**Title:** Tropicalization of temperate ecosystems in North America: The northward range expansion of tropical organisms in response to warming winter temperatures

**Running Title:** Tropicalization of temperate ecosystems

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**Abstract**

Tropicalization is a term used to describe the transformation of temperate ecosystems by poleward-moving tropical organisms in response to warming temperatures. In North America, decreases in the frequency and intensity of extreme winter cold events are expected to allow the poleward range expansion of many cold-sensitive tropical organisms, sometimes at the expense of temperate organisms. Although ecologists have long noted the critical ecological role of winter cold temperature extremes in tropical-temperate transition zones, the ecological effects of extreme cold events have been understudied, and the influence of warming winter temperatures has too often been left out of climate change vulnerability assessments. Here, we examine the influence of extreme cold events on the northward range limits of a diverse group of tropical organisms, including terrestrial plants, coastal wetland plants, coastal fishes, sea turtles, terrestrial reptiles, amphibians, manatees, and insects. For these organisms, extreme cold events can lead to major physiological damage or landscape-scale mass mortality. Conversely, the absence of extreme cold events can foster population growth, range expansion, and ecological regime shifts. We discuss the effects of warming winters on species and ecosystems in tropical-temperate transition zones. In the twenty-first century, climate change-induced decreases in the frequency and intensity of extreme cold events are expected to facilitate the poleward range expansion of many tropical species. Our review highlights critical knowledge gaps for advancing understanding of the ecological implications of the tropicalization of temperate ecosystems in North America.

**Keywords:** climate change, climate extreme, extreme cold events, poleward migration, range expansion, tropicalization, warming, winter climate change

**Introduction**

In the face of accelerating climate change, ecologists are increasingly challenged to better understand the ecological impacts of changes in the frequency and intensity of extreme climatic events (Parmesan et al., 2000; Smith, 2011; USGCRP, 2018). Near tropical-temperate transition zones in North America, decreases in the frequency and intensity of extreme cold events are expected to enable the poleward range expansion of cold-sensitive tropical organisms (Figure 1; Table S1), sometimes at the expense of temperate organisms (Weiss & Overpeck, 2005; Carter et al., 2018; Cavanaugh et al., 2019). Although astute scientists and naturalists have long recognized the critical influence of winter temperature extremes on tropical species’ range limits (e.g., Shreve, 1911; Davis, 1940; Lonard & Judd, 1991), the ecological effects of extreme cold events have been understudied (Boucek et al., 2016).

In this communication, our aim is to highlight, within the context of climate change, the critical ecological role that extreme cold events play within tropical-temperate transition zones in North America. We begin with sections that consider the frequency and return interval of ecologically relevant extreme cold events, recent winter warming trends, and the general physiological effects of winter cold temperature extremes. Next, in taxa-specific sections focused on terrestrial plants, coastal wetland plants, coastal fish, sea turtles, terrestrial reptiles, amphibians, manatee, and insects, we examine cold sensitivity and assess the influence of extreme cold events on northern range limits and ecosystem properties. These taxa-specific sections also examine climate change effects and gauge the potential for ecological tropicalization, which is a term used to describe the transformation of temperate ecosystems by poleward-moving tropical organisms in response to warming temperatures (Vergés et al., 2014). Following the taxa-specific sections, we include sections focused on the critical role of range expansion pathways, microclimates, and thermal refugia. Then, we finish with a section that highlights knowledge gaps for advancing understanding of climate change impacts.

**Extreme cold events are infrequent but ecologically important**

Across the globe, there is much variation in the frequency, intensity, and ecological significance of extreme cold events within tropical-temperate transition zones. In some transition zones—especially those in the Southern Hemisphere (e.g., South America, Africa, and Australia, where cold Antarctic air is modulated by large stretches of ocean before reaching other continents)—winter temperature means, rather than extremes, appear to govern the poleward range limits of tropical organisms (Osland et al., 2017b). However, tropical-temperate transition zones in the Northern Hemisphere (e.g., North America and Asia, where cold Arctic air can rapidly descend across large, high latitude land masses) are more strongly controlled by extreme cold events rather than mean winter temperatures (Stuart et al., 2007; Osland et al., 2017b). Cold air outbreaks from the poles into tropical-temperate transition zones are more frequent and intense in the Northern Hemisphere compared to the Southern Hemisphere (Smith & Sheridan, 2020).

Tropical-temperate transition zones in North America occur primarily in the southern United States and northern Mexico (see warm-to-cold color transitions in Figure 1). Across these zones, extreme cold events act in a similar manner as other ecologically important but infrequent disturbances such as hurricanes, fires, or floods, which can cause mass mortality, lead to large losses of aboveground biomass, reset successional dynamics, and, in the most severe cases, produce ecological regime shifts (Turner, 2010; Peters et al., 2011; Pickett & White, 2013). For example, extreme cold events can lead to ecological regime shifts in coastal wetlands, where cold-induced mangrove forest mortality events are followed by landscape-scale mangrove-to-marsh transitions, and conversely, the absence of cold events can lead to marsh-to-mangrove transitions (Osland et al., 2017a; Cavanaugh et al., 2019).

Within the USA, Florida, Alabama, Mississippi, Louisiana, Texas, New Mexico, Arizona, and California are states that possess tropical-temperate transition zones (Figure 1). In northern Mexico, Tamaulipas, Nuevo Leon, Coahuila, Chihuahua, Sonora, and Baja California are states with tropical species whose northern range limits are governed by cold temperature extremes. In northern Mexico and the southwestern and south-central United States, extreme cold temperature controls on tropical species’ distributional limits are also apparent at higher elevations along altitudinal gradients (Niering et al., 1963; Brusca et al., 2013; Bojórquez et al., 2019).

The Florida peninsula and the Sonoran Desert have been two notable hotspots for research on the effects of extreme cold events on tropical species’ range limits in North America. The Florida peninsula spans a comparatively conspicuous tropical-temperate transition zone, with more cold-sensitive tropical species in south Florida and more cold-tolerant temperate species in north Florida. Ecologists working in Florida have long studied the critical ecological role of extreme cold events (Davis, 1940; Olmsted et al., 1993; Boucek et al., 2016). The Sonoran Desert region is the most tropical of North America’s four great deserts (Shreve & Wiggins, 1964; Turner et al., 1995). The structurally diverse vegetation of the Sonoran Desert is unique in that it includes many species of columnar cacti and leguminous trees, both of which are rare or nonexistent in the other three shrub-dominated North American deserts (i.e., the Great Basin, Mojave, and Chihuahuan Deserts). Occasional catastrophic cold events (i.e., cold-induced mass mortality events) have historically been common and comparatively well studied in the northern Sonoran Desert along the Arizona-Sonora border (e.g., Shreve, 1911; Bowers, 1981; Steenbergh & Lowe, 1983).

Most winters in North-America’s tropical-temperate transition zones are comparatively mild, without major frequent cold events that lead to landscape-scale mortality or physiological damage. Major ecologically relevant cold events may occur just once every 20–30 years. To illustrate this point, we present 100 years of air temperature data from a location within the tropical-temperate transition in central Florida (Figure 2). In this example, the return time for ecologically relevant cold events is several decades (see red line within grey box), and most winters are comparatively warm, ecologically benign, and disturbance free (see blue peak outside of grey box). Due to accelerating climate change, the number of years with ecologically relevant cold events is expected to decrease (USGCRP, 2017; Carter et al., 2018; USGCRP, 2018). In other words, the red and blue lines in Figure 2 are expected to shift to the right, which would be above the threshold for cold damage or mortality. Although the temperatures along the y-axis and the position of the ecological threshold are species and location-dependent, the general relationships describing the return time and frequency of ecologically relevant extreme cold events apply to many other tropical-temperate transition zones across North America (i.e., all areas with warm-to-cold color transitions in Figure 1). Tropical-temperate ecotones in North America are highly sensitive to small changes in winter temperature regimes, and the northern limits of climatic zones that support tropical cold-sensitive species are expected to move northward under 2°C, 4°C, and 6°C increases in winter temperature extremes (Figure 3b,c,d, respectively).

Across North America, scientists use different terms to describe extreme cold winter temperature events, and these terms are often taxa or region specific. Some of these terms are introduced in the taxa-specific sections of this review. For example, sea turtle ecologists use the term “cold stunning event” (Griffin et al., 2019) and desert ecologists have used the term “catastrophic freeze” (Bowers, 1981). In Florida, citrus physiologists have used the terms “hard freeze” and “impact freeze” (Attaway, 1997). Other synonomous terms for winter cold temperature extreme events include “severe cold event” (Stevens et al., 2016), “cold snap” (Mazzotti et al., 2011), and “extreme cold spell” (Boucek et al., 2016). In the taxa-specific sections, we often use the term appropriate to the particular taxon we are describing to remain consistent with that literature.

**Recent decades have warmer winters with fewer extreme cold events**

To evaluate winter trends over recent decades, temperature data were analyzed from four representative sites in North America’s tropical-temperate transition zone. From west to east, these sites included San Francisco (California, on the Pacific Ocean), Tuscon (Arizona, in the Sonoran Desert), New Orleans (Louisiana, along the northern Gulf of Mexico coast), and Tampa (Florida, on the east coast of the Gulf of Mexico). These sites were selected to span the regions discussed in this review, but also based on the length and quality of climate records. All four sites had a common period of record going back to at least 1948.

At each of the sites, three different winter parameters were investigated to determine how winters have been changing as the climate warms (USGCRP, 2017). These parameters include: (1) mean winter temperature, (2) the single coldest temperature recorded each winter, and (3) the number of days each winter with temperatures of 0°C or less (i.e., the number of subzero days). The clear tendencies in the time series (Figure 4; Table S2) show that: (1) average winter temperatures are increasing, (2) the extreme minimum temperatures recorded each winter season are becoming milder, and (3) the number of annual subzero days is declining.

The decline in the number of subzero days is especially apparent in San Francisco (Figure 4a). Prior to 1980, most winters in San Francisco had subzero events, with the winter of 1949 having a total of 17 days with subzero temperatures. However, for the latter 40 years of the time series (1981–2020), only 14 days in San Francisco had subzero temperatures. There has only been one subzero event since 1999 in San Francisco and no subzero events since 2008. For Tucson, there was a warming trend for all three winter parameters (Figure 4b) (see also: Weiss & Overpeck, 2005). For New Orleans, the time series shows a relatively low number of subzero events in the late 1940s and 1950s, followed by a very high number of subzero events in the 1960s–1980s, and fewer subzero events over the past three decades (Figure 4c). Examination of the last 30 years (1991–2020) in New Orleans shows a total of 165 days with subzero temperatures, while the previous 30-year period (1961–1990) had 480 subzero days. Data from Tampa show a similar temporal pattern to New Orleans with 23 subzero days over the past 30 years (1991–2020) compared to 107 subzero days in the 30 years prior (1961–1990) (Figure 4d).

