

## Chapter 10.7. Phylogeographic patterns of the Southern Ocean crinoids

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Census of Antarctic Marine Life SCAR-Marine Biodiversity Information Network

# BIOGEOGRAPHIC ATLAS OF THE SOUTHERN OCEAN

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# **10.7. Phylogeographic patterns of the Southern Ocean crinoids (Crinoidea: Echinodermata)**

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#### 1. Introduction

Crinoids are well represented in the Southern Ocean. They locally may constitute one of the major mega-epibenthic components and therefore play a crucial role in the functioning of some ecosystems. Because some species are very abundant and well distributed over the whole Southern Ocean (Eléaume 2006; Hemery 2011; Eléaume *et al.* Chapter 5.25, this volume), they have been recognised as model organisms for studying the spatial variance of genetic diversity in the Southern Ocean.

The main hypothesis that explains the patterns of distribution of Antarctic benthic genetic diversity is linked to the cyclical advance of ice sheets on the Antarctic continental shelf. In this hypothesis, the Antarctic shelf ecosystems have undergone cyclical disturbance events during the last 35 My (see review in Turner et al. 2009), including as many as 38 sedimentary cycles of ice sheet advance during the last 5 My (Naish et al. 2009). It is thought that ice advance has regularly eradicated the benthic shelf fauna. Thatje et al. (2005, 2008) suggested that the benthic fauna now flourishing on the shelf arose from refuges either in areas left free of ice on the shelf (e.g. polynyas or areas not impacted by ice advance), on the adjacent slopes and deep-sea floor, or on the shelves of sub-Antarctic islands. Allcock & Strugnell (2012) summarised the expected molecular patterns for each of these hypotheses. As a result, widely distributed populations were fragmented into smaller populations that have diverged, and sometimes developed barriers to reproduction. This may have been followed by range expansion and, in the case of broadcast spawners, rapid recolonisation of habitats left free of ice, and secondary contact of refugial populations. Benthic taxa lacking a dispersal phase are often structured in haplogroups segregated in narrow geographical ranges (Rogers 2007). Clarke & Crame (1992, 2010) proposed that the cyclical nature of the vicariance events that may have resulted in genetic divergence and sometimes in allopatric speciation enhanced the Antarctic "biodiversity pump" that resulted from "the regular pulses of migration in and out of Antarctica driven by climate variability" (Clarke & Crame 1992, p. 299). Cycles of ice extension and retreat are often seen as catastrophic disturbance events leading to large-scale environmental instability, leading in turn to large-scale benthic eradication. However, the megabenthic fauna on the Antarctic shelf is thriving and seems to have rapidly recovered from the cyclical disturbance events, the latest being the Last Glacial Maximum (LGM) some 20,000 years ago. Variation of food availability (which may be linked to ice concentration but also to other factors such as current velocity), interspecific competition for space and food, pelagic larval duration or larval mortality, may also have contributed to the observed patterns, and should receive greater attention in the future. Here we explore, using the Cytochrome c Oxydase subunit I (COI) mitochondrial DNA, the spatial variance of the genetic diversity of the most common Southern Ocean crinoid species, which include broadcast spawners as well as brooders, and examine congruence with predictions of the biodiversity pump hypothesis.

#### 2. Datasets used

The datasets analysed here are already available from previous publications and registered in GenBank and BOLD (accession numbers are given in each publication). The sequences used for the phylogenetic reconstruction are available through Hemery *et al.* (2013a) and comprise data from COI, 16S, 18S and 28S genes. DNA extraction, gene amplification techniques and sequences used for the phylogenetic and phylogeographic analyses are described in Wilson *et al.* (2007), Eléaume *et al.* (2011), Hemery (2011), Hemery *et al.* (2012), Hemery *et al.* (2013a) and Hemery *et al.* (2013b).

#### 3. Antarctic crinoids in context

Hemery *et al.* (2013a) published a phylogenetic study based on a high-resolution taxon sampling, which serves as the basis for this study. Here, we use the same DNA markers, the same sampling effort, and the same reconstruction techniques. Hemery *et al.* (2013a) used four mitochondrial and nuclear genes and 105 taxa, which are differently presented here (Fig. 1). The classification and a nomenclature of extant crinoids used here predate the molecular analyses, and are mainly based on Roux *et al.* (2002) for stalked crinoids, and Messing (1997) for comatulids. The position of Antarctic species is highlighted in the tree to clearly demonstrate their phylogenetic context.

