Opinion

Paradigm lost, or is top-down forcing no longer significant in the Antarctic marine ecosystem?

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Abstract: Investigations in recent years of the ecological structure and processes of the Southern Ocean have almost exclusively taken a bottom-up, forcing-by-physical-processes approach relating various species' population trends to climate change. Just 20 years ago, however, researchers focused on a broader set of hypotheses, in part formed around a paradigm positing interspecific interactions as central to structuring the ecosystem (forcing by biotic processes, top-down), and particularly on a "krill surplus" caused by the removal from the system of more than a million baleen whales. Since then, this latter idea has disappeared from favour with little debate. Moreover, it recently has been shown that concurrent with whaling there was a massive depletion of finfish in the Southern Ocean, a finding also ignored in deference to climate-related explanations of ecosystem change. We present two examples from the literature, one involving gelatinous organisms and the other involving penguins, in which climate has been used to explain species' population trends but which could better be explained by including species interactions in the modelling. We conclude by questioning the almost complete shift in paradigms that has occurred and discuss whether it is leading Southern Ocean marine ecological science in an instructive direction.

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"New opinions are always suspected, and usually opposed, without any other reason but because they are not already common." John Locke (1690), "An Enquiry Concerning Human Understanding"

Introduction

According to Thomas Kuhn (1962), most science, which he called "normal science," is conducted within existing, well-accepted paradigms. Anomalous findings are set aside until they become so numerous that they overwhelm an existing paradigm's explanatory power, after which a breakthrough is realized and for a short time "revolutionary science" reigns. Through an intellectual tussle, the anomalies are

consolidated into a coherent body of thought and the older paradigm gives way, sometimes requiring the passing of a generation before full adoption of the new paradigm.

In Southern Ocean (SO) marine science, such a shift in paradigm (or at least a central unifying concept) occurred during the early 1990s, but without the passing of a generation and with few attempts to forestall it (Croxall 1992, Murphy 1995). Before then, and reaching a climax in large-scale, international programs like BIOMASS and FIBEX (El-Sayed 1994, Siegel 2005), the accepted paradigm proposed that between-species interactions were considered of great importance in structuring the "Antarctic marine ecosystem" (AME), particularly with regard to the removal of more than a million baleen whales during the 1950–60s, thought in turn to have resulted in a huge surplus of their prey, Antarctic krill *Euphausia superba* Dana (e.g. Laws 1977, Fraser *et al.* 1992 and references cited therein). Krill was viewed as a central species in the AME, i.e. the pelagic-continental slope ecosystem (e.g. Beddington & May 1982), and international survey efforts were organized to quantify krill abundance and importance to non-cetacean species.

If interspecific interactions (such as competition for prey) were important in the AME, certain predictable patterns should have emerged. This has been well substantiated where a top-trophic species or a trophic competitor has been experimentally removed or re-introduced, as is the case in terrestrial, limnetic and certain other marine ecosystems (e.g. Carpenter & Kitchell 1988, Mills et al. 1993, Verity & Smetacek 1996, Terborgh et al. 2006, Jackson 2006). Ensuing studies testing this expectation in the SO found some evidence of a krill surplus but also found many inconsistent or conflicting patterns (see also Ballance et al. 2006). For example, while investigating the expected compensatory response of first or second order krill predators, researchers found an inconsistent decrease in age of maturity among crabeater seals Lobodon carcinophagus (Hombron & Jacquinot) (Bengtson & Laws 1985); increasing or recovering populations of various fur seals Arctocephalus spp but simultaneously decreasing populations of southern elephant seals Mirounga leonina L. (Payne 1977, Priddle 1992, Reid & Croxall 2001, McMahon et al. 2005); increasing chinstrap Pygoscelis antarctica Forster but decreasing Adélie P. adeliae (Hombron & Jacquinot) penguin populations (both should have increased as both are first-order krill predators; Woehler et al. 2001, Lynnes et al. 2002); and, finally, while recovery in populations of the baleen whales is proceeding most remain severely depleted, even to this day (Best 1993, Branch et al. 2004). Despite lack of clear support, the species-interaction paradigm persisted long enough (especially the krill surplus portion) that a fishery treaty emerged from an effort to understand krill better. This was the Convention for the Conservation of Antarctic Marine Living Resources (CCAMLR), a formal attempt to manage resource extraction as fishery nations began to look closely at the supposed krill surplus. Ironically in retrospect (see below), CCAMLR hoped to avoid the situation where human fisheries rather than natural competitors would replace the whales and affect top-down structuring of the AME in their stead (Mitchell & Sandbrook 1980, Bengtson & Laws 1985, Murphy 1995).