**Physiological effects of cold temperature extremes: why are warming winters relevant?**

Warmer winters with fewer extreme cold events can lead to the range expansion of tropical cold-sensitive species due to the decreases in the frequency of events that lead to cold-induced physiological damage and mortality. In the taxa-focused sections, we provide specific examples of the damage and mortality caused by extreme cold events. This section covers some of the more general (i.e., cross taxa) physiological effects of extreme cold temperatures, which, even at temperatures above 0°C, can have significant effects on organismal function due to both direct damage to macromolecules as well as the indirect effects of reductions in enzyme activity. Low temperatures can lead to protein denaturation (Todgham et al., 2007), induction of the damaging gel phase of membranes (Wilson et al., 2003), and breakage of double- and single-stranded DNA (Yao & Somero, 2012). In addition, low temperatures can reduce the activity of the sodium-potassium ATPase enzyme, which is essential for the maintenance of ion balance in all organisms, and can induce chill coma in animals—a reversible loss of neuromuscular function that can lead to chilling injury (MacMillan & Sinclair, 2011; Overgaard & MacMillan, 2017). The vast majority of species are sensitive to chilling and die or are severely damaged long before the risk of ice formation in their body fluids (Lyons, 1973).

While deionized water freezes at 0°C, freezing points in organisms are set by a combination of factors including body water content, solute content, and the presence of ice nucleators and ice binding proteins and may or may not be relevant for any given species (Wilson et al., 2003). For example, the tropical fruit fly *Drosophila melanogaster* has a freezing point of around–20°C, but dies of chilling injury at around –5°C (Strachan et al., 2011). Tissue ice formation induces a separate series of stresses than chilling. Extracellular ice formation is frequently lethal to most organisms because it can induce osmotic damage as solutes are excluded from the growing ice lattice, cause mechanical damage to cell membranes and walls, and potentially cause ischemia-reperfusion damage (Khan & Vincent, 1996; Krivoruchko & Storey, 2010; Toxopeus & Sinclair, 2018). However, extracellular ice formation can be common in plants, where ice nucleation occurs in the saturated extracellular air spaces of leaves or other tissues. Upon extracellular ice formation, water is distilled from the relatively high water content of plant cells (~70 to 80%), resulting in their freeze-dehydration. This is because the water potential of ice is -1.16 × Temperature (°C) and is thus always lower than the water potential of adjacent liquid water within cells (Nobel, 2005). Taken together, these differences in the physiological effects of cold temperature extremes show that it is essential to consider cold tolerance as individual-, life stage-, and species-specific phenomena when conducting comparative studies or evaluating climate change responses.

**Terrestrial plants**

Agronomists, botanists, foresters, and naturalists have long observed the importance of cold and freezing air temperatures for the survival, fitness, and productivity of a wide range of plant types.

There are many terrestrial tropical and subtropical plant species in North America whose northern or altitudinal range limits are governed by extreme freezing air temperature events (Holdridge, 1967; Sakai & Larcher, 1987; Woodward, 1987). Many tropical plants undergo chilling stress when temperatures fall below about 10°C (Levitt, 1980). However, there are some subtropical plant species that can tolerate temperatures a few degrees below 0°C but are then severely damaged or killed by temperatures between approximately –5°C and –15°C (Lonard & Judd, 1991; Osland et al., 2020a; Osland & Feher, 2020).

In the desert ecosystems of the arid southwestern United States and northwestern Mexico, extreme freeze events control the northern and altitudinal distribution of foundation plant species like the giant saguaro cactus (*Carnegiea gigantea*) (Figure S1) (Shreve, 1911; Niering et al., 1963; Steenbergh & Lowe, 1976), desert ironwood tree (*Olneya tesota*) (Turnage & Hinckley, 1938), organ pipe cactus (*Stenocereus thurberi*) (Turnage & Hinckley, 1938; Bowers, 1981; Parker, 1988), senita (*Pachycereus schottii*) (Turnage & Hinckley, 1938; Felger & Lowe, 1967; Nobel, 1980c),creosote bush(*Larrea tridentata*) (Pockman & Sperry, 1997; D'Odorico et al., 2010; Ladwig et al., 2019), Joshua tree (*Yucca brevifolia*) (Smith et al., 1983; Loik et al., 2000; Dole et al., 2003), little-leaf paloverde (*Parkinsonia microphylla*), and many others. Landscape-scale pulses of adult plant death in the Sonoran Desert are caused by catastrophic freezes (Steenbergh & Lowe, 1977; Bowers, 1981; McAuliffe, 1996). Low air temperatures affect small individuals of cactus species more than large individuals because stems of small diameters reach lower minimum apical temperatures than larger stems under the same environmental conditions (Steenbergh & Lowe, 1969; Nobel, 1980a; Nobel, 1980b). On the other hand, the energy budgets of small-statured cacti can be coupled to microhabitat features, and warmed or insulated by objects such as rocks, other vegetation, litter, and snow (Nobel et al., 1991; Geiger et al., 2012). The distributional limits of many Sonoran Desert plants are regulated by winter freezes, and catastrophic freezes are known to cause large-scale die-offs of plants in this region (Shreve, 1914; Felger & Lowe, 1967; Bowers, 1981). For example, the northern limit of senita cactus (*P. schottii*), just north of the Arizona-Sonora border, is set by winter freeze-induced seedling mortality (Parker, 1989). The northern boundary of the Sonoran Desert itself is typically defined by the northern extent of such characteristic plants as saguaro (*C. giganteus*), triangle-leaf bur sage (*Ambrosia deltoidea*), canyon ragweed (*Ambrosia ambrosioides*), little-leaf palo verde (*P. microphylla*), and blue palo verde (*Parkinsonia florida*). This northern border coincides with the isotherm beyond which freezing temperatures have occurred lasting longer than 24 hours (Hastings, 1963; Bowers, 1981). Notably, some lineages of southwestern origin have escaped constraints from freezing and have spread widely into the North American temperate zone. For example, the diminutive prickly pear cactus, *Opuntia fragilis*, has evolved substantial freeze tolerance that allows persistence in many locations in North America, including elevations up to 3048 m in Colorado and to 56°N latitude in British Columbia, Canada (Loik & Nobel, 1993b).

In the Sonoran Desert, freeze events that lead to mass plant mortality are rare. Bowers (1981) analyzed weather patterns for Tucson (Arizona) from 1894 to 1979, noting that there is much variation in plant-relevant winter cold events ranging from mild freezes with little effect on native plants, to severe freezes that inflict damage to frost-sensitive plants, to catastrophic freezes that kill or injure many species of plants over large areas. Catastrophic freezes usually occurred within 17 days of the winter solstice and are characterized by the co-occurrence of low minimum temperatures and many consecutive hours of freezing. Between 1946 and 1979, there were four such catastrophic freezes that caused widespread frost damage to Sonoran Desert plants (Bowers, 1981). In Saguaro National Park, near Tucson, elevated adult mortality of saguaros since 2010 is probably related to a single event, a severe freeze in February 2011 that was the first major extended period of below-freezing temperatures in southern Arizona since the 1970s (Orum et al., 2016). The 2011 freeze event killed approximately 36% of older adult plants in Saguaro National Park and also killed substantial numbers of saguaros at Tonto and Organ Pipe Cactus National Monuments (Swann et al., 2018).

Along the southwestern coast of the United States, patterns of warmer winter temperatures are greatly influenced by elevation and proximity to the Pacific Ocean (Figure 1). Temperature gradients can be quite steep over short distances or elevations, resulting in an irregular mosaic of thermal stress environments across the region. This pattern is especially pronounced in comparison to the north-south temperature zonation of topographically uniform Florida (Figure 1). Although air temperatures rarely fall below 0°C along the Pacific coast, minimum temperatures can fall to –10°C just a few kilometers inland. Thus, plants in California can be exposed to simultaneous drought and high temperature stresses in summer, while drought may persist into winter and be accompanied by rare freeze events (Davis et al., 2007). The restriction of some species in California to coastal habitats may be related to this spatial zonation. For example, the coastal barrel cactus (*Ferocactus viridescens*) is limited in occurrence to within ~15 km of the coast in San Diego County, California, which may be due to its limited ability to survive episodic freezing (Loik & Nobel, 1993a). Rare freezing events in southern California can also lead to zonation of plant communities along elevation gradients (Davis et al., 2007; Kelly & Goulden, 2008). The ability of southern California chaparral shrubs to persist in habitats visited by rare freezing events depends on the ability of xylem cells to tolerate or avoid embolism formation, in which air bubbles form within trachea or vessels of the xylem and block water flow (Davis et al., 1999). Such freeze-induced emboli can be fatal, and tradeoffs between xylem flow and resistance to cavitation are important for the survival of freezing and the geographic distribution of coastal species such as *Ceanothus megacarpus* (big pod ceanothus) and *Malosma laurina* (laurel sumac)(Langan et al., 1997). The integrity of membranes and the photosynthetic apparatus within mesophyll leaf cells are also susceptible to freeze-induced damage, and patterns of damage are dependent on age, season, and location, yet consistent with observations of spatial and elevational distributions of the chaparral species *M. laurina*, *C. megacarpus*,*Rhus ovata* (sugar sumac), and *Ceanothus spinosus* (redheart) (Boorse et al., 1998). Considering the low frequency and high impact of subzero air temperature events in coastal California, more information on the role of rare episodic freezing would help better understand how this region became a hotspot of biodiversity.

In semi-arid ecosystems of the south-central United States (e.g., south Texas) and northern Mexico, extreme freeze events dictate the northern distribution of many tropical plant species, including native woody plant species like great leadtree (*Leucaena pulverulenta*), anacahuita (*Cordia boissieri*), and coyotillo (*Karwinskia humboldtiana*) (Lonard & Judd, 1985; Lonard & Judd, 1991; Bojórquez et al., 2019). In the more humid and seasonal wet-dry ecosystems of south and central Florida, extreme freeze events govern the northern distribution of entire forest biomes (Greller, 1980) as well as the range limits of plant species such as poisonwood (*Metopium toxiferum*), gumbo limbo (*Bursera simaruba*), cocoplum (*Chrysobalanus icaco*), and pigeon plum (*Coccoloba diversifolia*) (Myers, 1986; Box et al., 1993; Olmsted et al., 1993). In response to warming winter temperature extremes, many of these tropical freeze-sensitive native plant species are expected to move northward, within arid, semi-arid, and humid climates of western, central, and eastern parts of the continent, respectively (Box et al., 1999; Weiss & Overpeck, 2005; Carter et al., 2018) (Figure 3). Species-specific rates of poleward range expansion are, however, highly variable and greatly influenced by the presence of plant traits that foster rapid movement into novel climates, and also interactions with other global change factors that may hinder or accelerate range expansion (Parker, 1993; Davis & Shaw, 2001; Zhu et al., 2012). Long-distance seed dispersal, high seed production and viability, rapid growth and establishment, and the ability to outcompete incumbent species are some of the traits useful for species to migrate in response to climatic shifts.

