Connections to the Deep: Deep Vertical Migrations, an Important Part of the Life Cycle of *Apherusa glacialis*, an Arctic Ice-Associated Amphipod

Magnus Drivdal*, Erin H. Kunisch, Bodil A. Bluhm, Rolf Gradinger, Stig Falk-Petersen and Jørgen Berge

1 Akvaplan-Niva, Fram Centre, Tromsø, Norway, 2 Department of Arctic and Marine Biology, Faculty of Biosciences, Fisheries and Economics, UIT The Arctic University of Norway, Tromsø, Norway, 3 Department of Arctic Biology, The University Centre in Svalbard (UNIS), Svalbard, Norway, 4 NTNU Centre for Autonomous Marine Operations and Systems, Marine Technology Centre, Trondheim, Norway

Arctic sea ice contains a substantial amount of living biota of which part is lost through melt and export out of the Arctic Ocean every year. It is unclear how populations can be maintained within the Arctic Ocean. A representative ice inhabitant, the amphipod *Apherusa glacialis* was previously assumed to spend its entire life in the sea ice habitat, hence being dependent on sea ice to complete its life cycle. However, several recent studies report pelagic occurrences and suggest that seasonal vertical migrations might be an adaptive life history trait enabling a viable population size in the Arctic Ocean.

In this study we use a particle-tracking model to investigate to what extent vertical migration might affect the species’ retention in the Arctic Ocean and the sea ice habitat. The modeled trajectories of *A. glacialis* were calculated based on ice drift and ocean currents from a coupled ocean – sea ice model covering the Arctic Ocean. We test two scenarios: (1) trajectories of *A. glacialis* that stay attached to the ice or follow the surface currents if they melt out of the ice and (2) trajectories of *A. glacialis* that undertake a seasonal vertical migration to drift with the currents at depth for parts of the year. In the multi-year model simulations it is assumed that after an initial period of 2 years *A. glacialis* that are located outside sea-ice cover each spring will perish while those located within the ice-covered region will reproduce. The model results show that a seasonal vertical migration both increases the total number of individuals and leads to a population distribution within the Arctic Ocean more in line with previous findings than the results from the non-migrating *A. glacialis*. Our results support the hypothesis that a seasonal migration may be an adaptive life history strategy in this species.

Keywords: amphipod, Arctic, sea ice, life cycle, particle tracking, vertical migration, *Apherusa glacialis*
INTRODUCTION

In general, the Arctic pelagic marine ecosystem contains very few endemic species, mostly due to the absence of barriers and isolation between the Arctic Ocean and its surrounding seas. Rather, strong biogeographic connections and interchange of water masses with the North Atlantic sub-Arctic seas, and to a lesser extent with the North Pacific are characteristic features. Yet, Arctic sea ice provides a unique habitat absent from sub-Arctic seas, with a high biodiversity including a number of endemic species [examples in Bluhm et al. (2011)]. More than 1,000 species, ranging from unicellular bacteria and algae up to larger crustaceans, mammals and sea birds are known to live on, in, or in close connection to the sea ice during at least parts of their life cycle (Poulin et al., 2011; Bluhm et al., 2017). The ice habitat also serves as platform for breeding seals (Laidre et al., 2008; Kovacs et al., 2011), as protective habitat from predators for young polar cod (Gradinger and Bluhm, 2004), as feeding habitat for metazoan meiofauna in the brine channel systems (Gradinger and Bluhm, 2020) and the underside as habitat for e.g., sea ice associated algae and amphipods and for pelagic flora and fauna such as various copepods, amphipods, comb jellies, etc. (Ehrlich et al., 2020). Questions remain on how the sea ice taxa, which mainly drift passively with ice and currents, can maintain their populations within the Arctic Ocean when sea ice is continuously exported from the Arctic.

The ice-cover and its drift are closely related to Arctic Ocean hydrography and circulation. The Arctic Ocean is a semi-enclosed ice-covered ocean, with extensive shelves and a deep basin with maximum depths of more than 4,000 m, as first described by Nansen (1902). The main large-scale wind-driven surface circulation and sea ice drift features of the Arctic Ocean are the Beaufort Gyre and the Transpolar Drift (Figure 1). The fundamental features and current systems of the Norwegian Sea including the Greenland-Iceland-Norwegian (GIN) Seas and the connection to the Arctic Ocean were later described by Helland-Hansen and Nansen mainly based on the data collected between 1900 and 1904 (Helland-Hansen and Nansen, 1909). A prominent feature of this circulation is the inflow of warm surface water from the North Atlantic that submerges under colder Arctic water when entering the Arctic, and continues to flow through the Arctic and along the pan-Arctic slope; modified Atlantic water in fact fills all deep Arctic basins (Rudels et al., 2000, 2013; Figure 1). Our knowledge of these interconnected systems has been updated and documented in several fundamental studies (Hansen and Østerhus, 2000; Rächold et al., 2004; Falk-Petersen et al., 2007; Rudels, 2015; Semper et al., 2020; Wassmann et al., 2020). The ocean current system (Figure 1) provides exchange of water, particulate matter, phytoplankton, zooplankton, ice biota and other matter including sediments from the Siberian shelf and rivers, between the North Atlantic and the Arctic (Timmermans and Marshall, 2020; Daase et al., 2021). Connectivity to the Pacific is limited to northward inflow into the Arctic through the shallow Bering Strait (Woodgate et al., 2005). Sea ice is a unique feature of the Arctic Ocean, consisting of drifting pack ice and land fixed fast ice. Approximately 50% of the ice cover has been seasonally melting in the past, leaving the central Arctic ice covered year-round by multi-year ice floes, yet with less and less multi-year ice remaining in recent decades. Both annual and perennial sea ice formed inside the Arctic are exported out following the surface currents and prevailing winds, with the major export happening through western Fram Strait. This transport accounts for an estimated average export of 14% of the sea ice volume every year, and has a strong seasonality, with most of the export happening in the winter months from November to April (Spreen et al., 2020).

