

# Genetic analysis on stock structure of fin whales in the Antarctic based on mitochondrial and microsatellite DNA

Mutsuo Goto, Naohisa Kanda and Luis A. Pastene

*Institute of Cetacean Research, 4-5 Toyomi-cho, Chuo-ku, Tokyo 104-0055, Japan*

Contact e-mail: [goto@cetacean.jp](mailto:goto@cetacean.jp)

## ABSTRACT

Genetic samples (catches and biopsies) of fin whales obtained by JARPA and JARPAII were analyzed with two genetic markers, mtDNA control region sequencing and microsatellite DNA, to investigate stock structure of this species in the Antarctic feeding grounds. Genetic samples were available from Areas IIIE (n=6), IV (n=23), V (n=24) and VIW (n=2). No statistical significant difference in mtDNA haplotype frequencies was found between Areas IIIE+IV and Areas V+VIW. However, because the haplotype distributions were quite different among the areas, statistical comparison was made by re-grouping the haplotypes into two categories: those specific to each of the areas and those shared by more than one area. Results of the heterogeneity test showed statistically significant differences among whales in Areas IIIE, IV and V. The microsatellite analysis showed statistically significant differences in allele frequencies of 16 loci among whales from Areas IV and V, but not IIIE and IV, as well as IIIE and V, most likely due to the small sample size of Area IIIE. Results of the genetic analyses therefore suggested the possibility of genetic structuring of fin whales in the JARPAII research area, which should be further explored with the analyses of a large number of samples in the future.

**KEYWORDS:** ANTARCTIC, GENETICS, FIN WHALE, SCIENTIFIC PERMITS

## INTRODUCTION

Little information is currently available on the stock structure of fin whales in the Antarctic. As in the case of the blue whale, earlier mark-recapture analysis showed that most whales return to the same part of the Antarctic year after year (Brown, 1954). Subsequent mark-recapture studies conducted by Brown (1962) suggested that the six whaling areas are probably more valid for blue and humpback whales than for fin whales (see also Mackintosh, 1965). The past information suggested there was certain segregation of fin whales in the feeding ground between certain longitudes in four sectors which lie: South of the Atlantic Ocean, South of the Indian Ocean, South of Western South Pacific Ocean and South of Eastern South Pacific Ocean (Mackintosh, 1965). South of the Indian Ocean correspond approximately to JARPAII Areas IIIE and IV and South of Western South Pacific to JARPA and JARPAII Areas VIW and V. It is important to investigate whether such geographical segregation is supported by genetic differences.

Wada and Numachi (1991) conducted allozyme analysis using North Pacific, Spanish coastal and Antarctic fin whales. They showed significant allele frequency differences between Hemispheres. However they could not detect evidence of more than one stock within the Antarctic or within the North Pacific fin whales.

Only a single study based on JARPA biopsy samples and mtDNA has been conducted to examine genetic differences between the whales from IIIE+IV and VW (Pastene *et al.*, 2005). Although the authors found no evidence of the genetic differences between IIIE+IV and VW, the sample size was too small (8 and 15, respectively) to make a firm conclusion on stock structure of this species in the Antarctic.

Additional genetic samples were obtained during the JARPAII surveys. The analysis of those samples is important for testing the hypothesis of segregation in the feeding grounds of the IWC management areas. In this study we conducted mtDNA and microsatellite analyses using additional biopsy and research samples obtained by JARPA and JARPA II to investigate further the stock structure of fin whales in Areas

III, IV, V and VI.

## MATERIALS AND METHOD

### Samples

Genetic samples were available from fin whales caught by JARPAII surveys between 2005/06 and 2010/11 and from biopsies obtained from the sighting surveys of the JARPA and JARPAII, on an opportunistic basis. Table 1 and Fig. 1 show the number and geographical position of the genetic samples used in this study, by year and Area.

### Laboratory procedures

Total DNA was extracted from 0.05g of skin tissue using the protocol of Sambrook *et al.* (1989). Extracted DNA was stored in TE buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8.0).