In North America, there are many invasive non-native plant species that are expected to also expand northward in response to winter climate change. Many of these species are currently abundant in tropical parts of Florida, which is a state that has become a prominent example of the negative ecological impacts of invasive non-native species. In the past century, many of Florida’s native ecosystems have been affected, and in some cases transformed, by invasive non-native organisms (Simberloff et al., 1997; South Florida Ecosytem Restoration Task Force, 2015; Rodgers et al., 2018). In 2019, the Florida Exotic Pest Plant Council identified 84 invasive non-native plant species as Category I Invasive Plant Species, which are species that are “altering native plant communities by displacing native species, changing community structures or ecological functions, or hybridizing with natives” (FLEPPC, 2019). Many of these plants are tropical freeze-sensitive species that are expected to expand northward into other parts of the southeastern United States in response to warming winters. Brazilian pepper (*Schinus terebinthifolius*) is one of Florida’s most abundant and problematic invasive plant species and a prime example of the potential negative effects of invasive non-native range expansions (Osland & Feher, 2020). Millions of dollars are spent each year in Florida to mitigate the negative ecological impacts caused by Brazilian pepperinvasions (Hiatt et al., 2019). In response to warming winters, this freeze-sensitive species is expected to migrate northward and transform ecosystems in north Florida and across much of the Gulf of Mexico and south Atlantic coasts of the United States (Osland & Feher, 2020). Melaleuca (*Melaleuca quinquenervia*) (Turner et al., 1997), Old World climbing fern (*Lygodium microphyllum*) (Hutchinson & Langeland, 2014), and Australian pine (*Casuarina equisetifolia*) (Morton, 1980) are other problematic freeze-sensitive non-native invasive species that could move northward from Florida into other parts of North America in response to warming winters.

Warming winters are also expected to facilitate the northward spread of invasive non-native plant species in the arid southwestern United States. Abatzoglou and Kolden (2011) analyzed a suite of climate projections for the mid-21st century and concluded that changes in the length of the freeze-free season in this region will favor cold-intolerant, invasive, annual grasses [e.g., cheatgrass, *Bromus tectorum* (Griffith et al., 2014); red brome, *Bromus rubens*; buffelgrass, *Pennisetum ciliare*]. These grasses have been increasing in abundance and extent in the southwestern United States over the past three decades (Esque & Schwalbe, 2002; Abatzoglou & Kolden, 2011). Buffelgrass is an aggressive perennial grass species introduced from southern Africa that is transforming desert ecosystems in the Sonoran Desert by outcompeting native species, modifying ecohydrological processes, and altering fire regimes (Williams & Baruch, 2000; Marshall et al., 2012). Because buffelgrass is sensitive to freezing temperatures (Cox et al., 1988; Hussey & Bashaw, 1996; Stair et al., 1998), warming winter temperature extremes are expected to allow buffelgrass to continue expanding northward and affect desert ecosystems to the north of its current distribution (Martin et al., 2015; Jarnevich et al., 2018). Buffelgrass is also expanding its altitudinal distribution up mountain slopes, and in doing so carrying wildfires from the grass-invaded desert scrub to higher-elevation woodlands, and vice versa (Williams & Baruch, 2000; Esque & Schwalbe, 2002; Abatzoglou & Kolden, 2011).

**Coastal wetland plants**

Coastal wetland ecosystems in the tropical-temperate transition zone are highly sensitive to changes in the frequency and intensity of extreme cold events, which can produce landscape-scale ecological transformations (Lugo & Patterson-Zucca, 1977; Sherrod & McMillan, 1985; McKee et al., 2012). Coastal wetlands are abundant in North America, especially along the low-lying and comparatively flat coasts of the Gulf of Mexico and Atlantic Ocean, where tropical mangrove forests and temperate salt marshes support critical habitat and provide many valuable ecosystem goods and services (Barbier et al., 2011). Both coasts span tropical-temperate transition zones that generate comparatively large gradients in coastal wetland ecosystem structure and function, and there are strong linear and nonlinear relationships between winter temperature regimes and coastal wetland plant community composition, vegetation height, aboveground biomass, and productivity (Feher et al., 2017; Gabler et al., 2017; Osland et al., 2018b).

Extreme freeze events greatly influence whether a tidal saline wetland in eastern North America is dominated by woody trees or grass-like plants (i.e., mangrove forest or salt marsh graminoid plants, respectively) (Stevens et al., 2006; Cavanaugh et al., 2019; Osland et al., 2019a). Salt marsh graminoid plants are freeze tolerant and dominate coastal wetlands in climates with colder winters (Pennings & Bertness, 2001). In contrast, extreme freeze events lead to mangrove physiological damage or mortality (Ross et al., 2009; Lovelock et al., 2016; Osland et al., 2020a), which means that mangrove forests are most abundant in tropical and subtropical climates (e.g., south and central Florida, Mexico’s Gulf of Mexico coast, and Mexico’s Pacific coast) (Duke et al., 1998; Saenger, 2002). The three most common mangrove species in North America are the black mangrove (*Avicennia germinans*), red mangrove (*Rhizophora mangle*) (Figure S1), and white mangrove (*Laguncularia racemosa*) (Felger et al., 2001; Tomlinson, 2016; Osland et al., 2018a). Two other species often considered as mangroves or mangrove associates are the buttonwood mangrove (*Conocarpus erectus*) and the sweet mangrove (*Maytenus phyllanthoides*), which can be sporadically common in tropical coastal wetlands, especially at higher intertidal elevations near landward transitional margins (Felger et al., 2001; Whitmore et al., 2005; Mendelssohn et al., 2017). The poleward range limits of mangroves along the Gulf of Mexico and Atlantic coasts are strongly governed by extreme freeze events. *Avicennia germinans* is the most freeze-tolerant species and its range extends furthest north—into salt marshes in north Florida, Louisiana, and Texas (Armitage et al., 2015; Cook-Patton et al., 2015; Osland et al., 2020b). On the Pacific coast of Mexico, all five mangrove species are severely frost-damaged during exceptional cold events in the Gulf of California (Felger et al., 2001; Felger, 2004).

In eastern North America, the adaptive capacity of mangroves to changing winter temperature regimes is very high. Long-distance dispersal mechanisms (Van der Stocken et al., 2019a; Van der Stocken et al., 2019b) and rapid peat development (McKee, 2011; Krauss et al., 2014; Osland et al., 2020c) enable mangrove forests to adapt to climate change. Mangroves produce floating propagules that can be carried long distances by oceanic currents, especially when propagule release coincides with tropical storms that produce storm surges and rapidly moving northward surface currents (Kennedy et al., 2016; Van der Stocken et al., 2019b; Kennedy et al., 2020). Paleoecological records indicate that during the last glacial maximum (~19,000 years ago) mangrove range limits were much further south in Central America (Sherrod & McMillan, 1985). Due to a warming climate, the northern range limit of mangroves has moved poleward during the Holocene (Woodroffe & Grindrod, 1991; Kennedy et al., 2016), with local, decadal-scale range expansion and contraction phases occurring in the absence or presence of extreme freeze events, respectively (Osland et al., 2017a; Cavanaugh et al., 2019). Mangroves have been expanding in the southeastern United States for approximately 30 years, since the last major freeze events occurred in the late 1980s (Cavanaugh et al., 2014; Osland et al., 2017a; Cavanaugh et al., 2019). In response to accelerating anthropogenic climate change in the Anthropocene, warming winters are expected to accelerate the pace of poleward range expansion along the Gulf of Mexico and Atlantic coasts of North America, as mangrove forests move northward and replace grass-dominated salt marshes in much of Texas, Louisiana, and north Florida (Osland et al., 2013; Gabler et al., 2017; Cavanaugh et al., 2019). Many temperate salt marsh ecosystems in eastern North America are expected to become tropicalized by poleward-moving mangrove forests. However, rising sea levels are expected to simultaneously lead to the landward migration of tidal saline wetlands at the expense of upslope and upriver ecosystems (Doyle et al., 2010; Enwright et al., 2016; Borchert et al., 2018); thus, mangroves forests could also be replacing inland terrestrial and freshwater forests as they expand northward.

In North America, mangrove range limits along the Gulf of Mexico and Atlantic coasts are highly responsive to changing winter temperature regimes. However, mangrove range limits along the Pacific coast of North America, in Mexico within Sonora and Baja California Sur, are governed by a different combination of macroclimatic drivers (Felger et al., 2001; Whitmore et al., 2005; Glenn et al., 2006). Near these Pacific coast range limits, extreme freeze events are rare and only occasionally affect mangroves and other coastal plants (Felger et al., 2001). Mangrove range limits in western North America have been understudied; however, the abundance and distribution of mangroves in those areas are thought to be most strongly influenced by aridity (Brusca et al., 2006; Glenn et al., 2006) and dispersal constraints (Van der Stocken et al., 2019a; Bardou et al., 2020). Highly arid conditions (i.e., very low rainfall, high evapotranspiration, and minimal freshwater inputs) promote the development of hypersaline conditions that are inhospitable to mangroves and many other coastal wetland plants (Flores-Verdugo et al., 1992; Arreola-Lizárraga et al., 2004; Osland et al., 2019a). The northernmost mangrove forests in western Mexico occur in hypersaline coastal lagoons that entirely lack freshwater riverine input, and compared to more southerly, taller mangroves that occur in less saline estuarine habitats, the growth form of these northern mangroves is greatly stunted (Felger, 2004; Brusca et al., 2006; Glenn et al., 2006). Moreover, predominantly southward-moving oceanic currents (e.g., the coastal California Current) combined with large distances between suitable estuarine habitat are thought to hinder mangrove range expansion into climate-appropriate estuaries beyond current Pacific coast range limits (Cavanaugh et al., 2018; Van der Stocken et al., 2019a; Bardou et al., 2020). For example, winter air temperature regimes indicate that mangroves may be able to survive in San Francisco Bay’s coastal wetlands (Osland et al., 2019a), which are approximately 1500 km north of the current mangrove range limit in Baja California.

**Coastal fishes**

One of the early uses of the term “tropicalization” was to describe the effects of range-expanding tropical fish species into temperate reef ecosystems (Vergés et al., 2014). Within tropical-temperate transition zones across the globe, warming winter ocean temperatures are expected to increasingly lead to the poleward range expansion of tropical fish species (Figueira & Booth, 2010; Nakamura et al., 2013). Along the Pacific coast of North America, the distributions of fishes are strongly affected by ocean currents and upwelling events (Carlisle et al., 2015; Páez-Osuna et al., 2016); however, winter temperature effects on the distribution of some tropical species have been reported in shallow waters of the Gulf of California (Thomson & Lehner, 1976; Lehner, 1979). The warm-temperate ocean waters of southern California, and the subtropical waters of the Gulf of California (Mexico), show a warming trend during the past century (Lluch-Belda et al., 2009). As this trend is likely to continue, poleward expansion of tropical fish species is expected, and this phenomenon appears to already be taking place (González-Cuéllar et al., 2013; Fernández‐Rivera Melo et al., 2015; Booth et al., 2018).

The influence of winter temperatures on subtropical fishes is more apparent along coasts of the Gulf of Mexico and Atlantic Ocean, where extreme cold events periodically lead to mass mortality events (i.e., conspicuous fish kills that receive media and public attention). For example, an extreme cold event in 2010 resulted in fish kills across Florida’s Gulf of Mexico and Atlantic coasts (Boucek & Rehage, 2014; Stevens et al., 2016; Purtlebaugh et al., 2020), which span ~6 degrees of latitude and ~2200 km of coastline. Another prime example stems from severe freeze events in 1983 and 1989, which produced fish kills across the region, from Texas to Florida. In Texas alone, the 1983 and 1989 events killed at least 31 million fish, comprised of 103 species (Martin & McEachron, 1996). Along these eastern North American coasts, the poleward distributions of subtropical fishes—for example, common snook (*Centropomus undecimalis*, Figure S2) (Howells et al., 1990), smalltooth sawfish (*Pristis pectinate*) (Poulakis et al., 2011), gray snapper (*Lutjanus griseus*) (Hare et al., 2012), goliath grouper (*Epinephelus itajara*), and tarpon (*Megalops atlanticus*) (Mace et al., 2017)—are ultimately limited by their cold tolerance. This is also the case for freshwater fishes introduced from the tropics, some of which can tolerate moderate salinity and occupy the upper reaches of estuaries (Idelberger et al., 2011; Greenwood, 2017).