Historically, organisms living in or on the underside of the sea ice habitat were classified into two groups: the allochthonous and autochthonous ice-associated species. The former group consists of species that spend parts of their life cycle in connection to the sea ice, the latter classified organisms that were assumed to be fully dependent on sea ice throughout their life cycle. In terms of sea ice fauna biomass, the latter group is dominated by crustacean amphipods (Lonne and Gulliksen, 1991a,b) originally with four species (Gammarus wilkitzkii, Apherusa glacialis, Onisimus nansen, and Onisimus glacialis), later expanded to also include Pleusyntes karstensi (Macnaughton et al., 2007). Since their association with sea ice has been considered obligatory, these organisms would lose both their sea ice habitat during ice melt and be advected out of the Arctic Ocean by the Transpolar Drift (Gulliksen and Lonne, 1991; Lonne and Gulliksen, 1991b; Arndt and Swadling, 2006). When the sea ice melts, these organisms are subjected to horizontal transport in the water column by ocean currents. Unless they can remain in an area where ice forms in the subsequent winter or they are transported back into the ice within their lifespan, these organisms are inevitably lost from the population. Several studies have investigated the biomass transport of ice biota into the Greenland and Barents Seas and estimated it to be on the order of 3.55 × 10^5 t wet weight (4.2 × 10^5 tC) year^{-1} and 1.5 × 10^5 t wet weight (1.8 × 10^4 tC) year^{-1}, respectively (Hop et al., 2006; Wassmann et al., 2006). These calculations and estimates, however, are based on the general assumption that the organisms in question are truly dependent on sea ice for their survival.

For A. glacialis (Figure 2), the most abundant of the five assumed obligate ice-associated amphipod species, the obligate sea ice association may be less strong than assumed. It had been originally considered a pelagic species (Barnard, 1959), but extensive and regular findings of the species living on the underside of the Arctic drift ice changed the general understanding of its ecology to that of an autochthonous ice associated herbivorous species (Scott et al., 1999; Beuchel and Lonne, 2002; Werner and Auel, 2005). Recently, however, Berge et al. (2012) and Kunisch et al. (2020) have questioned the obligate nature of the ice association for this species. Based on findings of A. glacialis in the deep layers of inflowing Atlantic water into the Arctic Ocean, Berge et al. (2012) suggested a conceptual model in which deep migration was an integrated part of A. glacialis’ 2-year life cycle (Poltermann et al., 2006; Beuchel and Lonne, 2002) in at least parts of the Arctic. It was suggested that A. glacialis grazes on ice algae during the productive spring, summer and autumn as the sea ice drifts south into the Greenland Sea, then migrates down to depths following the current system across the Norwegian Sea and re-enters the Arctic Ocean via the
north flowing Atlantic Current. Kunisch et al. (2020) conducted an extensive review of all reported pelagic findings of *A. glacialis* in the Arctic and concluded that it was regularly found in the water column throughout the year.

In this study we examine a hypothesis based on the findings by Berge et al. (2012) and Kunisch et al. (2020), namely that deep vertical migration after grazing on the ice-algal bloom would enhance the return and retention of *A. glacialis* in the Arctic Ocean. This can occur in summer and until the polar night. Such deep migration may enhance their fitness by both avoiding predation at a time when they carry young in a large brood pouch, and secondly by allowing them to relocate themselves either within or even back into the Arctic Ocean by advection as proposed by Berge et al. (2012). To test the advective component of the hypothesis, we use a particle trajectory model. We consider two parallel model scenarios where initially *A. glacialis* (as seeded particles) are located in identical areas covering large parts of the ice-covered Arctic Ocean. The amphipods in the two scenario populations follow the same strategy for the periods when sea ice-based food is available, staying attached to the ice (if ice is available) or close to the ocean surface. For the rest of the year the two populations behave differently, with amphipods in one migrating downwards to the core of the relatively warm Atlantic layer (here chosen at a depth of about 250 m; Lind et al., 2016; Bluhm et al., 2020), while in the other they remain attached to the ice (if available) or otherwise in the surface water layer.

MODEL SETUP AND METHODS

The coupled ocean-sea ice model used in this study is a 4 km pan-Arctic setup (Arctic-4) of the Regional Ocean Modeling System (ROMS, Shchepetkin and McWilliams, 2005) with a coupled sea-ice component (Budgell, 2005). For Arctic-4 the initial hydrodynamic fields and boundary conditions are provided by a monthly global ocean reanalysis (Storkey et al., 2010); surface
atmospheric forcing fields, such as near-surface wind, sea-level pressure and temperature are provided by a global atmospheric reanalysis (ERA-Interim: Dee et al., 2011). Tidal forcing is added at the open ocean boundaries from the tidal model TPXO (Egbert and Erofeeva, 2002). The Arctic-4 model has previously been run from 1993, with the 10 first years acting as a spin-up from ice-free conditions. Here the ocean current and ice drift from these simulations have been used as input for a separate particle tracking model. A more detailed description of the Arctic-4 model can be found in Hattermann et al. (2016). In the present study the particle (amphipod) trajectories were calculated from the Arctic-4 model fields from the years 2008–2017. The currents and ice drift from Arctic-4 respond to the continuously shifting atmospheric and boundary forcing conditions. Prevailing atmospheric conditions and current patterns may lead to significant year to year variations in ice cover and drift patterns, which again lead to annual variations in particle distribution and transport in and out of the Arctic Ocean.

Trajectories of the modeled A. glacialis were computed with TRACMASS (Döös et al., 2013, 2017) forced by the daily averaged model velocities from the Arctic-4 model. By using the daily averages, high frequency variability such as tides were filtered out. To simulate the trajectories of the particles in the ice, the model forcing was modified so that particles at the surface follow ice-velocities, which are largely forced by the wind (e.g., Spreen et al., 2011), if the ice cover in the region is above 10%. Otherwise, the particles follow the water masses transported by the ocean currents. This 10% threshold for ice drift to replace surface current drift is slightly lower than the 15% often used to describe the sea ice extent (e.g., Stroeve et al., 2012), but was chosen as we assumed A. glacialis to have a high preference to stay in the ice when food is available, even at low sea ice concentrations. To model vertical migration, the model was run according to a specific procedure, where particles were removed and subsequently reseeded at different depths. This simplified procedure assumes that the migration itself is a short process and the drift patterns during the vertical migration itself were not considered. The dominating ocean circulation patterns and associated sea ice drift are sketched in Figure 1. However, there are temporal fluctuations in the current and ice-drift due to changing weather patterns that can influence the particle transport in the model. For example, the Norwegian Atlantic Current can be ~20% stronger in its mean flow in winter (Andersson et al., 2011), and wind forcing accounts for a large fraction of the variation of summer-fall sea ice extent in the Arctic (Ogi et al., 2010). Yet in the present study our focus is on the broad-scale differences in amphipod distributions with and without vertical migrations, recognizing that absolute magnitudes will differ between years.

