REVIEW ARTICLE

The challenges of a small population exposed to multiple anthropogenic stressors and a changing climate: the St. Lawrence Estuary beluga

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Abstract

Until 2012, the St. Lawrence Estuary beluga population was considered stable with about 1100 individuals. An abnormally high number of calves reported dead that year triggered a population status reassessment. This review article summarizes the findings from this reassessment and various studies subsequent to it and provides an updated analysis of carcass recovery rates up to 2019. The 2013 review indicated a decreased incidence of cancer in adults, suggesting positive impacts from the regulation of toxic substances (e.g., PCBs and PAHs). However, the review also revealed that the population initiated a decline of ca. 1% per year in the early 2000s and had reached a size of ca. 900 individuals by 2012. This decline was accompanied by high inter-annual variability in calf survival and pregnancy rates and by more frequent peripartum complications among dead females. The change in population dynamics coincided with a shift in the St. Lawrence ecosystem structure and warmer environmental conditions, suggesting a link through effects on reproductive success and adult female body condition. This was supported by the continued high calf mortality after 2012 and a documented decline of fat reserves in beluga blubber from 1998 to 2016. Other factors, such as the exposure to chronic vessel noise, increasing whale-watching activities, high contaminant levels and episodic harmful algal blooms, may also be contributing to the long-term non-recovery and current decline of the population. The strong natal philopatry and complex social system of the beluga likely increase its vulnerability to extinction risk by limiting dispersal.

Keywords
Delphinapterus leucas; climate change; population dynamics; noise; food availability; cumulative effects

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Abbreviations
AIC: Akaike information criterion
CI: confidence interval
COSEWIC: Committee on the Status of Endangered Wildlife in Canada
DDT: dichlorodiphenyltrichloroethane
DFO: Fisheries and Oceans Canada
NAFO: North Atlantic Fisheries Organization
PAH: polycyclic aromatic hydrocarbon
PBDE: polybrominated diphenyl ether
PCB: polychlorinated biphenyl
POP: persistent organic pollutant
SLE: St. Lawrence Estuary

Historical context

The beluga (Delphinapterus leucas) has a discontinuous circumpolar distribution in the Arctic and Subarctic regions, with eight populations occurring in Canadian waters (Stewart & Stewart 1989; COSEWIC 2016). The lack of a dorsal fin and a relatively thick dermis (5–12 mm) make the species particularly well adapted to environments with seasonal and extensive ice cover. The population in the SLE, Canada, is at the southern limit of the species’ global distribution and probably established itself thereafter the Wisconsin glaciation (Harington 1977, 2008). The beluga persistence in the SLE is probably largely due to the combination of an extensive and seasonal sea-ice cover and the cold and productive environmental conditions that are maintained in this region in part from the influx and upwelling of Arctic waters of the Labrador Current (El-Sabh & Silverberg 1990).

The SLE beluga, like several other populations, undertakes seasonal movements, but its extent appears limited to a few tens or hundreds of kilometres (Mougin et al. 2010). The core of its distribution remains in the SLE year-round, but an unknown and likely variable proportion of the population moves eastward each fall to winter in the western Gulf of St. Lawrence (Fig. 1). While immigration from other populations might have been notable a century ago (Vladykov 1944), the SLE beluga population now has the lowest haplotype diversity of all beluga populations and shares none with other populations (Postma 2017; Skovring et al. 2019).
Their nuclear genetic diversity is similar to insular populations of mammals (Patenaude et al. 1994), suggesting insignificant contributions from neighbouring populations and a distinctly isolated population (Postma 2017).

The SLE beluga population declined from a pristine size estimated at about 8000 individuals in 1866 to a few hundreds of individuals at the end of the 1970s as a result of hunt and bounty programmes that thrived from the 1500s to the mid-1900s (Vladykov 1944; Reeves & Mitchel 1984; Hammill et al. 2007). Interest in the fate of this population progressively rose with governments and the public, leading to a complete halt of the hunt in 1979 and the population designation as Endangered in 1983 by the Committee on the Status of Endangered Wildlife in Canada (Pippard 1985). There were concerns at the time about high contamination of sediments and

Fig. 1 Relative density of marine traffic in the summer range of SLE beluga and their designated critical habitat. Traffic includes merchant ships, ferries and some of the whale-watching vessels equipped with automatic identification systems. Recreational vessels and two ferries operating downstream of the Saguenay River and connecting the north and south shores are not included. White areas indicate areas with no traffic. Adapted from Chion et al. (2009).
habitat loss from the development of a marina in Tadousac Bay and hydroelectricity-related damming of three rivers considered part of the SLE beluga critical habitat (Pippard 1985).

A carcass monitoring programme was implemented in 1982 (fully in 1983) to monitor the demography and causes of death, pathologies and other illness in this population through tissue sampling. Every effort was made to ensure that reporting of carcasses and data collection were consistent over the study period. For instance, the existence of the programme was advertised each spring through a publicity campaign requesting the public to report dead beluga. Each reported case was investigated closely to confirm species identity using photographs or visual confirmation by an expert. Field data and samples were collected by three or four experts over the 30-year study period, while the necropsies were conducted by five veterinary pathologists from the same institution (for details see Lesage, Measures et al. 2014; Lair et al. 2015).

