## **BULLETIN No. 118**

# The Biology of the Ringed Seal (*Phoca hispida* Schreber) in the Eastern Canadian Arctic

### By

### I. A. McLAREN

Fisheries Research Board of Canada Arctic Unit, Montreal, P.Q.

# PUBLISHED BY THE FISHERIES RESEARCH BOARD OF CANADA UNDER THE CONTROL OF THE HONOURABLE THE MINISTER OF FISHERIES

**TTAWA, 1958** 

rice: \$1.00

## **BULLETIN No. 118**

# The Biology of the Ringed Seal (*Phoca hispida* Schreber) in the Eastern Canadian Arctic

By

I. A. McLAREN Fisheries Research Board of Canada Arctic Unit, Montreal, P.Q.

# PUBLISHED BY THE FISHERIES RESEARCH BOARD OF CANADA UNDER THE CONTROL OF THE HONOURABLE THE MINISTER OF FISHERIES

**OTTAWA, 1958** 

W. E. Ricker N. M. Carter *Editors* 

#### EDMOND CLOUTIER, C.M.G., O.A., D.S.P. QUEEN'S PRINTER AND CONTROLLER OF STATIONERY OTTAWA, 1958

Cat. No. Fs 94-118

BULLETINS OF THE FISHERIES RESEARCH BOARD OF CANADA are published from time to time to present popular and scientific information concerning fishes and some other aquatic animals; their environment and the biology of their stocks; means of capture; and the handling, processing and utilizing of fish and fishery products.

In addition, the Board publishes the following:

An ANNUAL REPORT of the work carried on under the direction of the Board.

The JOURNAL OF THE FISHERIES RESEARCH BOARD OF CANADA, containing the results of scientific investigations.

ATLANTIC PROGRESS REPORTS, consisting of brief articles on investigations at the Atlantic stations of the Board.

PACIFIC PROGRESS REPORTS, consisting of brief articles on investigations at the Pacific stations of the Board.

The price of this Bulletin is \$1.00 (Canadian funds, postpaid). Orders should be addressed to the Queen's Printer, Ottawa. Canada. Remittance made payable to the Receiver General of Canada should accompany the order.

All publications of the Fisheries Research Board of Canada still in print are available for purchase from the Queen's Printer. Bulletin No. 110 is an index and list of publications of the Board to the end of 1954 and is priced at 75 cents per copy postpaid.

For a listing of recent issues of the above publications see inside of back cover.

### "CALANUS" SERIES NO. 12

### CONTENTS

	Page
Abstract	vii
Introduction	1
Methods and Materials	3
AGE DETERMINATION Ageing by dentinal annuli Other methods of ageing Factors affecting the formation of dentinal annuli and claw bands	5 8 11
GROWTH AND MORPHOMETRY	13
FEEDING Food preferences Patterns of feeding The fasting season Feeding and seal condition	16 17 21 20
The Moult	26
REPRODUCTION Ages at sexual maturity The breeding season and the annual sexual cycle Implantation and development of the embryo Birth and care of the young The sex ratio	28 38 52 55 59
Local Movements, Distributions, and Factors of Abundance Reproductive ecology Movements Population samples Distribution factors in the eastern Canadian arctic	60 70 75 77
Summary	79
Acknowledgments	82
References	83
Appendix	86

#### ABSTRACT

The ringed seal (*Phoca hispida* Schreber) was studied in the eastern Canadian arctic, particularly in southwest Baffin Island and northern Foxe Basin.

Canine teeth, which deposit incompletely calcified dentine in the spring season of low nutrition, are shown to be reliable indicators of age. The claws are useful for the first ten years. The seal grows from 65 cm. (2.1 feet) to a final adult length of about 135 cm. (4.4 feet). At least 72 food species were identified. Mysis oculata, decapods, and Boreogadus saida were most commonly taken in inshore waters, while most offshore seals feed on plankton, particularly Themisto libellula in southwest Baffin Island. Food does not limit the range or abundance of this seal. The basking and fasting season, in June and early July, results in much blubber loss, which is regained by summer feeding. Late departure of ice, giving longer opportunity for basking, may increase blubber loss. The moult occurs mainly in June. The sex ratio is about 53.5 males to 46.5 females. The male matures at about seven years, and the first ovulation of the female, which may be outside the breeding season and not result in pregnancy, usually occurs in the seventh year. The males are in rut from March to mid-May and the female ovulates during lactation, shortly after parturition. The blastocyst implants after about 3<sup>1</sup>/<sub>2</sub> months delay, in early August, and the corpus luteum changes its structure and increases in size at this time. The pup is produced in one of two basic types of birth-lair on the fast ice, generally in early April, and may be attended until the ice breaks up. Coastal configuration and latitude govern the amount of fast ice, its stability, and its snow cover, all of which determine the suitability of an area for pupping. More starvelings and small pups are produced on simple coasts, and the adult population of complex coasts is older and the seals are larger in size. Seals of higher latitudes are also larger. There is an annual, small-scale migration, and a dispersal of young seals from productive complex coasts to simple coasts. Ice conditions govern the distribution and abundance of this seal throughout the eastern Canadian arctic.

#### INTRODUCTION

The ringed seal, *Phoca hispida* Schreber, is a widely ranging hair seal inhabiting arctic and subarctic waters (as defined by Dunbar, 1951) of the entire northern In North American waters it is found from the Gulf of St. Lawrence hemisphere. (rarely, Fisher, 1950) across the Canadian arctic to the Bering Sea. No overall biological study has been made of this seal in North America. Much of our knowledge comes from Eskimo observations and much that is presumed from these sources has not been checked by careful observation and collection. Allen, in his monograph on the Pinnipedia (1880), offers the best early summary of the biology of this animal. Dunbar (1949) brings in more recent material and reviews the knowledge of the species to that date. Chapsky (1940) apparently gives a thorough review of the species in the Siberian arctic, but this paper has not been available to the author. Particular studies have included those of Sleptsov (1943) on reproduction, Dunbar (1941) and Pikharev (1947) on feeding in the Canadian and Siberian arctic respectively, and Anderson (1942) on the taxonomy of the Canadian arctic groups. Manning (1944) describes well the methods and implements used by the Eskimos in hunting this animal. Other published material consists largely of incidental observations made on several expeditions to Greenland and the Canadian arctic.

The ringed seal is exploited by most of the Eskimo groups inhabiting the Canadian arctic. In some areas it contributes much of the food, clothing and fuel to the Eskimo welfare and is, in fact, the backbone of the native economy. In recent years some concern has been felt over the utilization of arctic sea mammal stocks. The problem, in the case of the ringed seal, has two main aspects:

1. Primitively, the Eskimo filled many of his needs with this animal; today, his needs have increased through the development of a trapping economy. Also, the seal itself has become an item of trade (for example, 2,506 skins were turned in to the Hudson's Bay Company at Cape Dorset, Baffin Island, in 1953-1954). It is not known whether the hunting pressure in such areas constitutes a danger to the maintenance of the seal stocks.

2. In some areas, local and seasonal scarcities result in depression of the native economy.

This paper deals with some basic features of the biology of the ringed seal in the eastern Canadian arctic. Economic problems are not treated directly here, but it is hoped that some of the biological findings, particularly the factors of distribution, will be of assistance in the management of the species as a valuable resource in the Canadian north. The development of an accurate ageing technique has stimulated research on seals in recent years. This technique has not hitherto been applied to the ringed seal and has been of great value in this study. The application of the ageing method to the study of reproduction and patterns of distribution constitutes the core of the work. Two phases of the study, parasitology and taxonomy, are not included in this paper and will be published at a future date.

#### METHODS AND MATERIALS

The study of the ringed seal was initiated with the analysis of stomach contents of seals from Ungava Bay, collected from 1947 to 1950. In the summer of 1953, the Fisheries Research Board ketch, M/V Calanus, worked near Cape Dorset, southwest Baffin Island, where collections from 44 ringed seals were made; these served as the basis for an overall biological study. The success of the 1953 operations suggested that southwest Baffin Island would be a suitable area for the study of the ringed seal in 1954. The author arrived at Cape Dorset, by chartered aircraft, on March 2 and departed, by ship, on October 2, 1954. It soon became apparent that many features of ringed seal biology revolve around local and seasonal distributions and movements, and that populations differ in their make-up throughout southwest Baffin Island. Thus, extensive travelling was required in the area between Cape Dorset and Markham Bay, about 135 miles to the east. The author was completely dependent on the Eskimos for transportation and securing seals and about two-thirds of the season was spent travelling and living with these people, the remainder of the time being spent in working out of the trading post at Cape Dorset. The bulk of the material in this paper is the result of collections made during that seven-month stay on southwest Baffin Island. Substantial collections were also made by the author between mid-September and mid-November from the *Calanus* in northwest Foxe Basin, in 1955. Very limited material was collected between 1951 and 1955, from Frobisher Bay, Cumberland Sound, and Churchill, Manitoba. Valuable collections have been made available through the efforts of other individuals in the pursuit of other arctic field-problems. Seals from Southampton Island, northern Baffin Island and northern Foxe Basin were obtained in some numbers, and smaller collections were offered from Ellesmere and Prince Patrick Islands, southwest Baffin Island, and Nettilling Lake on Baffin Island.

The lower jaws of all seals examined were collected for the purpose of ageing and preserved dry or rotted; whole skulls were taken when possible. The extracted canine teeth were placed in glycerine and water, with thymol as a preservative, and were sectioned and ground according to the techniques described by Fisher and Mackenzie (1954). The teeth sections were examined, in reflected and transmitted light, under a binocular microscope. In many cases an optical micrometer was used to measure tooth thickness, size of pulp cavity and thickness of dentinal bands in section. One or two claws were removed from many individuals and generally preserved rotten.

Stomach contents were examined in the field or collected for laboratory analysis. Those collected were identified to species, where possible. Organisms were counted, except where the stomach contents consisted of large numbers of small crustacea; in such cases the numbers were estimated by volumetric subsampling. The dominant food (in bulk) was estimated by eye, or, in doubtful cases, by displacement methods.

The reproductive organs of males and females were examined or collected; those of immature seals were not always examined. Measurements were generally taken of the fresh tracts and glands in the field. Portions of testes and epididymides, chiefly those of adults, were preserved in Bouin's fixative or 10%formalin, sectioned, and stained with haematoxylin and eosin or Mallory's stain. Ovaries were sectioned by hand, with a razor blade, into slices one to three millimetres in thickness. These sections were examined for corpora lutea, corpora albicantia and extent of follicular development, and the diameters of these were measured. Selected corpora lutea were sectioned, and stained with haematoxylin and eosin, or azocarmine, phloxine and fast-green. Embryos were preserved when found. Attempts were made, without success, to find unimplanted blastocysts in the uteri. Bacula, or penis bones, were collected from the males and preserved dry or rotted. After a superficial cleaning, these were boiled in a dilute solution of ammonium hydroxide, then weighed and measured. The cleaning was not rigidly standard, but it is felt that the small variation in residual fat content has little effect on the weights of bacula.

Notes were kept on the moult and other individual characteristics of the specimens.

Field numbers were given to all specimens. In this paper, reference to various years and localities is as follows: "S-" was used in southwest Baffin Island in 1953; "SD-" was used in the same area in 1954; "S-55-" applies to specimens from northwest Foxe Basin in 1955; "S-56-" designates northern Foxe Basin seals in 1956; "A-" is for Southampton Island seals.

#### AGE DETERMINATION

#### Ageing by Dentinal Annuli

The technique of ageing by dentinal annuli has been thoroughly discussed by one of the discoverers of the method (Laws, 1953a) and by Fisher (1954a). The work of Fisher, on the harp seal (*Phoca groenlandica*), is more applicable to studies of the ringed seal, since the construction and nature of the dentinal rings is similar in these two species. The validity of the method, the physiological basis of dentine deposition, and much background information has been given by both authors. In this section it is proposed to deal chiefly with the evidence for the annual nature of annulus formation. The nutritional basis for differential dentine deposition will be discussed on page 11.