In the early 1990s a few authors (e.g. Taylor & Wilson 1990, Fraser *et al.* 1992) theorized that climate change and its affect on sea ice, rather than a krill surplus, explained some of the inconsistent trends and patterns evident in certain ecosystem components. This idea was closely followed by additional studies and a quickly rising tide of acceptance of what became the new central

paradigm - bottom-up forcing by physics and climate change as the single most important ecosystem driver - and it is within this that the majority of researchers now interpret their data (see summaries in Croxall et al. 2002, Siegel 2005, Nicol 2005). The paradigm shift was perhaps partly facilitated by the increasing ease with which physical data could be acquired owing to the technological revolutions also underway (e.g. powerful computers, satellite imagery and remote sensing, moorings of electronic instruments, etc). Under the new paradigm not only was there never a krill surplus but, in effect, whales were replaced not by krill but by gelatinous salps Salpa thompsoni (Foxton)! Atkinson et al. (2004) showed that the increase in salps was correlated with decreased sea ice extent during the 1960s-early 1970s; many other statistically significant correlations also have been shown to relate species' trends with sea ice at varying temporal scales (e.g. Loeb et al. 1997, Fraser & Hofmann 2003, Jenouvrier et al. 2005, Forcada et al. 2006).

Progressing beyond the description of correlations, but perhaps emboldened by the plethora of the fine work about species' relationships to climate change, as referred to above, other researchers have looked for and believe that they have found biotic proxies indicating major change in climate and the large-scale sea ice regime, also during the 1960s, well before the era of remote sensing (De La Mare 1997, Curran et al. 2003, Hilton et al. 2006). Such results, in accordance with the new paradigm, have been cited widely and with little questioning are used to confirm the current "normal science," emphasizing the supreme, direct importance of physical factors in structuring the AME (e.g. Smetacek & Nicol 2005, Barbraud & Weimerskirch 2006). Problematically, but in keeping with the rules of normal science, contrary views about climate and sea ice change during that period have been ignored and relegated to the status of anomalies. For example, in contrast to an analysis of whale catch records (De La Mare 1997, postulating a reduction of mean sea ice extent during the 1960s), Ackley et al. (2003), examining historical, direct observations of sea ice from ships as well as comparing more recent observations with satellite imagery, found no evidence of a large-scale shift from higher to lower mean ice extents from the 1950s to the 1970s. Zwally et al. (2002), using satellite imagery, have also found no evidence for a dramatic multi-year decrease in ice extent anywhere other than the Antarctic Peninsula region in recent decades, a finding acknowledged recently by Siegel (2005) and Nicol (2005) but ignored by many other biologists. We do not dispute that climate change has been occurring around Antarctica, affecting changes in sea ice extent, ice thickness and the sea ice season (also Parkinson 2002) as well as some populations - especially in the Antarctic Peninsula region (e.g. Smith et al. 1999) but also elsewhere (see above, also Barbraud & Weimerskirch 2001, Ainley et al. 2005).

We do find it perplexing, though, that recent studies have rarely considered the extent of the total biotic variation explained by physical factors, or that factors other than physical ones could also be involved in explaining temporal and regional variation in the AME foodweb structure.

It cannot all be climate

While momentum for this physical forcing/climate change paradigm accelerated, overtaking the species-interactions paradigm, Pauly et al. (1998) published an analysis that introduced the concept of "fishing down the food web." This concept has gained acceptance for marine environments other than the SO as more and more researchers detail the loss of top-down trophic structuring owing to over-fishing of top- and middle-trophic level species (e.g. NSF 1998, Schindler et al. 2002, Soulé et al. 2005, Scheffer et al. 2005, Frank et al. 2005). Included in the Pauly et al. analysis, among examples from around the world, was a tally of the SO (non-cetacean) fishery landings that showed the now characteristic successively downward depletion of trophic levels in order to maintain economic viability of burgeoning fishing fleets (see also Myers & Worm 2003, for independent, corroborating analysis for the Scotia Sea). According to this analysis, ostensible fishery-driven food-web shift in the SO (not to