In 1988, another programme was undertaken to monitor trends in distribution and abundance using a systematic and repeatable study design (for details see Gosselin et al. 2014). Specifically, photographic aerial surveys were flown every 3–5 years along equally spaced transect lines perpendicular to the main axis of the SLE using two planes, allowing a high-intensity (ca. 50%) coverage of the entire SLE beluga summer range in a single day.

These two ongoing programmes confirmed that the population numbered only a few hundred individuals (Kingsley & Hammill 1991) and that it was amongst the most contaminated marine populations on the planet (Martineau et al. 1987; Muir et al. 1990; Wagemann et al. 1990; Martineau et al. 1994), with an incidence of tumours much higher than observed in any other wild population (Bélond et al. 1993; De Guise et al. 1994).

From the 1980s through the 2000s, health effects from high contaminant loads were seen as the most imminent threat to the SLE beluga population (Bailey & Zinger 1995). As a result, research efforts were focused on monitoring population size and documenting pathologies, levels of toxic substances and their potential effects on the beluga immune and endocrine systems (reviewed by Kingsley 2002; Brousseau et al. 2003; Lebeuf 2009). Toxicological studies revealed that SLE beluga had a variety of persistent organohalogen pollutants in their tissues, which were mainly from agricultural and industrial origins. Among these pollutants were PCBs, DDT and their metabolites, heavy metals and PAHs (e.g., Martineau et al. 1987; Wagemann et al. 1990; Muir et al. 1996; Lebeuf 2009). Some of these toxic substances were suspected of inducing immunosuppression and of being responsible for the severity, high prevalence and diversity of lesions observed in the population (De Guise et al. 1994; Martineau et al. 1994; De Guise et al. 1995; De Guise 1998). PAHs, in particular, originating from aluminium smelters, were thought to be responsible for the high rates of cancer documented in the population (Martineau et al. 2002). High burdens of some of these persistent organic pollutants were also suspected of having negative effects on the population’s reproductive rate (Martineau et al. 1987). However, scientific evidence supporting correlations between pollutants and pathologies or high rates of cancer in SLE beluga and whether the sample of recovered carcasses was representative of the living population were debated at the time (Addison 1989; Theriault et al. 2002; Hammill et al. 2003).

The long-term programmes monitoring SLE beluga carcasses and abundance revealed a stable number of recovered carcasses over time at around 15 individuals per year (median), including up to three calves annually (reviewed by Lesage, Measures et al. 2014) and a population growing at an abnormally slow pace, if growing at all (Hammill et al. 2007). A demographic analysis conducted in 2007 concluded that the adult mortality rate was comparable to Arctic beluga populations, the emigration rate was likely minimal and the recruitment rate was low and likely responsible for the apparent lack of recovery of the population (Hammill et al. 2007).

Reasons for the non-recovery and low recruitment rate of the population could be a result of one or multiple anthropogenic stressors acting on SLE beluga or their habitat, either independently or in synergy (Beauchesne et al. 2020). SLE beluga live downstream of highly industrialized and urbanized regions discharging toxic chemical substances which eventually reach their habitat (Martel et al. 1986; Gearing et al. 1994; Gobeil et al. 1995; Viglino et al. 2004; Lebeuf & Nunes 2005). The St. Lawrence River represents the main seaway to interior North America (Fig. 1). Several thousands of ships transit through the SLE beluga habitat each year, chronically elevating ambient noise levels (Simard et al. 2010; Simard et al. 2014; Gervaise et al. 2015). Adding to this noise and potential disturbance are the multiple ferries and the multi-million dollar whale-watching industry operating year-round or seasonally in the SLE beluga critical habitat (Gervaise et al. 2012; Ménard et al. 2014). The collapse of commercially exploited fish stocks in the 1990s (Worm & Myers 2003; Savenkoff et al. 2007; Cairns et al. 2014) and the population increase of potential competitors, such as grey (Halichoerus grypus) and harp (Phoca vitulina groenlandicus) seals (Hammill et al. 2015; Hammill et al. 2017), have modified ecosystem trophodynamics (Savenkoff et al. 2007) with potential consequences on prey availability to SLE beluga.
The canaries in the mine

In 2012, an alarmingly high number of newborn calves \((n = 16)\) were reported dead in the SLE through the carcass recovery programme. This calf stranding rate was five times higher than the annual maxima of three recorded since the beginning of the programme in 1983 and 16 times higher than the annual median calf stranding rate of one during the period 1983–2019 (Lesage, Measures et al. 2014). Four years earlier (in 2008), eight calves and 14 adults had been found dead, an event attributed at the time to a particularly intense and well-documented harmful algal bloom of the toxic dinoflagellate *Alexandrium tamarense* in the SLE, which had also killed hundreds of seabirds, seals and fish (Starr et al. 2017). A re-examination of beluga deaths between 1983 and 2012 revealed that an unusually high number of calves (10) had also washed ashore in 2010. This series of high calf mortality events over a short period (2008–2012) led to a coordinated review in 2013, examining long-term information available on multiple aspects of the biology, ecology and the environment of SLE beluga. In years subsequent to the review, additional research efforts were undertaken (e.g., diet, noise exposure and impacts, and health effects of contaminants), along with a close monitoring of anomalies across multiple demographic parameters. The following sections summarize the findings from the 2013 review (see also DFO 2014) and the various studies subsequent to it. This review article also updates the analysis of sex- and age-specific carcass recovery rates presented as part of the 2013 review (Lesage, Measures et al. 2014) by incorporating seven additional years of data: 2013–19.