GENERAL APPEARANCE. A thin cross-section of a ringed seal canine tooth usually shows alternating bands of dense and thin dentine. Dense dentine is more opaque under transmitted light and is solidly constructed. Thin dentine may have one of two basic forms; it may be clear and continuous, transmitting more light than does the dense dentine, or it may be reflective, vacuolar and reticulated in structure. A typical series of bands, especially in the earlierformed, outermost annuli, consists of dense dentine followed by thin, followed by reticulated, followed again by dense dentine; sometimes the thin or the reticulated dentine is absent. The density of the dense band varies considerably in comparison with its associated thin band. The relative widths of the bands within different series may be variable, as small, misplaced thin bands occasionally break the continuity of a dense band. In spite of all this variation, the pattern is distinct; denser and more fully calcified dentine alternates with thinner and less well-formed dentine. The deposition rate of the dentine appears to be regular within one year, but decreases as the number of annuli increases. In verv old teeth, the annuli are very narrow. In the most typically annulated teeth the widths of adjacent annuli are similar. The adventitious nature of occasional variations and anomalies can be shown up by the use of an optical micrometer on the binocular microscope.

Both Laws (1953a) and Fisher (1954a) have offered convincing evidence that the deposition of the two major types of dentine occurs at different times of the year in the harp seal and the elephant seal (*Mirounga leonina*), and that this justifies the use of dentinal annuli in determining the age of these seals. The same reasoning applies to the ringed seal.

EVIDENCE FOR THE ANNUAL FORMATION OF DENTINAL ANNULI. Teeth from over 750 ringed seals have been sectioned for the study of ageing. From some of these teeth a histogram (Fig. 1) has been constructed showing, in two-week periods, the deposition of dense dentine at the pulp cavity. Many teeth obtained from the Eskimos were not dated, and not all the dated teeth could be entered into this histogram. Where the innermost annuli were smaller and crowded the detection of a dentine change would be difficult until some time after its occurrence. Thus, the histogram represents only those seals with fewer than four complete annuli. Probably there is a lag in the detection of dentinal change in all teeth, but this can be kept at a minimum by excluding older teeth. All suitable teeth collected from southwest Baffin Island in 1953 and 1954, as well as those collected from Foxe Basin in September and October, 1955, and February to April, 1956, are included in the histogram.

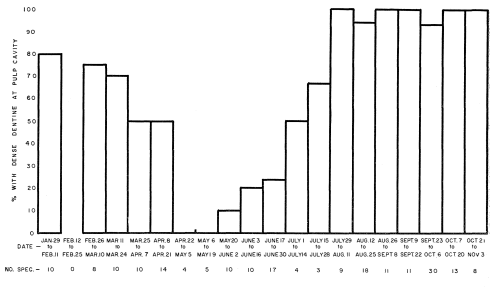


FIGURE 1. Seasonal changes in the deposition of dense dentine at the pulp cavity of the canine tooth of *Phoca hispida*, in 205 specimens.

Reference to Figure 1 shows that of ten teeth taken between January 29 and February 11, eight showed no interruption of the most recently formed dense dentine. One had a very narrow, thin band and another showed thin dentine in a fairly broad band next to the pulp cavity. The first tooth with reticulated, as opposed to merely thin, dentine was taken on March 11. Dense or thin dentine bordering the pulp cavity was found in seals taken up to April 19. Only thin or reticulated dentine was found in the next members of the series, killed on April 29, and in all seals taken after this date until late May. At this time one seal, which may have been anomalous, showed a broad, dense band bordering the pulp cavity with no evidence of recent thinning. The first teeth to show an obvious change from thin to denser dentine were two taken shortly before The last reticulated dentine was being formed on July 3, and theremid-June. after the proportion of teeth depositing dense dentine next to the pulp cavity increased. Only 3 out of 100 teeth taken between August 6 and October 30 had thin bordering dentine. Two seals taken in November were depositing dense dentine, although another with 8 dense bands on December 17 showed what appeared to be slight thinning of the dentine at this date. The majority, however, must continue to deposit dense dentine through the winter, judging from the suggestion of the first few histogram units of Figure 1. In those seals which showed the broadest bands of thin dentine in the spring, there was a tendency for the most recently formed dentine to be even thinner, or reticulated. The evidence for the deposition of the "poorest" dentine in the spring, from about mid-March to the beginning of July, is thus based on a series of teeth which covers this entire period. The annual nature of annulus formation may be taken as established and the structure of the dentine may be used with confidence to deduce the age of the seal. Thus, a seal taken in late summer with no annular pattern will be in its first year. Occasionally, however, a tenuous line of thin dentine, which may represent a line of birth or weaning, may be present. A seal taken in May with a single annulus of thinner or vacuolar dentine next to the pulp cavity, will have been born in March or April of the previous year. Figure 2 illustrates a variety of tooth sections, showing the seasonal changes as elaborated above. Some seals could not be aged accurately by this method after the tenth year and few could be aged after 20 years of age. This is due to the decrease in

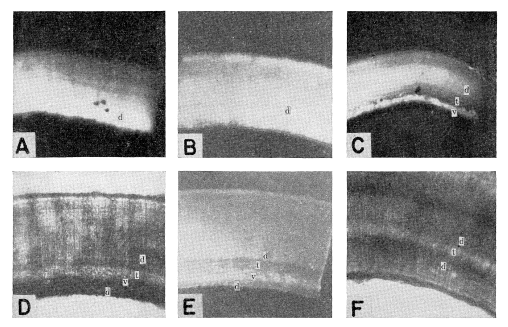


FIGURE 2. Tooth sections of young seals in various seasons. On the sections: d = dense dentine, t = thin dentine, and v = vacuolar or reticulated dentine.

- A. S-50, August 31, 1953. Reflected light. This tooth, dense to the pulp cavity, is taken from a seal of the year.
- B. S-56-12, February 11, 1956. Reflected light. A seal of the previous year. No thinning of the dentine is apparent. The narrow dark border is the face of the pulp cavity.
- C. SD-177, July 3, 1954. Reflected light. A moulting seal, born in the previous year. Note that the vacuolar dentine, on the inner border, is highly reflective.

- D. S-55-57, October 16, 1955. Transmitted light. One thin band, deposited in the previous spring, indicates that this animal is in its second year. The dentine bordering the pulp cavity is dense.
- E. S-55-57, October 16, 1955. Reflected light. The same section as in D. Note that the vacuolar dentine, which is not visible in the transmitted light of D, is here revealed as a reflective band (as in C.)
- F. S-56-15, February 12, 1956. Transmitted light. Dense dentine is being deposited at the pulp cavity, although it is not as dense as the dentine of the previous summer, which had been laid down immediately following the spring thin band. Second year.

dentine deposition rate, which results in crowding of annuli after many years of age. In very old seals the annular structure of the cementum served as an added check. Dense cementum bands show a one-to-one correlation with dense dentine bands in younger seals and do not decrease in size with age. By this technique one male, the oldest seal obtained, was found to be about 43 or more years in age.

#### Other Methods of Ageing

The Eskimos of southwest Baffin Island are able to distinguish what they suppose to be the younger age-classes of the ringed seal. The ringed seal (*netserk* s.l.) is divided into seven age-groups with the following Eskimo names (spelling based on Thibert, 1954);

Netsiak	the white-coat
Netsiavinerk	a seal of the year, before the first moult
Pivinerk	a seal of the second year, before the second moult
Saggalak	a seal of the third year, before the third moult
Togrotinerk	a seal of the fourth year, before the fourth moult
Netsilak	a seal of the fifth year, not always distinguishable

All other seals are considered to be adult (*netserk* s.s.); the adult female is often called *nunerk* and the adult male *tiggak*. Thus the natives consider that maturity is reached after the fourth or fifth year. The names actually refer to the length of the hair and its state of growing; the same names are also applied to stages of the moult and formation of the new coat. Thus a seal of the second year may be called *pivinerk* on the basis of age, but if taken shortly after the moult, it may also be referred to as saggalak (translated approximately as "new-grown hair"). Naturally, the determination of age by this method, if it is valid, is most simple after the hair is fully grown and before the moulting season. The author was unable on this basis to distinguish any of the young age-classes except the seal of the year, which, in softness and length of hair, is quite distinct. All twelve seals which were entered in the field-notes as young of the year were correctly aged. Three out of four *pivinerk* were in the second year and one was in the third year, but had not yet moulted. Four third-year seals, saggalak, were correctly aged by the natives; another seal, actually in its first moult, was put into this class by the Eskimos. The class torgrotinerk was less reliable. Four out of seven of this class were correctly aged; however, two in their fifth year should have been called netsilak and one, in the seventh year, was actually a netserk in the strict sense.

The Eskimos also make use of the claws in determining the age of the seal. They consider that a large number of alternating light and dark claw annuli in some way indicates an old seal. Plekhanov (1933) noted annular bands on the claws of the harp seal. He presumed that these represented annual increments of growth. Doutt (1942) found the same pattern on ringed seal claws.

The claws of 105 ringed seals were collected between April 28 and October 27, in order to assess their value in age determination. Dried or rotted claws are easily removed from the phalanges, but those preserved in formalin cannot easily be removed from the attached bone and flesh. The claws show alternating dark and light bands. The light bands are variable in width, but are usually marked in the middle by a narrow, encircling ridge. Claws showing few bands are capped at the tip by a section of more or less translucent structure. This translucent tip, when not worn off, is usually separated from the rest of the claw by a narrow constriction which may represent the limit of prenatal growth. The band at the growing base is very narrow and light in the early spring, broadening through the summer, and darkening before the autumn. The slight ridges in the light band appear to be laid down in the spring. These features are shown in a typical claw on Figure 3. The hypothetical age of a seal may be determined by assuming that the dark bands are the result of autumn and winter growth and that the light bands are formed in the spring and summer. The ages derived in this manner are checked against ages determined from dentinal annuli in Table I. The claws, which generally have a regular and clear pattern, are very useful in checking tooth-determined ages and for preliminary ageing in the field. Wear at the tip obliterates first the 'constriction of birth' and then the first year's growth. Ageing by claws is invalid beyond about the tenth year, judging from comparisons of age-estimations of seals of 9 to 17 years in the Table.

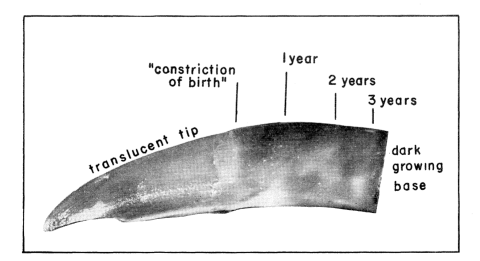


FIGURE 3. A typical claw, of a fourth-year seal killed on October 4, 1955.

51238 - 4 - 2

Deduced age	Number of specimens	Tooth age	Claw age
First year	4	1	1
Second year	11 2	2	2 3?
Third year	7 1 1	3 2? 3	3 3 2?
Fourth year	8 1 1	4 5? 4	4 4 5?
Fifth year	3 2	5 4?	5 5
Sixth year	9	6	6
Seventh year	2 1	7 6?	7 7
Eighth year	3 1	8 7?	8 8
Ninth year	2 1	9 9	9 7+
Tenth year	1 1	10 10	$10 \\ 8+$
Eleventh year	1 1	11 11	10+ 8+
Twelfth year	1	12	10+
Thirteenth year	1 1	13 13	10+ 11+
Fourteenth year	1 1	14 14	10+12+
Seventeenth year	1	17	10+

TABLE I. Comparison of ages of *Phoca hispida* deduced from teeth and claws. A query following an age indicates that this age is the less accurate of the two, due to anomalies or obscurities. A plus sign indicates that wearing of the claw has eliminated the first year's growth.