Antarctic crinoids are polyphyletic (see blue taxa in Fig. 1). They do not constitute a single clade that could have originated from a single ancestor, but are found scattered in the tree in most of the major clades (e.g. clades A, B1, B2, B3, B4, see Hemery *et al.* 2013a for details). This suggests that crinoids have colonised the Southern Ocean several times at different periods, or that crinoids have colonised the Southern Ocean independently from different ocean basins at roughly the same time. Within the Hyocrinidae, Antarctic and sub-Antarctic taxa are not monophyletic and are found associated with North

and South Pacific, and North Atlantic species. Within the Pentametrocrinidae, one Pentametrocrinus and one Thaumatocrinus species (see Eléaume et al. Chapter 5.25, this volume) are known from the Antarctic and sub-Antarctic regions. Thaumatocrinus has representatives in the North Atlantic, and Pentametrocrinus is found in the Indo-West Pacific and North Atlantic deep basins. In addition, Rouse et al. (2013) have demonstrated that the Indo-Pacific stalked crinoid genera Guillecrinus and Vityazicrinus are closely related to Pentametrocrinus. Within Bathycrinidae, a clade composed of Antarctic Bathycrinus australis and an Indo-West Pacific Monachocrinus sp. (both stalked) is sister to Atelecrinidae, a widespread bathyal family of feather stars. The Antarctic brooding genera Eumorphometra, Eometra and Isometra are related to North-Atlantic and Indo-West Pacific species (clade Antedonidae #1). Heliometrinae, a subfamily of Antedonidae that includes numerous Antarctic species, returns as two separate clades (see Eléaume 2006 for morphological arguments). Heliometrinae #1 is purely Antarctic and may result from in situ diversification events. Heliometrinae #2 comprises one Caribbean species, East Pacific and Arctic taxa, and at least one true Antarctic species, Anthometrina adriani.

Antarctic crinoids seem to have originated from ancestors from various ocean basins, including the Southern Ocean. Some taxa seem to have colonised the Southern Ocean repeatedly (e.g. Hyocrinidae, Pentametrocrinidae), other taxa seem to have radiated in situ (e.g. Heliometrinae #1, Notocrinidae, Isometrainae, and the hyocrinid genus *Dumetocrinus*).



**Figure 1** Cladogram of extant crinoids (modified from Hemery *et al.* 2013a) using molecular markers (COI, 16S, 18S and 28S). A total of 105 taxa are included and 3848 base positions were analysed. Maximum Likelihood (ML) and Bayesian Inference (BI) were performed on the four genes combined as an unlinked-partition dataset. For ML analyses, the model GTR+F (General Time Reversible) was used for each unlinked-partition, and bootstrapping was carried out with 1000 replicates using the same model. For BI analyses, the model GTR+I+F was used for each unlinked-partition. Two iterations of fifty million generations were run with eight chains, sampling every 1000 generations. ML and BI tree topologies were identical. Bayesian posterior probabilities values are given close to nodes; family and subfamily names are in bold; numbers in brackets indicate number of genera, species and specimens, respectively, included in the clade; names of clades discussed in Hemery *et al.* (2013) are given at nodes; Antarctic taxa are highlighted in blue; arrows indicate taxa that contain Southern Ocean species; stars indicate brooding species; HN = Holopus + Neogymnocrinus; CAP = Calometridae + Asterometridae + Ptilometridae; ZHMC = Zygometridae + Himerometridae + Mariametridae + Colobometridae.