Population trends and dynamics

Abundance estimates obtained from visual line-transect surveys (2001–09) and photographic strip-transect surveys (1988–2009) indicated that the 2009 estimate—the latest available for this population at the time of the 2013 review—was the lowest in both survey time series (Gosselin et al. 2014). A photogrammetric analysis to detect calves and yearlings (zero and one year in age) in aerial images from the photographic surveys (Desrosiers 1994) indicated a decrease in these age class proportions within the population from 15–18% in the 1990s to 3–8% in the 2000s (Gosselin et al. 2014). The incorporation of this information in an age-structured population model, along with abundance estimates from the photographic survey time series (1988–2009; summarized by Gosselin et al. 2014) and numbers of dead calves and non-calves reported through the carcass monitoring programme (Lesage, Measures et al. 2014), indicated that several aspects of SLE beluga demography had changed over time (Mosnier et al. 2015). Modelling showed that over the period 1983–2012, the population had shifted from a relatively stable to an unstable state starting around 1999 (Fig. 2). From 1983 to 1998, the calf mortality rate was relatively stable (median values: 14–27%), as was the pregnancy rate (about 30%, with small peaks every three years) and the population age structure (50.5% adults, 42% juveniles and 7.5% calves). Starting in 1999 and at least until 2012 (last year modelled), an increase in inter-annual variability was noted in calf mortality and pregnancy rates, along with a decline in the proportion of juvenile individuals (1–7 years of age) to 33% in 2012 (Mosnier et al. 2015). There was no clear pattern in adult mortality over the modelled period, 1983–2012 (Fig. 2). However, the model predicted a change in female reproduction during the last few years of the modelling exercise, when calf mortality was above the long-term median. The reproductive cycle shifted from a three-year cycle, with a third of mature females pregnant each year, to a two-year cycle, with about half of the females pregnant (Mosnier et al. 2015). These predicted changes in mortality and reproductive schedules resulted in the population shifting from a stable or slightly increasing trend at about 0.13% per year until 2002 to a declining trend at an average annual rate of −1.13% per year and an estimated population size of 889 individuals (95% CI: 672–1167) in 2012 (Mosnier et al. 2015). An additional photographic aerial survey was conducted in 2019 to update population abundance estimates, trends and demographic parameters; results are pending.

These model-derived demographic parameters were consistent with independent empirical observations made in the context of a long-term photo-identification programme of live SLE beluga from 1989 to 2012 (Michaud 2014). Specifically, an index for the proportion of immature individuals in the population (grey animals), which included juveniles and young adults, increased slightly from 1989 to the mid- and late 2000s. Consistent with model predictions, this period was followed by a decline in the representation of immature individuals in the population until the last year modelled in 2012 (Michaud 2014). Field data collections also indicated that years predicted to be associated with high pregnancy rates in the model for the period 2004–2012 were followed by years of high calf production in the field, building our confidence in model predictions.

Carcass monitoring and cause of death

The number of beluga recovered each year through the carcass monitoring programme was variable between
years for the different age and sex classes but appeared relatively stable from 1983 to at least 2007 (Fig. 3). Trends emerged for some age and sex classes when carcasses from the most recent period (2013–19) were added to the time series examined during the review (1983–2012). Specifically, the number of dead calves, which had increased in 2008–2012, continued to remain higher than the long-term maxima observed from 1983 to 2007 (three individuals) in 10 of the 12 most recent years, that is, 2008–2019 (Fig. 3). In contrast, adult animals (aged eight years or more) tended to be found less often after 2010. A sex-specific analysis revealed that this decrease in adult mortality was entirely attributed to the decline in adult male representation among carcasses (Fig. 4a). The exact timing of this decline, however, remains uncertain. The most parsimonious model suggests that it began in the late 1990s (Fig. 5; generalized additive model with penalized cubic splines smooth, best model based on lowest AIC: 3 knots, edf = 2.0, df = 4, deviance explained = 33.4%). However, the downward trend is heavily driven by the 2010–19 data; allowing more knots to the generalized additive model or using a different statistical approach (e.g., broken stick or sequential analysis) would have resulted in a decline at a later date (not shown). The decrease in male representation among adult carcasses was strongly correlated (Pearson’s r = –0.86) with an increase in the sex ratio among deaths towards adult females (Fig. 4b; best model: linear regression, df = 3, p < 0.001) even if the absolute number of adult females among carcasses, although variable among years, did not change over the study period (best model: linear regression, df = 3, p = 0.70; Fig. 4a). Similar to adult females, the number of carcasses of juvenile individuals (i.e., 1–7 years old) remained stable over the study period (Fig. 3b; best model: linear regression, df = 3, p = 0.09, AIC for linear and non-linear regressions were identical).