10

FACTORS AFFECTING THE FORMATION OF DENTINAL ANNULI AND CLAW BANDS

The general appearance, variability and times of deposition of the dentinal annuli have been discussed. Essentially, incompletely calcified dentine is laid down in the spring season, and solid, well-formed dentine is deposited in the summer and generally through the winter. The springtime is featured by a relaxation of feeding, followed by more intensive fasting as the season progresses. Fewer than 60% of seals were feeding in each of the two-week periods between April 22 and June 30, which corresponds to the period in Figure 1 when fewer than 25% of the seals were depositing dense dentine at the pulp cavity. Thus, there is a distinct correlation between seal feeding or fasting, and the deposition of dentine. Dense dentine is deposited from early summer until the following spring when feeding intensity is greater. Mellanby (1928) found that a diet deficient in cod-liver oil, but complete in other respects, would result in poor dentine formation in puppies, and other workers have found that the necessary factor is Vitamin D. Fisher (1954a) and Laws (1953a) discuss this effect of Vitamin D in greater detail. Certain individual seals show thinning of the dentine before the season of more intensive fast and some of these, as suggested above, may have been depositing such dentine through much of the winter. It is possible that a sufficient Vitamin D intake is assured only by voracious feeding, for even though a seal may be eating enough to maintain its condition of fatness, as seems to be usual in the winter, it may not be ingesting sufficient Vitamin D to ensure the formation of the densest form of dentine. Moreover, adults experience the metabolic strain of the breeding season during the spring and this may well effect the formation of dentine. The moult normally occurs from mid-May until mid-July, with a peak in the month of June, but the physiological cause of the process may well extend back to an earlier period and effect the balance of dentine deposi-Several animals were taken in August and September which were in very tion. late or prolonged moult. In these abnormal cases the seals were all feeding and dense dentine was being deposited at the pulp cavities of their teeth. Seals of the year, whose moult is not associated with a prolonged fast, do not deposit poorly formed dentine until the following spring. Thus the moulting process, possibly through further drain on the resources of Vitamin D, may be a supporting factor in thin dentine formation, but it cannot be the sole cause. Some teeth may show narrow bands of adventitious thin dentine within the dense dentine and it is probable that this a reflection of individual variations in feeding. Teeth from seals taken at Herschel Island, in the western arctic, are often difficult to interpret for this reason, and these anomalies may be due to abnormal feeding circumstances imposed by the geography and productivity of the area.

The pattern of annular deposition in the ringed seal is in accord with the findings of Fisher (1954a) on the harp seal, although Laws (1953a) has reached different conclusions from his work on the elephant seal. He considers that dense, fully formed dentine is laid down while the animal is fasting and points out that Vitamin D may be formed in optimum amounts by irradiation, when the

 $51238 - 4 - 2\frac{1}{2}$ 

animal is lying in the sun; the same factor may be available in sub-optimum amounts when the animal is feeding in the water. Fisher (1954a) discusses the opposition of the two views, but the present author can add nothing more in explanation of the two different conclusions.

The claws are less subject to individual variation than the teeth and their growth is probably more closely dependent on the moulting cycle, which is a more precisely annual phenomenon than the fasting season.

#### GROWTH AND MORPHOMETRY

The ringed seal is probably the smallest of the hair seals (Phocidae). The largest specimen taken for this study was a male, 156 cm. (5.1 feet) from nose to tip of tail and 222 lb. in weight; the longest female measured 152.5 cm. (5.0 feet). Soper (1928) recorded a male of slightly greater length, 162.5 cm. (5.3 feet), and Stefansson (1944) weighted one of 225 lbs. or more, including estimated blood loss. Five and one-half feet (168 cm.) and 250 lb. are probably close to the limits for the male of this species. There is great individual variation in adult size, which appears to be independent of age; the factors governing this variation will be discussed in a later section.

The relationship between standard length and age of the ringed seal is outlined in Figure 4. The symbols represent the mean of size and age of seals

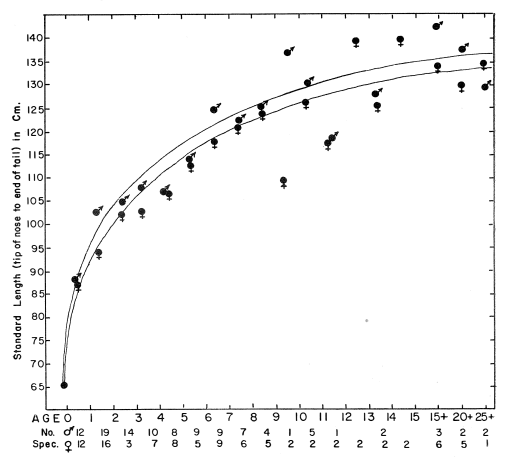


FIGURE 4. Growth of *Phoca hispida*. Each symbol represents the average, in length and age, of each age-group. The curves are fitted by eye with the aid of several straight lines of least squares.

from each age-group. The size at birth (65 cm. or 25.5 inches, derived from three full-term foetuses and two new-born pups and averaged for April 1) is not fixed with accuracy, but the first-year size-mean, derived from 24 specimens, is more firmly based. The high degree of individual variation and the different geographical origins of the specimens make it difficult to fit exact growth curves. The curves on Figure 4 must be considered to be approximate only, although they were fitted with the aid of several successive straight lines of least squares. During its life, the ringed seal appears to grow from approximately 65 cm. at birth to a final adult size of about 135 cm. (4.4 feet), the males averaging about 3 cm. longer than the females after maturity. The shape of the growth curve is similar in both sexes. The slight negative deviation of the third, fourth and fifth year size-means is considered to be a result of sampling error, and not related to the double sigmoid growth-curve found in the males of polygynous species (Laws, 1953b; Scheffer and Wilke, 1953).

Laws (1956a) has related growth rates of Pinnipedia and Cetacea to their ages at puberty. He found that about 86% of the final length is attained by the females at the age of sexual maturity. This rule holds well for the ringed seal which ovulates for the first time, on the average, when about 6 years old. At this time, from Figure 4, the female would measure about 117 cm., or 87.6% of the final adult length of 133.5 cm. at 25 + years.

Other body measurements are given in Appendix Tables I and II. Since field-measurement techniques and criteria differ, the measurements are defined here. The "same" measurements often produced quite different results when applied by more than one person to the same specimen. These Tables are made up of the author's data alone.

1. *Standard length* was measured in a straight line from tip of nose to the end of the tail, with the head and neck in a natural position.

2. Axillary girth was taken with the aid of a piece of string around the body, under the fore-flippers, at the level of the axillae.

3. *Maximum girth* was taken around the largest circumference of the chest or abdomen; this did not always exceed the axillary girth in very thin seals.

4. *Navel to anus* distance was measured from the centre of the umbilical scar to the anterior notch of the anus of the male and the vestibule of the female, along the body contour.

5. *Penis to anus* distance was measured from the centre of the penis orifice to the anterior notch of the anus, along the body contour.

6. *Foreflipper length* was measured in a straight line along the radial border of the forelimb, from the axilla to the tip of the longest digit (not claw). The forelimb was held flat and extended posteriorly along the side of the body.

7. Foreflipper width was taken from the tips of the first and last digits (not claws) of the stretched flipper, in a straight line.

8. *Hindflipper length* was measured from the base of the tail, along the inner edge of the extended flipper, to the level of the longest digit.

9. *Hindflipper width* was taken from the tips of the first and last digits with the flipper stretched to its fullest extent.

10. A probe was inserted into a small incision over the sternum. The *blubber thickness* was then marked on the probe from the bared bone of the sternum to the inner surface of the hide.

11. Weight was taken by a spring balance of 200 lb. capacity. No estimate was made for loss of blood.

A few data on tail length and distance between mammae are not included in the Table. The tail length appears to increase from about 6.5 cm. in 80 cm. seals, to about 9 cm. in 150 cm. seals( from line of regression of tail length to standard length of 29 specimens). The distance between mammae (from 22 specimens) appears to range from about 5 cm. in seals of 85 cm. in length, to about 9 cm. in seals about 140 cm. long.

For comparative purposes, some of the measurements have been reduced to percentages of the standard length and each such percentage plotted against its standard length. Straight lines of least squares have been fitted to the measurements expressed in this fashion, with the sexes separated. Appendix Table II gives the results of this analysis. The percentage of standard length of each measurement is given for seals of 60 cm. and 160 cm. in standard length and serves to fix the line of regression, and one standard error of the estimate is given to express the variation around this line. The seasonally variable girths, weight, and blubber thickness are excluded. Heterogonic growth is suggested in several of these comparisons. The limbs seem to grow proportionately less rapidly than the animal as a whole. The hind end of the body appears to grow only slightly more rapidly than the anterior end, as demonstrated by the comparison of the distance from navel to anus with standard length; thus there is little evidence for an antero-posterior growth gradient in this species.

A large series of skulls was collected from southwest Baffin Island. Morphometric studies of these have led to the consideration of taxonomy, and in particular to the status of the two subspecies described by Anderson (1942). Taxonomic matters will be left for a later paper and the morphometry of the skulls will be discussed then.

#### FEEDING

#### Food Preferences

Of the 216 seal stomachs examined in the field by the author, 36 were empty and 104 were sampled, in whole or in part, for laboratory analysis. The stomach contents of 48 seals, collected by other groups and individuals, were also available. At least 72 species were detected in these stomachs. Appendix Tables III to IX list the food species, along with data on their incidence, according to localities.

The ringed seal makes use of a wide variety of planktonic, nektonic, and benthonic food species. It is apparently widely adaptable in its feeding habits, but does not seem to eat many of the slow-moving, sedentary or burrowing animals (molluscs, echinoderms, tubeworms, large prawns, etc.) which are the chief components of the diet of the bearded seal (*Erignathus barbatus*). The few molluscs taken were small, with the shells present, and were probably more or less accidentally ingested. Among the decapod food species, only one crab (*Hyas coarctatus*) and none of the heavily armoured *Sclerocrangon boreas*<sup>2</sup> were detected. The smaller individuals of the natantic decapods were common and such food may be selected just off the bottom, not benthonically. The most abundant food elements were *Themisto libellula*, *Mysis oculata*, and the polar cod, *Boreogadus saida*.

The ringed seal probably feeds by individual selection of each food organism. It is known to feed only on the larger *Themisto* of a population having a bimodal size distribution (Dunbar, 1941). The possibility of a sifting or filtering method of feeding cannot be excluded, however. For example, two seals from Foxe Basin were feeding almost exclusively on the small (10-12 mm.) amphipod A pherusa glacialis, of which there were about 1,100 individuals in each stomach. These amphipods were noted in thick surface swarms at the time and almost certainly, due to their small size, could not have been taken individually. Small terrestial leaves and bird feathers were occasionally found in plankton-filled stomachs, especially in seals feeding inshore on mysids, indicating that selection was not too critical. Bearded seal stomachs which have been examined by the writer often contained stones, sand and mud. Such evidence of indiscriminate feeding was rarely found in the ringed seals which were feeding benthonically. Pebbles were found in the otherwise empty stomachs of three seals taken during the fasting season. These may have been swallowed purposely, perhaps to get rid of the nematode worms with which the stomachs were infested (see also Hahn, 1884).

Little evidence is avilable on the depth at which ringed seals may feed, for hydrographic charts of the areas studied are not generally available, or are not complete. One ringed seal taken near Churchill, Manitoba, in February, 1954,

<sup>&</sup>lt;sup>2</sup> Squires (1957) lists 39 *Sclerocrangon boreas* in ringed seal stomachs from Ungava Bay, and other differences are found between the decapod food species in Appendix Table III of the present paper and those listed by Squires from the same collection of seal stomachs. Mr. Squires has kindly informed the author that these discrepancies are due almost entirely to the mistaken inclusion in his list of the findings from a single bearded seal stomach.

was feeding on benthonic decapods in 14 to 15 fathoms (26–28 m.) of water and decapods were also eaten by seals in 15 to 20 fathoms (28–37 m.) of water in Foxe Basin in the autumn of 1955. Several seals killed in over 20 fathoms of water in the winter of 1956 were feeding benthonically and at least one of these, judging from the few soundings in this northern area of Foxe Basin, might have been in around 50 fathoms (90 m.) of water. The fur seal (*Callorhinus ursinus*) is known to dive to at least 40 fathoms (70 m.) (Scheffer, 1946) and the walrus (*Odobaenus rosmarus*) to 44 fathoms (80 m.) (Vibe, 1950). One undisturbed seal, observed in a fairly shallow bay of southwest Baffin Island, stayed down for an average period of 3 minutes and 13 seconds, and surfaced for an average of 1 minute and 29 seconds, during four successive dives.

No differences were apparent in the choice of food species through the season and no differences could be detected in feeding with respect to seal age. It is evident that the choice of food depends on its availability, and that feeding falls into a pattern which is partly determined by depth of water and distance from the coast.