With the applied setup, the trajectories could be followed both as particles leave and re-entered the ice-covered regions. In the experiments, a large sector of the Arctic was seeded with approximately 350,000 particles (Figure 3) on 1 April 2008, and the individual trajectories were followed for several years (first for the period 2008–2013, and then in a separate run for 2008–2017). For modeling purposes, the yearly cycle of A. glacialis was split into two periods: the biologically active season (approximately when A. glacialis are assumed to be feeding on fresh algal material and algal detritus at/near the sea ice; Scott et al., 1999; Poltermann, 2001) 1 April – 31 July, and the biologically passive period 1 August – 31 March (little primary production (Leu et al., 2015), with little or no feeding). The trajectory model was first run for 2 years for two different scenarios: one with no vertical migration, and the other with seasonal migration as was proposed
by Berge et al. (2012). For the non-migrating scenario, we allowed *A. glacialis* to follow the ice drift if located in an area with ice cover greater than or equal to 10%. In contrast, *A. glacialis* was allowed to drift with the surface current when they reached areas with less ice cover (<10%). For the migrating scenario, *A. glacialis* had the same behavior in the active period (1 April – 31 July) but migrated down to 250 m (or the sea bottom if shallower) and followed the ocean currents passively at that depth for the period 1 August – 31 March before they migrated back up to the surface every 1 April. The water depth of 250 m was chosen because it is a reasonable approximation of the Atlantic layer core in the study area where temperatures are above 0°C (and even warmer in the Atlantic inflow area of the Fram Strait and Barents Sea) and salinity is at 34.8–35.0 (Loeng, 1991; Lind and Ingvaldsen, 2012; Bluhm et al., 2020) and current velocities northward can reach upward of 15 cm s\(^{-1}\) (Aksenov et al., 2011; Menze et al., 2019).

For simplicity, vertical migration in the model was assumed to occur at the same time every year for all individuals. The choice of timing of the arrival after migration up to the sea ice is based on the observation of the predominance of small size classes in the sea ice habitat in spring months (Melnikov, 1997; Kunisch et al., 2020). The timing of migration to depth is inferred from observations of mating during early winter, the finding of gravid females in deeper water layers, and the virtual absence of gravid females in surface layers from spring to fall (Melnikov, 1997; Poltermann et al., 2000; Berge et al., 2012; Kunisch et al., 2020).

The vertically migrating particles thus avoid following the ice drift during the winter months, when the major sea-ice transport through the Fram Strait occurs (Spreen et al., 2020). Although the sea ice transport may vary from year to year, this behavior was expected to lead to a significant reduction in the transport of particles from the Arctic to the GIN seas.

For practical reasons the representation of mortality and reproduction in the particle tracking model was simplified. From the start of the simulation, 1 April 2008 until 1 April 2010, the number of particles representing *A. glacialis* was kept constant (unless they exited the model domain). Subsequently, starting in 2010 and each 1 April from then on, all particles were counted and split into two groups: group one included those remaining or having re-entered the ice-covered region (defined as greater than or equal to 10% ice cover) and group two included those located in the open ocean. The model was then reseeded with 2 particles for each particle in group one (representing reproduction) and run forwards until next 1 April. The particles in group two were removed and accounted for mortality/population loss every year. This procedure was used to represent a simplified life cycle with mortality and reproduction, where an average of two offspring survived each year (for particles in the sea ice). With this simplistic 1:2 reproduction rate there was population growth for both scenarios (migrating and non-migrating) and the number of particles became impractical for the model to handle after 5 years (2013). In addition, we assumed that individuals may survive in the water column during the entire winter period but die if they are located in a totally ice-free area the following 1 April. Hence, the modeled *A. glacialis* may leave the sea-ice after 1 April and still be able to survive and reproduce later if they re-enter the sea-ice habitat before 1 April the following year. During such a pelagic stage, one possible return-route to the sea-ice habitat for the migrating particles could be to follow the Atlantic Current northwards, as proposed by Berge et al. (2012).

To analyze general longer-term distribution patterns and inter-annual variability, a separate run was performed for both scenarios until 2017, only accounting for mortality (using a 1:1 reproductive rate for individual particles in sea-ice). The reason for using a 1:1 reproductive rate in this model run is a practical one, namely that the model could not handle the population growth with the previously used 1:2 rate in the long term. These simulations diverge from the first ones from 2010. For each 1
April following this, particles that are located outside the sea ice are removed from the simulation. This leads to a year-to-year decline in particles for both migrating and non-migrating particles but at different rates.

RESULTS

Particle Distribution Before Migration
Since both the migrating and non-migrating model particles stayed at the ocean surface and followed the ice drift pattern during the first period (1 April 2008 – 31 July 2008), the trajectories of the two model scenarios were identical, and the end positions for the particles are shown in Figure 4. Clearly, the transport through the Fram Strait was significant, and particularly high concentrations of particles could be found in and south of the Fram Strait and in the marginal ice zone along the Barents and Kara Sea shelves. The ice drift in the Bering, Chukchi, and East Siberian Seas led to a northwards transport, resulting in large parts of the marginal ice zone in those shelf regions being devoid of A. glacialis during this period.

Particle Distribution After First Vertical Migration
From the 1 August 2008 and the onset of amphipod vertical migration in scenario 2, the particle densities of the two scenarios differed, (Figure 5 shows the concentrations 1 April 2009 – 1 April 2010). It is clear that after the first period with deep vertical migration (Figure 5A) a visibly larger amount of the non-migrating particles was transported out through the Fram Strait with the Transpolar Drift, exiting the Eurasian Basin north of Svalbard. For the migrating particles that avoided the upper ocean transport toward the Fram Strait for large parts of the year, densities north of Svalbard and the Arctic Basin remained high, especially along the shelf break to the Kara Sea and the Siberian shelf, as well as the St. Anna Trough (Figure 5, right hand side). In contrast, the number of non-migrating particles in these areas are greatly reduced by 1 April 2010 (Figure 5, left hand side). Recounting the number of particles after 2 years of simulations (Figure 5C) for both scenarios revealed that about 88% of the migrating particles ended up in ice-covered waters, whereas only 62% of the non-migrating particles remained under ice.