#### 4. Phylogeographic patterns and demographic history

Only a subset of the Southern Ocean species collected and presented in the cladogram above were represented by a sufficient number of individuals to be suitable for further phylogeographic analyses. These species, the number of sequences analysed, and their reproductive strategies are as follows: *Anthometrina adriani* (146, broadcast spawner), *Feracrinus heinzelleri* (34, possibly a broadcast spawner), *Florometra mawsoni* (513, broadcast spawner), *Isometra graminea* (47, brooder), *Notocrinus virilis* (175, brooder), *Promachocrinus kerguelensis* (1429, broadcast spawner), and *Ptilocrinus amezianeae* (66, possibly a broadcast spawner). The sampling method was opportunistic and dependent on ship-based and station-based operations in Antarctica. Many areas of interest have not been sampled yet because of ice conditions or remoteness of these areas. Many spatial gaps remain, even though the sampling effort used here is unprecedented and in most cases is likely to encompass the whole distribution range of species under study.

from neighboring population by several hundred nautical miles. Ten different populations have been identified: Amundsen Sea (AS), Burdwood Bank (BB), Bouvet Island (BI), Davis Sea (DS), Dumont d'Urville Sea (DDU), Eastern Weddell Sea (EWS), Kerguelen Plateau (KP), Ross Sea (RS), Admiralty and Scott Island Seamounts (ASIS), Scotia Arc (SA), West Antarctic Peninsula (WAP). In the case of P. kerguelensis for which enough specimens are available, the Scotia Arc area has been separated into four distinct populations: South Shetland Islands (SSh), South Orkney Islands (SO), South Sandwich Islands (SSa), and South Georgia (SG). Hemery et al. (2012) described the methods used to estimate population diversity and the genealogical relationships among haplotypes. COI sequences were aligned using BioEdit Sequence Alignment Editor v7.0.9.0 (Hall 1999); haplotype networks were constructed using Network software (version 4.5.0.0; http://www.fluxus engineering.com); TCS1.21 (Clément et al. 2000) was used to test the connection threshold at which groups of haplotypes disconnect. Population diversity indices (Hd, haplotype diversity;  $\pi$ , nucleotidic diversity; Fu's F<sup>s</sup>) were calculated using ARLEQUIN v.3.5.1.2 (Excoffier & Lischer 2010) and are given in Table 1.

Defining a population is not trivial (Avise 2000). Here we define a population as a group of sequences collected in the same area, typically separated

**Table 1** Diversity indices for seven species largely distributed in the Southern Ocean. An = number of haplotypes; Fu's FS = result of the Fu's FS test; Hd = haplotype diversity; N = number of specimens;  $\pi$  = nucleotidic diversity; p = significance level of the Fu's FS with \* = p<0.05, \*\* = p<0.01, \*\*\* = p<0.005. AS = Amundsen Sea; ASIS = Admiralty and Scott Island Seamounts; BB = Burdwood Bank; BI = Bouvet Island; DDU = Dumont d'Urville Sea; DS = Davis Sea; EWS = Eastern Weddell Sea; HEA = Heard Island; KP = Kerguelen Plateau; RS = Ross Sea, and SA = Scotia Arc, the latter divided for *Promachocrinus kerguelensis* into SSa = South Sandwich Islands, SSh = South Shetland Islands, SO = South Orkney Islands, SG = South Georgia, and WAP = West Antarctic Peninsula.