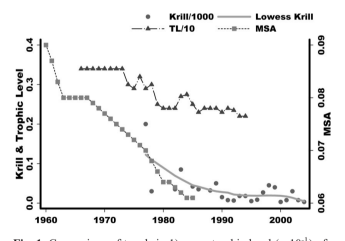


Fig. 1. Comparison of trends in 1) mean trophic level ($\times 10^{-1}$) of fishery landings from FAO areas 48, 58 (Scotia Sea) and northern 88 (does not include cetaceans already depleted in 1950s–early 1960s; from Pauly *et al.* 1998), 2) krill density from net tows taken in Scotia Sea region, with salps having a mirror-image trend [no. m⁻² 1000⁻¹; line shows predicted values from locallyweighted regression (lowess smoothing), data from Atkinson *et al.* 2004], and 3) 20 yr running mean of methanesulphonic acid (MSA, µM), a derivative of phytoplankton-produced dimethylsulphide (DMS), found in glacial ice-core annual layers at Law Dome, (western) East Antarctica (Curran *et al.* 2003). Less DMS and MSA are produced if salps rather than krill are grazing the phytoplankton (Katamatsu *et al.* 2004).

mention the large-scale whale extraction), also took place during the 1960s, much later than elsewhere on the globe. And, more critical to the old SO paradigm and perhaps providing reasons for inconsistent support of a krill surplus, it was not just cetaceans that had been depleted. Also removed were large predatory fish, a fact that is critical to consider in an analysis of food-web structure as fish are the most important predators in most marine ecosystems (Sheffer et al. 2005, and others). What is not surprising within the current confines of normal science is that we have yet to find even one paper in the SO literature that has cited Pauly et al., even though that paper provides crucial support for the importance of interspecific interactions required for the credibility of the betweenspecies-interactions paradigm. Moreover, many Antarctic papers published over the past decade that have shown correlations between a long-term biological variable and a physical one, such as sea-surface temperature or sea ice extent (or their proxies), probably would find an equally compelling correlation if observations were compared instead against the declining trophic level/commercial fisheries data of Pauly et al. The latter are also characterized by a steep decline beginning in the 1960s (cf. De La Mare 1997, Curran et al. 2003, Atkinson et al. 2004; Fig. 1). The ocean around the colonies of polar penguin species (whose diet trophic level has also been declining - a metric that has been suggested as a proxy for decreased phytoplankton production (Hilton et al. 2006)) historically has been one of the most heavily fished, and depleted, waters on the globe (Pauly et al. 2005). It seems, therefore, that the Pauly et al. analyses are included in those that most researchers have relegated, under the climate change paradigm, to the category of anomaly, even though they certainly appear to be relevant to understanding the AME.

Assuming that the rate of commercial take of resources can be a proxy for their availability (see references above regarding top-down forcing; also Myers & Worm 2003), these data also indicate that a major shift has occurred in the food-web structure of the AME (Pauly et al. 1998), and this shift has little to do with climate. Many feed-back loops and competitive and synergistic relationships are now probably gone (Pauly & Maclean 2003) due to the loss of major portions of upper- and middle-trophic levels especially around most sub-Antarctic and lower latitude Antarctic islands and northern Antarctic continental shelves, although food webs in some more southerly areas, e.g. the Ross Sea, remain to date intact. This loss occurred within the professional lifetimes of still-practicing SO ecologists and their students, and occurred just before the dramatic advances of modern technology that now define SO marine science (mid-1970s onward), as noted above. Thus, the vast majority of data supporting the physicallyforced view of the AME have been collected well after the depletion of top predators in the system was well underway.