#### MtDNA

The first half of the control region of the mitochondrial genome was amplified by using the polymerase chain reaction (PCR). For the amplification of approximately 500bp of the mtDNA control region, primers MT4 (Árnason *et al.*, 1993) and Dlp 5R (5'-CCA TCG AGA TGT CTT ATT TAA GGG GAA C-3'), were used. Reactions were carried out in 25  $\mu$ L volumes containing 10-100ng of DNA, 2.5 pmole of each primer, 0.5 units of Ex Taq DNA polymerase (Takara), 2mM of each dNTP, and 10x reaction buffer. After an initial denaturation step at 95°C for 5 minutes, a PCR amplification cycle of 30 seconds at 94°C, followed by 30 seconds at 50°C and 30 seconds at 72°C are repeated 30 times. The amplification is completed with a final extension step of 10 minutes at 72°C. Subsequent cycle sequencing reactions are performed with 100ng of products generated in the above PCR amplifications using the BigDye® Terminator Cycle Sequencing Kit (Applied Biosystems). The oligo-nucleotides used to prime the cycle sequencing reaction were the same as employed in the initial PCR amplification listed above. A total of 25 cycles for 10 seconds at 96°C, 20 seconds at 56°C and four minutes at 60°C are performed. The nucleotide sequence of each cycle sequencing reaction is determined using Applied Biosystems 3500 Genetic Analyzer (Life Technology) under standard conditions. Both strand samples are sequenced in their entirety for all samples.

#### Microsatellite

Genetic variation at microsatellite loci were analyzed using 16 sets of primers as follow: EV1, EV14, EV21, EV94, EV104 (Valsecchi and Amos, 1996), GT011 (Bérubé *et al.*, 1998), GT23, GT211, GT271, GT310, GT575 (Bérubé *et al.*, 2000), GATA28, GATA53, GATA98, GATA417, GGAA520 (Palsbøll *et al.*, 1997), and DlrFCB17 (Buchanan *et al.*, 1996). Primer sequences and PCR cycling profiles generally followed those of the original authors. PCR amplifications were performed in 15ml reaction mixtures containing 10-100ng of DNA, 5 pmole of each primer, 0.625 units of Ex Taq DNA polymerase (Takara Shuzo), and 2mM of each dNTP, and 10x reaction buffer containing 20mM MgCl<sub>2</sub> (Takara Shuzo). Amplified products with internal size standard (GENESCAN400HD, Applied Biosystems Japan) were run on a 6% polyacrylamide denaturing gel (Long Ranger) using BaseStation100 DNA fragment analyzer (Bio-Rad). Although alleles were visualized using Cartographer software specifically designed for the BaseStation, allelic sizes were determined manually in relation to the internal size standard and fin whale's DNA of known size that were rerun on each gel.

### Statistical analysis

#### MtDNA

Sequences were aligned initially using Sequence Navigator (Applied Biosystems, Inc). The aligned sequences were then corrected by eye for minor inconsistencies.

Haplotype and nucleotide diversity were calculated following Nei (1987). Conventional hypothesis testing procedure was conducted based on the randomized chi-square test of independence (Roff and Bentzen, 1989) to assess genetic differences among Areas. *F<sub>st</sub>* was calculated using the AMOVA (Excoffier *et al.*, 1992). Furthermore heterogeneity tests were conducted as described in Hudson *et al.* (1992), using the *H<sub>st</sub>* and the *K<sub>st</sub>\** statistics. The level of statistical significance was estimated from 10,000 Monte Carlo simulations as the proportion of simulations in which a similar or more extreme value of *H<sub>st</sub>* or *K<sub>st</sub>\** was observed.

#### Microsatellites

The number of alleles per locus and expected heterozygosity per locus was calculated using FSTAT 2.9.3 (Goudet, 1995). Statistical tests for the deviations from expected HardyWeinberg genotypic proportions

were conducted using GENEPOP 4.0 (Rousset, 2008).