Species/ phylogroup	Population	Ν	An	Hd	π	Fu's FS	р
Promachocrinus kerguelensis A	KP	113	16	0.7162	0.0019	-18.519	***
	DS	1	1	_	_	_	
	DDU	32	6	0.6855	0.0043	-1.397	
	RS	86	9	0.6621	0.0048	-3.256	
	SSh	26	2	0.6092	0.0040	1.349	
	SSa	15	3	0.7619	0.0018	-2.841	**
	SG	2	2	_	_	—	
	BI	36	2	0.1619	0.0002	-2.590	***
	EWS1	3	2	—	—	—	
Promachocrinus kerguelensis B	DS	1	1	—	—	—	
	DDU	54	4	0.6988	0.0017	-3.221	*
	RS	40	1	0.5859	0.0015	-0.935	
	SSh	9	0	0.2222	0.0004	-0.263	
	EWS1	2	2	—	—	—	
	EWS2	1	1	—	—	—	
Promachocrinus kerguelensis C	DS	54	2	0.6108	0.0013	-0.987	
	DDU	179	9	0.7423	0.0040	-2.056	
	RS	10	0	0.3778	0.0020	0.683	
	AS	3	1	_	—	—	
	WAP	18	4	0.8301	0.0040	-2.202	
	SSh	13	2	0.6923	0.0016	-2.036	*
	SSa	35	6	0.5664	0.0013	-6.758	***
	SO	34	1	0.4688	0.0010	-0.920	
	EWS2	30	0	0.6644	0.0027	0.430	
	EWS1	71	11	0.8072	0.0059	-3.258	
Promachocrinus kerguelensis D	DS	38	2	0.6088	0.0031	-0.660	
	DDU	82	5	0.6197	0.0044	-1.706	
	RS	53	7	0.8694	0.0074	-0.261	
	AS	13	2	0.7308	0.0029	0.514	
	WAP	10	1	0.5333	0.0047	1.176	
	SSh	18	7	0.8497	0.0083	-1.623	
	EWS1	13	3	0.8590	0.0054	-2.291	
	EWS2	8	0	0.4643	0.0035	1.493	
Promachocrinus kerguelensis E1	DDU	11	0	0.0000	0.0000	—	
	RS	8	0	0.0000	0.0000	—	
	SSh	5	0	0.0000	0.0000	—	
	SO	25	6	0.4300	0.0012	-4.900	***
	SG	6	0	0.0000	0.0000	—	
	EWS1	1	1	—	—	—	
Promachocrinus kerguelensis E2	DS	22	0	0.0000	0.0000	-	
	DDU	51	3	0.1153	0.0002	-4.339	***
	RS	68	1	0.0294	0.0001	-1.894	***
	WAP	2	1	-	-	-	
	EWS1	22	2	0.7056	0.0016	-0.184	
	EWS2	2	2	-	_	_	



Species/ phylogroup	Population	Ν	An	Hd	π	Fu's FS	р
Promachocrinus kerguelensis F	DS	68	1	0.5083	0.0009	0.580	
	DDU	9	0	0.0000	0.0000	—	
	RS	4	1	—	—	—	
	AS	1	1	—	—	—	
	WAP	9	0	0.0000	0.0000	—	
	SSh	1	1	—	—	—	
	SO	2	1	—	—	—	
	SSa	1	1	—	—	—	
	EWS1	8	1	0.6071	0.0014	-0.224	
Anthometrina adriani	DS	45	7	0.7636	0.002104	-1.606	
	DDU	33	7	0.6818	0.001716	-2.781	*
	RS	50	8	0.6833	0.001627	-3.461	*
	EWS	18	4	0.6078	0.001314	-0.841	
Florometra mawsoni	KP	62	20	0.7039	0.0020	-21.957	***
	DS	90	9	0.2844	0.000705	-8.394	***
	DDU	240	30	0.8009	0.003114	-22.585	***
	RS	27	5	0.4416	0.000879	-2.709	**
	AS	3	1	—	—	—	
	WAP	12	2	0.1667	0.000605	0.432	
	SA	9	3	0.4167	0.0008	-1.081	*
	BB	2	2	—	—	—	
	EWS	68	12	0.6817	0.003611	-2.966	
Notocrinus virilis A	SA	1	1	—	—	—	
	EWS	35	5	0.2185	0.007143	-4.015	***
Notocrinus virilis B	DS	5	1	—	—	—	
	DDU	67	5	0.2687	0.011578	-1.973	
	RS	56	4	0.1052	0.00267	-4.521	***
	SA	6	1	—	—	—	
	EWS	5	1	—	—	—	
Isometra graminea	DDU	31	5	0.5441	0.001282	-1.61702	
	RS	13	2	0.5128	0.000968	1.1512	
	WAP	3	2	0.6667	0.001258	0.20067	
Feracrinus heinzelleri	DS	4	4	1.0000	0.007202	-0.61511	
	RS	28	12	0.8730	0.00674	-1.74279	
	AS	2	2	—	—	—	
Ptilocrinus amezianeae	KP	23	6	0.4585	0.001116	-3.23397	***
	ASIS	40	9	0.5026	0.000932	-7.74253	***
	SA	3	1	—	—	—	