Particle Distribution After Reproductive and Mortality Events
From 1 April 2010 the scenarios were run with the 1:2 reproduction rate and the mortality assessment (particles in open water by 1 April) as previously described. This led to an increase in total population numbers in both cases, but the vertically migrating population grew faster since a higher share of these particles were located within the ice every 1 April. The particle densities for both cases for each 1 April in the years 2010 – 2013 (Figure 6) showed clear differences between these two cases. The migrating particles were more evenly spread within the Arctic Ocean, except for low concentrations over the Siberian shelf. The non-migrating particles on the other hand largely exited the Eurasian Basin and were mostly located in high concentrations within the Beaufort Gyre domain in the Amerasian Basin. Also, the total amount of particles differed between the two populations, with the migrating particle number increasing at a higher rate. By 1 April 2013 the migrating population was nearly twofold larger, consisted of more than 1.4 million particles whereas the non-migrating population had reached about 750,000 particles. For the non-migrating particles, the seasonal reproduction was not enough to compensate for the
rapid transport away from the Eurasian Basin, leading to this area being largely devoid of particles by 1 April 2013. Particles that performed a vertical migration avoided the rapid transport of ice with the Transpolar Drift toward the Fram Strait for the winter period and could recolonize new ice the next spring when migrating back toward the surface.
Mortality Effect
To get a longer-term perspective of mortality effects, both scenarios were rerun for a longer period (from 2010 to 2017), but assuming a lower reproductive rate where each particle in the ice was reseeded 1:1 every 1 April. This scenario in both cases (migrating and non-migrating) led to decreasing particle density.
numbers. For these simulations, the distribution pattern was similar to what is shown in Figure 6 for the years 2010 – 2013, but with lower concentrations. The subsequent development for the years 2014 – 2017 is shown in Figure 7. For the non-migrating particles, the total population size initially dropped rapidly as most A. glacialis in the Eurasian Basin exited through the Fram Strait within the first few years. From 2013, the population size decreased at a much lower rate, as most remaining particles...
were located in the Amerasian Basin and tended to circulate the basin with the Beaufort Gyre rather than following the Transpolar Drift toward and exiting through the Fram Strait. The number of migrating particles decreased at a steadier yet slower pace, as the transport out through the Fram Strait was slowed due to the seasonal migration. From 2016 to 2017 the spreading pattern of the non-migrating particles shifted quite rapidly in that also particles in the Beaufort Gyre/Amerasian Basin moved toward the Fram Strait, indicating a change in the drift during this period.

**DISCUSSION**

In this paper we studied the possible effects of seasonal migration of the amphipod *A. glacialis* on their population size and distribution within the Arctic Ocean using a particle tracking model. In our simulations we separately ran the model with two scenarios: (a) *A. glacialis* that stay close to the surface, following the ice drift or surface current year-round and (b) *A. glacialis* that behave in the same manner 1 April to 31 July but perform a deep vertical migration staying at 250 m for the rest of the year. After the first summer (31 July 2008) the particles were evenly distributed in the Arctic Ocean, the GIN Seas and the Barents Sea. However, after the initial 2 years, the number of non-migrating particles were reduced to nearly 60%, while the number of migrating particles remained high, nearly 90% of the original. Another interesting result was that, as years progressed to 2013, the non-migrating particles were mostly retained in the ice-covered Beaufort Gyre. Very few of the non-migrating particles were found in the central Arctic Ocean Basin, along the shelf break along Siberia, the shelf areas as well as the northern Barents Sea, in Svalbard waters, the Fram Strait and the Greenland Sea. The non-migrating particles were largely transported away from these areas and lost as the ice is melting. This is not surprising since the upper ocean currents in this region can lead to transport from the Siberian shelf to the Fram Strait within approximately 1 year (Timmermans and Marshall, 2020). On the other hand, the migrating particles were well distributed throughout the Arctic Ocean, along the shelf breaks, the Fram Strait and the northern Barents Sea.

The Transpolar Drift, along with the seasonal melting of sea ice in the Fram Strait, Greenland and Barents Seas, represents a large one-way transport of the core population of ice amphipods out of the Arctic Ocean. Accordingly, the ice amphipods (including *A. glacialis*) released as the ice melts over deep oceans, have been assumed to be permanently lost from their populations being subjected to predation in the pelagic system and/or become part of the vertical carbon flux (Lønne and Gabrielsen, 1992; Werner et al., 1999; Hop and Pavlova, 2008). Our findings, however, support the hypothesis of Berge et al. (2012) that *A. glacialis* instead reaches a more stable population size through performing a seasonal vertical migration. After they are transported by the ice drift into the GIN Seas, they migrate to depths in autumn and into the inflowing Atlantic Currents, being able to be transported back to the core population within the then ice-covered parts of the Arctic Basin. Further, our results suggest that without a seasonal vertical migration *A. glacialis* would be absent from most of the Arctic Ocean and a core population would only be maintained in the Beaufort Gyre. We argue that the seasonal vertical migration is a generally useful trait in lipid-rich, high-latitude herbivores as has been shown for the high-latitude *Calanus* copepods (Falk-Petersen et al., 2007, 2009).