#### PATTERNS OF FEEDING

In southwest Baffin Island (Fig. 5) all seals which had been feeding in deep offshore waters were subsisting mainly on *Themisto libellula*, a planktonic amphipod. Seals taken in bays or in shoal water were feeding predominantly on the

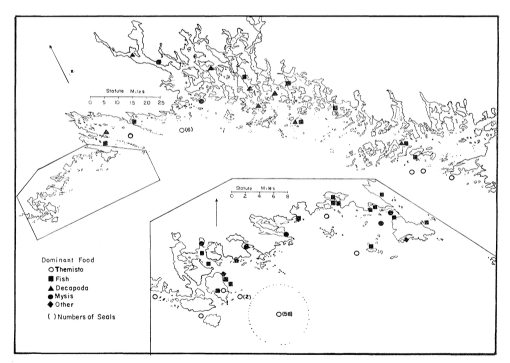


FIGURE 5. The pattern of feeding of Phoca hispida in southwest Baffin Island

polar cod (Boreogadus saida), Mysis oculata, and on various decapods; Themisto was dominant in the stomach of only one inshore seal. This stomach contained fresher decapod material, including a single crab (Hyas coarctatus), and the seal had probably come in recently from the offshore waters. Two seals taken near land fronted by the open sea had been feeding largely on the amphipods Haploops setosa and Gammarus setosus. Other seals killed near the open coast were feeding on polar cod, or Themisto, or both.

In northern Foxe Basin (Fig. 6) the delimitation between inshore and offshore feeding is not so apparent. Foxe Basin is generally shallow and almost all the seals were taken well within the 25-fathom limit (see Campbell and Collin, 1956). Also, *Themisto* does not appear to be as abundant here as in Hudson Strait. The polar cod appears to be the most common food choice in shoal water and *Mysis oculata* also figures largely at such depths. The insert in Figure 6 suggests that *Mysis* is not taken in the shallowest water and this is in keeping with the known habits of this crustacean. Large numbers of polar cod were taken in nine out of eleven seals feeding over the shallow bar of Turton Bay (Figure 6).

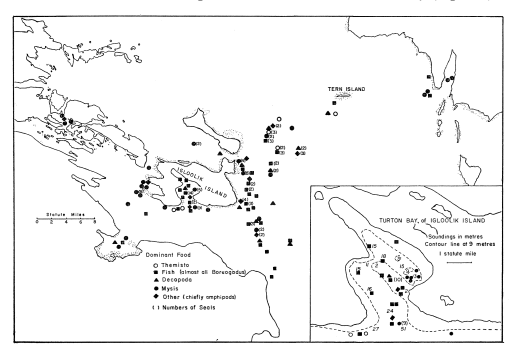


FIGURE 6. The pattern of feeding of Phoca hispida in northern Foxe Basin.

This bar was covered with grounded ice during much of the season, and the Eskimos reported having seen great schools of this fish there earlier in the season. All the seal stomachs shown in Figure 6 which are distributed to the southeast and east of Igloolik Island, and between Igloolik and Tern Islands, were examined in the field during the winter of 1956 (*Calanus* field notes, 1956). *Themisto* was

dominant in four stomachs from the waters between the northern tip of Igloolik Island and Tern Island, where the depths range around 50 fathoms. It is perhaps of interest that the two other stomachs in which *Themisto* was dominant were taken from the southwest corner of Turton Bay (Fig. 6) which is probably the deepest water from which seals were taken, excluding the area to the southwest of Tern Island. Thus, in spite of the obscurity of the pattern in Foxe Basin, a similar correlation between choice of food and depth is suggested.

Many of the stomachs in Ungava Bay were collected by Eskimos and were not accompanied by precise data on origin. Nevertheless, most were probably taken in inshore waters, for the dominant food species, *Mysis* spp., *Thysanoessa* spp., and the fish are all inshore forms; *Themisto* was a minor food element there. Seals from bays and fiords of Frobisher Bay were feeding largely on *Mysis* and fish. *Mysis*, pelagic amphipods such as *Gammarus* spp. and *Apherusa*, and the benthonic genera *Atylus*, *Socarnes* and *Anonyx*, and fish were found in inshore stomachs from Ellesmere and Prince Patrick Islands. The geographical pattern of feeding is not clear in Southampton Island, where *Themisto*, *Mysis* and *Thysanoessa* were the most commonly found food species. A seal off the mouth of the Churchill River, in 14–15 fathoms (26–27 m.) of water, was feeding largely on Decapoda.

The literature suggests that a similar feeding pattern may apply in other parts of the arctic. Johansen (1910) states that seals shot in the pack ice off East Greenland were feeding on pelagic amphipods, but the polar cod was the principal food for fiord-dwelling seals in May to August. The old fiord-seals of East Greenland feed on the larger species of Crustacea (Decapoda?) according to Pedersen (1930). Offshore and inshore seals of the same region may feed on polar cod (*idem*, 1942). The adult seals of Cumberland Sound feed largely on unspecified Crustacea (Kumlien, 1879); in the winter they live inshore, under the fast ice, and subsist to a considerable extent upon fish. Vibe (1950) considers that the polar cod is the principal food when the seals are foraging at great depths and that amphipods and decapods are the chief food elements near the coast of northwest Greenland. This is just the reverse of the feeding pattern displayed in southwest Baffin Island and other localities in the eastern Canadian arctic. Vibe found polar cod almost exclusively in stomachs from Nege, which, judging from his map of the area, is actually an inshore region. At Lake Harbour, on southern Baffin Island, Dunbar (1941) found Themisto dominant in seals taken during August; the stomachs were taken in offshore waters (M.J. Dunbar, personal communication). Other references in the literature, lacking in geographical specificity, do not illustrate the pattern of feeding and depth, but do suggest that the ringed seal has some latitude in choice of food.

Thus, in shallow, inshore waters the ringed seal may feed benthonically or near the bottom, chiefly on fish, *Mysis*, and decapods. It can turn quite easily to strictly planktonic feeding, particularly on Amphipoda, and in the offshore waters of southwest Baffin Island its chief food is *Themisto libellula*. The ringed seal has the ability to utilize organisms at several points of the food chain (as outlined in Dunbar, 1953) and although there may be individual, seasonal and geographical differences in their nutritional histories, food is not likely a limiting factor in the local distribution and numbers of the seal. Important food species, such as *Mysis* and *Boreogadus*, are available in areas where seals are scarce (for example, parts of Hudson Bay) and food availability is evidently not a critical factor in determining the abundance of ringed seals over various parts of their wide range.

#### THE FASTING SEASON

Figure 7 shows, in histogram form, the season of fasting. The degree of fullness of stomachs is not incorporated into this histogram, which is made up of seals sampled from southwest Baffin Island, Southampton Island and northern Foxe Basin. Although there is some suggestion of relaxation of feeding in late winter and early spring, the period of more intensive fasting would appear to extend from April to late June. The fast seems to be strongly associated with the habit of hauling out onto the fast ice, for Freuchen (1935), Pedersen (1930,) and Kumlien (1879) mention the relative emptiness of stomachs from seals which have been basking on the ice. In southwest Baffin Island in 1954, seals began

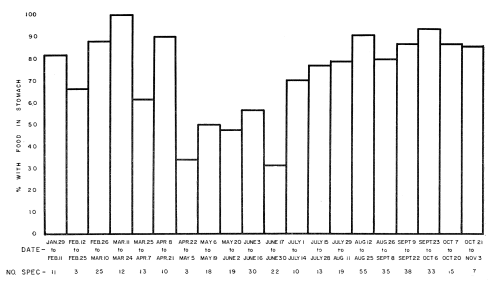


FIGURE 7. The annual cycle of feeding intensity of Phoca hispida, from 381 specimens.

to appear on the fast ice as early as late March, but the numbers were very small until May. On May 14, a day of low cloud-cover, only 3 seals were seen in about 21 miles of travel on the fast ice. On May 16, a day of bright sunshine, eleven seals were seen during 30 miles of travel. The numbers were still relativelysmall in late May, for this period was generally cold and windy. During the hot days of mid-June to early July, large numbers of seals were seen on the ice of Andrew Gordon Bay (see Fig. 21). On June 18, 16 seals were observed lying-

alongside a tide-crack in the fast ice, about 150 yards (140 m.) in length. On July 13, 27 animals were sunning on a sheet of ice, about  $\frac{3}{4}$  square mile in area. After mid-June large numbers of immature animals, which had spent the winter in the open water, moved into the fast ice of southwest Baffin Island. Seals were very common on the fast ice of Southampton Island before mid-June of 1955, when the season was hot and sunny (A. W. Mansfield, personal communication). In northwest Foxe Basin in 1956, on the other hand, the sunny weather occurred much later in the season and seals were not very common on the ice before late June and early July. Table II lists the proportions of food-containing and empty stomachs in relation to the fast ice, for the crucial period of late April to mid-July. During this period, 29% of seals killed in the open water outside the fast ice had empty stomachs, whereas 79% of those taken on the fast ice were not feeding. It appears, then, that there is a general relaxation of feeding in the early spring, perhaps more pronounced in seals living under the fast ice, most of which are adults concerned with reproduction in late March until mid-May. The season of most intensive fasting coincides with the periods when most seals are sunning on the fast ice, and it may be that the timing and intensity of this pronounced fast is dependent on the conditions for hauling out in the sun.

2		Date						
Origin of stomach	Contents	April 22 to May 5	May 6 to May 19	May 20 to June 2	June 3 to June 16	June 17 to June 30	July 1 to July 14	Totals
Outside fast ice	Empty With food	1 1	1 1	<b>4</b> 8	5 14	2 6	1 4	14 34
Inside fast ice (cracks and tide-rips)	K		4 7	•••				4 7
On fast ice	Empty With food	1	4 1	6 1	8 3	13 1	2 3	34 9

TABLE II. Seal feeding in relation to fast ice.

#### FEEDING AND SEAL CONDITION

Some of the routine measurements taken in the field are useful in determining the condition or degree of fatness of the seals. From these measurements, the following indices of seal condition may be derived:

*Index 1.* Blubber thickness may be compared directly with standard length. The thickness of the blubber of the ringed seal is small and this comparison is subject to error of measurement.

*Index 2.* The cube root of the weight may be compared with the standard length. This index shows a very wide range of values and the seasonal trends are not too clear. Since loss of blood cannot be controlled, the wide range of values is not unexpected.

Index 3. This index of condition has been applied by Sivertsen (1941) to the harp seal and by Laws (1953b) to the elephant seal. The circumference of the seal, which reflects blubber thickness by a ratio of change of about 6 to 1  $(2\pi r)$ , is compared with the length of the seal. The size of the seal enters into both numerator and denominator of the formula:

Seal condition =  $\frac{\text{Maximum girth}}{\text{Standard length}} \times 100$ 

This formula was applied to 326 seals, excluding seals of the year, since these have not fasted in the spring, and illustrates the seasonal trends of condition (Fig. 8). The different geographical origins of the seals and the wide range of variation in condition suggest that some data on individual seals and circumstances are required to clarify the picture of seasonal changes.

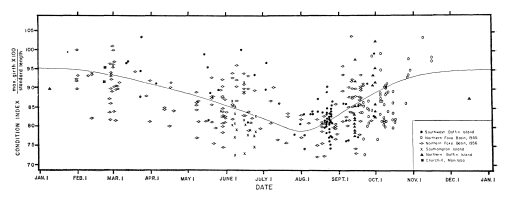


FIGURE 8. Seasonal changes in seal condition. The curve has been fitted by eye, with the aid of semi-monthly means, and has been raised at the left and lowered at the right, in the belief that seal condition does not change much during the winter months.