Hypotheses to explain this sex-related difference in carcass recovery rate are multiple. While a brief overview is provided here, a thorough investigation of each hypothesis will be required before any conclusion can be
drawn from the observed patterns. At least during part of the year, adult males and females are spatially segregated, leading to differential exposure to environmental stressors and differential access to prey items (Michaud 1993; Mosnier et al. 2016; Lesage et al. 2020). The lower representation of adult males in the recovered carcasses could arise from a progressive decrease in detection probability, for instance, as a result of a change in adult male distribution (e.g., due to ecosystem change) towards areas that are less urbanized or less accessible to humans. Males could also have decreased in abundance over time compared to females, resulting in fewer relative deaths. Male survival may also have increased compared to adult females, for instance, through a progressively lesser exposure to habitat degradation or threat factors, or through access to more profitable prey compared to females. Alternatively, the health and survival of both males and females might have improved because of a reduction of potential stressors (e.g., contamination) or higher prey availability but additional sources of mortality prone to affect females might have eliminated these benefits for them. While one of these hypotheses is elaborated below, all of them are plausible. The timing of the change in adult male representation in the recovered carcasses will be a key in assessing the likelihood of each of these hypotheses.

Complete necropsies of 222 SLE beluga conducted between 1983 and 2012 revealed that the main cause of death for this population was an infectious disease.
which accounted for half of the cases in juveniles and one-third (32%) of all cases (Lair et al. 2016). Malignant neoplasia (tumours) was the cause of death for 14% of the beluga and affected only beluga of 28 years or older, whereas dystocia or post-partum complications caused the death of 19% of sexually mature females. A small fraction of deaths were attributed to vessel strikes (4%), primary starvation (2%), fishing-gear entanglement and (1%) and intoxication (one case). The absence of neoplasia was also noted for SLE beluga born after 1971 (Lair et al. 2016).

Lair et al. (2016) postulated the existence of a relationship between the reduction of gastrointestinal cancer and the cessation in 1976 of PAH emissions from changes in the aluminium production process and stricter environmental regulations. A recent study examining PAH–DNA adducts in beluga intestine lent support to this hypothesis (Poirier et al. 2019). Banning PCBs in 1979, which reduced the levels of PCBs in beluga over time (Lebeuf, Measures et al. 2014), may also have contributed to the progressive disappearance of some neoplastic diseases in this population (Martineau et al. 2002; Lair et al. 2016). A reduction in non-sex-specific cancers should have improved the health of, and increased survival among, both males and females (i.e., reducing numbers in reported carcasses). Whether the improvement in survival occurred equally and simultaneously for males and females, and whether it was progressive or instead occurred when contaminant levels fell below a certain level in beluga tissues, is unknown. Parturition-associated mortality, a cause specific to females and which occurred only occasionally from 1983 to the early 2000s, increased in frequency in the 2000s (Lair et al. 2016). Assuming a persistence of this phenomenon in the 2010s, it could have masked an increase in the survival rate of adult females from an overlapping reduction in cancer-related mortalities and led to the stable pattern in female mortality observed over time (Fig. 4a).

**Potential stressors and their effects**

There are also competing, although not exclusive, hypotheses for the observed high calf mortality and change in the demography of the SLE beluga population. These include health problems or impaired reproductive functions from the toxic effects of specific chemical compounds on endocrine and immune functions; perturbation of critical activities (e.g., parturition and foraging) from chronic vessel exposure or isolated or repeated disturbance; fishery- or climate-related changes in ecosystem structure leading to a decrease in habitat quality and prey availability or quality; and recurring stochastic events such as harmful algal blooms (DFO 2014).

SLE beluga have a long history of exposure to a variety of POPs, such as DDT, PCBs, Mirex, toxaphene, PAH, dioxins and furans (reviewed by Lebeuf 2009; see also Lebeuf, Raach et al. 2014; Simond et al. 2017). Decreasing in the environment since at least 1987, these pollutants are unlikely to be the primary factor for the recent increase in mortality and population decline (Muir et al. 1996; Lebeuf, Raach et al. 2014); however, new contaminants from the 1980s, such as toxic flame retardants (PBDEs), have accumulated at an exponential rate in beluga and their environment in the 1990s, with PBDEs remaining at maximum levels in beluga since that time (De Wit 2002; Lebeuf, Measures et al. 2014; Lebeuf, Raach et al. 2014; Simond et al. 2017).

PBDEs and several other POPs can have endocrine-disrupting effects in a variety of species including humans and other mammals, with possible impacts on reproduction, immune system, behaviour and offspring development (reviewed by Lebeuf 2009; Lebeuf, Raach et al. 2014; see also Costa et al. 2014; Yu et al. 2015). An analysis comparing PBDE concentrations in adult females and calves among three periods (1995–2001, 2002–07 and 2008–2011) was unsuccessful at identifying the role of PBDEs in the recently elevated frequency of complications.
at parturition and mortality of calves (Lebeuf, Measures et al. 2014). This is not to say that PBDEs have no detrimental effects on health but that the study, with its specific design, does not reveal a direct relationship between PBDE burden and beluga health linked to mortality. A negative effect of PBDE homologs on blood thyroid hormone levels has been documented in beluga in Svalbard, Norway (Villanger et al. 2011). The negative effects of hypothyroidism on parturition and foetal health and survival, and of foetal exposure to PBDE on calf survival and cognitive abilities, have been reported in humans and other animals (reviewed by Lair et al. 2016). While there is yet no evidence for PBDE-related hypothyroidism in SLE beluga, a correlation between thyroid-related gene expression and levels of some emerging halogenated flame retardants has been documented in this population (Simond et al. 2019). However, a mismatch between the period of increased calf mortality and parturition complications (the late 2000s) and of high concentrations of PBDEs in SLE beluga—which have plateaued since 1998—indicates a need for further investigation (Lair et al. 2016).