Probability test (or Fisher's exact test) implemented in GENEPOP 4.0 (Rousset, 2008) was used to conduct the heterogeneity tests. When simultaneous multiple tests were conducted, correction for the multiple tests was performed (Rice, 1989). Statistical significance was also determined using the chi-square value obtained from summing the negative logarithm of p-values over the 16 microsatellite loci (Sokal & Rohlf 1995). A p-value smaller than 0.05 was used as a criterion to reject the null hypothesis of panmixia.

## RESULTS AND DISCUSSION

### MtDNA

#### *Diversity*

A segment of 479bp of the mtDNA control region was sequenced for all samples. A total of 32 polymorphic sites identified a total of 45 unique sequences (haplotypes) in the total sample of 55 fin whales (Table 2). Except for two transversion sites, all substitutions were transitions.

Level of mtDNA diversity based on nucleotide and nucleon diversity is shown in Table 3. Both indices showed a high and similar degree of mtDNA diversity through the Areas. Nucleotide diversity estimate for the total sample was 0.0121 (Table 3).

#### *Heterogeneity test*

No significant differences were found in the statistical comparison between Areas III+IV and Areas V+VI (Table 4). Values of  $F_{st}$ ,  $H_{st}$  and  $K_{st}^*$  were negative. It was possible that no evidence of the statistically significant difference was because most of the haplotypes were singletons in each area. Because the haplotype distributions were quite different among the areas, an additional statistical comparison was made by re-grouping haplotypes into two categories: those specific to each of the areas and those shared by more than one area (Table 5). In this case, statistically, highly significant differences were observed among the areas (Table 6).

### Microsatellite

#### *Diversity*

Table 7 shows the number of alleles, expected heterozygosity and test result for expected Hardy-Weinberg genotypic proportions at 16 microsatellite loci. Total number of alleles at the loci in all the samples combined ranged from six at EV104 to 25 at EV94 with an average of 13.7. Expected heterozygosity ranged from 0.487 at FB14 to 0.955 at EV94 with an average of 0.843. Three loci (GAT98, EV21, TAA31) after correction for the simultaneous multiple tests and overall value showed significant deviation from the expected Hardy-Weinberg genotypic proportions.

#### *Heterogeneity test*

Pair-wise comparisons among the areas showed the statistically significant differences between whales from Areas IV and V, but not between III+IV and V, as well as III+IV and VI. (Table 8).

### Stock structure

Both of the genetic markers detected the genetic heterogeneity between Areas IV (Indian Ocean) and V (Pacific Ocean). In contrast to the past genetic studies (Wada and Numachi, 1991; Pastene *et al.*, 2005), our genetic study raised the possibility of genetic structuring in the JARPAII research area. This corresponded to the past observation of the feeding segregation between South of the Indian Ocean (IV) and South of Western South Pacific Ocean (V) (Mackintosh, 1965).

The catch distribution of fin whales in the Antarctic based on the IWC catch database (1913-1973) is shown in Fig. 2. Fin whales were widely distributed from the middle latitudes to south of 60°S along all longitudes. They were concentrated in Area III (between 10°E-60°E), in Area IV (between 80°E-110°E), Area V (between 140°E-170°E) and Area VI (between 170°W-150°W). A low density zone was observed around 130°E in the JARPAII research areas. Therefore this distribution pattern also matched well with the results of our genetic analyses.

The mtDNA analysis also detected the difference in the haplotype distributions between Areas III+IV and V, although the microsatellite analysis failed to detect such differences most likely due to the small sample size of III+IV. Our study thus suggested the possibility of an additional structuring in the JARPAII research area. In Area III+IV, an additional stock may be distributed or multiple stocks may be mixed. However, it

was difficult for us at this time to distinguish between these two possibilities. A similar pattern of genetic structure was observed in humpback whales in the JARPAII area (Pastene *et al.*, 2013; Kanda *et al.*, 2014). Pastene *et al.* (2013), by analyzing mitochondrial DNA variations on humpback whales from both the feeding and breeding grounds, showed that IV from 80°E to 120°E was occupied by one stock (Western Australia stock) and V from 140°E to 160°E by another stock (Eastern Australia stock) and that the rest of the areas were mixing areas of the adjacent stocks.