**Patterns** – Haplotype networks exhibit different topological patterns across all species (Fig. 2). *Promachocrinus kerguelensis* is composed of seven groups of haplotypes (A, B, C, D, E1, E2, F), each displaying a star-like topology, with one central and common (ancestral) haplotype surrounded by a crown of a varying number of derived secondary and less frequent haplotypes or singletons. *Florometra mawsoni*, which displays a star-like topology as well, connects to this network at a larger distance (Fig. 2A). Hemery *et al.* (2012) interpreted this pattern as congruent with the hypothesis of multiple refugia on the high-Antarctic and sub-Antarctic islands shelves during past glaciations, followed by population expansion, rapid recolonisation and secondary contact. Nuclear marker networks (Hemery *et al.* 2012) display two distinct lineages that indicate that the divergence in the mitochondrial genes was not found in the nuclear genes. These results have been interpreted as congruent with a single or two different species in *Promachocrinus*.

Anthometrina adriani displays a very different, much simpler, pattern with two major haplotypes surrounded by a crown of derived less frequent haplotypes and singletons (Fig. 2B). Anthometrina adriani is only found on the high-Antarctic shelf (Eléaume *et al.* Chapter 5.25, this volume). This less complex phylogeographic pattern is in agreement with the interpretation that this species is well mixed today or survived the LGM in fewer refugia on the High Antarctic shelf.

Isometra graminea, Notocrinus virilis, and Ptilocrinus amezianeae exhibit clear geographical structure. Isometra graminea is divided in three distant haplogroups (Fig. 2D), one restricted to the Ross Sea, a second to the Dumont d'Urville Sea, and the third to the West Antarctic Peninsula (Marguerite Bay). Notocrinus virilis also is divided into five distant haplogroups (Fig. 2C): two very distinct from each other restricted to the Eastern Weddell Sea, another two very distinct restricted to the Scotia Arc, and one restricted to a larger area comprising the Ross, Dumont d'Urville and Davis seas. The haplogroups containing more than one haplotype consist of one or two central haplotypes surrounded by a small number of singletons. Ptilocrinus amezianeae displays a dumbbell-shape topology with two haplogroups separated by nine unsampled haplotypes. One haplogroup is restricted to the Kerguelen Plateau and the second to the Admiralty and Scott Island Seamounts and Scotia Arc (Fig. 2E). Populations in the first two areas display a star-like topology, with one central and common (ancestral) haplotype surrounded by a crown of derived singletons. The recent discovery of *P. amezianeae* on the Kerguelen Plateau and Scotia Arc, in addition to the population known from the Admiralty and Scott Island Seamounts (Bowden *et al.* 2011; Eléaume *et al.* 2011), seems to suggest that this species may be well-distributed in the Southern Ocean and that the gap observed between the two haplogroups may be due to a sampling bias.

A fourth type of haplotype network pattern is the bush-like topology displayed by *Feracrinus heinzelleri* (Fig. 2F) which is a deep-sea, more-or-less eurybathic species. The genetic pattern observed could be a signature of a slope-dwelling species that may have found refuge during glacial periods in the deep basins around the continental shelf.

Demography - Based on the number of samples available, a total of seven species and up to ten populations within a species were analysed (Table 1). Most of the populations display medium to high haplotype diversity and medium to high nucleotidic diversity, indicating an overall rather high withinand among-sequences diversity. Within P. kerguelensis, phylogroups A, C and D are composed of generally highly diverse populations showing signatures of a bottleneck (or genetic sweep) followed by population expansion. A reduced gene flow between several populations also characterises these phylogroups. Other phylogroups are less diverse and show no sign of expansion, and some level of connectivity between populations is suspected (Fig. 7 in Hemery et al. 2012). Within each phylogroup, the populations seem to have either been variously impacted or reacted differently to past glaciations. The DS, RS, SA, and WAP populations of F. mawsoni, and DDU and RS populations of N. virilis, display a low haplotype diversity and low nucleotidic diversity suggesting a recent bottleneck or genetic sweep. In addition, significant negative Fu's Fs results support the idea that Anthometrina adriani DDU and RS populations. and Florometra mawsoni DDU, DS, KP, RS and SA populations have undergone a recent expansion, purifying selection or selective sweep. If the recent expansion hypothesis is confirmed, these results would suggest that DS, RS and WAP populations may be recovering from benthic eradication due to one of many glaciation events, as previously observed for P. kerguelensis (Hemery et al. 2012). Other populations seem to have recovered from more diverse and less impacted populations. Anthometrina adriani DS and EWS populations appear stable with a smaller number of singleton haplotypes, suggesting that these populations have been less impacted by, for example, past glaciation

events. *Feracrinus heinzelleri* displays stable population structure and no sign of population decrease or expansion, a result congruent with populations not impacted by drastic events, and capable of surviving glaciation periods on the slopes or deeper environnements.