Chronic exposure to noise and its potential effects on marine mammals is a concern worldwide (Clark et al. 2009; Boyd et al. 2011; Williams et al. 2015). In the SLE, chronic commercial shipping and ferry operations, as well as seasonal whale-watching and recreational activities, raise ambient noise in the beluga habitat (Simard et al. 2010; McQuinn et al. 2011; Gervaise et al. 2012). Between 5000 and 6000 transits from merchant ships are reported each year through the critical habitat of SLE beluga (Fig. 1; McQuinn et al. 2011). There are 11 marinas near SLE beluga critical habitat, four ferry services operating within its bounds, resulting in tens of thousands of ferry transits each year, and a multi-million dollar whale-watching industry, mainly targeting species other than beluga but operating within its critical habitat (Ménard et al. 2014). Vessel traffic from tourism and whale-watching activities peak in July–August, when SLE beluga give birth (Sergeant 1986). While SLE beluga appear more tolerant to noise than beluga populations in the Arctic, where there is comparatively little shipping (Finley et al. 1990), SLE beluga are not immune to deleterious effects from noise exposure. Noise can have negative effects on animals in a number of ways, including masking vocalizations and important signals, reducing acoustic space, diverting attention, disrupting natural behaviour, habituation (i.e., learned deafness) and causing chronic stress (Erbe et al. 2018). Noise can also limit the energy and time allocated to critical activities like foraging or impair social interactions by, for example, interfering with the acoustic bond between mother and calf (Tyack & Clark 2000). Vocal responses to noise have been documented in SLE beluga (Lesage et al. 1999; Scheifele et al. 2005), and their abandonment of Tadoussac Bay established after a marina is suspected to be a consequence brought by disturbance from increased vessel traffic and noise exposure (Pippard 1985). Ferries and other large ships can reduce beluga acoustic space considerably. At the mouth of the Saguenay River where as many as three ferries operate simultaneously, acoustic space has been reported at a fraction (30%) of its expected value under natural conditions during half of the time, regardless of the beluga call frequency (Gervaise et al. 2012). Beluga emit contact calls, which can be heard over ranges that are 18 times larger for adults and subadults than for newborn calves (Vergara et al. 2021 [this special cluster]). In a key habitat of the SLE (Baie Sainte-Marguerite in the Saguenay River), vessel noise reduced the communication range from a median of 6.7–2.9 km for adults and subadults and from a median of 360–170 m for newborn calves. These 57% and 53% reductions in communication range for calls with a known function are evidence that anthropogenic noise from marine traffic may impair communication, and especially mother–call contact.

The exposure of SLE beluga to noise varies between habitats and is highest near the shipping lane and at the Saguenay River mouth, where a marina and most of the whale-watching companies are based (Fig. 1; McQuinn et al. 2011). Noise exposure is lowest in habitats located along the south shore, where traffic of all sources is currently light (McQuinn et al. 2011; Lesage, McQuinn et al. 2014; Roy & Simard 2015). On a daily basis, merchant ships in the SLE repeatedly expose a substantial proportion of the beluga population (15–53%, depending on traffic direction) to noise levels likely to interfere with their behaviour, and the vast majority of exposed animals (72–81%) are females with calves or juveniles (Lesage, McQuinn et al. 2014).

Shipping traffic in the SLE has likely been lighter in the 1990s and 2000s compared to the 1970s or 1980s (Dionne 2001). While interannual variations are noted, there is no evidence for any substantial increase in traffic volume at least until 2012 (McQuinn et al. 2011; Ménard et al. 2014). Whale-watching operations in traditional sectors of activity (i.e., in the SLE downstream of the Saguenay River where large whales are located; Fig. 1) have also remained relatively constant over time, even decreasing slightly in the 2010s (Ménard et al. 2014). However, newly established companies specifically targeting beluga have increased whale-watching operations from 2004 to 2012 in non-traditional sectors located in the upstream portion of the SLE and within the beluga critical habitat (Ménard et al. 2014). Recreational activities have also increased in this sector since 2004 (Ménard et al. 2014). Indicators for the volume of recreational activities in the beluga summer
range indicated a coincidence of the high calf mortality events from 2010 and 2012 with a high number of boat-nights in marinas, high per cent days with low-infration reports for interactions with beluga and clement weather conditions, that is, low precipitation and high air temperature (Ménard et al. 2014). Physical or acoustic interferences from motorized or non-motorized vessels (e.g., kayaks) with parturition, lactation or mother–calf acoustic bond can induce parturition-related complications or abandonment of the calf. The continuous presence of a human observer during parturition has been reported to induce such complications in cows (Mee 2008). Although it is unknown if disturbance during the calving season was specifically higher during the high mortality events from 2010 and 2012 than in previous years, the persistence of high calf mortality after 2012 in spite of variable weather conditions suggests that this factor might not be the main cause for the increased calf mortality observed over the past decade.