**Is a Deep Migration Feasible?**

Our model output supports the concept that vertical migration behavior of *A. glacialis* as suggested based on observations of *A. glacialis* in deeper water layers in the Arctic Ocean (Berge et al., 2012; Kunisch et al., 2020) acts as a strategy that enhances survival and maintains a stable population in the ice-covered Arctic. Furthermore, since the suggested migrations are assumed to occur in the autumn after the peak of primary production it is a strategy that would not require additional energy reserves for the period the species would spend at depth. *Apherusa glacialis* is a mainly herbivorous species feeding on ice algae as well as phytoplankton when those sources are available (Scott et al., 1999; Kohlbach et al., 2016). During that time, it builds up lipid resources resulting in high lipid content from 45 to 60% lipids of dry mass (Berge et al., 2012; Kohlbach et al., 2016). These are lipid levels on par with those found for other key Arctic herbivorous species such as the copepods *Calanus finmarchicus*, *C. glacialis*, and *C. hyperboreus*. These three *Calanus* species are known for their unique life history strategies, feeding on high latitude blooms until they have filled their lipid depots, then migrating to depth for overwintering (Conover and Huntley, 1991; Falk-Petersen et al., 2000, 2009; Lee et al., 2006). While the benefits of diel vertical migrations are linked to predator avoidance, those of seasonal vertical migration have rather been linked to enduring the food-poor season (for herbivores) with lowered metabolic rates outside the often colder surface while subsiding on stored energy (Hagen, 1999). *Apherusa glacialis* also has the energy reserves needed to survive a long winter without feeding (Werner and Auel, 2005). Arctic blooms, consisting of ice algae and pelagic phytoplankton, are spatially variable and can occur at any time between March and September, dependent on light, nutrients, ice conditions, leads and opening in the sea ice, and upwelling of nutrients through the halocline in the Arctic Ocean and along the shelf break (Zenkevich, 1963; Daase et al., 2013; Ardyna et al., 2014; Falk-Petersen et al., 2015; Darnis et al., 2019; Randelhoff et al., 2020). The three *Calanus* species have life history strategies which are geared to different bloom situations. While *C. finmarchicus* mainly feeds on the traditional spring – summer bloom, the two other species being larger in size with large lipid stores, can utilize the bloom at any time during the primary production period between the spring and autumn equinoxes (Falk-Petersen et al., 2000). As soon as the *Calanus* species have filled up their lipid reserves they will migrate down to depth being transported by the interconnected current system inside and outside of the Arctic Ocean (Falk-Petersen et al., 2007). The seasonal migration to deep waters can take place as early as May – June and as late as September (Falk-Petersen et al., 2009). The present study, together with records of pelagic occurrences (Kunisch et al., 2020) and high lipid content (Berge et al., 2012; Kohlbach et al., 2016) suggest that similar strategies are
conceivable for a sympagic species such as *A. glacialis*. Whether or not *A. glacialis* is actively feeding during the phase at depth is unclear, though recent trophic marker studies suggest they may continue feeding (Kunisch et al., unpubl. data).

Another important part of the life history of *A. glacialis* that needs to be considered in evaluating the feasibility and ecological role of deep vertical migration is its life cycle. *Apherusa glacialis* has a lifespan of 2 years (Melnikov and Kulikov, 1980; Poltermann, 1998). Though it has initially been assumed to have a high fecundity compared to most gammarid amphipods with 370–727 eggs per female year$^{-1}$ (Poltermann, 2000), our recent observations show the number of eggs to be much lower (pers. obs. by author group, Figure 2). The here evaluated deep vertical migration is consistent with *in situ* observations that gravid *A. glacialis* females are more or less absent from spring to fall when these amphipods can be abundant under sea ice. Gravid females have, however, lately been regularly observed in deeper layers of the Arctic Ocean north of Svalbard in polar night months (Berger et al., 2012; Kunisch et al., 2020). In terms of required energy reserves and development, a deep migration in the polar night to layers consisting of water masses with temperatures several degrees higher than those found close to the sea ice might be advantageous for the developing juveniles carried in the brood pouch of the female. In spring, high numbers of freshly hatched juvenile stages (1–2 mm body length) directly under sea ice indicate upward migration of females with fully developed juveniles (or even of juveniles themselves) by that time. This timing is thought to take advantage of the ice algal spring bloom in sea ice (Kunisch et al., 2020) that precedes the pelagic bloom by weeks to months (Leu et al., 2015; Wassmann et al., 2020).

**Model Set-up, Performance, and Caveats**

With the model approach used in this study we simulated the trajectories of *A. glacialis* following both the sea-ice drift and ocean currents for several years. The current model set-up is clearly a simplistic first step to link the life strategy of *A. glacialis* to the Arctic hydrographic conditions, and future modifications to the model could reveal more details. Future runs could be modified to use observation-based abundance estimates of *A. glacialis* including their spatial variation. This species is the most abundant of sympagic amphipods with densities that vary from 0 to >150 ind. m$^{-2}$ with large spatial variation at multiple scales (Kunisch et al., 2020). Highest values reported are from around Svalbard and the Beaufort shelf, and lowest from the Beaufort Gyre (synthesized in Bluhm et al., 2017). *Apherusa glacialis* have not been reported from the Bering and southern Chukchi Seas (ibid.). In addition, experimental work focused on reproductive success could constrain a reproductive ratio closer to reality. Here our model estimates that a reproductive rate between 1:1 (loss) and 1:2 (increasing populations) would result in near constant abundances.

We chose a migration depth of 250 m, based on a combination of highest current velocities thereabouts in the core Atlantic layer in the Atlantic inflow region (Bluhm et al., 2020; Polyakov et al., 2020) and known vertical migration behavior of Calanoid copepods (Soreide et al., 2010; Daase et al., 2013). Modeled outputs (especially years 2012 and 2013, Figures 6C,D) showed the effect of this choice in enhanced *A. glacialis* particle concentrations at the shelf break in the Kara and Siberian Seas where the Arctic Circumpolar Boundary Current (ACBC) is the likely vector and reaches velocities on the order of 20 cm s$^{-1}$ (Bluhm et al., 2020). We did not adjust particle depth although we are aware that the ACBC eventually becomes deeper during its transit into and throughout the Amerasian Basin (Bluhm et al., 2020). Adjusting the migration depth according to the ACBC could result in the particles following the current trajectory more closely. So far, however, the observational evidence rather suggests that *A. glacialis* may occupy several depths (Kunisch et al., 2020). Considering this, the choice of a constant migrating depth probably underestimates the spread of the migrating particles. The more important finding, however, is that the higher numbers of migrating than non-migrating particles retained inside the Arctic Ocean largely stems from avoiding the major export with the ice in the winter months through the Fram Strait, which indicates that a pelagic state at any depth is advantageous for the retention in the Arctic. In the model simulations the migrating *A. glacialis* all undertake a deep migration at fixed dates and are assumed to passively stay at depth for the entire period from 1 August to 31 March the following year. This is clearly a simplification that possibly leads to an overestimated difference in the spreading pattern between the migrating and non-migrating particles. Algal productivity in the Arctic Ocean can occur also past August 1, as recently shown through remotely sensed algal autumn blooms (Ardyna and Arrigo, 2020). A more likely vertical movement pattern is that the deep migration is restricted to a shorter period. Furthermore, it seems reasonable to assume that the timing of the migration is more flexible and could be triggered at an individual and regional level, by e.g., lipid reserves being filled or by reduced sea ice extent or food concentrations. This is also more in line with the fact that *A. glacialis* can be found in the entire water column throughout the year (Kunisch et al., 2020). Adding individual behavior such as feeding to the model would require information about the quantity of food sources. A possible approach to this could be to run a coupled hydrodynamic, sea ice and biogeochemical model system that simulates e.g., ice algae blooms. However, running such a coupled system would greatly increase the computational cost. An advantage of the simpler approach in this study is the computational efficiency and hence the possibility to run simulations with a large number of particles. We consider therefore our approach as a first step to investigate the possible effect of vertical deep migration of *A. glacialis*. Although the simplified migration pattern introduces uncertainty, the clear results from the simulations show that a vertical deep migration will have a large impact on the spreading of this species.