Figure 2 Haplotype networks for seven of the most common species of crinoids in the Southern Ocean. Haplotypes are derived from COI sequences. The genealogical relationships among haplotypes were estimated using the median-joining algorithm, allowing for the definition of clades (based on divergence up to 0.5%), and using statistical parsimony. A - Promachocrinus kerguelensis and Florometra mawsoni networks are shown as connected because F. mawsoni has been shown to be closely related to P. kerguelensis and may be considered another lineage within the P. kerguelensis complex. All lineages are represented and indicated with an uppercase letter from A to F. B - Anthometrina adriani network showing two major haplotypes surrounded by secondary less frequent haplotypes. C - Notocrinus virilis network showing the relationship between haplogroups A and B. Haplogroup A displays one major haplotype in the Eastern Weddell Sea and 3 secondary ones found in the Eastern Weddell Sea and in the Scotia Arc. Haplogroup B is more diverse and distributed in both East and West Antarctica. These patterns are not typical of brooders and suggest some dispersal capabilities for this species. D – Isometra graminea network showing strict geographical segrega-tion of all three haplogroups. This pattern is typical of a non-dispersive species, i.e., a brooder. E - Ptilocrinus amezianeae network showing two major haplotypes apparently geographically segregated but with gene exchange between the Scotia Arc and the Admiralty and Scott Island Seamounts some 5000 km away. F - Feracrinus heinzelleri network showing no particular spatial patterning. Pk = Promachocrinus kerguelensis; Fm = Florometra mawsoni; Nv-A = Notocrinus virilis haplogroup A; Nv-A = N. virilis haplogroup B. All circles reprsent a unique COI sequence (haplotype), and their diameter is proportionnal to the number of specimens sharing this haplotype. Numbers in red indicate the number of unsampled haplotypes between two closely related haplotypes

#### 5. Geographic distributions of haplogroups

The following maps show the distribution and haplotype composition of populations of *Promachocrinus kerguelensis* phylogroups A to F (Maps 1 to 7), *Florometra mawsoni* (Map 8), *Anthometrina adriani* (Map 9), *Isometra graminea* (Map 10), *Notocrinus virilis* haplogroups A and B (Maps 11, 12), *Ptilocrinus amezianeae* (Map 13) and *Feracrinus heinzelleri* (Map 14). On each map, each color represents a different haplotype and the pie size is proportionnal to the total number of specimens analysed per population.

#### 6. Conclusion

Sample bias is only to be expected in the Southern Ocean, because of the logistic challenges that remoteness, ice and cold confer. However, we have tried to cover, as best as possible, the distributional range and habitat of selected crinoid species. Still, deep-sea habitats are obviously undersampled, as well as a large area between the Eastern Weddell Sea and the Davis Sea, between the Ross Sea and the West Antarctic Peninsula, and the southern part of the Atlantic sector (i.e., Burdwood Bank, Bouvet Island). Unsampled haplotypes (see Fig. 2) from these regions, if added to our dataset, might change some of the conclusions drawn here.

Brooders and broadcast spawners showed strongly contrasting population structures. As expected, brooders showed high levels of geographic structure, whereas broadcast spawners did not. This is probably due to their contrasting life history traits. Planktonic larvae are usually produced in larger numbers and are able to cover longer distances. Among the brooders, *I. graminea* haplogroups have never been found sympatrically on the Antarctic shelf. This species seems to lack the ability to disperse and rapidly colonise new habitats. By contrast, *Notocrinus virilis*, another brooder, seems to be capable of larval dispersion (S. Schiaparelli, pers. com.). Among boadcast spawners, all phylogroups are found sympatrically except within *P. kerguelensis* at Bouvet Island and on the Kerguelen Plateau, where only the clade A is present.