The frequency at which the lower lethal water temperature (~10°C) of subtropical fishes is reached greatly influences whether a species occurs in a region, and if so, its population status. Recent population dynamics of a well-studied gamefish, common snook, highlight the critical ecological role of winter temperatures. In 2010, a severe cold event resulted in greater than 75% reductions of snook populations in some Florida estuaries (Stevens et al., 2016). These findings led to the temporary closure of the recreational snook fishery along the Gulf of Mexico coast, and recovery of the populations took up to four years. Since then, a decade of mild winters has allowed for expansion of the species approximately 200 km beyond the historic distribution of snook (Anderson et al., 2020; Purtlebaugh et al., 2020). Adaptive behaviors (e.g., movement to thermal refugia during winter) have likely made this range expansion possible (Stevens et al., 2018) and illustrate some plasticity in the life history traits of subtropical fish species.

Just as for other taxa, changes in the frequency, intensity, and duration of cold temperature extremes will greatly influence fish distributions and migrations. The frequency of extreme events is important because combined effects of consecutive events over short periods of time can elicit different responses by communities (Stevens et al., 2006; Boucek & Rehage, 2014). The intensity of a cold event, and the context of its onset, determine the time available for fishes to respond. For example, in 2008, a rapid decrease in temperature that occurred in Florida during an otherwise mild winter resulted in mortality of subtropical species (Blewett & Stevens, 2014). If given more time, these species can respond to cold temperatures by moving to deep, wind-protected shorelines (Blewett & Stevens, 2014; Scharer et al., 2017). However, extended periods of cold weather allow near-surface water temperatures to equilibrate to air temperatures, and any thermal refuge provided by stratification at deeper sites may eventually be lost due to mixing. A well-studied cold event in 2010 had a large impact on fish due in part to its long duration, nearly a full week (Boucek et al., 2016). Although the event had only moderate effects on subtropical coastal vegetation such as mangroves, the effects on subtropical fish populations were extreme. The long duration of cold likely reduced the efficacy of deeper sites that would have otherwise provided thermal refugia during a shorter-term event (Blewett & Stevens, 2014; Boucek et al., 2017).

The ecological and economic implications associated with changes in winter temperatures and their effects on fisheries are complex and industry dependent (e.g., different for estuary vs. nearshore coastal fisheries). The ranges of tropical/subtropical estuarine fishes will likely expand poleward if winter temperatures continue to warm (Fodrie et al., 2010), and these species could support novel fisheries to the north of their current distribution. Some subtropical species already have adult ranges that extend to mid-Atlantic states—for example, gray snapper (Wuenschel et al., 2012), bull shark (*Carcharhinus leucas*) (Matich & Heithaus, 2012), smalltooth sawfish (Norton et al., 2012), goliath grouper (O’Hop & Munyandorero, 2016), and tarpon (Mace et al., 2017), but their juvenile habitat is largely limited to subtropical estuaries. Poleward expansion of subtropical fish nurseries may prompt efforts to identify and protect these habitats if new fisheries targeting subtropical species are encouraged by managers. Warming winter temperatures are also likely to alter the migrations (or behavior) of nearshore coastal fishes. Winter movements of coastal migrants to the south (e.g., the winter run of sailfish, *Istiophorus platypterus*) and return trips back north (e.g., the spring run of cobia, *Rachycentron canadum*) have economic and cultural significance to coastal communities (Colburn et al., 2016). Large and collaborative passive acoustic telemetry networks established along the Atlantic (Young et al., 2020) and Gulf of Mexico (Lowerre-Barbieri et al., 2019) coasts are allowing the timing and geographic extent of coastal migrations to be better quantified for many managed sharks, rays, and sport fish. The migrations of a representative coastal migrant, lemon shark (*Negaprion brevirostris*), was found to vary by ~200 km depending on the severity of winter (Reyier et al., 2014). Fewer coastal fishes moving along the coasts in response to winter temperature drivers would have dramatic effects on recreational and commercial fisheries that depend on the migrations. Range expansions and altered migration patterns could eventually require greater management coordination across U.S. states, especially for certain fishes that are managed at the state, not federal level. Mechanistic models can be used to proactively develop fisheries regulations and habitat protection efforts to support range-expanding species in novel areas.

**Sea Turtles**

As ectothermic (cold-blooded) reptiles, winter temperatures affect nearly all aspects of sea turtle ecology, and warming winter temperature regimes could affect their distribution and abundance in North America. Sea turtles have a complex life cycle, which includes several ontogenetic shifts in habitat use (Bolten, 2003; McClellan & Read, 2007; Seminoff, 2010). Near northern range limits, females usually deposit multiple clutches per season on nesting beaches during summer where incubation and hatching occur. In the tropics, they may have winter nesting seasons. Each clutch is separated by a period termed the internesting interval. After nesting is complete, the adults migrate to distant foraging areas where they spend several years before returning to mate and nest again, with the time between nesting migrations termed the remigration interval. Juveniles use shallow coastal habitats where they forage on benthic invertebrates, marine macroalgae, and seagrasses (Wallace et al., 2009; Jones & Seminoff, 2013; Williams et al., 2014). Cold temperature extremes in any of these habitats and life stages can have significant effects on sea turtle populations.

Cold temperatures affect reproduction in many ways including phenology, fecundity, and incubation (Davenport, 1997; Hamann et al., 2003; Hawkes et al., 2009). All follicles that will become eggs are developed at foraging areas; thus, cold temperature extremes at winter foraging sites can result in the development of fewer follicles and smaller clutch sizes (Hamann et al., 2003; Lamont & Fujisaki, 2014). Cold temperatures at foraging grounds can also affect remigration intervals by constricting the foraging season, which limits nutrient acquisition (Hays, 2000). During the nesting season, cold temperature extremes can increase internesting intervals and result in fewer clutches and lesser reproductive output (Schofield et al., 2009).

Perhaps the most significant and striking effect of extreme cold on sea turtles is direct mass mortality following stranding. While most freshwater and terrestrial turtle species that inhabit temperate environments hibernate to survive extreme cold (Claussen et al., 1991; Ultsch et al., 1999), sea turtles typically migrate to warmer waters to escape winter temperature extremes. Many sea turtle species utilize high latitude, resource-rich bays during summer and migrate to deeper or warmer waters in winter, especially along the U.S. Atlantic coast (Hawkes et al., 2011; Seney & Landry Jr, 2011; Lamont & Iverson, 2018). However, when extreme cold impacts an area suddenly and turtles have not already moved, they can suffer immobility, which can lead to stranding and eventually death. In the sea turtle literature, these mass stranding occurrences are called “cold stunning events” (Still et al., 2005; Foley et al., 2007; Anderson et al., 2011). During cold events in 2010 and 2018, for example, more than 2,900 sea turtles stranded in St. Joseph Bay, a foraging site in northwestern Florida (Lamont et al., 2018) (Figure S3). The 2010 event resulted in the mortality of 434 individuals (Avens et al., 2012). Water temperatures below 8°C to 10°C can significantly affect sea turtle physiology (Davenport, 1997). At these temperatures, blood chemistry can be altered (Anderson et al., 2011) and turtles can become lethargic and float at the water’s surface (Milton & Lutz, 2003; Avens et al., 2012). Mortality rates increase significantly as temperatures drop below 5°C to 6°C (Schwartz, 1978).

Some sea turtles along the Pacific coast of North America and in the Gulf of Mexico tend to winter in place (Madrak et al., 2016; Lamont et al., 2018), and green turtles have been observed active and foraging in water temperatures as low as 15.8ºC (Hanna et al., In review). The low temperatures elicit a languid response during which turtles spend extended periods inactive on the seafloor, which may result in increased risk of direct harvesting (Felger et al., 1976). For example, turtles impacted by extreme cold in the Gulf of California (Mexico) can fall prey to illegal harvesting by local artisanal dive fisheries for spiny lobster (*Panulirus interruptus*) and sea cucumber (*Apostichopus californicus*) (Felger et al., 1976; Seminoff & Wallace, 2012).

In addition to affecting population abundance, cold-induced mortality can also affect population structure (Lamont et al., 2018). In two of the largest, recent cold stunning events in the Gulf of Mexico (2010 and 2018), a disproportionate number (> 45%) of small turtles (<40 cm straight carapace length) stranded (Avens et al., 2012; Lamont et al., 2018). This disproportionate mortality can affect the structure of juvenile turtle populations at temperate foraging sites. A 20-cm turtle will be 30 cm in 3–4 years and 40 cm in 5–7 years; thus, removal of small turtles during extreme cold events will affect larger size classes for many years after the event. This appears to result in a variable size class structure with peaks in abundance of larger individuals (40–49.9 cm straight carapace length) in the years immediately following an event, followed by a shift to smaller size classes as new recruits enter the population (Lamont et al., 2018). In addition to size structure, this mortality most likely impacts other aspects of demography such as growth and survival. Finally, cold temperatures themselves can reduce somatic growth rates (Avens et al., 2012).

As sea turtles move farther northward in response to warming air and ocean temperatures, the potential for sea turtle mortality events in novel, northerly locations may also increase and affect range expansion dynamics. For incubating embryos, nesting farther north increases the risk of exposure to cold temperature extremes. For example, on the Atlantic coast of North America, green turtles have typically been considered a tropical/subtropical species with rare nesting occurring in Georgia and South Carolina. However in 2011, a green turtle nested as far north as Delaware (Shamblin et al., 2018). Although that nest successfully hatched, a late nest deposited on a temperate beach risks exposure to early winter temperature extremes. Similarly, warmer summer waters at northern foraging areas, such as Cape Cod Bay in Massachusetts, allows juvenile turtles to travel farther north to resource-rich habitats. However, these areas are also more susceptible to rapid decreases in temperature during fall, thereby exposing turtles to potential cold stun stranding. In Cape Cod Bay, higher cold-stunning years occurred when the Gulf of Maine had warmer sea surface temperatures in late October through early November (Griffin et al., 2019). Similarly, green turtles in the northeastern Pacific occasionally nest as far north as the Cape Region of the Baja California Peninsula, Mexico (Tiburcio-Pintos & Cariño-Olvera, 2017), but they forage throughout the Gulf of California and north as far as southern California (Seminoff & Wallace, 2012). Warming winters may allow this threatened turtle to extend its nesting sites northward into the Gulf of California.

As warming winters increasingly lead to the tropicalization of coastal and marine ecosystems in North America, sea turtles are expected to increasingly migrate northward of their current historical range limits. However, given the complexity and sensitivity of so many stages of the sea turtle life cycle to winter temperature extremes, we expect that future changes in the distribution and migration of sea turtles will be highly dynamic and heavily influenced by interactions with other aspects of global change. For example, sea turtles have temperature-driven sex determination, such that the proportion of female embryos increases with incubation temperature (Yntema & Mrosovsky, 1982). Thus, many sea turtle populations are in danger of population feminization (i.e., production of females only) as global temperatures increase (Jensen et al., 2018). However, nesting range expansion onto more northerly, and thus cooler, beaches may help maintain population sex ratios.