Our study implies that effective management of fin whales in Antarctic should be conducted Area by Area although it is premature yet to finalize their stock structure. In order to better understand the stock structure of fin whales, further genetic analyses should be conducted in the future based on larger sample sizes.

#### ACKNOWLEDGEMENTS

We would like to thank the researchers and crew members of the JARPA and JARPAII surveys for their efforts in collecting the samples. Our gratitude goes to H. Oikawa and S. Azumi for their laboratory assistance.

#### REFERENCES

- Arnason, U., A. Gullberg, and B. Widegren. 1993. Cetacean mitochondrial DNA control region: sequences of all extant baleen whales and two sperm whale species. *J. Mol. Biol. Evol.* 70: 960-970.
- Bérubé, M., Aguilar, A., Dendanto, D., Larsen, F., Notarbartolo-di-Sciara, G., Sears, R., Sigurjónsson, J., Urban-Ramirez, J., and Palsbøll, P.J. 1998. Population genetic structure of North Atlantic, Mediterranean Sea and Sea of Cortez fin whales, *Balaenoptera physalus* (Linnaeus, 1758); analysis of mitochondrial and nuclear loci. *Mol. Ecol.* 7:585-599.
- Bérubé, M., Jørgensen, H., Mcewing, R., and Palsbøll, P.J. 2000. Polymorphic di-nucleotide microsatellite loci isolated from the humpback whale, *Megaptera novaeangliae*. *Mol. Ecol.* 9:2181-2183.
- Brown, S.G. 1954. Dispersal in blue and fin whales. *Discovery Reports*. Vol. XXVI:355-384.
- Brown, S.G. 1962. The movement of fin and blue whales within the Antarctic zone. *Discovery Reports*. Vol XXXIII:1-54.
- Buchanan, F.C., Friesen, M.K., Littlejohn, R.P., and Clayton, J.A. 1996. Microsatellites from beluga whale *Delphinapterus leucas*. *Mol. Ecol.* 5:571-575.
- Excoffier, L., Smouse, P.E. and Quatro, J.M. 1992. Analysis of molecular variance inferred from metric distances among DNA haplotypes: application to human mitochondrial DNA restriction data. *Genetics* 131:479-91.
- Goudet, J. 1995. FSTAT, version 1.2: a computer program to calculate F-statistics. *J. Hered.* 86:485-486.
- Hudson, R.R., Boos, D.D. and Kaplan, N.L. (1992). A statistical test for detecting geographic subdivision. *Mol Biol Evo* 9:138-151.
- Kanda, N., Goto, M. and Pastene, L.A. 2014. Stock structure of humpback whales in the Antarctic feeding grounds as revealed by microsatellite DNA data. Paper SC/F14/J31 presented to the JARPAII review meeting. February 2014. (unpublished). 5pp.
- Mackintosh, N.A. 1965. *The stocks of whales*. Fishing News (Books) Ltd., London. 232pp.
- Matsuoka, K., Hakamada, T., Kiwada, H., Murase, H. and Nishiwaki, S. 2006. Distributions and standardized abundance estimates for humpback, fin and blue whales in the Antarctic Areas III, IV, V and VIW (35°E -145°W), south of 60°S. Paper SC/D06/J7 submitted to the JARPA REVIEW meeting. December 2006. (unpublished) 37pp.
- Nei, M. 1987. *Molecular Evolutionary Genetics*. Columbia University Press, New York. x+512pp.
- Palsbøll, P.J., Bérubé, M., Larsen, A.H., and Jørgensen, H. 1997. Primers for the amplification of triand tetramer microsatellite loci in baleen whales. *Mol. Ecol.* 6:893-895.
- Pastene, L.A., Goto, M., Kanda, N. and Nishiwaki, S. 2005. Genetic analyses on stock identification in the Antarctic humpback and fin whales based on samples collected under the JARPA. Paper [JA/J05/JR16](#) submitted to the JARPA review meeting. January 2005. (unpublished) 13pp.
- Pastene, L.A., Kitakado, T., Goto, M., and Kanda, N. 2013. Mixing rates of humpback whales of Stocks D, E and F in the Antarctic feeding grounds based on mitochondrial DNA analyses. Paper SC/65a/SH13 presented to the IWC Scientific Committee, June 2013, Cheju Island, Korea (unpublished).
- Rice, W.R. 1989. Analyzing tables of statistical tests. *Evolution* 43:223-225.
- Roff, D.A. and Bentzen, P. 1989. The statistical analysis of mtDNA polymorphisms: chi-square and the problem of small samples. *Mol. Biol. Evol.* 6:539-45.
- Rousset, F. 2008. Genepop'007: a complete re-implementation of the genepop software for Windows and