Demographic approaches indicate that populations display contrasting histories, depending on the species or phylogroup. Population expansion after a bottleneck or genetic sweep can be detected, but not in all populations under study. Population expansion after a bottleneck is an expected signature after a drastic demographic event such as a large-scale benthic eradication resulting from grounded ice expansion during a glacial period. The "biodiversity pump" scenario, which states that population fragmentation and isolation in refugia during glacial periods may have induced allopatric speciation in Antarctic taxa, is therefore congruent with these results.

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Phylogeography Crinoida Maps 1-4 Map 1 Promachocrinus kerguelensis clade A is well represented on the Kerguelen Plateau, East Antarctic Ross Sea and Dumont d'Urville Sea shelves, tip of the Peninsula and in the sub-Antarctic Islands but the number of haplotypes is higher on the east Antarctic Ross Sea and Dumont d'Urville Sea shelves. In contrast, this clade is rare on the shelves extending from the Weddell Sea to the Davis Sea. The haplotype composition of each population may be indicative of a past refuge on the Kerguelen Plateau, and subsequent recolonisation from different source haplotype in the East and West Antarctic. Map 2 Promachocrinus kerguelensis clade B is well represented on the East Antarctic Ross Sea and Dumont d'Urville Sea shelves. It is rare from other part of the Antarctic shelf, and absent from all sub-Antarctic islands. Map 3 Promachocrinus kerguelensis clade C is well represented on the high-Antarctic shelves. It displays a higher number of haplotypes in the West Antarctic: Weddell Sea, Peninsula, South Orkney and South Shetland Islands. It is less diverse on the East Antarctic Dumont d'Urville Sea and Ross Sea shelves. It is completely absent from other sub-Antarctic Islands. Promachocrinus kerguelensis clade D is well represented on the East Antarctic Dumont d'Urville Sea and Ross Sea shelves. It is completely absent from the sub-Antarctic Islands. Proportion of haplotypes in the East Antarctic Davis Sea and Dumont d'Urville Sea and Dumont d'Urvil



Phylogeography Crinoida Maps 5-8 Map 5 Promachocrinus kerguelensis clade E1 is rare overall with very low haplotype diversity. It is better represented with a higher diversity on the West Antarctic South Orkney shelf, and seems absent from the sub-Antarctic islands. Map 6 Promachocrinus kerguelensis clade E2, like E1, is also very rare with low haplotype diversity. It is only found on the high-Antarctic shelf and is absent from all sub-Antarctic islands. Map 7 Promachocrinus kerguelensis clade F is also rather rare and shows very reduced haplotype diversity. It is found on the high-Antarctic shelf and from the West Antarctic South Orkney and South Sandwich Islands. It is absent from other sub-Antarctic islands. Map 8 *Florometra mawsoni* is closely related to *P. kerguelensis*. It is well represented on the East Antarctic shelf. It is found in lower abundance in the Peninsula, Scotia Arc, and Burdwood Bank. It seems to be absent from other sub-Antarctic localities except the Kerguelen Plateau. The Kerguelen, Dumont d'Urville Sea and Weddell Sea populations are composed of high numbers of different haplotypes and show high levels of diversity. As in the case of *P. kerguelensis* clade A, *F. mawsoni* may have found refuge on the Kerguelen Plateau during past glaciation, and recolonised high-Antarctic areas from there.