**Terrestrial and freshwater reptiles**

Like their marine counterparts (e.g., sea turtles and sea snakes), terrestrial reptiles are highly sensitive to freezing and chilling temperatures. The thermal physiology of reptiles affects their behavioral patterns, movements, and survival. Extreme cold events can impose strong natural selection on wild populations (Campbell-Staton et al., 2017). Near their northern range limits in North America, winter temperature regimes greatly influence the physiology, distribution, and abundance of tropical snake, lizard, turtle, and crocodilian species.

For terrestrial reptiles, the tropical-temperate transition zone in North America is especially striking in south Florida, where both temperate and tropical reptile species are present. For example, south Florida is the only part of North America where cold-tolerant American alligators (*Alligator mississippiensis*) can be found within the same watersheds as tropical, cold-sensitive American crocodiles (*Crocodylus acutus*). The northern distribution of American alligators extends much farther into higher latitudes than American crocodiles, in part due to differences in their behavioral responses to extreme cold events. While the American alligator seeks refuge during cold events, the American crocodile attempts to compensate for cold air temperatures by basking in the sun, which actually increases its exposure to lethal air temperatures (Mazzotti et al., 2016).

In addition to controlling the northern distribution of native tropical reptiles, extreme cold events also constrain the distribution of invasive non-native species. Prolonged periods of extremely cold winter temperatures in south Florida are uncommon, yet occasional cold ‘snaps’ occur. Within the past century, the introduction of tropical non-native reptiles to Florida has had a tremendous impact on native ecosystems. Florida has had more non-native herpetofaunal species introductions than anywhere in the world (Engeman et al., 2011; Krysko et al., 2011; Fujisaki et al., 2015). Of the estimated 180 reptile and amphibian taxa that have been introduced to Florida, 63 species have become established, including 54 reptiles (48 lizards, 5 snakes, and 1 crocodile) (Krysko et al., 2016). Most of these introductions (86%) have occurred due the pet trade (Krysko et al., 2016), and the northern distributions of many of these non-native species are governed by winter cold temperature extremes.

Burmese pythons (*Python bivittatus*) are an especially harmful and well-known example of the negative ecological impacts of pet trade-driven tropical reptile introductions (Figure S4). This large constrictor snake is an opportunistic apex predator, whose expansion within and around Everglades National Park has decimated mammal, bird, and other prey populations (Dorcas et al., 2012; McCleery et al., 2015; Sovie et al., 2016). Burmese pythons are sensitive to cold temperature extremes, and mortality can occur during exposure to several days of extremely cold, wet weather (Mazzotti et al., 2011; Mazzotti et al., 2016). Pythons are projected to spread northward in response to warming winter temperature regimes (Jacobson et al., 2012). Two experiments tracking pythons through various freeze events in captive or lab settings revealed cold tolerance down to 0°C (Avery et al., 2010; Dorcas et al., 2011). Further, recent evidence of directional selection in genomic regions enriched for genes associated with thermosensation, behavior, and physiology was detected in pythons sampled before and after a 2010 south Florida freeze event (Card et al., 2018). With this new evidence that cold adaptation genes are evolving measurably faster, it is likely that pythons can extend their range further north in response to warming winters. Further, Card et al. (2018) reported that several of these genes are linked to regenerative organ growth, which is an adaptive response that controls organ size and function with feeding and fasting in pythons. Thus, there may be selective pressure that favors survivors with higher tolerance for cold and appropriate refuge-seeking behaviors. Green iguanas (*Iguana iguana*) (Krysko et al., 2007), anole species (e.g., *Anolis sagrei, A. equestris*) (Dalrymple, 1980; Kolbe et al., 2014), chameleon species (e.g., *Furcifer oustaleti*, *Chamaeleo calyptratus*), and Nile monitor lizards (*Varanus niloticus*) are other examples of problematic invasive non-native tropical reptiles that are cold sensitive and would have the potential to spread northward in warmer winters.

Beyond south Florida, extreme cold temperatures also play an important role in controlling the northern distribution of tropical reptile species in tropical-temperate transition zones along the Gulf of Mexico (i.e., Texas, Louisiana, Mississippi, and Alabama) and Atlantic (i.e., Georgia) coasts as well in arid and semi-arid parts of North America. For example, the freeze-sensitive brown anole is currently also established in Texas, Louisiana, Georgia, and California (EDDMapS, 2020). Green iguanas have expanded from their northern range limit in Mexico into south Texas (Hibbitts & Hibbitts, 2015). In the Sonoran Desert, the neotropical vine snake (*Oxybelis aeneus*) is just one example of a terrestrial native reptile whose distribution and habitat use are governed by winter temperatures (Van Devender et al., 1994).

Dispersal constraints may affect the potential for some reptile species to expand northward in response to climate change. However, due to the potential for long-distance dispersal via the pet trade, there are many reptile species in Florida that have the potential to rapidly move northward and westward in response to warming winters.

**Amphibians**

As ectotherms, amphibians have evolved physiological and behavioral traits to allow them to survive in a range of environmental temperatures, but most are disabled or even killed by extreme cold (Lillywhite et al., 2017). Many amphibians have evolved traits to survive freezing through avoidance or freeze tolerance (Costanzo & Lee, 2013). Some species of amphibians (e.g., wood frogs, *Lithobates sylvaticus*) are so well adapted to cold temperatures that they can survive ice formation in tissues through physiological mechanisms (Costanzo et al., 2013). However, most amphibian species are much less immune to the detrimental effects of extreme cold temperatures. Tropical amphibians tend to have higher thermal minima than amphibian species from temperate climates (Brattstrom, 1968), and their only mechanism for survival in cold temperatures is to find a suitable refuge where they can endure until warmer temperatures return.

There are several North American examples of amphibians that appear to have recent northward range expansions in the tropical-temperate transition zone. In the southeastern United States, there are no native amphibian species that are exclusively found in subtropical regions (Means & Simberloff, 1987). However, there are several recently introduced invasive amphibian species that originated in tropical regions (Krysko et al., 2011; Meshaka, 2011). For example, greenhouse frogs (*Eleutherodactylus planirostris*) have spread widely throughout Florida and continue to expand their range (Meshaka et al., 2004), and Rio Grande chirping frogs (*Eleutherodactylus cystignathoides*) have been expanding in Texas and Louisiana (Battaglia et al., 2015). Cuban treefrogs (*Osteopilus septentrionalis*) have spread from the site of their original introduction in the Florida Keys in the early 1930s (Barbour, 1931) to occur in most of peninsular Florida (Dodd, 2013) and are increasingly being found in nearby states (Glorioso et al., 2018) (Figure S5). There is a growing body of evidence that Cuban treefrogs can tolerate freezes through the use of microclimatic retreats (i.e., warmer shelter sites that provide protection from surrounding freezing temperatures) and are therefore likely to continue to expand their range (Meshaka, 1996; Haggerty & Crisman, 2015). Another species that has periodically established in Florida and may be less freeze tolerant than Cuban treefrogs is the Puerto Rican common coquí (*Eleutherodactylus coqui*). Coquís have been documented in southern Florida from at least two separate establishment periods (Austin & Schwartz, 1975; Loftus & Hernon, 1984), but there is plausible evidence that this species was extirpated during a hard freeze in 1977 (Wilson & Porras, 1983). In response to winter warming, more of Florida is expected to become suitable for coquí, and this species would have a better chance of establishing there (Rödder, 2009).

Dispersal constraints may affect the ability of some amphibian species to move northward in response to warming winters. However, there are species (e.g., Cuban treefrogs and coquís) that have successfully established populations in new areas following long distance transport (>100 km) with ornamental tropical plants. Some species have inadvertently traveled on horticultural shipments from Florida and other more tropical locations to nurseries across the United States (Kraus & Campbell, 2002; Glorioso et al., 2018; Morningstar & Daniel, 2020), which is a long-distance dispersal pathway that may enable more rapid poleward range expansion of some amphibian species in response to climate change.

The consequences of tropicalization of the amphibian fauna will vary depending on the species involved. Cuban treefrogs are a major threat to native amphibian communities because they can reduce the abundance (Rice et al., 2011) and occupancy (Waddle et al., 2010) of native treefrogs through competition and predation of native species (Meshaka, 2001). Cuban treefrogs produce a toxic secretion that can be noxious to humans and may be harmful to frog predators such as snakes (Goetz et al., 2018). In addition, Cuban treefrogs may have societal and economic impacts by causing power outages and other damages while seeking refuge from cold temperatures in buildings (Johnson et al., 2010). In Hawaii, invasive non-native coquí have become highly disruptive by altering the soundscape with loud choruses of vocalizations that have had an economic impact (i.e., decreasing property values and purchase interest) (Kraus & Campbell, 2002; Rödder, 2009), and their potential ecological impacts include altering the native fauna and disrupting nutrient cycles in ways that could benefit invasive plant species (Beard & Pitt, 2005; Sin et al., 2008). The ecological impacts of other species like greenhouse and Rio Grande chirping frogs are currently unknown. Few studies have been conducted on these small, ground-dwelling amphibian species.

**Manatee**

The West Indian manatee (*Trichechus manatus*) is a large marine mammal that lives at the freshwater-marine interface in low-energy estuaries, tidal creeks, and shallow near-shore waters where it feeds on seagrass and submerged aquatic vegetation. The Florida subspecies (*T. m. latirostris*) on the Atlantic and Gulf of Mexico coasts of the southeastern United States defines the northern limits for the species (Lefebvre et al., 2001). All extant manatee species and dugongs (order Sirenia) are tropical in distribution and share similar thermal tolerance to cold water temperature with physiological and behavioral traits adapted to the tropics (McNab, 2002; Marsh et al., 2012). As obligate herbivores, their low-energy diet leads to low metabolic rates and low capacity for thermogenesis compared to other marine mammals of similar size (Irvine, 1983; O'shea, 1988). Additionally, high thermal conductance results in rapid heat transfer to and from the environment. Homeostasis of core body temperature for manatees can be controlled to some extent by a countercurrent exchange circulatory system that conserves heat under cold stress or radiates heat under heat stress (Rommel & Caplan, 2003).

For the Florida manatee at the tropical-temperate transition in North America, mortality events from winter freezes and continental cold fronts, which can extend down the entire Florida peninsula, have been a focus of research and a concern for federal and state managers charged with recovery of the protected West Indian manatee. Manatees are subject to acute hypothermia or chronic cold stress pathologies that can be debilitating or lethal (Bossart et al., 2003). When winter water temperatures begin to drop and stay below approximately 20°C, manatees have to move to warmer water (Hartman, 1979; Irvine, 1983). In summer, individuals are dispersed to a wide range of forage habitats, but in fall and winter, they converge and form large aggregations in areas near warm-water refugia (Deutsch et al., 2003). Three types of warm water refugia have been important to manatees (Laist et al., 2013): (1) natural artesian springs where warm geothermal groundwaters come to the surface (Hartman, 1979); (2) artificial warm-water discharges produced by thermoelectric power plants and other industrial plants (Moore, 1956) (Figure S6); and (3) canals, boat basins, and drainage ditches with limited mixing of surface and bottom waters, which can maintain water temperatures passively by thermal inertia (Stith et al., 2011; Stith et al., 2012).