The model simulations showed a change in the drift pattern during 2016–2017, as non-migrating particles that were in the Beaufort Gyre and Amerasian Basin drifted quite rapidly toward the Fram Strait. This feature is possibly related to the unusual sea ice cover pattern in 2016, which exhibited a record-low sea ice extent at the start of the year followed by a higher-than-expected sea ice extent in July–September and a record late freeze-up in the central Arctic (Petty et al., 2018).
CONCLUSION

Seasonal vertical migration considerably enhanced both the survival of *A. glacialis* and its retention within the Arctic Ocean region in our model. With a simplified parameterization of reproduction/mortality, the population increased from a factor slightly above 2 to a factor of 4 over the course of 5 years when vertical migration was included. We suggest that this behavior will increase survival in a warmer Arctic with little sea ice. If this pattern holds true for a fully ice-free Arctic, however, remains questionable, and additional targeted sampling will be beneficial. Targeted sampling would help our understanding if *A. glacialis* conducts vertical migrations en masse, similar to *Calanus* spp. One way to further test this hypothesis is by conducting a population genetics study in which we would hypothesize less genetic structure if *A. glacialis* was indeed conducting seasonal vertical migration, while we would expect more genetic differences if the species was entirely sympagic and more constrained to either the Transpolar Drift or the Beaufort Gyre systems.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

REFERENCES

Aksenov, Y., Ivanov, V. V., Nurser, A. J. G., Bacon, S., Polyakov, I. V., Coward, A. C., et al. (2011). The Arctic circumpolar boundary current. *J. Geophys. Res.* 116:09017. doi: 10.1029/2010JC006637

Anderson, M., Ovirk, K. A., LaCasce, J. H., Koszalka, I., and Mauritzen, C. (2011). Variability of the Norwegian Atlantic current and associated eddy field from surface drifters. *J. Geophys. Res.* 116:C08032. doi: 10.1029/2011JC007078

Ardyna, M., and Arrigo, K. R. (2020). Phytoplankton dynamics in a changing Arctic Ocean. *Nat. Clim. Chang.* 10, 892–903. doi: 10.1038/s41558-020-0905-y

Ardyna, M., Bahin, M., Gosselin, M., Devred, E., Rainville, L., and Tremblay, J. -É (2014). Recent Arctic Ocean sea ice loss triggers novel fall phytoplankton blooms. *Geophys. Res. Lett.* 41, 6207–6212. doi: 10.1002/2014GL061047

Arndt, C. E., and Swadling, K. M. (2006). Crustacea in Arctic and Antarctic sea ice: distribution, diet and life history strategies. *Adv. Mar. Biol.* 51, 197–315. doi: 10.1016/S0065-2881(06)51004-1

Barnard, J. L. (1959). Epipelagic and under-ice Amphipoda of the Central Arctic Basin. Scientific studies at Fletcher’s Ice Island, T-3. *Geophys. Res. Pap.* 63, 115–153.

Berge, J., Varpe, O., Moline, M. A., Wold, A., Renaud, P. E., Daase, M., et al. (2012). Retention of ice-associated amphipods: possible consequences for an ice-free Arctic Ocean. *Bioll. Lett.* 8, 1012–1015. doi: 10.1098/rsbl.2012.0517

Beuchel, F., and Lonne, O. J. (2002). Population dynamics of the sympagic amphipods *Gammarus wilkitzkii* and *Apherusa glacialis* in sea ice north of Svalbard. *Polar Biol.* 25, 241–250. doi: 10.1007/s00300-001-0329-8

Bluhm, B. A., Beuk, A. V., Gradinger, R., Hopcroft, R. R., Huettmann, F., Kosobokova, K. N., et al. (2011). An update of species richness and examples of biodiversity change. *Oceanography* 24, 232–248. doi: 10.5670/oceanog.2011.75

Bluhm, B. A., Hop, H., Melnikov, I. A., Poulin, M., Vihatarki, M., Collins, R. E., et al. (2017). “Sea ice biota,” in *CAFF State of the Arctic Marine Biodiversity Report*. *Conservation of Arctic Flora and Fauna*, eds T. Barry, C. Price, M. Olsen, T. Christensen, and M. Frederiksen (Akureyri: International Secretariat).

Bluhm, B. A., Janout, M. A., Davidson, S. L., Ellingsen, I., Gavrilov, M., Grebmeier, J. M., et al. (2020). The pan-Arctic continental shelf: sharp gradients of physical processes affect pelagic and benthic ecosystems. *Front. Mar. Sci.* 7:886. doi: 10.3389/fmars.2020.544386

Budgell, W. P. (2005). Numerical simulation of ice-ocean variability in the Barents sea region towards dynamical downscaling. *Ocean Dyn.* 55, 370–387. doi: 10.1007/s10236-005-0008-3

Conover, R. J., and Huntley, M. (1991). Copepods in ice-covered seas – distribution, adaptations to seasonally limited food, metabolism, growth patterns and life cycle strategies in polar seas. *J. Mar. Syst.* 2, 1–41. doi: 10.1016/0924-7963(91)90011-1

Daase, M., Berge, J., Serøide, J. E., and Falk-Petersen, S. (2021). “Ecology of Arctic pelagic communities,” in *Arctic Ecology*, ed. D. N. Thomas (Hoboken, NJ: Wiley Blackwell), 219–259. doi: 10.1002/9781118846582.ch9