**Phylogeography Crinoida Maps 9-12** Map 9 *Anthometrina adriani* is confined to the high-Antarctic shelf and is absent from the South Shetland Islands and the Scotia Arc as well as from the sub-Antarctic islands. The number of haplotypes is higher on the East Antarctic shelf. The number of specimens analysed is much smaller in the Antarctic Peninsula area. This is probably not a sampling bias; *A. adriani* populations from the Peninsula consist of fewer individuals (M. Eléaume, personal observations). Map 10 *Isometra graminea* is restricted to the high-Antarctic. The higher diversity detected on the East Antarctic Dumont d'Urville Sea shelf is probably linked to the greater number of specimens collected there. Each locality displays a set of endemic haplotypes, suggesting absence of gene flow in the past. However, the current situation remains unknown. It is possible that each of these populations represent a separate species. Map 11 *Notocrinus virilis* haplogroup A is restricted to the West Antarctic Veddell Sea and Scotia Arc. It shows a high level of geographic segregation with two populations displaying no haplotype in common. It is possible that the Weddell Sea and the South Shetland Islands populations may represent two distinct species. Map 12 *Notocrinus virilis* haplogroup B shows a high level of geographic segregation. The West and East Antarctic populations have no haplotypes in common. The East Antarctic populations display a reduced number of haplotypes in common, suggesting some degree of connectivity in the past. The current situation is unknown. It is, however, possible that the West and East Antarctic populations may represent two or three separate species. Colors and pies as in Map 1.



Phylogeography Crinoida Maps 13-14 Map 13 The Kerguelen and Admiralty and Scott Island Seamounts Ptilocrinus amezianeae populations display no haplotype in common. The Ross Sea and South Sandwich populations have one haplotype in common, suggesting some degree of gene flow in the past. This pattern is suggestive of two disconnected populations that may have diverged into separate species. However, a detailed morphological analysis (Eléaume et al. 2011), and the fact that only a portion of the potential habitat of this species has been explored (sample bias), argue in favor of a single, undersampled species possibly a ring species. Map 14 Feracrinus heinzelleri shows a greater number of haplotypes in the Ross Sea population. This is probably due to the much greater number of specimens analysed in this population. New samples from the Kerguelen Plateau indicate that this species is also well represented there

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#### THE BIOGEOGRAPHIC ATLAS OF THE SOUTHERN OCEAN

Biogeographic information is of fundamental importance for discovering marine biodiversity hotspots, detecting and understanding impacts of environmental changes, predicting future distributions, monitoring biodiversity, or supporting conservation and sustainable management strategies The recent extensive exploration and assessment of biodiversity by the Census of Antarctic Marine Life (CAML), and the intense compilation and validation efforts of Southern Ocean biogeographic data by the SCAR Marine Biodiversity Information Network (SCAR-MarBIN / OBIS) provided a unique opportunity to assess and synthesise the current knowledge on Southern Ocean biogeography

The scope of the Biogeographic Atlas of the Southern Ocean is to present a concise synopsis of the present state of knowledge of the distributional patterns of the major benthic and pelagic taxa and of the key communities, in the light of biotic and abiotic factors operating within an evolutionary framework. Each chapter has been written by the most pertinent experts in their field, relying on vastly improved occurrence datasets from recent decades, as well as on new insights provided by molecular and phylogeographic approaches, and new methods of analysis, visualisation, modelling and prediction of biogeographic distributions. A dynamic online version of the Biogeographic Atlas will be hosted on www.biodiversity.aq.

#### The Census of Antarctic Marine Life (CAML)

CAML (www.caml.aq) was a 5-year project that aimed at assessing the nature, distribution and abundance of all living organisms of the Southern Ocean. In this time of environmental change, CAML provided a comprehensive baseline information on the Antarctic marine biodiversity as a sound benchmark against which future change can reliably be assessed. CAML was initiated in 2005 as the regional Antarctic project of the worldwide programme Census of Marine Life (2000-2010) and was the most important biology project of the International Polar Year 2007-2009.

The SCAR Marine Biodiversity Information Network (SCAR-MarBIN) In close connection with CAML, SCAR-MarBIN (www.scarmarbin.be, integrated into www.biodiversity.aq) compiled and managed the historic, current and new information (i.a. generated by CAML) on Antarctic marine biodiversity by establishing and supporting a distributed system of interoperable databases, forming the Antarctic regional node of the Ocean Biogeographic Information System (OBIS, www.iobis.org), under the aegis of SCAR (Scientific Committee on Antarctic Research, www.scar.org). SCAR-MarBIN established a comprehensive register of Antarctic marine species and, with biodiversity.aq provided free access to more than 2.9 million Antarctic georeferenced biodiversity data, which allowed more than 60 million downloads.

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AnT-ERA)

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