The average condition index of 6 seals taken in southwest Baffin Island between March 11 and April 4 was 94.9. The average for 5 seals killed on the fast ice during May was 90.9. One of the May seals had been badly wounded a few days previously and its poor condition (index 81.7) may have resulted from this abnormal stress. Thus, there appears to have been little loss of blubber in May and this is in accord with the relative scarcity of seals basking on the ice at this time. At Southampton Island in May, numbers of seals were basking on the fast ice, and in June, all the seals from this locality were killed on the ice; their condition is correspondingly poor, giving an average index of 80.0. The average condition index of the Foxe Basin specimens changed from 90.8 in the period of February to March, to 86.0 in the period of mid-May to mid-June. Both these values are lower than those for the equivalent periods in southwest Baffin Island, but the decrease in condition is about the same in each region. The figures may represent actual differences in seal condition between southwest Baffin Island and Foxe Basin, though they may result from different measuring techniques, since the measurements of the Foxe Basin seals were not made by the author. In southwest Baffin Island on June 12 and 23 two offshore animals

were feeding heavily and were in winter condition (indices 100.0 and 93.6); in living outside the fast ice, presumably neither had experienced much fasting. Seals taken on the fast ice of this area showed a wide range of condition. It is believed that those seals taken near the outer edge of the fast ice in June and early July had come in recently from the open water. Those taken well inside the fast ice were probably winter residents, which had had a longer opportunity for basking and associated fasting. Six seals taken at the head of Andrew Gordon Bay (see Fig. 21) were all quite thin in mid-June. On the other hand, seals taken near the edge of the fast ice in this area were either fat or thin. Two measured from this area in early July exceeded in condition index all but one of the 43 August specimens from southwest Baffin Island and were expected to lose much more blubber before the summer season. In northern Foxe Basin the seals were not common on the ice until late June and early July and, with the late departure of the ice in that area, were common on the ice throughout July. The condition index of the population continued to drop during this period, being 82.1 in the first half of July and 78.8 in the first half of August. The southwest Baffin Island seals, which had probably resumed feeding somewhat earlier, were slightly fatter than the Foxe Basin seals in mid-August. The curve of Figure 8 shows that blubber thickness increases throughout the summer and fall and probably reaches winter condition around the time for freezeup. The northern Foxe Basin specimens best show this trend in late August and early September.

Thus Figure 8 does not adequately express the blubber changes of an individual seal. In brief, a seal probably maintains a high condition index through the winter, undergoes a gradual decline in blubber thickness through the early spring, and loses condition more rapidly during the intensive basking and fasting period of late spring. The blubber increases with resumed feeding in the early summer until a maximum thickness may be reached in the late fall.

The specific gravity of a seal is lowest when the blubber is thickest, and the percentage of sinking seals is greatest immediately after the fast, when the seals are thin. Table III gives the percentages of sinking seals, in two week intervals, for the period of July to the end of September in southwest Baffin Island. Many seals, hunted from rowboat, whaleboat or schooner are retrieved at the last possible moment; the table cannot account for these potential sinkers, nor for variations in the harpooners' abilities. The hunters are much more diligent when the seals are most likely to sink and these factors have an effect on the proportions retrieved.

There may be differences in the feeding activity of inshore and offshore seals of southwest Baffin Island. Thus, 9 out of 40 seals killed in inshore waters after July 15 had empty stomachs whilst all but one of 57 killed during the same period in the offshore waters southeast of Cape Dorset (Fig. 5, insert) had been feeding. Inshore stomachs were also inclined to be less full, whereas those from the offshore area were almost always crammed with *Themisto*. This does not necessarily imply that inshore animals were experiencing sub-optimal feeding conditions. *Themisto* probably offers less food, weight for weight, than do fish

and perhaps decapods, for the intestinal contents of offshore seals often contain much indigestable exoskeletal material; the faeces of fish-eating seals are considerably more liquid and further digested. It is possible that the intensity of feeding is dependent on the nature of the food, and seals eating *Themisto*, or other plankton, must apply themselves more diligently to feeding.

	July 1-15	July 15-31	Aug. 1-15	Aug. 15-31	Sept. 1-15	Sept. 15-30
Killed	15	1	24	34	23	32
Sank	9	1	4	5	1	1
Percentage sinking	60		16.6	14.7	4.4	3.1

TABLE III. Seasonal sinking percentages of offshore *Phoca hispida* in southwest Baffin Island in 1953 and 1954.

There is no evidence of poorer condition of inshore seals in the Cape Dorset area in August, and seals taken from the fast ice in the period of March to late May were certainly not more undernourished than those taken offshore during the same period. Nevertheless, seals living deep within the archipelago to the east of Cape Dorset in September may have been in poorer condition than those living in offshore waters at the same time. From September 6 to 9, 1954, the author accompanied some Eskimos on trips within this region, some 20 to 30 miles from the open sea. Fifteen seals were killed here, but only five were re-This loss is much larger than that found near Cape Dorset in August trieved. and early September (66.7% as opposed to 12.4%; the standard error of the difference in percentage is 11.4% and the chance that these were random samples from the same population is less than 0.006 in 100). The flotation of a seal is delicately balanced and it is clear that the fresher superficial water layer, resulting from the pouring of rivers into the coastal bays and lagoons, would not support the seals as well as the more saline and denser offshore waters. However. this is probably not the whole answer. The condition indices of 43 seals from near Cape Dorset in August ranged from 74.1 to 90.3, with a mean index of 81.8. Four seals from within the archipelago in early September ranged from 76.2 to 85.9 and had a mean condition index of 80.1. These means are not significantly different, but since the seal with the lowest index from the deep-inshore waters was actually recovered from the bottom, the ten which were not recovered may have been as thin as or thinner than this seal. This may indicate that the seals inhabiting deeply indented coasts are thinner than offshore seals at the same It has been pointed out in the last paragraph that it cannot be presumed time. that there is a shortage of available food in the inshore waters. However, it is possible that the long duration of ice in these deeply indented coasts results in a more prolonged period of spring and early summer fasting among the resident seals, and this will be reflected in a greater lag in the return to floating condition after feeding has been resumed.

In 1955, according to A. H. Macpherson (personal communication), seals in the Cape Dorset area were sinking in unusually large numbers throughout the early summer. The spring season was unusually hot and the ice departed rather later than in 1954. It seems likely that the high sinking rate was caused by better conditions for spring basking, which resulted in a lower average index of condition in the summer seals.

#### THE MOULT

The moult in Pinnipedia is annual and always coincides with a period when the seals are able to haul out on the land or ice. Table IV summarizes the moulting season from observations made in southwest and northern Baffin Island and Southampton Island. The moulting season may extend from mid-May to mid-July, though the phenomenon is most marked in June. At this time the seals are able to take advantage of the basking season and the animals are observed to scratch and rub the moulting skin while lying in the hot sun. The Eskimos of southwest Baffin Island believe that hauling-out behaviour, in enabling the seal to free itself of the old hair and epidermis, is necessary for the completion of the moult, and their reasoning is as follows: they suppose that those seals in which the moulting process is prolonged throughout the summer months to the fall were unable to crawl out onto the ice during the basking season, perhaps because they were caught too far offshore in areas lacking suitable ice; but the evidence for their not having been able to come out onto the ice is in the protracted moult, and the argument is thus circular.

Dates (inclusive)	Moulting	Not moulting	Percentage moulting
	No.	No.	%
May 1-7 May 8-14 May 15-21 May 22-28 May 29-June 4 June 5-11 June 12-18 June 19-25 June 26-July 2 July 3-9 July 10-16 July 17-23		$     \begin{array}{r}       3 \\       2 \\       9 \\       3 \\       5 \\       12 \\       4 \\       2 \\       5 \\       \dots \\       4 \\       5 \\     \end{array} $	$ \begin{array}{c} 0 \\ 0 \\ 10 \\ 0 \\ 62 \\ 40 \\ 60 \\ 75 \\ 38 \\ \dots \\ 20 \\ 0 \\ \end{array} $
July 24–30	••••	3	0

TABLE IV.Number, by weekly periods, of moulting Phoca hispida of southwest and northernBaffin Island and Southampton Island in 1954, 1955 and 1956.

When moulting seals are taken much later in the season, the condition is considered to be abnormal. Individuals in this state were taken on August 6 (southwest Baffin Island), September 1 (Southampton Island) and on October 8 and 28 (Foxe Basin). The September 1 and October 28 specimens were second

year females, and were extremely small and delicate for seals of this age (81 and 79 cm. in length respectively, the average length of females in this age-group being about 93 cm.). The September 1 specimen, with an almost bare skin, is illustrated in Figure 9. Another abnormal animal, which was completely devoid of hair, was reported to have been killed in early September, 1954, in southwest Baffin Island, and Kumlien (1879) reports similar hairless seals in Cumberland Sound.

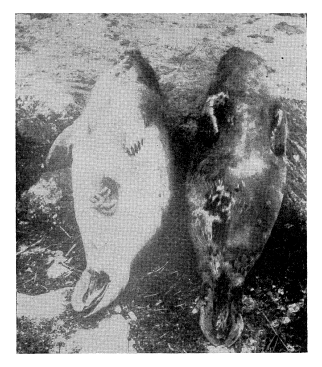


FIGURE 9. An abnormally late and complete moult on September 1. This unusual specimen, on the right, is compared with a non-moulting first-year seal. Note that very little hair remains, chiefly on the right flank, and that the new hair is very short.

#### REPRODUCTION

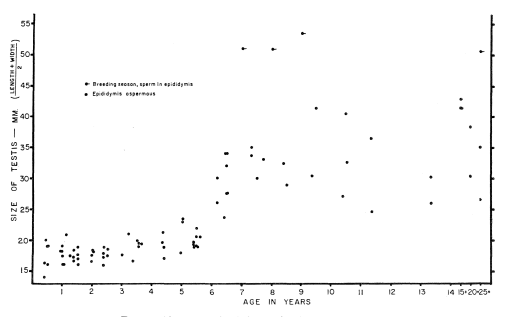
#### Ages at Sexual Maturity

Until the development of an accurate ageing technique, the question of age at sexual maturity for various pinnipeds could not be answered satisfactorily. Data on age at sexual maturity of the ringed seal are scarce and are based on general considerations, rather than on absolute ageing methods. Sleptsov (1943) assumes that maturity is reached after two years. Pedersen (1926) and Freuchen (1935) suggest that the ringed seal is not fully developed until the fifth year, although neither gives any direct evidence for this assumption. It is interesting to note that the latter two authors, both of whom lived with the Eskimos, give estimates which coincide with Eskimo conclusions on southwest Baffin Island and Foxe Basin (see page 8).

THE MALE. The advent of sexual maturity in the male may be considered to coincide with the appearance of spermia in the ejaculatory ducts for the first time. A large sample of seals taken during the breeding season would be required to fix the peak and range of age at sexual maturity, by this definition. Due to the difficulties of hunting at this season (March to mid-May), no such large sample could be obtained.

Two males of 5 years of age and all younger males lacked spermia in the ducts during the breeding season. Another 5-year-old seal on June 5 showed no evidence of having contained spermia (degenerate remains) in the past breeding season, and the epididymis of a 6-year-old on May 3 was aspermous. The youngest male taken with spermia in the epididymis was 7 years in age (SD-27, April 7, 1954).

Other criteria are useful in establishing age of sexual maturity in the male. As with all other mammals, adult seals have a larger test is than immature animals, even outside the breeding season (see Scheffer, 1950). Figure 10 indicates that an increase in test is size occurs during the seventh year and there is a corresponding increase in the diameters of the tubules in the test and also the epididymis (Table V). Apparently there is an increase of sexual activity during the breeding season in immature males as well as in adults and this results in the wide range of histological appearance and tubule size, as demonstrated in the Table. Spermatids were detected in the one fifth-year seal shortly after the breeding season. All sixth and seventh-year testes bore spermatids, and small numbers of spermia were found in the testis, but not in the epididymis, of a sixth-year seal in early June.



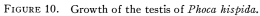


TABLE V. Tubule diameter and spermatogenetic activity in young Phoca	hispida. Tubules were measured from the basement membranes across the
narrowest diameter; the mean of 25 tubules was taken for each specimen.	Measurements are in microns and the ranges are bracketed after the means.

A	Number	Diamete	er of tubule3		Spermatoger	netic stages		
Age group	of specimens	Testis	Epididymis	Spermatogonia only	Spermatocytes	Spermatids	Spermia	
0-1	5	52.8 (45.1-66.2)	72.0 (59.5-87.4)	1	4			
1-2	2	49.8 (45.8-53.8)	58.5 (55.7-61.2)	1	1			
2-3	2	48.2 (43.6-52.8)	51.9 (51.2-52.5)	1	1			
3-4	4	55.4 (47.0-63.0)	74.9 (58.2-91.5)		4			
4-5	4	69.1 (68.2-70.1)	101.4(66.6-157.1)		4	1		
5-6	6	76.8 (58.9–126.1)	100.2 (65.3-165.4)		6	2	1	
6-7	4	109.1 (98.9–121.6)	145.0 (98.2-206.4)		4	4		

Both baculum length and baculum weight (Fig. 11, 12) show a steep rise in the seventh year, though the increase may begin before the seal is 6 years old. Growth appears to level off during the eighth year. The baculum continues to grow throughout the life of the seal, but most of the weight increase appears to result from increase of thickness, and perhaps density, of the bone.