Daase, M., Falk-Petersen, S., Varpe, O., Darnis, G., Serøide, J. G., Wold, A., et al. (2013). Timing of reproductive events in the marine copepod *Calanus glacialis*: a pan-Arctic perspective. *Can. J. Fish. Aquat. Sci.* 70, 871–884. doi: 10.1139/cfas-2012-0401

Darnis, W. O., Falk-Petersen, S., Graeve, M., and Fortier, L. (2019). Could offshoring predation offset the successful reproduction of the marine copepod *Calanus hyperboreus* under reduced sea-ice cover conditions? *Progr. Oceanogr.* 170, 107–118. doi: 10.1016/j.pocean.2018.11.004

Deep, P. U., Uppala, S. M., Simmons, A. J., Berrisford, P., Poli, P., Kobayashi, S., et al. (2011). The ERA-interim reanalysis: configuration and performance of the data assimilation system. *Q. J. R. Meteorol. Soc.* 137, 553–597. doi: 10.1002/qj.828

Doös, K., Jonsson, B., and Kjellson, J. (2017). Evaluation of oceanic and atmospheric trajectory schemes in the TRACMSS trajectory model v6.0. *Geosci. Model. Dev.* 10, 1733–1749. doi: 10.5194/gmd-10-1733-2017

Doös, K., Kjellson, J., and Jonsson, B. (2013). “TRACMSS—a Lagrangian trajectory model,” in *Preventive Methods for Coastal Protection*, eds T. Sœmøre and E. Quak (Heidelberg: Springer), doi: 10.1007/978-3-19-00460-2_7

Egbert, G. D., and Erofeeva, S. Y. (2002). Efficient inverse modeling of barotropic flow. *Geophys. Res. Lett.* 29, 1707. doi: 10.1029/2001GL013826

Egbert, G. D., and Erofeeva, S. Y. (2002). Efficient inverse modeling of barotropic flow. *Geophys. Res. Lett.* 29, 1707. doi: 10.1029/2001GL013826

Egbert, G. D., and Erofeeva, S. Y. (2002). Efficient inverse modeling of barotropic flow. *Geophys. Res. Lett.* 29, 1707. doi: 10.1029/2001GL013826

Egbert, G. D., and Erofeeva, S. Y. (2002). Efficient inverse modeling of barotropic flow. *Geophys. Res. Lett.* 29, 1707. doi: 10.1029/2001GL013826

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AUTHOR CONTRIBUTIONS

MD performed the particle tracking modeling, the model post-processing and visualization of results. SF-P and MD wrote the first draft of the manuscript. All authors contributed to the manuscript revision, read and approved the submitted version and conception and design of the study and analysis of model results, and wrote sections of the manuscript.

A. glacialis remains questionable, and additional targeted sampling will be beneficial. Targeted sampling would help our understanding if *A. glacialis* conducts vertical migrations en masse, similar to *Calanus* spp. One way to further test this hypothesis is by conducting a population genetics study in which we would hypothesize less genetic structure if *A. glacialis* was indeed conducting seasonal vertical migration, while we would expect more genetic differences if the species was entirely sympagic and more constrained to either the Transpolar Drift or the Beaufort Gyre systems.
Falk-Petersen, S., Hop, H., Budgell, W. P., Hegseth, E. N., Korsnes, R., Loyning, T. B., et al. (2000). Physical and ecological processes in the marginal ice zone of the northern Barents Sea during the summer melt period. J. Mar. Syst. 27, 131–139. doi: 10.1016/S0924-7963(00)00064-6

Falk-Petersen, S., Mayzaud, P., Kattner, G., and Sargent, J. R. (2009). Lipids and life strategy of Arctic Calanus. Mar. Biol. Res. 5, 18–39. doi: 10.1080/17451009002512267

Falk-Petersen, S., Pavlov, V., Berge, J., Cottier, F., Kovacs, K. M., and Lydersen, C. (2015). At the rainbow's end: high productivity fueled by winter upwelling along an Arctic shelf. Polar Biol. 38, 5–11. doi: 10.1007/s00304-014-1482-1

Falk-Petersen, S., Pavlov, V., Timofeev, S., and Sargent, J. R. (2007). “Climate variability and possible effects on arctic food chains: the role of Calanus,” in Arctic Alpine Ecosystems and People in a Changing Environment, eds J. B. Orbeck, R. Kallenborn, I. Trombe, E. N. Hegseth, S. Falk-Petersen, and A. H. Hoel (Berlin: Springer).

Gradner, R., and Bluhm, B. A. (2004). In-situ observations on the distribution and behavior of amphipods and Arctic cod (Boreogadus saida) under the sea ice of the high Arctic Canada Basin. Polar Biol. 27, 595–603. doi: 10.1007/s00300-004-0630-4

Gradner, R., and Bluhm, B. A. (2020). First analysis of an Arctic sea ice meiofauna food web based on abundance, biomass and stable isotope ratios. Mar. Ecol. Prog. Ser. 634, 29–33. doi: 10.3354/meps13170

Gulliksen, B., and Lønne, O. J. (1991). Sea ice macrofauna in the Antarctic and the Arctic. J. Mar. Syst. 2, 53–61. doi: 10.1016/0924-7963(91)90013-K

Hagen, W. (1999). Reproductive strategies and energetic adaptations of polar zooplankton. Invertebr. Reprod. Dev. 36, 25–34. doi: 10.1080/074229999.9652674

Hansen, B., and Osterhus, S. (2000). North Atlantic–Nordic Seas exchanges. Progr. Oceanogr. 45, 109–208. doi: 10.1016/S0079-6611(99)00052-X

Hattermann, T., Isachsen, P. E., von Appen, W. J., Albreten, J., and Sundfjord, A. (2016). Eddy—driven recirculation of Arctic water in far strait. Geophys. Res. Lett. 43, 3406–3414. doi: 10.1002/2016GL068323

Helland-Hansen, B., and Nansen, F. (1909). The Norwegian Sea, It’s Physical Variability and possible effects on arctic food chains: the role of Calanus,” in Arctic Alpine Ecosystems and People in a Changing Environment, eds J. B. Orbeck, R. Kallenborn, I. Trombe, E. N. Hegseth, S. Falk-Petersen, and A. H. Hoel (Berlin: Springer).