Unfortunately this depletion probably continued coincident with apparent or real shifts in physical factors (Zwally et al. 2002, Parkinson 2002, Jacobs et al. 2002, Jacobs 2006). Because fishing down the food web can affect ecosystems just as dramatically as climate change (Pauly & Maclean 2003), causes of patterns have merged and what is concluded about the AME may primarily be distorted by researcher bias and the lack of long-term data on food-web structure. In the Bering and North seas and Benguela Current, for instance, where researchers accept the idea of major climate change as well as heavy fishing pressure, the dramatic rise in the abundance of gelatinous organisms has been explained by the fishery pressure that has reduced grazing on their larvae (Brierly et al. 2001, Brodeur et al. 2002, Heymans et al. 2004, Lynam et al. 2005), with similar arguments being made for other, smaller systems (e.g. Mills 2001, Xian et al. 2005). On the other hand, in the mind-set of most scientists working under the climate change paradigm in the "relatively pristine" AME (e.g. a concept used in the summaries by Croxall et al. 2002, Smetacek & Nicol 2005), the current hypothesis is that it is simply changed sea ice that is responsible for a similar dramatic rise in gelatinous creatures (Atkinson et al. 2004, Kawaguchi et al. 2005). However, because both heavy fishing and salps are concentrated and/or dominant in waters of non-existent, reduced or declining ice cover in the AME (see Nicol et al. 2000), a positive correlation between heavy fishing and increased salps could well be involved (see also Lynam

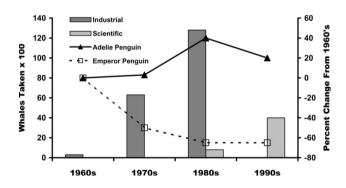


Fig. 2. Numbers of minke whales (×100) removed from IWC areas V and VI (Adélie Land and Ross Sea sector) by decade in the industrial and scientific whaling eras, compared to percent differences relative to 1960s numbers of Adélie penguins breeding at capes Royds and Bird, Victoria Land, and emperor penguins at Pointe Géologie, Adélie Land. The Cape Royds colony, the longest monitored in the Antarctic, was the same size in 1959 as it was in 1909, with no evidence that it had been larger in historical times (Ainley 2002a). These penguins and whales spend the late summer, autumn and winter in the same habitat and region; minke whales and Adélie penguins have the same diet (Ainley 2002b). Data on penguins are from Wilson *et al.* (2001) and Weimerskirch *et al.* (2003); those for whales from Brown & Brownell (2001).

et al. 2006). In the case of the gelatinous salps, although notoriously difficult to detect in the highly acidic stomachs of fish, they have been found in the diet of certain SO piscine species (Casaux *et al.* 1990, Pakhomov 1997, Barrera-Oro 2003, Bushula *et al.* 2005), though not yet in the little-researched diets of pelagic juveniles of commercially important benthic fish (e.g. Pakhomov & Pankratov 1992, Koch & Eversen 1997, Barrera-Oro *et al.* 2005). Finally, it is very likely that salps eat eggs and early larvae of krill (reviewed by Siegel 2005), a scenario consistent with the decline in krill as salps proliferate (cf. Atkinson *et al.* 2004).

Whale extraction may well have had a major effect

Some of us, exercising "normal science," recently presented ideas about how changes in climate, weather and sea ice may have affected the long-term population trends of certain Antarctic penguin species breeding in the Ross Sea/ western Pacific Ocean sector and elsewhere in the SO (Ainley et al. 2005). By contrast, we compare the Ross Sea penguin trends to the rate of extraction of a major trophic competitor of Adélie penguins, the Antarctic minke whale Balaenoptera bonaerensis Burmeister, from the same region (Fig. 2). The population of this pagophilic cetacean, which completely overlaps the penguins' habitat and food preferences (an important coincidence; Murphy 1995), was aggressively targeted two decades later than the other, more open-water baleen whales (Bengtson & Laws 1985). The effects to the population of the more than 116 600 taken (Clapham & Baker 2001), about 20% from the region of the Ross Sea (Brown & Brownell 2001) is unknown. Adélie penguins, seemingly released from trophic competition, show the appropriate demographic lag as increased penguin productivity and survival result in population growth for this slow-to-mature (up to seven years) upper-level predator. The fact that coastal polynyas were increasing in size during this time as well would facilitate the penguin increase (cf. Ainley 2002a, Parkinson 2002, Ainley et al. 2005). As the take of minke whales slackened beginning in 1987 (Brown & Brownell 2001), and best demonstrated in the Ross Sea, the penguin increase slackened.