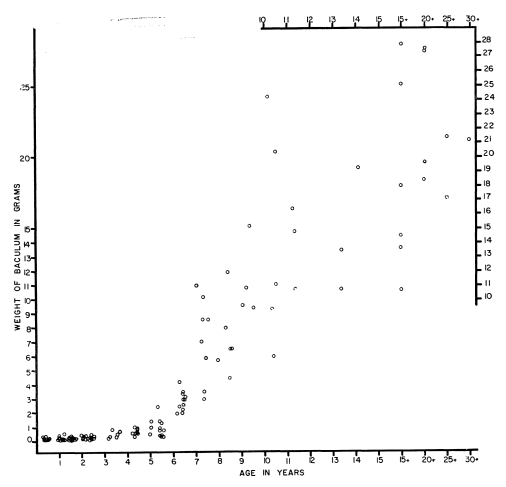
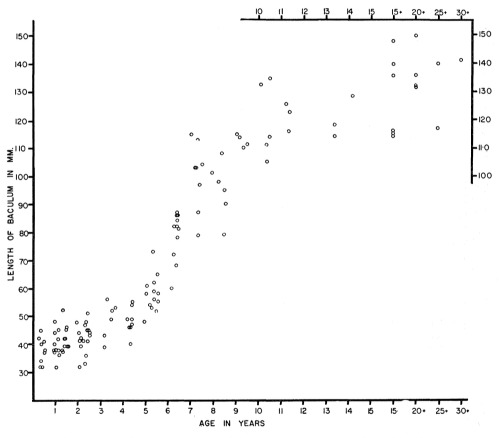
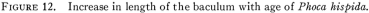


FIGURE 11. Increase in weight of the baculum with age in Phoca hispida.

During the breeding season, the adult male is peculiarly malodorous. Such seals are referred to as *tiggak* by the Eskimos, and the youngest seal recorded in this class proved to be 7 years old. It is also of interest to note that two seals of the early sixth year were the oldest taken outside the limit of fast ice of southwest Baffin Island in the winter and early spring. Residence in the fast ice is a response to breeding urges (see page 38), and this is probably analogous to the incomplete representation of the immature harp seals in the southward breeding migration (see Fisher, 1954a).

All these features suggest that, on the average, the male ringed seal is capable of breeding for the first time when about 7 years of age, though some seals may mature a year earlier. Thus, of 2 seals (SD-106 and 107), both 6 years and about 2 months in age, one had larger testes and these were associated with a baculum of almost twice the weight of the other. The epididymis of the smaller testis was empty except for cellular detritus, whilst the other epididymis contained what appeared to be small numbers of degenerate spermia, together with the cellular detritus. The more advanced of the two individuals may thus have been in breeding condition when about 6 years of age. It is also probable that some seals, for example those in Figure 12 whose bacula are closer to the 7th-year size-mean than to the size-mean of their companion bacula of the 8th year, may not have been capable of breeding when 7 years old.





Fisher (1954a) found an average maturation at 8 years in the harp seal male. However, spermia were found in the testes, although not in the ejaculatory ducts, of harp seals as young as 4 years old, and on this basis Fisher considered seals of 4 to 7 years old to represent a class of adolescents. In the

ringed seal, a very few spermia were found in only one of the 6 breeding-season testes from the age-group 4 to 6+ years (A-15, 5 years and 2 months old); thus, the adolescent class is not well defined.

The age at maturity of the female may be defined as the age The Female. when ovulation first occurs. Productive maturity, the age when the female is first capable of conceiving, may or may not coincide with the age of first ovulation. Ovulation may be detected, after its occurrence, by the development of the corpus luteum at the site of ovulation. In many mammals, especially in pinnipeds, the corpus luteum regresses to a fibrous scar, the corpus albicans, which may persist for some time after parturition. If these scars were retained throughout the life of the seal, they would be of great value, in conjunction with the known age of the seal, in determining the ages of first ovulation. Bertram (1940), assuming them to be permanent, attempted to use corpora albicantia for age determination of the Weddell seal (Leptonychotes weddelli), but the assumption has been found to be unsound for this species (Mansfield, in press). Laws (1953b) suggests that corpora lutea of missed pregnancies in the elephant seal do not result in persistant corpora albicantia as do those of completed pregnancies. Thus there are limits to the value of the corpora albicantia in determining ovulation history.

Macroscopically visible follicles were first noted in one of the 8 fourth-year seals examined. All but 2 of the 10 fifth-year animals showed follicles. Five of the 7 sixth-year ovary pairs and all ovaries from older seals were folliculating.

The youngest seal with a corpus luteum was a fifth-year animal taken in northern Baffin Island between September 9 and 14. The corpus luteum was very small and showed definite signs of regression; at this late date an embryo should have been apparent. Another small corpus luteum was observed in a nulliparous 5-year-old taken on March 25, in southwest Baffin Island. The gland showed some vascular intrusion and was not of recent ovulation. This seal was large and measurements of its reproductive tract indicated that it was considerably advanced sexually beyond 7 other seals of its age-class. It may be concluded that these fifth-year seals were pseudopregnant<sup>2</sup>, the corpora lutea reflecting neither pregnancy in the former animal, nor recent parturition in the latter.

Two out of 7 sixth-year females had ovulated. One, killed on August 24 in southwest Baffin Island, had a less well-developed reproductive tract than the 5-year-old described above. The left ovary contained a small corpus luteum, in which connective tissue was found to be replacing the luteal elements, and the seal was undoubtedly pseudopregnant. The other specimen, a nulliparous female with a well-developed tract, was killed on July 26. The corpus luteum was large, and appeared quite functional. This seal might well have been pregnant, but an unimplanted blastocyst would have been difficult to detect at this time.

<sup>&</sup>lt;sup>2</sup> This term is used here and later to describe several possible types of ovulation without fertilization. Although the corpus luteum in the ringed seal is retained for some time, there are not the associated changes in the reproductive tract which are found in pseudopregnancy of some mammals.

Nine of the seventh-year seals had ovulated and showed corpora lutea. Five of these seals were pregnant and the other 4 were taken at a time when the embryos might not have been apparent. One seal had a normal corpus luteum, measuring 9 by 10 mm. in section on June 18, and another killed on August 9 had a substantial corpus luteum of 10 by 13 mm. and probably contained an unimplanted blastocyst. The other 2 seals, killed on August 6 and 21, had corpora lutea of 8.5 by 8.5 mm. and 8.5 by 9 mm. respectively. The average size of 6 other pre-implantation corpora lutea in August was 10.7 by 12.3 mm., the smallest being 9 by 12 mm. in section. It is probable that the smaller corpora lutea were not supporting blastocysts, though these would be very difficult to detect in the preserved material. Two seventh-year females had not ovulated.

All 7 seals in the eighth-year group had ovulated and 2 contained foetuses. One seal taken on August 6 had a quite large  $(10 \times 12.5 \text{ mm.})$  corpus luteum and was probably pregnant. An attempt was made to locate a blastocyst in the cornu of a female taken on August 25 (corpus luteum 11  $\times$  11.5 mm., and probably pregnant) with no result, but this does not invalidate the assumption that the normal-appearing corpus luteum reflected pregnancy. Three other eighth-year females were taken in September and early October, long after the normal time of implantation of the blastocyst. Their corpora lutea were small  $(7 \times 7 \text{ to } 9 \times 9 \text{ mm.})$ ; two resembled those of pre-implantation seals and the other was regressing, with development of connective tissue and vascular intrusion. These three seals would not be expected to produce young in the following pupping season.

The development of the corpus luteum will be dealt with in a later section. The above discussion is necessary for a consideration of the ages of sexual maturity, here defined as age of first ovulation, in the female.

Table VI summarizes the ages of first ovulation in the female, based on counts of corpora lutea and corpora albicantia. It is assumed, for example, that a female in the tenth year with one corpus luteum and four albicantia will have ovulated for the first time in age-group 5-6; a female in her ninth year, with one luteum and one albicans, will have ovulated for the first time in age-group 7-8. The ovaries of 35 seals younger than three years old were examined in the field or laboratory, but since none of these had ovulated, they are not included in the Table. Laws (1953b) suggests that corpora lutea are absorbed within a year in most elephant seals, while Fisher (1954a) found that a balance between addition and loss of corpora albicantia develops after ten years of life in the average female Figure 13 suggests that the latter phenomenon holds true in the harp seal. ringed seal and Table VI is thus limited to the fourth to tenth-year age-groups. A discrepancy is found between the percentage of mature seals within each agegroup (second column from the right of Table VI) and the percentage ovulating for the first time in each age-group (lower line of Table). The latter percentages, based on the assumption that corpora albicantia of the first ovulation are retained up to the tenth year, are derived from greater population samples. But, since no evidence of first ovulation in the age-group 4-5 is found in the higher age-

	Number of	1	Number o	Percentage	Percentage					
Age-group	specimens	3-4	4–5	5–6	6-7	7-8	8–9	9–10	mature	mature (adjusted)
									%	%
Fourth year	8								0	0
Fifth year	10		2						20	12
Sixth year				2					28	22
Seventh year	11			2	7				81	78
Eighth year	7				5	2			100	100
Ninth year				1	1	3			100	100
Tenth year	3	• • • •		1	2				100	100
Totals	51		2	6	15	5				
Numbers of specimens in age-groups 5-10, etc Percentage ovulating for first time in ag		51	43	33	26	15	8	3		
4–5, etc		0.0	4.7	17.9	57.7	33.3	0.0	0.0		

 TABLE VI.
 Summary of deductions on ages of first ovulation in Phoca hispida from counts of corpora lutea and corpora albicantia.

 Explanation in the text.

35

groups, the assumption that *all* albicantia are retained until the tenth year may well be invalid. It is quite possible that the smaller corpora lutea of pseudopregnancy, described above, are not retained for long, and this is supported by Laws (1953b), who suggests that those of missed pregnancies are not retained at all in the elephant seal. It is felt that a closer approximation of the true picture may be gained by assuming that a given ovulation is apparent until at least the end of the following year (one year after parturition, if this occurs), but that it may or may not be apparent after this time. Thus, limiting the ovulation history of each age-group to the end of the following year, the adjusted percentages of the last column of Table VI are derived. For example, of 7 seals in the sixth year, 2 had ovulated; of 11 seals in the seventh year, 2 showed a single corpus albicans each, and had thus ovulated in the previous year, in age-group 5–6. Thus of 18 seals in the sixth *and* seventh year age-groups, 4 or 22.2%, were mature in the sixth year.

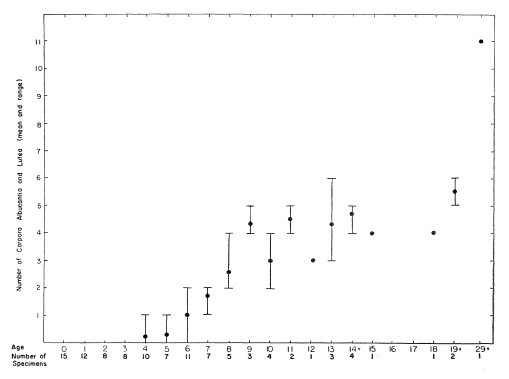


FIGURE 13. The relationship between number of corpora lutea and corpora albicantia, and the age of the female of *Phoca hispida*.

It appears that the left ovary is generally the first to ovulate, for of 13 first corpora lutea, 9 were produced in the left ovary, and if the evidence from corpora albicantia of seals younger than 10 years old is included, then 18 out of 26 first ovulations were in the left ovary. H. D. Fisher (personal communication) found that early collections of harp seal ovaries revealed the same tendency for first ovulation in the left ovary; however, this pattern was not found in later and larger samples. It may be that the phenomenon is due to small samples, but Laws (1956b) found very good evidence that one ovary, in this case the right, tends to ovulate first in the elephant seal.