Cold winters have been less frequent in recent years and changes in manatee movements and habitat use are emerging. In cold winters, densities are higher at traditional warm water refugia where scientists monitor populations with photo-identification and aerial surveys. However, in warm winters, fewer numbers use these refugia because they can find adequate thermal protection at sites closer to prime foraging areas. However, expansion of the winter range beyond Florida due to winter warming may not be realized in the foreseeable future, given the magnitude of temperature changes required to reduce the need for refugia. The quality and quantity of natural artesian spring refugia are not found outside Florida, and current spring flows may be reduced with rising sea levels and increasing groundwater withdrawals. Furthermore, warm water discharges from power plants may decrease in the future as companies switch to more efficient technology and add alternative energy sources. Water conservation is an additional factor driving electricity production away from once-through cooling systems (Dieter et al., 2018). Nonetheless, further tropicalization of the climate would reduce morbidity and mortality probabilities from chronic cold stress within the current winter range.

Summer manatee sightings are increasing in numbers beyond historical range limits along the northern Gulf of Mexico (Fertl et al., 2005) and Atlantic coasts (Rathbun et al., 1982; Cummings et al., 2014). However, the role of climate change has been understudied. Manatee population growth due to recovery efforts may play a larger role in these increases. Recently, the U.S. Fish and Wildlife Service reclassified the species from endangered to threatened status (U.S. Fish and Wildlife Service, 2017). Regardless, the large number of manatee individuals and now annual sightings of known individuals in Alabama and on coastlines in Georgia, South Carolina, and North Carolina suggest that these are expansions of the summer foraging range. Due to exploratory migrations, manatee sightings are also increasing farther north, including in the Chesapeake Bay (Reid, 1995), Delaware River, and Cape Cod.

**Insects**

Across the globe, winter temperature regimes greatly influence the poleward distribution and abundance of insects (Sinclair et al., 2003; Robinet & Roques, 2010; Williams et al., 2015) and winter warming can facilitate poleward range shifts (Crozier, 2003; Battisti et al., 2005; Fält-Nardmann et al., 2018). Cold tolerance limits frequently parallel minimum temperature isotherms, supporting a widespread role for winter low temperatures in setting range limits (Sinclair et al., 2012; Lynch et al., 2014; Overgaard et al., 2014). Tropical insect species are typically sensitive to cold temperature extremes, which can lead to mortality at any life stage (Lee Jr., 2010). Tropical species are less cold tolerant than their higher latitude relatives, which constrains the poleward distributions of tropical insect species (Overgaard et al., 2011; Halbritter et al., 2018). Thus, warming winter temperatures associated with climate change are expected to allow some tropical species to move northward into ecosystems that have typically supported temperate insect-dominated communities. Because insects often have disproportionate effects on the structure and functioning of ecosystems (Ayres & Lombardero, 2000; Weed et al., 2013), and many also are vectors of vertebrate (including human) pathogens, the ecological and societal consequences of tropical insect range expansions are large.

From a human health perspective, the northward movement of tropical mosquitoes in response to warming winters has the potential to facilitate the spread of harmful viruses (Morin et al., 2013; Monaghan et al., 2016). The tropical “southern house mosquito,” *Culex quinquefasciatus*, a vector of many human and wild/domestic animal pathogens (e.g., St. Louis encephalitis, West Nile virus, Western equine encephalitis, possibly Zika virus) occurs from California to Florida but has been gradually expanding northward (Turell et al., 2001; Hahn et al., 2015). Although this mosquito species will occasionally feed on human blood, it prefers to feed on animals (Pruszynski et al., 2020). Over the past several decades the two most important mosquito vectors for dengue, Zika, chikungunya, and yellow fever—*Aedes aegypti* and *Aedes albopictus*—have been expanding their ranges worldwide (Gubler, 2002; Lambrechts et al., 2010; Roth et al., 2014). In fact, the domesticated subspecies *Aedes aegypti aegypti* has become the most cosmopolitan species among all insect vectors (Christophers, 1960; Powell & Tabachnick, 2013; Powell et al., 2018) (Figure S7). Both *A. aegypti* and *A. albopictus* are tropical/subtropical species that are range-limited by winter low temperatures (Joy et al., 2012; Johnson et al., 2017). Both are established in Florida and Texas where local transmission of dengue, chikungunya, and Zika occurs (Johnson et al., 2017). Cold winter temperatures kill *A. aegypti* eggs, which currently constrains the species to southern tropical regions of the United States (e.g., southern Florida and Texas) (Monaghan et al., 2016). However, the eggs of *A. albopictus* are able to diapause, increasing its ability to survive in colder winters and spread farther north than *A. aegypti* (Hawley et al., 1989; Brady et al., 2014; Johnson et al., 2017), and this species has now been reported from over 30 U.S. states. In response to warming winters, *A. aegypti* is now found in southern California and Arizona (Joy et al., 2012), and it is expected to increasingly move northward and become established in other parts of the southern United States (e.g., north Florida, Georgia, Alabama, Mississippi, Louisiana, north Texas, New Mexico), which means that Zika (Monaghan et al., 2016), dengue (Morin et al., 2013; Ebi & Nealon, 2016), and other mosquito-borne viruses (Iwamura et al., 2020) could potentially also expand northward and become locally transmitted in new areas. Numerous studies have produced global predictions of climatic suitability for *A. aegypti* and *A. albopictus* and all have concluded that warming winters are a primary factor driving mosquito range expansion poleward (Khormi & Kumar, 2014; Campbell et al., 2015; Monaghan et al., 2018). Rochlin et al. (2013) estimated that regions with suitable environmental conditions for the development of *A. albopictus* will increase by 50% by the end of the century, placing an additional 30 million people at risk.

For natural and managed forests, the northward movement of forest insect herbivores could have large ecological and economic implications (Ayres & Lombardero, 2000; Dale et al., 2001; Duehl et al., 2011). For example, the southern pine beetle (*Dendroctonus frontalis*) is the most destructive insect to pine forests in the southeastern United States. The southern pine beetle is sensitive to extreme cold winter temperatures, which control its northern range limit; thus, warming winters are expected to allow *D. frontalis* to expand into new areas and affect pine forests north of its current distribution (Ungerer et al., 1999; Trân et al., 2007; Lesk et al., 2017). Range limitation by winter cold is common in forest pest insects, so winter warming is leading to large-scale poleward range shifts in many insect species (Paradis et al., 2008; Sambaraju et al., 2019).

Many non-indigenous insect species that were initially constrained by cold temperatures are expanding their ranges poleward. The introduced sub-Saharan, arid-land African dung beetle (*Digitonthophagus gazelle*) has already expanded from the Gulf of Mexico coast to the southwestern United States, and it is expected to expand its range farther northward as winter climate warms (Harpootlian et al., 2001; Noriega et al., 2010). Similarly, the fire ant (*Solenopsis invicta*), which is native to the tropical regions of Argentina but has been introduced in the southern United States, Australia, and New Zealand, is moving poleward with climate warming. This species was estimated to be responsible for approximately US $600 million annually in damage to livestock, public health, and wildlife in the United States alone (Pimentel et al., 2000). While initially thought to have low capacity for cold tolerance, recent work has demonstrated local adaptation in cold in this species, suggesting that further poleward expansion is possible (Lytle et al., 2020).

Continental-scale insect migrations can also be affected by changing winter temperature regimes. For example, most monarch butterflies (*Danaus plexippus*) in eastern North America migrate from temperate regions (Canada and the northern United States) to the tropics (Mexico) to avoid winter-time temperature extremes (Brower & Malcolm, 1991). However, warming winters have the potential to allow migratory monarchs to overwinter in areas north of their historical overwintering grounds (e.g., along the northern Gulf of Mexico coast, instead of Mexico) (Howard et al., 2010).

**Pathways to range expansion**

Since the last glacial maximum about 19,000 years ago, warming temperatures have allowed cold-sensitive tropical organisms to move northward to their current northern range limits in North America (Pielou, 1991; Woodroffe & Grindrod, 1991; Zink & Gardner, 2017). Many of these species have traveled large distances (~2000–3000 km) as they adjusted to a warming world. Despite this tremendous capacity for adaptive movement, the unprecedented pace of climate change in the coming centuries is expected to be much faster than the past, which will require even faster rates of range expansion. To complicate matters, there are many other aspects of global change that will interact with warming winters to affect species’ northward movement. Thus, there is need to more closely examine potential barriers to and opportunities for range expansion.

In the taxa-specific sections, we provide some examples of different range expansion pathways. For example, long-distance range extension is most likely for species that can fly or swim (e.g., birds, fish, manatee, sea turtles, insects), or those easily transported by humans. Some plants are capable of long-distance dispersal by water transport. For example, mangroves, whose water-dispersed propagules can be moved by tropical cyclones and oceanic currents (Van der Stocken et al., 2019b). Other plants can travel large distances when their seeds are carried by winds or animals. For example, the non-native Brazilian pepper tree can be widely dispersed by birds that have consumed its fruit (Ewel et al., 1982). Plants can also be dispersed by humans; for example, ornamental tropical plants sold at nurseries have often traveled long distances via truck transport on highways. In general, amphibian and reptile movements are slower than for birds or fish. However, there are many reptile and amphibian species that have moved large distances due to human transport; for example, Cuban treefrogs have crossed states while concealed in ornamental plant shipments (Glorioso et al., 2018), and Burmese pythons and other reptiles have traveled across oceans via the pet trade (Krysko et al., 2016). The massive introduction of invasive non-native tropical reptile (Engeman et al., 2011) and plant species (Simberloff et al., 1997) in Florida provides a cautionary example of the potential for transformative ecological change due to human-assisted range extensions. Beyond just dispersal and movement capacity, habitat constraints can limit range expansion for some species (Zhu et al., 2012; Corlett & Westcott, 2013). For example, urban development or geological barriers northward of current range limits can reduce the habitat available for organisms to expand into. Collectively, these examples show that the pathways for range expansion are complex and warrant close consideration.

**Microclimates and thermal refugia**

Thermal refugia play a critical role in the tropical-temperate transition zone by providing warmer microclimates, which enable tropical cold-sensitive species to survive extreme cold events. For example, during extreme cold events: manatees congregate near warmer waters (Laist et al., 2013); coastal fish (Stevens et al., 2018) and sea turtles (Lamont et al., 2018) move to warmer offshore, deeper, or spring-fed waters; reptiles (Mazzotti et al., 2016) and amphibians (Meshaka, 1996) seek shelter closer to warmer soil or water; and cold-sensitive plants are typically located in landscape positions that provide warmer microclimates (D'Odorico et al., 2013). Northern mangrove range limits along the northern Gulf of Mexico, for instance, are often located on barrier islands, which are far from the colder mainland and surrounded by comparatively warmer oceanic waters (Osland et al., 2019b). Human-built infrastructure can also provide thermal refuge to cold-sensitive species. For example, the Cuban treefrog seeks shelter in buildings (Johnson et al., 2010), manatees take advantage of the warmer water provided by thermoelectric plants (Moore, 1956), and the mangrove tree crab (*Aratus pisonii*) seeks warmer conditions provided by docks (Cannizzo et al., 2020). As tropical cold-sensitive species move beyond their current range limits, the warmer microclimates provided by thermal refugia will continue to shape range expansion dynamics.