Stochastic events such as harmful algal blooms were also among the factors examined in the context of the recent increase in calf mortality. Red tides caused by blooms of the dinoflagellate *Alexandrium tamarense* and their associated release of phycotoxins are responsible for outbreaks of paralytic shellfish poisoning and can pose serious health risks for humans and marine organisms (Geraci et al. 1989; Landsberg 2002). Three major red tides were documented over the past two decades in St. Lawrence: in 1996, 1998 and 2008 (Scarratt et al. 2014; Starr et al. 2017). The 2008 red tide was especially well-documented, and a relationship between high cell concentrations of *A. tamarense* near Tadoussac, in the heart of SLE beluga habitat, and the high mortality of beluga was established for that year (Scarratt et al. 2014). There was no evidence for such red tides during the high mortality events from 2010 and 2012 (Scarratt et al. 2014).

**Diet and ecosystem change**

Changes in the St. Lawrence community structure have occurred over time as a result of climate variability and anthropogenic factors such as the collapse of demersal fish stocks from overexploitation and ocean warming (Worm & Myers 2003; Savenkoff et al. 2007). These environmental changes likely have modified prey availability, with potential effects on predator population dynamics (e.g., Plourde, Grégoire et al. 2014). A time-series analysis of 28 ecosystem indices of habitat quality, prey availability and physical environmental variability of the Gulf of St. Lawrence over the period 1990–2012 highlighted a major shift in the ecosystem in the late 1990s, that is, approximately at the time when beluga population dynamics started to change (Plourde, Galbraith et al. 2014). The ecosystem shifted from a period of above long-term averages, characterized by a relative abundance of prey and cool environmental conditions, to a period of below long-term averages when Gulf of St. Lawrence stocks of Atlantic spring herring (*Clupea harengus*), Atlantic cod (*Gadus morhua*) and other large demersal fish (NAFO Division 4T) had collapsed and were at their lowest biomass, and when the environment was subject to below-average ice conditions and above-average water temperatures. Trend analysis of these variables since 1971 also identified unprecedented extremes in environmental conditions from 2010 to 2012, with particularly high temperatures and low indices for ice duration and volume (Plourde, Galbraith et al. 2014). These extreme conditions have persisted at least to 2019 (Galbraith et al. 2020).

An optimum ice condition for calf survival is likely to exist based on the U-shaped relationship predicted between ice duration and calf mortality for SLE beluga (Williams et al. 2017). In the Arctic, females tend to avoid very high or very low ice cover in the fall (Barber et al. 2001), supporting model predictions indicating that extreme ice cover may be unfavourable to calf survival. While the physical presence of ice could reduce energy output for adults and calves by offering shelter against storm (Barber et al. 2001), its relative benefits to beluga energetics remain uncertain. Ice also influences the potential for prey to concentrate in the marginal ice zone (Fréchet 1990) and the biomass and timing of spawning of some important beluga prey, such as capelin (*Mallotus villosus*; Buren et al. 2014; Lesage et al. 2020). As a result, extremes in ice conditions might affect calf survival by reducing energy intake of pregnant females during winter (Truchon et al. 2013) and during critical periods such as the spring when intense feeding on prey such as spawning capelin and herring likely occurs before calving (see Lesage et al. 2020). Elevated sea surface temperatures in the summer months have also been predicted to have a negative effect on calf survival (Williams et al. 2017).

Calf mortality was also inversely related to biomass indices for some beluga prey such as spring herring (NAFO Division 4T) and demersal species (NAFO Division 4Tw), which, together with sea-ice indices and sea-surface temperature, accounted for 82% of the deviance of the model predicting calf mortality (Williams et al. 2017). Conversely, prey availability and variations in environmental conditions were not good predictors of adult female mortality and fecundity. Such an effect on reproductive success or survival would be likely to arise from extreme changes in adult female body condition.

The coincidence of the shift in the St. Lawrence community structure and environmental conditions with the
change in beluga population dynamics, including higher calf mortalities in periods of extreme conditions (starting around 2010), supports the hypothesis highlighting the role of ecosystem change in beluga population dynamics, potentially through effects on reproductive success and adult female body condition. While data on diet remain fragmentary for this population, studies from the 1930s and from more recent years (1987–2019) suggest that pelagic species such as capelin and herring, sand lance (Ammodytes spp.) and large demersal species such as cod (Gadus morhua and G. ogac) and redfish (Sebastes sp.) may be seasonally important in the SLE beluga diet (Vladykov 1946; Lesage 2014; Ferchou 2019; Lesage et al. 2020). Evidence for a potential shift in SLE beluga diet has been reported through an assessment of SLE beluga dietary sources and trophic position from 1988 to 2012, which identified a progressive decrease (approximately one part per mil, equivalent to one trophic level), in a tracer of carbon sources starting around 2003 (Lesage 2014). A decline in a tracer of trophic position (nitrogen isotope ratios), although overall modest at approximately 0.6‰ over 10 years, was also observed over the same period (Lesage 2014). Whether these changes reflect a change in beluga diet or a change in the isotopic signature of their prey is unknown.