Fisher (1954a) found fertilization to be 80% successful in mature female In the ringed seal, it appears that corpora lutea are not as likely to harp seals. indicate pregnancy in young seals as in older seals. The virgin uterine cornu is distinctive in its lack of folding and rugosity and is not vascularized as much as are cornua which have been pregnant. Table VII shows the percentages of virgin females in the younger year-classes. Virgin cornua were identified by the criteria mentioned above. Presence of embryos, or of corpora lutea which appeared normal for the pre-implantation period, were taken to indicate contemporary pregnancy. Absence of corpora lutea, or presence of regressing or abnormally small corpora lutea, were taken to reflect absence of an embryo. In order to increase the samples determining the percentages of virgin females, it may be assumed that seals of one age-group show the history of the previous age-groups, at least up to the level of one previous pregnancy. Thus, 7 out of the 11 seals in age-group 6–7 were pregnant, and 2 out of the 7 seals in the age-group 7-8 exhibited one previous pregnancy and were therefore pregnant in the agegroup 6-7. This means that 9 out of 18, or 50%, of seals in these age-groups were virgin in the age-group 6-7. The results of this adjustment are shown on the lower line of Table VII and are probably as close to the real picture as the size of the sample will permit.

		1	Age Group	)	
	4–5	5-6	6-7	7-8	8–9
Number of specimens Virgin Contemporary pregnancy only One previous pregnancy One or more previous pregnancies	10 10 	6 6 	11 4 5 2	7 3 2 2	5
Percentage virgin in each age-group Adjusted percentage virgin	100 100	100 92	36 50	43 25	4 0 0

TABLE VII. Virginity in young and newly matured Phoca hispida. Explanation in the text.

The approach toward maturity in the fifth to eighth year in the female is accompanied by an increase in the size of the genitalia, which is demonstrated in Table VIII. Only tracts from nulliparous females are included, since the cornua are considerably distorted after pregnancy. A series of measurements from 31 older seals show that adult ovary size is reached by the seventh or eighth year; these ovaries averaged 26.1 mm. in length (range, 17–39) and 13.3 mm. in width (range, 10–19).

TABLE VIII. Growth of the virgin female reproductive tract of *Phoca hispida*. Lengths of cornua are measured externally from the junction of cornua to the beginning of the fallopian tube. Average widths are derived from the mean of maximum and minimum widths. Measurements are in millimetres and the ranges are bracketed after the means.

		Ovaries		Cornua				
Age-group	Number of specimens Length		Average width	Number of specimens	Length	Average width		
		mm.	mm.		mm.	mm.		
First year	7	16.7 (14-18)	7.4 (5-10.5)	6	38.3 (27-57)	3.3 (3-4)		
Second year	6	18.7 (15.5-25)	7.5 (6-8.5)	5	47.4 (35-53)	3.6 (3-4)		
Third year	4	18.1 (12.5-25)	7.8 (6-10.5)	3	53.3 (36-71)	3.6 (3.25-4)		
Fourth year	6	17.0 (15-19)	6.9 (5.5-8)	4	58.8 (44-72)	3.8 (3.5-4)		
Fifth year	8	19.9 (16-27)	8.2 (6-13)	4	67.0 (60-73)	5.8 (5-8)		
Sixth year	6	19.9 (17-25.5)	8.8 (6.5-11)	5	71.4 (58-81)	5.3 (4-7)		
Seventh year	6	23.8 (18-28)	11.9 (8.5-15)	5	75.1 (70-87)	10.3 (9.5-12)		
Eighth year	5	24.2 (17-28)	12.7 (10-19)	4	76.9 (63-88)	8.8 (6-10.5)		

There was one senile female aged 35 or more years. No folliculation was detected and each ovary was a mass of scar tissue. At least 11 corpora albicantia stood out distinctly (Fig. 13).

#### THE BREEDING SEASON AND THE ANNUAL SEXUAL CYCLE

THE MATING SEASON. The literature dealing with the ringed seal has little to offer on the subject of the timing of the pairing season. Freuchen (1935) considered that mating occurred shortly after the break-up of the winter ice. Pedersen (1930) suggests that younger females are impregnated first and that copulation takes place from the beginning of May until the end of June. Neither author mentions actual observations of the mating act and neither offers any anatomical evidence for his conclusions. Sleptsov (1943) was probably the first to approach the problem from an anatomical point of view. He concludes that pairing of the Okhotsk Sea *Phoca hispida* occurs in late July through August. The validity of Sleptsov's results will be discussed later in this section.

The only sound methods of establishing the breeding seasons of animals whose mating activities are difficult to observe are histological examination of the testes and sperm ducts and determination of the time of ovulation in the female. Several assumptions on the timing of seal breeding seasons, even when supported by accounts of apparent mating activity, have been proved false by critical anatomical studies (see Fisher, 1954b, on the harbour seal).

A high level of spermatogenesis and, more specifically, presence of spermia in the epididymis, are the sure indications of sexual activity in the male. Eleven adult males fulfilling both these conditions were taken during the period of March 23 to May 15.

The epididymides of 12 adult males taken between May 19 and June 30 were all aspermous, though one on May 22 contained a very few scattered spermia

in some tubules. Most contained varying concentrations of cellular detritus, sloughed spermatogenetic cells, consisting mostly of spermatids, and what appeared to be degenerate spermia. Most summer and fall epididymides contained such detritus and sloughed cells.

Spermatogenetic activity declined after mid-May. Spermia were found in only 9 out of 100 testis tubules of a specimen killed on May 24. No spermia were noted in the testes of males taken on May 27 and May 28 and of 2 killed on June 7, although 3 testes taken on May 19, May 22 and June 7 were still quite active, with spermia present in most tubules. No spermatogenetic stages beyond spermatids were detected in the testes of 5 adults killed between June 16 and June 30. Spermatids were present in varying numbers in all testes from later dates.

Although the majority of males are sexually quiescent during the summer and fall seasons, 2 specimens from Foxe Basin disturb the pattern somewhat. Spermia were present in many of the testis tubules of one killed on August 28, 1956, although the epididymis was aspermous. Both testis and epididymis of another killed on October 10, 1955, contained spermia. Sixteen tubules out of 200 in the section of epididymis contained spermia and 5 of these, which were grouped and possibly represented a single convoluted tubule, were packed full. Both seals were very large, one being the largest and the other the fourth largest in size of males taken during the entire study. Their testes were also among the largest secured in this season. The significance of these active males is not known. They may be quite abnormal and cannot be considered to represent the general fertility of the male population in the summer and fall seasons. An adult from northern Baffin Island on December 19, 1954, was sexually inactive and stages only up to spermatids were represented. Two adult males from Foxe Basin in late February, 1956, less than one month before the date when an active male was taken in southwest Baffin Island, showed considerable activity in the testes, but spermia were not present in the epididymis.

Table IX summarizes the spermatogenetic activity of all adults, 7 or more years in age, as observed from histological sections. Photomicrographs of a selected series of these sections are displayed in Figure 15.

The size of a testis is governed by its internal activity. Breeding season testes are swollen and compact, while those of males taken outside this season are smaller and more flaccid. These differences are due in most part, or entirely, to the increase and decrease of the diameters of the testis tubules, as shown in Table IX. The average tubule diameter in the August 14 to October 10 specimens is 70.5% of the average diameter in the March 23 to May 15 group. This difference is highly significant (P < 0.01) and this has also been demonstrated in changes in the epididymis tubules. Mean external measurements of the testis show that the average size in the period of August to October is only 67% of the size from March to May. These measurements reflect the size of the enclosed tubules within a few per cent, though regression of the interstitial elements between the tubules may contribute slightly to a decrease in testis size. Harrison *et al.* (1952) and Laws (1956b) found a similar cycle of tubule size in antarctic seals.

TABLE IX. Seasonal spermatogenetic activity and tubule diameters of adult testes and epididymides of *Phoca hispida*. Tubules were measured from the basement membranes across the narrowest diameter; the mean of 25 tubules was taken for each specimen. Measurements are given in microns and ranges are bracketed after the means.

Dates (inclusive)	Numbe <b>r</b> of specimen <b>s</b>	Spermia in testis	Spermia in epididymis	Diameter testis tubules	Diameter epididymis tubules
	No.	No.	No.	microns	microns
March 23-April 30 May 8-28 June 7-30 August 14-24 September 8-29 October 2-10 December 19 February 27-28	7 9a 8 7 2 5 1	7 7 1 1 0 1 0 2	7 4 0 0 1 0 0	154.6 (126.1-178.3) 146.9 (130.6-168.3) 119.4 (102.7-147.5) 113.9 (90.9-153.3) 102.7 (99.5-105.9) 104.6 (79.4-144.0) 91.8 142.4 (113.3-171.5)	224 (180.8-272.0) 216.6 (178.9-256.6) 186.9 (128.3-249.6) 146.6 (121.6-185.6) 139.2 (137.9-140.5) 125.4 (113.9-139.5) 123.2 194.0 (143.7-244.3)

<sup>a</sup> Measurements taken on only 6 of these, due to poor preservation.

Figure 14 shows the annual cycle of testis size in scatter-diagram form. The eye-fitted curve gives an indication of the period of testis activity, the peak occurring shortly after mid-April. As will be pointed out below, pups are probably produced in greatest numbers near the beginning of April; thus impregnation might be expected to occur, on the average, about two weeks after parturition.

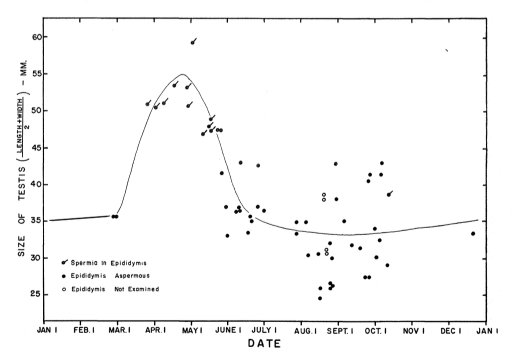


FIGURE 14. The seasonal variation in testis size in Phoca hispida.

Adult males taken in this early spring season have a peculiar musky odour. This is probably of sexual significance and is further field-evidence for the occurrence of the breeding season at this time.

In the central and western Canadian arctic, the ringed seal is commonly taken at its breathing holes. This technique is less often practiced in the eastern arctic, where hunting at the limit of fast ice, or at tide-rips, proves more profitable. The adult female is rarely taken by the native hunter after the beginning of March, until the season of basking on the spring ice. A few females are taken in the fast ice by lowering a firmly secured white-coat seal through the exit of its birth-cave, where it acts as a bait to lure the mother within striking distance of the harpoon. In the hope of determining the state of folliculation and ovulation before the females had weaned their pups, the author entrusted a native with materials for preserving the reproductive tract of a female with pup. Such a tract, well preserved and accompanied by the jaws of the female and pup, was brought to the author in August. It had been taken in mid-April. The right ovary contained the corpus luteum of the previous pregnancy, which, in fact, consisted largely of connective tissue with a few islands of luteal tissue. The opposite ovary contained a new, completely formed corpus luteum, though somewhat smaller than maximum size; it was probably more than one week old, by the criteria used in Fisher's harp seal study (1954a). Three lactating females taken on May 15, May 17 and May 20 all contained fully regressed corpora lutea of previous pregnancy (i.e., corpora albicantia) and new corpora lutea of pregnancy. Thus, the evidence indicates that the male is in full breeding condition before the peak of the pupping season and that the post-parturient female ovulates before the termination of lactation. It is suggested that the peak of mating comes shortly after the peak of pupping, perhaps in mid-April. Among other Phocidae, the harp seal and the harbour seal (Fisher, 1954a, 1954b), and the Weddell seal (Mansfield, in press) probably ovulate shortly after weaning; the elephant seal comes into oestrus during lactation (Laws, 1956b). Of the Otaridae, the Cape fur seal (Arctocephalus pusillus) ovulates about six days after parturition, during lactation (Rand, 1954). Post-partum female harp seals do not establish much contact with the male population until the abandonment of the young, and timing of ovulation is linked with this fact. In the case of the ringed seal, the adult males, which become increasingly potent as the season progresses, and the pregnant adult females live together under the fast ice throughout the winter. Here again, the timing of ovulation with respect to parturition seems to insure a high reproductive success. Pedersen's (1930) suggestion, that young females are impregnated first, may have some basis in fact, when it is considered that nonpregnant females would be the first available to the sexually active males.