**Knowledge gaps to better understand climate change impacts**

In each of the taxa-specific sections, we provide examples of the critical influence of winter temperature extremes on tropical species’ range limits. Most cold tolerance research has been focused on strongly cold-tolerant temperate species, leaving a large gap in our knowledge of the cold tolerance of tropical species. Many of the species that we highlight have been understudied, and to better assess the potential tropicalization of temperate ecosystems in response to climate change, there are many critical knowledge gaps to be filled. In this section, we identify a subset of these critical research areas within the context of the three key components of climate change vulnerability assessments: sensitivity, exposure, and adaptive capacity (Glick et al., 2011; Stein et al., 2014).

Knowledge gap #1: How sensitive are species, ecosystems, and biomes within tropical-temperate transition zones to extreme cold temperatures? Much of our knowledge on this topic has developed from sporadic and opportunistic observations that quantify the damage or mortality caused by a specific extreme event to a species, ecosystem, or biome. These ad hoc observations are extremely valuable, but there is also a need for laboratory and field-based manipulative experiments that elucidate physiological effects and better quantify temperature thresholds that lead to physiological damage or mortality. Moreover, opportunistic post-freeze observations from single locations could be made more strategic, organized, and valuable via their integration into regional coordinated networks of scientists that are collectively prepared to rapidly respond to and quantify the effects of extreme events along ecologically relevant abiotic gradients (e.g., Osland et al., 2020a). For many species, historical patterns of range expansion and contraction have been understudied, and historical temperature data, in combination with temporal ecological data and knowledge of critical physiological temperature thresholds, can be used to better understand historical changes in ecological distribution, structure, and function due to cold sensitivity.

Knowledge gap #2: What is the expected exposure of species, ecosystems, and biomes to changes in the frequency and intensity of extreme cold temperatures? In prior decades, climate change vulnerability assessments focused primarily on changing means rather than changing extremes. However, given the demonstrated ecological importance of winter temperature extremes, future climate projections that better characterize changes in the frequency and intensity of winter temperature extremes could improve estimates of the biological effects of changing winters (Dillon et al., 2016). In addition to long-term projections (e.g., end of 21st century), vulnerability assessments could be improved by better understanding of the shorter-term potential for extreme winter events in the coming decades due to Arctic warming and jet stream dynamics that affect midlatitude climates (Cohen et al., 2019; Blackport & Screen, 2020). To gauge exposure, future air temperature projections can be used for many cold-sensitive terrestrial organisms (e.g., plants); however, for marine and aquatic organisms (e.g., sea turtles, fishes, manatee), there is a need for customized models that quantify linkages between changing air temperatures and aquatic and coastal water temperatures. For coastal species (e.g., mangroves), models that incorporate ocean-driven microclimates (i.e., temperature variation across land-ocean gradients) are critical for assessing potential changes in species distribution and abundance (Osland et al., 2019b; Osland et al., 2020b). Conversely, for montane species, models that incorporate mountain-driven microclimates (i.e., temperature variation due to landscape aspect, topographical variation, and elevation gradients) can improve estimates of upslope migration. Likewise, during drought years in arid and semi-arid regions, cloudless nights present greater risk of freezing exposure and have been understudied. For species that use refugia (e.g., frogs, pythons, fish, manatee), models that quantify the microclimatic protection offered by those refugia are critical.

Knowledge gap #3: What is the adaptive capacity of species, ecosystems, and biomes to changes in the frequency and intensity of extreme cold temperatures? Adaptive capacity via range expansion depends on many factors including a species’ potential for long-distance dispersal, growth, reproduction, abiotic stress tolerance, and biotic competition for resources in novel environments (e.g., Riley & Griffen, 2017). Figure 5 provides a generalized illustration of four alternative range expansion pathways for different species. Some of the species covered in this review (e.g., mangroves, Burmese pythons, Brazilian pepper, Cuban treefrogs, snook) have a tremendous capacity to adapt to climate change and clearly have the potential to expand their ranges northward in response to warming winters (Figure 5, lines A-C). In contrast, there are other species that lack the capacity to rapidly adapt or move from their current locations, often due to dispersal constraints or other biotic limits (Figure 5, line D). For foundation plant species, there is some indication that ecological regime shifts and northward range expansion could be accelerated by vegetation-microclimate positive feedbacks (D'Odorico et al., 2013) or from metabolic interactions with elevated CO2 (Loik et al., 2000; Dole et al., 2003); however, these feedback mechanisms are understudied. Plastic responses to thermal extremes (including beneficial acclimatization or cumulative damage) will strongly influence adaptive capacity and are key priorities for future study (Williams et al., 2016). The effects of changing return intervals on range expansion rates are also poorly understood. As species move northward, how will changing return intervals—specifically, larger time periods between extreme cold events (i.e., longer return intervals)—affect physiological processes, population growth, and ecosystem development near moving range limits? For example, in Figure 5, line C depicts an expansion pathway that is initially dampened by recurring extreme cold events near northward moving range limits (see expansion and contraction cycles) but then unaffected by cold events as extreme cold events disappear (see line smoothening and the absence of contractions).

**Conclusions**

In recent decades, ecologists have increasingly shown that ecological transformations due to climate change are often driven by climate extremes and extreme climatic events, rather than just changes in climatic means (Jentsch et al., 2007; Smith, 2011; Pecl et al., 2017). In this review, we have used a diverse assemblage of examples to highlight the critical role that extreme cold temperatures play in controlling the distribution, abundance, and structure of species, ecosystems, and biomes near tropical-temperature transition zones in North America. We have also examined the potential for tropicalization as tropical species move poleward in response to warming winters. Climate change-induced decreases in the frequency and intensity of extreme cold events are expected to facilitate the poleward range expansion of many tropical species. Assessing the ecological implications of the tropicalization of temperate ecosystems in North America would provide critical knowledge to inform natural resource managers.

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Table 1. Tropical cold-sensitive organisms whose northern range limits are governed by winter cold temperature extremes. These examples are a small subset of the organisms discussed in the taxa-specific sections. The numbers correspond to the photos in Figure 1.

|  |  |  |
| --- | --- | --- |
| **Number** | **Common name** | **Scientific name (s)** |
| 1 | Burmese python | *Python molurus bivittatus* |
| 2 | Joshua tree | *Yucca brevifolia* |
| 3 | Saguaro | *Carnegiea giganteus* |
| 4 | Red mangrove | *Rhizophora mangle* |
| 5 | Melaleuca | *Melaleuca quinquenervia* |
| 6 | Brazilian pepper | *Schinus terebinthifolius* |
| 7 | Cuban treefrog | *Osteopilus septentrionalis* |
| 8 | American crocodile | *Crocodylus acutus* |
| 9 | Buffelgrass | *Pennisetum ciliare* |
| 10 | Goliath grouper | *Epinephelus itajara* |
| 11 | Sawfish | *Pristis pectinata* |
| 12 | Cobia | *Rachycentron canadum* |
| 13 | Bull shark | *Carcharhinus leucas* |
| 14 | Manatee | *Trichechus manatus* |
| 15 | Loggerhead sea turtle | *Caretta caretta* |
| 16 | Kemp's ridley sea turtle | *Lepidochelys kempii* |
| 17 | Greenhouse frog | *Eleutherodactylus planirostris* |
| 18 | Coqui frog | *Eleutherodactylus coqui* |
| 19 | *Aedes aegypti* | *Aedes aegypti* |
| 20 | *Culex quinquefasciatus* | *Culex quinquefasciatus* |
| 21 | Monarch butterfly | *Danaus plexippus* |
| 22 | Organ pipe cactus | *Stenocereus thurberi* |
| 23 | Chapparal plants | *Ceanothus megacarpus*, *Malosma laurina* |

Figure 1. A map illustrating tropical-temperate climate and ecological transition zones in North America. While the more temperate zones are depicted with blues, the more tropical zones are depicted with warm colors (i.e., red, orange, and yellow). In response to warming winters, cold-sensitive tropical organisms are expected to move northward, sometimes at the expense of cold-tolerant temperate organisms. The photos provide examples of tropical cold-sensitive organisms whose northern range limits are governed by winter cold temperature extremes (photo numbers correspond to the species in Table 1; see Table S1 for sources). Temperatures represent the absolute coldest temperature recorded for the period extending from 1980 to 2009, obtained from Daymet data (https://daymet.ornl.gov).



Figure 2. Near tropical-temperate transition zones in North America, winter cold temperature extremes play a critical ecological role. Data from 1900–1999 were used to create this generalized depiction of the return time and frequency of ecologically relevant cold events for a location in central Florida. The vertical line represents a generalized threshold for cold damage or mortality, and the grey box represents the zone where cold damage or mortality would occur in this scenario. The return time for ecologically relevant cold events in this depiction is ~20 years (see red line in grey box). This generalized scenario was developed using knowledge of cold temperature thresholds for two plant species in this region (Osland et al., 2020a; Osland & Feher, 2020).



Figure 3. Maps comparing the northward position of North American climatic zones supporting tropical cold-sensitive species (i.e., red, orange, and yellow areas) under (a) recent climatic conditions (1980–2009) and three alternative future climate scenarios—a 2°C, 4°C, and 6°C increase in winter temperature extremes (b-d, respectively). For (a), temperatures represent the absolute coldest temperature recorded during the 30-year period, obtained from Daymet data (https://daymet.ornl.gov).



Figure 4. Recent decades have had warmer winters with fewer extreme cold events. Winter temperature metrics from four representative locations in North America’s tropical-temperate transition zones: (a) San Francisco, California; (b) Tucson, Arizona; (c) New Orleans, Louisiana; and (d) Tampa, Florida. The upper green lines show mean winter temperatures [Mean Avg Temp (DJF)]. The middle blue lines show the absolute coldest annual winter minimum temperature [Lowest Min Temp]. The lower grey bars show the number of subzero days each winter [# of days <= 0°C]. See Table S2 for trend test results. Temperature data were obtained from the Applied Climate Information System ([http://scacis.rcc-acis.org](https://gcc02.safelinks.protection.outlook.com/?url=http%3A%2F%2Fscacis.rcc-acis.org%2F&data=04%7C01%7Cmosland%40usgs.gov%7C171c1803aa84418f57d908d8cdd6ae73%7C0693b5ba4b184d7b9341f32f400a5494%7C0%7C0%7C637485669229716542%7CUnknown%7CTWFpbGZsb3d8eyJWIjoiMC4wLjAwMDAiLCJQIjoiV2luMzIiLCJBTiI6Ik1haWwiLCJXVCI6Mn0%3D%7C3000&sdata=GC1u9xXX0DiivfQ5IAG8VzsRNpSTV4LbYi0WOlyJWyM%3D&reserved=0)).