Some prey species and feeding periods may be critical for pregnant or lactating females to ensure successful reproduction and weaning of calves (Lockyer 1986; Jönsson 1997; Miller et al. 2011). This appears to be the case for the SLE beluga population during spring feeding, when beluga rapidly accumulate lipid in their blubber (Casgrain 1873; Vladykov 1944; see Lesage et al. 2020). Food intake increases 1.5- to twofold in late pregnancy and peaks at fourfold their normal intake during the first month after giving birth in captive beluga (Kastelein et al. 1994). There is currently little information available on SLE beluga body condition. However, a recent study using the sum of blubber fatty acids as an indicator of beluga body condition suggests that it has deteriorated from 1998 to 2016 in both males and females (Bernier-Graveline et al. 2021). Whether a deteriorating body condition in pregnant females played a role in the increased calf mortality and parturition-associated complications observed in recent years is currently unknown (Lair et al. 2016). In other mammals, such as cows, nutritional stress during the last semester of pregnancy has been associated with an increased risk of parturition-associated complications (e.g., dystocia) (Gruner 1973). Disturbance of parturient females and exposure to endocrine-disrupting chemicals such as PBDEs also represent risk factors linked to parturition-associated complications (Lair et al. 2016).

Relative importance of the multiple stressors

A recent population viability analysis incorporating environmental stressors provided insight into how the SLE beluga population was likely to respond to changes in environmental conditions and varying levels of three anthropogenic threats (Williams et al. 2017). The analysis incorporated PCB contamination as a proxy for other contaminants, the combined biomasses of demersal fish and spring herring, and noise-mediated effects on foraging efficiency as potential stressors acting on population dynamics, while accounting for potential effects of changing sea-ice conditions and seawater temperatures. With the caveat that this model was data limited and could only account for the relationship between these stressors or the environment and calf mortality, the study indicated that prey availability contributed the most to changes in calf mortality, followed by noise and PCBs. Among the management scenarios that explored potential reductions in threats separately and in combination, only the scenario in which all three threats were mitigated simultaneously was likely to create the necessary resilience for the population to persist in the context of the current climate change and warming conditions (Williams et al. 2017).

Conclusions and perspective

The very slow population growth rate observed during the 1980s and 1990s despite the total halt on hunting suggests that population growth was already constrained by anthropogenic stressors or sub-optimal environmental conditions associated with local climate, or both. While a harmful algal bloom event was likely responsible for the unusually high calf mortality in 2008, the timing of the change in population dynamics and of increased calf mortality suggests that warming environmental conditions and a decrease in sea-ice extent and duration, with their cascading effects on ecosystem structure (Smetacek & Nichol 2005), may be key drivers of the current population decrease and the persistently high calf mortality. The apparent deterioration of SLE beluga body condition over the past 20 years supports a hypothesis in which climate or fishery-driven impacts on energy balance likely play an important role, by reducing prey availability, quality or both. A similar, although weak, trend in body condition (blubber thickness) and body growth rate has been documented in Beaufort Sea beluga from 1997 to 2008, although causative factors remain unclear (Harwood et al. 2014). Poorer body condition can also arise solely from an increase in energy output caused by anthropogenic stressors, for example, repeated avoidance response
or disruption of foraging by vessel proximity, or noise-mediated reduction of foraging efficiency (Lusseau & Bejder 2007; Christiansen et al. 2015; Pirotta et al. 2015; Senigaglia et al. 2016). In food-limited conditions or in the case where females are accompanied by a dependent calf, the ability of an individual to cope with these stressors can be reduced (e.g., Williams & Loren 2009; Senigaglia et al. 2016). Therefore, while a food-mediated effect on body condition and reproductive output represents a highly plausible explanation for the poorer body condition and current demographic situation of the SLE beluga population, other stressors should not be dismissed as potential contributors to the challenges that this beluga population face in its changing environment.

The vulnerability of a population to changes in its environment is a function of its exposure, sensitivity, and adaptive capacity (Dawson et al. 2011). Exposure depends on the magnitude and rate of change; the sensitivity depends on life-history traits such as habitat specificity, dietary specialization, range size, population size and reproductive rate; and adaptive capacity depends on habitat loss and fragmentation, and also on intrinsic traits such as phenotypic plasticity, and dispersal and colonization capacity (Ofori et al. 2017).

The beluga is fundamentally a polar species that has adapted over evolutionary time to an environment with a permanent sea-ice cap as its most defining character (O’Corry-Crowe 2008; Kovacs et al. 2011). Beluga in the SLE are at the southern margin of the species distribution, in an environment where only annually forming ice is present, and its formation is only seasonal. Beluga have persisted in this environment over the past 10 000 years (Harington 2008) and are presumably well-adapted to an extreme and dynamic environment given their anatomy and primary Arctic distribution. Climate models are forecasting that the SLE will be a nearly ice-free environment within 50–80 years (Ruest et al. 2016). The rate of the current climate warming and the high degree of isolation of the SLE beluga population may hinder their capacity to adapt to these changes and to survive in an ice-free or nearly ice-free and warm environment (O’Corry-Crowe 2008).