- Linux. *Molec. Ecol. Resources* 8:103-106.
- Sambrook, J., Fritsch, E.F. and Maniatis, T. 1989. *Molecular Cloning: A Laboratory Manual. Second Edition*. Cold Spring Harbor Laboratory, New York.
- Sokal, R.R., and Rohlf, F.J. 1995. *Biometry: the Principles of Statistics in Biological Research*. Freeman and Company, New York.
- Valsecchi, E., and Amos, W. 1996. Microsatellite markers for the study of cetacean populations. *Mol. Ecol.* 5:151-156.
- Wada, S. and Numachi, K. 1991. Allozyme analyses of genetic differentiation among the populations and species of the *Balaenoptera*. *Rep. int. Whal. Commn* (special issue 13):125-54.

Table 1 Number of samples used in this studies by year on Antarctic fin whales collected during JARPA and JARPAII sueveys.

Areas	1999	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011	Total
IIIE					4							2		6
IV				4				19						23
V	3		6		6				6		1		2	24
VIW						2								2

Table 2. Variable sites defining 45 mtDNA haplotypes in the Antarctic fin whale. The column on the left are haplotype ID. The numbers above listed on the right side are the nucleotide positions of the polymorphic sites starting from the 5' end of the mtDNA control region. Haplotypes '2' through '45' are listed with reference to haplotype '1'. A dot indicates an identical nucleotide at the position relative to haplotype '1'. The left side of the table are the frequencies of the 45 haplotypes in the four Antarctic Areas.