We then began to question the rapid decline attributed to a mysterious short-term episode of adult mortality in emperor penguins Aptenodytes forsteri Gray in the same region (Weimerskirch et al. 2003), a mortality event correlated to climate effects (Barbraud & Weimerskirch 2001. Jenouvrier et al. 2005). Also considered subsequently have been effects of changed climate on this species' inability to recover from that mortality event (Jenouvrier et al. 2004, Ainley et al. 2005). In accord with massive adult mortality, the population decline in Adélie Land, a short distance west of the Ross Sea, was sharp (see Jenouvrier et al. 2005) and it occurred in the early part of the Adélie penguin increase (Fig. 2). Not coincidentally we suggest, the mortality event happened at the time when minke-whaleeating killer whales Orcinus orca (L.) type A (Pitman & Ensor 2003), inhabiting the waters frequented by these penguins (again, see Murphy 1995), were being challenged by a major, rapid large-scale removal of their main prey, i.e. the minke whale. Until that time it was thought that minke whale and killer whale populations, prey and predator, were in equilibrium (N.V. Doroshenko in Mikhalev et al. 1981). Did these top predators switch to alternate prey including emperor penguins, especially penguins whose vulnerability was increased by a possible short-term divergence in sea ice (as proposed by Jenouvrier et al. 2005), and a habitat frequented only at its periphery by type A killer whales (Pitman & Ensor 2003)? Emperor penguins have been found in killer whale stomachs (type unknown; Prévost 1961), and it is well known that these predators (type A) eat high numbers of the closely related king penguin A. patagonicus in the northern part of this SO region (e.g. Condy et al. 1978, Guinet 1992, Guinet & Bouvier 1995). Owing to confounding climate effects (warmer winter temperatures, stronger winds, thinner sea ice on which to breed), the emperor penguin population has not been able to recover from the initial massive mortality (cf. Barbraud & Weimerskirch 2001, Jenouvrier et al. 2004, Ainley et al. 2005). This scenario is reminiscent of a hypothesized killer whale prey switch, in this case from whales to sea otters Enhydra lutris L., an idea developed to explain changes in the fauna of the Aleutian Islands (Springer et al. 2003), and a hypothesized switch to sea lions (Otaria flavescens (Shaw)) and elephant seals in the Southern Ocean (Branch & Williams 2006). Moreover, an emperor penguin is far more of a meal for a killer whale than is the similar sized sea otter, both weighing 25-35 kg as the high fat content of the penguin would make it superior prey, probably equivalent to an energy-rich sea lion pup (Williams et al. 2004).

Modelling Southern Ocean ecosystem structure and change

We will end here with some questions, hoping that the reader has at least given some credence to the hypothesis that the AME is vastly different from the pre-1970s due not just to climate effects but equally to biotic extractions from the ecosystem. We point out that in ecology, but not the physical sciences, it is important to keep in mind how paradigms have evolved or been replaced (Dunlap 2006). Why did the paradigm involving the massive 1960s whale loss and its possible effect on the AME fall out of favour so quickly and with very little contest? Why have the possible ecological effects of the concurrent massive extraction of finfish from the SO not been acknowledged by ecologists? By keying on physical, bottom-up forcing, most SO marine biologists, in effect, are now

acknowledging that the AME no longer has the resiliency it once had when numerous biotic feedback loops were in place to ameliorate climate effects? That is, lacking the biotic feed-backs, is the AME now more susceptible to physical forcing (see Balance et al. 2006), as is now being argued for depleted non-marine ecosystems (e.g. Post et al. 2000, Wilmers et al. 2002, Wilmers & Getz 2005)? Indeed, the dramatic year-class cycling evident in the krill that dominate the Adélie penguin diet off the Antarctic Peninsula (Fraser & Hofmann 2003) - and where alternate prey (fish) are no longer available in quantity (see above, also Emslie & McDaniel 2002) - is reminiscent of the simple food webs of the terrestrial Arctic, where predator populations closely track their limited prey. No wonder that in this area strongly affected by both climate change and fishery depletion, unlike high-latitude portions of the Antarctic, the Adélie penguins are being replaced only in part by other, open water species (Fraser & Patterson 1997, Ainley et al. 2005, Forcada et al. 2006).

No doubt a strong mix of factors is involved in the restructuring of the AME food web, with both climate change and depletion of top- and middle-trophic level species playing a role (as noted earlier, e.g. by Croxall 1992), but surely it is time for some serious re-thinking on their relative contributions and what it is we are actually measuring or managing in the observed trends of SO organisms? Finally, will we not be able to answer some of these questions better if portions of the SO, such as the Ross Sea, are kept free of intensive biotic extractions? The latter, following the progression experienced by other ocean systems (Pauly *et al.* 1998), has now begun in that neritic system (Ainley 2002b).

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