Hop, H., Falk-Petersen, S., Svendsen, H., Kwasniewski, S., Pavlova, V., Pavlova, O., et al. (2006). Physical and biological characteristics of the pelagic system across far strait to Kongsfjorden. Progr. Oceanogr. 71, 182–231. doi: 10.1016/j.pocean.2006.09.007

Hop, H., and Pavlova, O. (2008). Distribution and biomass transport of ice amphipods in drifting sea ice around Svalbard. Deep Sea Res. II Top. Stud. Oceanogr. 55, 2292–2317. doi: 10.1016/j.dsr2.2008.05.023

Hof, H., Falk-Petersen, S., Svendsen, H., Kwasniewski, S., Pavlova, V., Pavlova, O., et al. (2006). Physical and biological characteristics of the pelagic system across far strait to Kongsfjorden. Progr. Oceanogr. 71, 182–231. doi: 10.1016/j.pocean.2006.09.007

Hop, H., and Pavlova, O. (2008). Distribution and biomass transport of ice amphipods in drifting sea ice around Svalbard. Deep Sea Res. II Top. Stud. Oceanogr. 55, 2292–2307. doi: 10.1016/j.dsr2.2008.05.023

Kohlbach, D., Graeve, M. A., Lange, B., David, C., Pape, E., and Flores, H. (2015). Arctic spring awakening – Steering principles behind the phenology of the Barents Sea. Polar Biol. 10, 5–18. doi: 10.3402/polar.v10i1.6723

Lønne, O. J. (1991). Sea ice macrofauna in the Antarctic and the Arctic. J. Mar. Syst. 2, 53–61. doi: 10.1016/0924-7963(91)90013-K

Melnikov, A. (1997). “Modern terrigenous organic carbon input to the Arctic ocean,” in Proceedings of the Arctic Ocean Circulation, Processes and Water Masses: a Contribution to the International Polar Year 2007–2009, eds R. Stein and R. W. Garwood (Berlin: Springer), 33–55.

Pettersson, I. A. (2008). “Modern terrigenous organic carbon input to the Arctic ocean,” in Proceedings of the Arctic Ocean Circulation, Processes and Water Masses: a Contribution to the International Polar Year 2007–2009, eds R. Stein and R. W. Garwood (Berlin: Springer), 33–55.

Pettersson, I. A. Mel’nikov (Moscow: Nauka), 97–111.

Menne, S., Ingvaldsen, R. B., Haugan, P., Beschynska-Moeller, A., Fer, I., Sundsfjord, A., et al. (2019). Atlantic water pathways along the north-western Svalbard shelf mapped using vessel-mounted current profilers. J. Geophys. Res. Oceans 124, 1699–1716. doi: 10.1002/2018JC014299

Falk-Petersen, S., Pavlov, V., Berge, J., Cottier, F., Kovacs, K. M., Lydersen, C., Overland, J. E., and Moore, S. E. (2011). Impacts of climate-induced habitat change. Ecol. Appl. 20, 134–138. doi: 10.1890/06-0546.1

Falk-Petersen, S., Pavlov, V., Berge, J., Cottier, F., Kovacs, K. M., Lydersen, C., Overland, J. E., and Moore, S. E. (2011). Impacts of climate-induced habitat change. Ecol. Appl. 20, 97–125. doi: 10.1890/06-0546.1
of the Arctic Ocean from the 1990s to the late 2000s. Ocean Sci. J. 9, 147–169. doi: 10.5194/os-9-147-2013
Scott, C. L., Falk-Petersen, S., Sargent, J. R., Hop, H., Lomne, O. J., and Poltermann, M. (1999). Lipids and trophic interactions of ice fauna and pelagic zooplankton in the marginal ice zone of the Barents Sea. Polar Biol. 21, 65–70. doi: 10.1007/s00300030335
Semper, S., Pickart, R. S., Våge, K., Torres, D. J., and McRaven, L. (2020). CTD Temperature and Salinity Profiles Along Five Transects in NE Iceland / Iceland-Faroe Ridge. PANGAEA. doi: 10.1594/PANGAEA.919516
Shchepetkin, A. F., and McWilliams, J. C. (2005). The regional oceanic modeling system (ROMS): a split-explicit, free-surface, topography-following-coordinate oceanic model. Ocean Model. 9, 347–404. doi: 10.1016/j.ocemod.2004.08.002
Søreide, J. E., Leu, E., Berge, J., Graeve, M., and Falk-Petersen, S. (2010). Timing of blooms, algal food quality and Calanus glacialis reproduction and growth in a changing Arctic. Glob. Chang. Biol. 16, 3154–3163. doi: 10.1111/j.1365-2486.2010.02175.x
Stroeve, J. C., Kattsov, V., Barrett, A., Serreze, M., Pavlova, T., Holland, M., et al. (2012). Trends in Arctic sea ice extent from CMIP5, CMIP3 and observations. Geophys. Res. Lett. 39:L19501. doi: 10.1029/2012GL049870
Stroeven, A., Blockley, E. W., Furner, R., Guiavarch, C., Lea, D., Martin, M. J., et al. (2010). Forecasting the ocean state using NEMO: the new FOAM system. J. Oper. Oceanogr. 3, 3–15. doi: 10.1080/1755876X.2010.11020109
Timmermans, M.-L., and Marshall, J. (2020). Understanding Arctic ocean circulation: a review of ocean dynamics in a changing climate. J. Geophys. Res. Oceans 125:e2018JC014378. doi: 10.1029/2018JC014378
Wassmann, P., Carmack, E. C., Bluhm, B. A., Duarte, C. M., Berge, J., Brown, K., et al. (2020). Towards a unifying pan-arctic perspective: a conceptual modelling toolkit. Progr. Oceanogr. 189:102455. doi: 10.1016/j.pocean.2020.10.2455
Wernberg, T., Larkum, A. W. E., and De'ath, G. (2010). How will reduced coral cover affect coastal ecosystem services? Trends Ecol. Evol. 25, 75–83. doi: 10.1016/j.tree.2009.10.013
Woodgate, R. A., Aagaard, K., and Weingartner, T. J. (2005). A year in the physical oceanography of the Chukchi Sea: moored measurements from autumn 1990–1991. Deep Sea Res. Part II 52, 3116–3149. doi: 10.1016/j.dsr2.2005.10.016
Zenkevich, L. A. (1963). Biology of the Seas of the U.S.S.R. New York, NY: Interscience Publishers.

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