Haplotype	Area IIIE	Area IV	Area V	Area VIW	10	11	20	30
1	1	0	0	0	1667788900	3790015555	6888890026	90
2	1	1	0	0	4082978748	4910144689	6347811245	80
3	1	0	0	0	GCAGTTTT	GCCCTAACG	CTTTATCAT	CA
4	1	0	0	0	.TG..C....	....C....	....C....	
5	1	0	0	0	.TG.....	....C....	.....G..	
6	1	0	0	0	..G.....	.....T.	.....C..	
7	0	1	0	0	..A.C....	.....	.....C....	
8	0	2	0	0	.G.....	....CG...	.C....C....	
9	0	1	0	0	....C....	....C....	.....G..	
10	0	1	0	0	.TG..C.CC.	....C....	..C...T..	
11	0	1	1	0	....	....	....C....	
12	0	1	0	0	..G..C....	.....	....C....	
13	0	1	0	0	..G.....	.....A	.....G.T..	
14	0	2	2	0	..G.....	.....	.....C....	
15	0	1	0	0	..G..C....	....C....	.....C....	
16	0	1	0	0	..G..C....	.TATC....	.C..C....	
17	0	1	0	0	..G.....	....C.G...	....CT...	
18	0	1	0	0	ATGT.....G	....C....	.....G	
19	0	1	0	0	.TG..C....	....C....	..C...CT...	
20	0	1	0	0	..G.....	A...C....	.....	
21	0	1	0	0	.TG.....	.....	.....C...G	
22	0	1	0	0	.TG.....	.....	.....C....	
23	0	1	0	0	..G.....	.....T.	.....	
24	0	1	1	0	.TG.....	.....	T....C...G	
25	0	1	0	0	.TG.....	....C....	.....CT...G	
26	0	1	1	0	..G.....	....TC....	.....G	
27	0	0	1	0	..G.....	....C.G..	.....G	
28	0	0	1	0	..G..C....	....C.G...	.....	
29	0	0	1	0	.TG.....C.	....C....	.....C...G	
30	0	0	1	0	.TG.....	.....	..CC..CT...	
31	0	0	2	0	..G.....	....CCG...	.....C....	
32	0	0	1	0	..G.....	.....	..CC....	
33	0	0	1	0	.TG..C....	....C....	.....CT...	
34	0	0	1	0	.TG..C....	.....	.....CT...	
35	0	0	1	0	..G..C....	....C....	..C...C....	
36	0	0	1	0	..G..C....	.....A	.....C....	
37	0	0	1	0	..G.....	.....	.....C....	
38	0	0	2	0	.TG..C....	....C....	.....CT...G	
39	0	0	1	0	..G.....	.....	.C.....T.	
40	0	0	1	0	..G.....	.....	.....G.T..	
41	0	0	1	0	....	....	.....T.	
42	0	0	1	0	.TG.....	.....	..C...C...G	
43	0	0	1	0	....C....	....C....	.....T.	
44	0	0	0	1	....C....	....C....	.....	
45	0	0	0	1	.TG.....G	....C...T.	.....G	
	6	23	24	2	2223222222	2222222222	2222222222	22

Table 3. Estimates of the nucleon and nucleotide diversities in the Antarctic fin whale.

	<i>H</i>	$\pi$	SE
III+IV	0.99261	0.01076	0.00086
V+VI	0.99077	0.01052	0.00049

Table 4. Results of the heterogeneity tests using randomized chi-square, Fst, Hst and Kst\* in pair-wise comparison.

	$\chi^2$	Fst	Hst	Kst*
P	0.413	0.524	0.574	0.935
value		-0.001	-0.0005	-0.0058

Table 5. Frequencies of area specific haplotypes and shared common haplotype groups in areas III, IV and V.

	III	IV	V
III specific	5	0	0
Share between III & IV	1	1	0
IV specific	0	17	0
Share between IV & V	0	5	5
V specific	0	0	19

Table 6. Statistical comparison among areas III, IV and V re-grouped in the Table 6.

	$\chi^2$	Fst	Value
III vs IV	<0.0001	<0.0001	0.599
III vs V	<0.0001	<0.0001	0.659
IV vs V	<0.0001	<0.0001	0.599

Table 7. The number of alleles (A), expected heterozygosity (He), and test results for expected Hardy-Weinberg genotypic proportions (HW) at 16 microsatellite loci.

	A	He	HW
EV37	17	0.913	n.s
EV1	24	0.941	n.s
GT310	14	0.921	n.s
GAT28	17	0.849	n.s
GT575	13	0.734	n.s
EV94	25	0.955	n.s
GT23	15	0.888	n.s
GAT98	9	0.775	sign.
EV104	6	0.639	n.s
GAT417	20	0.843	n.s
GT211	12	0.895	n.s
EV21	8	0.614	sign.
FB14	8	0.487	n.s
EV14	16	0.788	n.s
GT195	13	0.875	n.s
TAA31	8	0.512	sign.
All			High. Sign.

Table 8. Results of the heterogeneity test comparing allele frequencies of 16 microsatellite loci of fin whales, among Antarctic management Areas.