Beluga as generalist feeders are better adapted than specialists for coping with changes in prey assemblages (Kassen 2002; Dennis et al. 2011). In the Canadian Arctic, where temperate species such as sand lance and capelin are expanding their range as a result of warming conditions (Wassmann et al. 2011), beluga responded to this change in prey availability by incorporating these new prey items into their diet (Kelley et al. 2010; Marcoux et al. 2012), although with unknown consequences related to prey quantity and/or quality on their health. Whether a similar pattern will or has occurred in SLE beluga as a result of the expansion or higher abundance of more temperate fish or invertebrate species is unknown (Lesage et al. 2020). The apparent deterioration of their body condition over the past 20 years (Bernier-Graveline et al. 2021) suggests that adaptation to the changing ecosystem has been challenging. The reduction in ice cover may increase interspecific competition as other species extend the duration of their stay or expand their range (Kingsley 2002). The continuing decrease in oxygen levels (i.e., hypoxia) in waters of the SLE and Gulf of St. Lawrence, partly resulting from the decreased proportion of the oxygen-rich Labrador Current water entering this region (Gilbert et al. 2005), is likely to affect biodiversity and productivity (Diaz & Rosenberg 2008), with potential effects on the availability of some of beluga prey (Lavaud et al. 2019).

With 1938 as a baseline, SLE beluga occupy an area that represents 65% of their historical range (COSEWIC 2014). Whether this range contraction is the consequence of population decline, as appears to be the case for Cook Inlet beluga (Rugh et al. 2010), or whether it occurred as a result of changes to their habitat or over-hunting at some historical sites is unknown. The species is known for its natal philopatry to summer concentration areas and to migratory circuits within regions, with migration routes and site fidelity being culturally learned (Caron & Smith 1990; de March & Postma 2003; Turgeon et al. 2012; Colbeck et al. 2013; O’Corry-Crowe et al. 2018). This cultural inheritance likely also explains the resilience of beluga movements to inter-annual variation in sea-ice conditions, although anomalies in migration patterns can occur in highly anomalous ice years (O’Corry-Crowe et al. 2016). Fidelity to breeding or feeding sites, such as those documented in beluga and other species like southern right whale (Eubalaena australis) and humpback whales (Megaptera novaeangliae), may make these species more prone to extinction risk compared to other cetaceans or to pinnipeds (Valenzuela et al. 2009; Baker et al. 2013; O’Corry-Crowe et al. 2018). Stable matrilineal groups have long been seen as the cornerstone of beluga society when interpreting the aspects of their ecology and behaviour, but a recent study indicates that beluga groupings can also be organized around close paternal relatives (O’Corry-Crowe et al. 2020). The complex social system and the high degree of sociality of beluga and other odontocetes may further increase their vulnerability to extinction risk (Wade et al. 2012). Predictions related to these life-history traits, which appear to reduce the potential for dispersal to new areas more suitable for their survival, could explain in part the absence of recolonization of abandoned or over-hunted sites (Mosnier et al. 2010; Colbeck et al. 2013).
St. Lawrence Estuary beluga in a pressured and changing ecosystem

V. Lesage

Together, these characteristics make the beluga a species that is considered moderately vulnerable to climate change or environmental variability (Laidre et al. 2008). However, vulnerability is likely to be higher for the SLE beluga population given its location at the southern margin of the species range and the low probability of migrants from other populations replenishing its declining population (O’Corry-Crowe et al. 2016). The frequency, intensity and geographic extent of harmful algal blooms appear to be increasing worldwide as a result of climate change, coastal eutrophication and other environmental perturbations and are expected to increase the frequency of associated mortality events (Van Dohal 2000; Gulland & Hall 2007). These multiple predictions underscore the need for immediate actions to be taken to reduce the population’s exposure to non-climatic stressors.

Since the first recovery plan was proposed in 1995 for SLE beluga (Bailey & Zinger 1995), multiple actions have been implemented to improve the quality of its acoustic and physical environment (reviewed in DFO 2017). The regulation of toxic chemical substances such as PCBs and PAHs, although not in response to beluga decline—they were regulated in the 1970s—may have reduced the incidence of cancer in adult beluga (Lair et al. 2016), showing that appropriate specific actions can result in measurable positive impacts on the population. While recovery measures implemented to date have allowed the population to stabilize prior to 2000, they were insufficient to allow the population to grow at the targeted 2% growth rate and to cease decreasing (DFO 2017). Additional actions are necessary and have been proposed to address the various threats (DFO 2017, 2020). In the case of noise, a number of actions have been thoroughly examined for their potential benefits and capacity to decrease exposure across beluga habitats (e.g., Lesage, McQuinn et al. 2014; Chion et al. 2017). The variety of threats, the number of stakeholders that need to be involved in the implementation of recovery actions and inherent delays associated with regulatory processes will remain a challenge, but the timely implementation of additional and necessary recovery measures is possible.

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St. Lawrence Estuary beluga in a pressured and changing ecosystem

V. Lesage
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