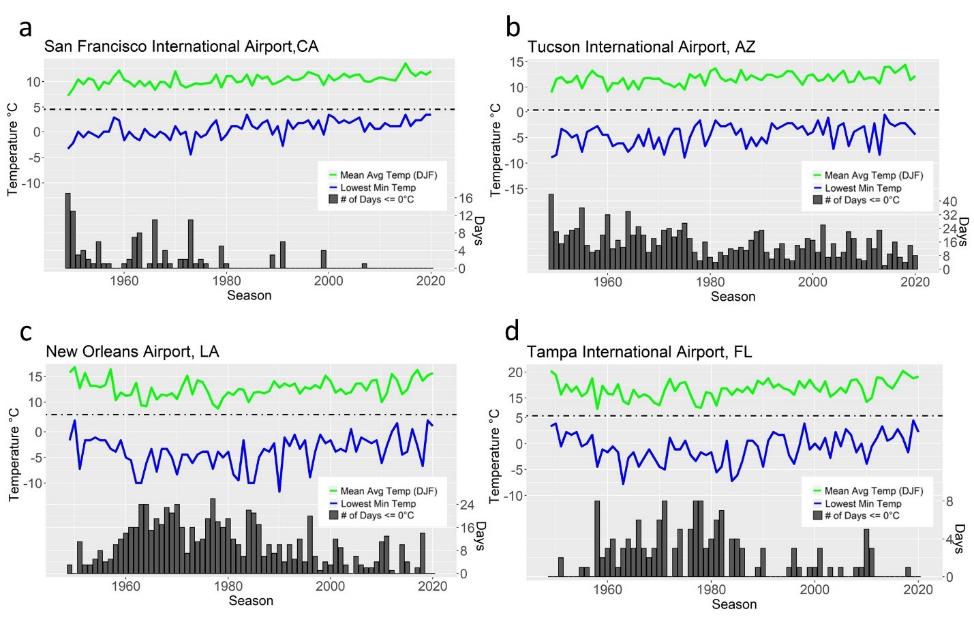


Figure 5. Four generalized depictions of alternative poleward range expansion pathways for tropical organisms in response to warming winters. A depicts a species that moves north in a step-like fashion by long-distance transport in horticultural or pet trade shipments. B depicts a species with continuous, exponential poleward range expansion, while C depicts a species whose poleward range expansion is discontinuous and punctuated by extreme cold events that result in short-term range contractions, but these extreme cold events eventually dissipate, producing a smoother, continuous range expansion. In contrast, D depicts a species that is not able to expand poleward, perhaps due to dispersal constraints or other biotic limits. The y axis represents the latitude of the species’ poleward range limit, where High represents a higher latitude north of the species' current range limit and Low represent a lower latitude near the species' current range limit.



Table S1. Sources for the photos in Figure 1. The numbers correspond to the photo numbers in Figure 1. All but one of the photos are from U.S. Government websites.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Number** | **Common name** | **Scientific name(s)** | **Source** | **Credit** | **Website** |
| 1 | Burmese python | *Python molurus bivittatus* | USGS | Bryan Falk | <https://www.usgs.gov/media/images/a-burmese-python-coiled-grass-everglades> |
| 2 | Joshua tree | *Yucca brevifolia* | NPS | Brad Sutton | <https://www.nps.gov/jotr/learn/news/newspaper.htm> |
| 3 | Saguaro | *Carnegiea giganteus* | USFS | Charlie McDonald | <https://www.fs.fed.us/wildflowers/plant-of-the-week/carnegiea_gigantea.shtml> |
| 4 | Red mangrove | *Rhizophora mangle* | USGS | Michael Osland | Not on a website |
| 5 | Melaleuca | *Melaleuca quinquenervia* | USDA | NA | <https://www.ars.usda.gov/southeast-area/fort-lauderdale-fl/iprl/docs/melaleuca/> |
| 6 | Brazilian pepper | *Schinus terebinthifolius* | USDA | NA | <https://www.ars.usda.gov/southeast-area/fort-lauderdale-fl/iprl/docs/schinus-terebinthifolius-brazilian-pepper-tree/> |
| 7 | Cuban treefrog | *Osteopilus septentrionalis* | USGS | Brad M. Glorioso | <https://www.usgs.gov/media/images/invasive-cuban-treefrog-new-orleans-la> |
| 8 | American crocodile | *Crocodylus acutus* | NPS | NA | <https://www.nps.gov/ever/learn/news/joe-bay-is-open-to-non-motorized-boating.htm> |
| 9 | Buffelgrass | *Pennisetum ciliare* | NPS | NA | <https://www.nps.gov/orpi/learn/nature/invasive-plant-species.htm> |
| 10 | Goliath grouper | *Epinephelus itajara* | NOAA | NA | <https://www.fisheries.noaa.gov/southeast/endangered-species-conservation/goliath-grouper> |
| 11 | Sawfish | *Pristis pectinata* | NOAA | NA | <https://www.fisheries.noaa.gov/species/smalltooth-sawfish> |
| 12 | Cobia | *Rachycentron canadum* | NOAA | NA | <https://www.fisheries.noaa.gov/species/cobia> |
| 13 | Bull shark | *Carcharhinus leucas* | NOAA | NA | <https://graysreef.noaa.gov/science/research/fish_tagging/visitors.html> |
| 14 | Manatee | *Trichechus manatus* | USGS | NA | <https://www.usgs.gov/centers/wetland-and-aquatic-research-center-warc/science/manatee-health-assessment-and-biomedical?qt-science_center_objects=0#qt-science_center_objects> |
| 15 | Loggerhead sea turtle | *Caretta caretta* | USGS | NA | <https://www.usgs.gov/news/after-hurricane-devastation-sea-turtle-scientists-rebound-help-rebuild> |
| 16 | Kemp's ridley sea turtle | *Lepidochelys kempii* | USGS | Margaret Lamont | <https://archive.usgs.gov/archive/sites/soundwaves.usgs.gov/2018/02/staff.html> |
| 17 | Greenhouse frog | *Eleutherodactylus planirostris* | USGS | Brad M. Glorioso | [https://armi.usgs.gov/gallery/species.php?itis=173568](https://armi.usgs.gov/gallery/result.php?search=Eleutherodactylus) |
| 18 | Coqui frog | *Eleutherodactylus coqui* | USGS | Chris Brown | [https://armi.usgs.gov/gallery/species.php?itis=173559](https://armi.usgs.gov/gallery/result.php?search=Eleutherodactylus) |
| 19 | *Aedes aegypti* | *Aedes aegypti* | CDC | NA | <https://www.cdc.gov/features/stopmosquitoes/index.html> |
| 20 | *Culex quinquefasciatus* | *Culex quinquefasciatus* | CDC | NA | <https://www.niaid.nih.gov/diseases-conditions/west-nile-virus> |
| 21 | Monarch butterfly | *Danaus plexippus* | USDA | Peggy Greb | <https://www.ars.usda.gov/oc/images/photos/oct19/d3980-1/> |
| 22 | Organ pipe cactus | *Stenocereus thurberi* | NPS | NA | <https://www.nps.gov/articles/nps-geodiversity-atlas-organ-pipe-cactus-national-monument-arizona.htm> |
| 23 | Chapparal plants | *Ceanothus megacarpus, Malosma laurina* | NPS | NA | <https://www.nps.gov/samo/learn/nature/chaparral.htm> |

Table S2. Test for trends in winter temperature (Temp) data from four representative locations in North America’s tropical-temperate transition zones, reporting Spearman’s correlation (rs) between year and: (1) mean winter temperature (Mean Avg Temp), (2) the absolute coldest annual winter minimum temperature (Lowest Min Temp), and (3) the number of subzero days each winter (# of days ≤ 0°C). See trend depictions in Figure 4.

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
|  | Mean Avg Temp | |  | Lowest Min Temp | |  | # of days ≤ 0°C | |
| rs | *p* |  | rs | *p* |  | rs | *p* |
| San Francisco (CA) | 0.55 | 0.000 |  | 0.62 | <0.001 |  | -0.66 | <0.001 |
| Tucson (AZ) | 0.39 | 0.000 |  | 0.41 | <0.001 |  | -0.46 | <0.001 |
| New Orleans (LA) | 0.19 | 0.104 |  | 0.23 | 0.047 |  | -0.39 | 0.001 |
| Tampa (FL) | 0.32 | 0.007 |  | 0.22 | 0.068 |  | -0.30 | 0.011 |

Figure S1. Within the tropical-temperate transition zone, extreme cold temperatures control the northern distribution of foundation plant species like the saguaro cactus (*Carnegiea gigantea*; left) and red mangrove (*Rhizophora mangle*; right). Photo credits: NPS (saguaro) and Michael Osland (mangrove).

Figure S2. Winter temperature extremes control the distributions of subtropical fishes and drive movements of coastal migrants. The photo on the left is of the common snook (*Centropomus undecimalis*), aggregating at a spring head in northern Florida during winter. The photo on the right shows a biologist tagging a coastal migrant, cobia (*Rachycentron canadum*), with an acoustic transmitter during its run along the Florida panhandle. Photo credits: Florida FWC [Phil Stevens (left) and Jessica Carroll (right)].



Figure S3. USGS scientists picking up cold-stunned sea turtles floating at the surface of St. Joseph Bay in northwestern Florida (USA) during an extreme cold event in 2018. Note the cold-stunned juvenile sea turtles in the boat. During cold stun events, mortality is often highest for juvenile, smaller sea turtles (Lamont et al. 2018). St. Joseph Bay is located in the northeastern Gulf of Mexico, approximately 50 km southeast of Panama City and 350 km northwest of Tampa Bay. Photo credit: USGS.

A ship in a body of water

Description automatically generated

Figure S4. Extreme cold events constrain the distribution of many invasive non-native tropical reptile species. Burmese pythons (*Python bivittatus*) are an especially harmful and well-known example of the negative ecological impacts of pet trade-driven tropical reptile introductions. This large constrictor snake is an opportunistic apex predator, whose expansion within and around Everglades National Park (southwestern Florida, USA) has decimated mammal, bird, and other prey populations. Burmese pythons are also sensitive to cold temperature extremes and expected to expand northward in response to warming winters. Photo credit: USGS.



Figure S5. Dispersal constraints may affect the ability of some species to expand their range northward in response to warming winters. However, there are species [e.g., the Cuban treefrog (*Osteopilus septentrionalis*), as shown in this photo] that have successfully established populations in new areas following long-distance transport (>100 km) in ornamental tropical plants. These species have inadvertently traveled on horticultural shipments from Florida and other more tropical locations, which is a long-distance dispersal pathway that may enable more rapid poleward range expansion of certain amphibian species in response to climate change. Photo credit: Brad M. Glorioso.



Figure S6. Near the northern limits of the distribution of the West Indian manatee (*Trichechus manatus*), individuals of the southeastern United States forage on seagrass and submerged aquatic vegetation. With advancing cold the population contracts toward Florida warm-water refugia where they often form large aggregations. This photo shows an aggregation at an artificial warm-water discharge produced by a thermoelectric power plant. Photo credit: Catherine Langtimm.



Figure S7. The northern range limits of most tropical insect species are governed by cold temperature extremes, which can lead to mortality of eggs, larvae, pupae, or adults.This photo is of *Aedes aegypti*, whichis a cold-sensitive mosquito species. This species is expected to increasingly move northward due to warming winters and become more established in other parts of the southern United States. Photo credit: CDC.