	P value
III E vs IV	0.6776
III E vs V	0.9997
IV vs V	0.0136

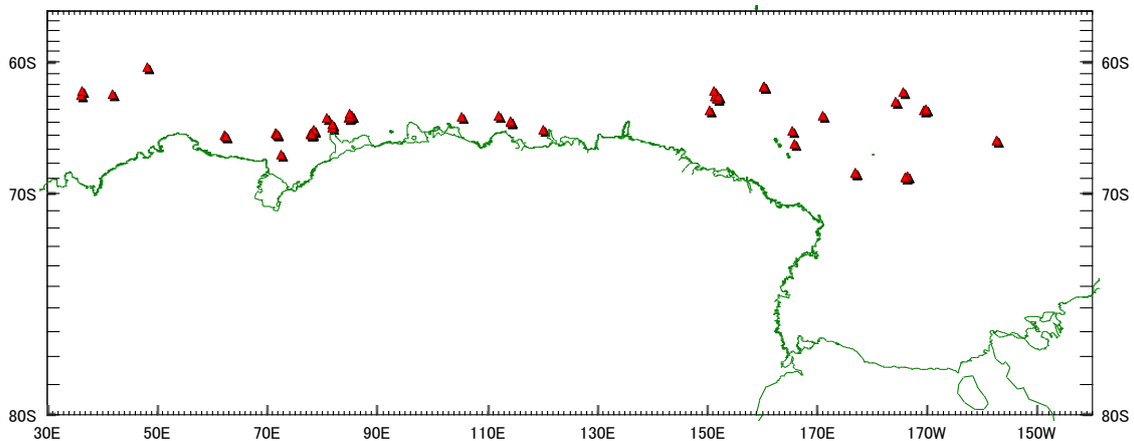


Figure 1. Geographic sampling localities of fin whales by catching and biopsy during JARPA and JARPAII.

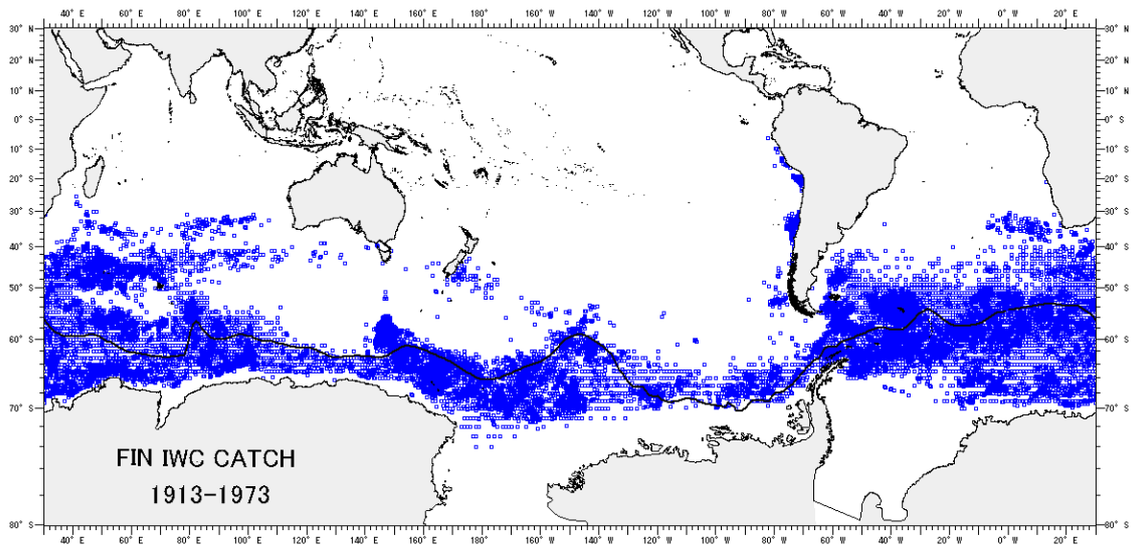


Fig.2. Distributions of fin whales from December to March (IWC catch database, noon positions (NP) of the days they were caught along the Southern Boundary of Antarctic Circumpolar Current) (after Matsuoka *et al.*, 2006)