Sleptsov (1943) gives a quite different picture of the breeding season in the Okhotsk Sea *Phoca hispida*. His material was collected by a pelagic sealing expedition, between May 11 and October 20, 1939. He states that the onset of ovulation occurs in late June and early July. The probability that the adult

51238 - 4 - 4

seals would not be found outside the fast ice before break-up suggests that this conclusion may be the result of sampling error, due to the absence of adult females in the catch before late June. His conclusions on spermatogenesis in the male are apparently based on field observations of exudate in the spermatic ducts, and not on histological examination. The former approach may be very misleading, for a milky exudate of cellular detritus and degenerate spermia may be present after the breeding season. Sleptsov's whole thesis is a refusal to acknowledge delayed implantation of the blastocyst, which had been suggested by several workers to occur in seals of other Russian waters. In the author's opinion, further work on the Okhotsk Sea pinnipeds will reveal inadequacies in Sleptsov's results.

THE FOLLICULAR CYCLE AND OVULATION. Hamilton (1939), Bertram (1940) and Fisher (1954a) have all commented on the primary importance of the male pinnipeds in determining the timing of the breeding season. In the several species which have been studied, the peak of testicular activity has been seasonally very precise.

One effect of the developing corpus luteum is in the suppression and control of folliculation in some species of seals. Harrison *et al.* (1952) suggest that there are two periods of follicular stimulation in the seals which they studied. The first, in the ovary with the corpus luteum, extends from probably a week after implantation of the blastocyst until the embryo is about 50 mm. in length, and the second occurs in late pregnancy, in the ovary without the corpus luteum, and continues throughout the remainder of gestation, during parturition and until ovulation. Laws (1956b), in his more representative study of the elephant seal, found a more complicated picture of follicular activity. In this species there is a stimulation of folliculation in the ovary destined to release an ovum just before parturition, and there is a period after ovulation in which follicles of both ovaries increase in size and number. After this latter period, a progressive decline is demonstrated in number and size of follicles until implantation occurs, after which there is again an increase in follicular activity, corresponding to the increase found by Harrison at this time. Folliculation is suppressed after the embryo reaches 3-4 cm. in length and this suppression is probably maintained throughout the remainder of gestation. A number of ringed seals are available from the period of pre-implantation and early pregnancy, and examination of the ovaries shows no evidence of very strong follicular stimulation in this period. Table X does suggest that folliculation is suppressed in the ovary containing the corpus luteum from some time in late pregnancy until after parturition, for no follicles of over 3 mm. were detected in the ovary of the post-parturient cornu of a female taken in mid-April. However the opposite ovary, containing the new corpus luteum of pregnancy, was very active and included one 10 mm. follicle. Other ovaries collected on May 15, 17, and 20 did not show marked suppression in the ovaries without corpora lutea. There is a slight increase during early pregnancy in the average number of follicles of ovaries without corpora lutea, and a slight decrease in ovaries with corpora lutea, but the range of variation implies that the changes are not invariable.

Time of	Number	Ov	VARY WITH C	ORPUS LUTI	EUM	Ova	Ovary without Corpus Luteum			Diameters of
year and status of pregnancy	of	Numbers of follicles			Numbers of follicles			Largest	corpus luteum in section	
of pregnancy	specimens	3 mm.	3-5 mm.	5 mm.	follicle	3 mm.	3-5 mm.	5 mm.	follicle	in section
	No.	No.	No.	No.	mm.	No.	No.	No.	mm.	mm.
Mid-April: attending young May 15-20: one or more months after breeding	1	4	1	5	10	6	0	0	2.5	$10.0 \times 8.5$
season	3	7.7 (2-17)	7.4 (5-9)	3.7 (3-4)	5.7 (5.5-6)	8.4 (1-14)	8.4 (5-12)	0.7 (0-2)	5.2 (4.5-6)	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$
June 8-15: about two months after breeding season.	4	12.8 (8-17)	5.5 (0-8)	2.0 (0-5)	4.9 (2-7.5)	5.8 (3-8)	1.8 (0-3)	0.8 (0-1)	4.3 (1-6)	$11.0 \times 9.6$ (10-12 × 9-10)
June 18-26: two or more months after breeding		( )	(/	. ,	(,	. ,	, í	. ,		
season	6	5.5 (3-7)	6.5 (2-16)	3.7 (1-11)	6.8 (5.5-8)	4.8 (3-7)	3.0 (0-5)	0.7 (0-1)	4.4 (1-7)	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$
July 20-31: three plus months after breeding										
season	3	8.0 (7-9)	5.0 (3-6)	1.0 (0-3)	5.5 (3.5-8)	5.9 (3-7)	2.7 (1-5)	0.3 (0-1)	4.2 (3.5-5)	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$
Aug. 6-25: no embryo found, but probably pre-									1.1	
implantation	6	4.4 (3-5)	2.5 (0-5)	1.8 (0-6)	6.5 (0-6)	6.5 (5-9)	5.2 (3-8)	2.0 (0-5)	5.6 (3.5-7)	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$
Aug. 16-24: embryo less than 25 mm. long	8	5.0 (1-14)	2.4 (1-8)	0.9 (0-3)	5.8 (4-8)	6.4 (2-17)	5.5 (1-14)	1.6 (1-3)	6.1 (4-8)	$11.5 \times 9.3$ (10-14.5 × 8-10)
Aug. 24-Sep. 27: embryo 28-42 mm. long	9	6.7 (2-15)	4.2 (0-9)	1.3 (0-4)	5.3 (3.5-7)	10.5 (5-48)	2.8 (0-7)	1.0 (0-4)	5.1 (3-9)	$\begin{array}{c c} 12.2 \times 10.9 \\ (11-14 \times 10-12) \end{array}$
Sep. 9-Oct. 7: embryo 63-110 mm. long	11	9.0 (2-24)	5.7 (1-10)	1.9 (0-6)	5.8 (5-7)	6.8 (4–15)	7.3 (2-11)	2.6 (0-7)	6.4 (5-8)	$12.7 \times 10.8$ (10-14 × 9-13.
Aug. 6-Oct. 9. virgius with small corpora lutea in			, í	. ,	. ,		. ,			
various stages and probably pseudopregnant	7	8.3 (3-16)	2.3 (0-9)	0.4 (0-2)	3.6 (1-6)	10.0 (2-22)	1.4 (0-6)	1.1 (0-6)	3.4 (1-7)	$7.8 \times 7.4$ $(5-9 \times 4-9)$
Sep. 9-14: missed pregnancy, corpus luteum			. ,	. ,		. ,				
regressing	1	14	2	0	3.5	26	0	0	2.5	$12.0 \times 10.0$
Oct. 4: no corpus luteum, but with large follicle,		4	0	1	8.5	0	0	0		
probably about to ovulate	1	4	0	1	0.5	0		U		
March 25: virgin with regressing corpus luteum	1	4	5	2	7.5	12	4	2	9.0	$8.5 \times 5.0$

## TABLE X. Summary of seasonal follicular and corpus luteum development in adult female Phoca hispida. (Ranges are bracketed below mean values.)

The control of folliculation around the time of ovulation tends to impart an alternation of pregnancy between right and left uterine cornua. The unbalance of corpora albicantia in some ovary pairs (e.g.: S-4, 3 to 1; S-27, 4 to 1; SD-528, 4 to 1) suggests that alternation does not always occur. It may be that presence of an embryo most effectively suppresses ovulation in the ovary of the pregnant cornu. Thus the female SD-58, taken on May 17, 1954, contained a new corpus luteum in the left ovary. This ovary also held what was probably the regressed corpus luteum of the previous pregnancy. The *right* cornu was flabbier and more heavily vascularized, and was probably the site of the previous pregnancy; here the ovum, or the blastocyst, probably passed from the site of its origin to the opposite horn. A definite case of this was found in a pregnant August specimen. It would seem, from the May example, that the influence of this transmigration has permitted ovulation to occur in the ovary which contained the corpus luteum. This may be one of the factors which upsets any regular alternation of ovulation in the ringed seal.

THE DEVELOPMENT OF THE CORPUS LUTEUM. The corpus luteum develops rapidly after ovulation in seals, staying relatively small until implantation of the blastocyst and then increasing to its full size. It then maintains this size, or regresses slowly, according to the species concerned (Harrison, 1948). The first part of this cycle, up to the early development of the embryo, is outlined in Table X. The Table reveals an increase in the diameters of the corpus luteum in August, both in pregnant seals and in seals in which implantation was not yet evident. The means of the average diameters of the two groups of corpora lutea, from mid-April to July 31 and from August 6 to October 7, differ by 1.354 mm. with a standard error of 0.31 mm.; thus there is a highly significant increase in size of the corpus luteum around the time of implantation in August.

Structural changes occur during the life of the corpus luteum; these are shown macroscopically in Figure 16 and microscopically in Figure 17. The new corpus luteum (SD-443, mid-April) is pale yellow in appearance, with little vascularization apparent macroscopically. Histologically, the gland is rather loosely packed with large, occasionally vacuolated cells and there is little connective tissue present, except along the trabeculae which project from the old follicle wall into the cell mass. The condition of corpora lutea in May and June is more or less similar; the luteal elements are more packed, but there is little development of connective tissue between the cells. Near the time of implantation, in late July and early August, the appearance of the gland is considerably changed. There is much vascular intrusion, particularly along the trabeculae, which become readily visible macroscopically. Nodules of connective tissue develop and almost all the luteal cells are separated by collagenous fibres, which stain very well with fast-green. Corpora lutea of later pregnancy, corresponding to foetuses up to 110 mm. in length, do not differ materially from this condition. No late pregnancy ovaries were secured in this study, but some of the regressing corpora lutea of pseudopregnancy exhibited increased connective tissue and vascular intrusion as well as the heavier rusty colour described by Fisher

(1954a) in late pregnancy harp seals. The corpus luteum of the previous pregnancy in the mid-April female was considerably changed. Luteal tissue remained isolated in small islands cut off by radiating masses of connective tissue, and individual luteal cells were shrunken and surrounded by collagenous fibres.

These changes in size and structure of the corpus luteum have been described by Harrison et al. (1952) and Laws (1956b). They found that the sudden increase in size and change of structure is associated with the implantation of the embryo in the several species studied, as it is in the ringed seal. In the ringed seal, the gland retrogresses rapidly after parturition and ovulation occurs during lactation, as Laws found in the elephant seal. The corpus luteum of the Weddell seal also degenerates rapidly after parturition, but this animal does not ovulate until the completion of the lactation period of about two months (Mansfield, in press). In the harp seal, on the other hand, the gland appears to maintain its size and activity until the short, two-week lactation period is complete (Fisher, 1954a). The harp seal does not ovulate until after lactation and the abandonment of the young, and Fisher suggests that the maintenance of the glandular activity of the corpus luteum in this species is linked with the timing of the breeding season. If the corpus luteum thus has a direct effect in suppressing oestrus, it is not known how the Weddell seal (Mansfield, in press) prevents ovulation from occurring until the end of lactation, long after the regression of the corpus luteum of the last pregnancy. It would seem that the whole question of timing of ovulation and development of the corpus luteum cannot be subjected to too much generalization, and that specific patterns are adapted to environmental and behavioural circumstances.

POLYOESTRUS IN THE YOUNG FEMALES. It is by no means an established fact that all female seals are monoestrus, ovulating only once annually, within the breeding season. Fisher (1954a) offers good evidence for a period of follicular stimulation during the breeding season, even in immature harp seals. On the other hand, the unimpregnated southern sea lion (Otaria byronia) can folliculate outside the normal breeding season (Hamilton, 1939). The newly matured cape fur seal may experience its first ovulation three months before the normal season of the older adults. Such premature oestrus may result in pregnancy in this species, as the male may be in rut at this time (Rand, 1954). The material from the ringed seal suggests that it too may be polyoestrus. On March 25, 1954, a five-year-old female was found to have a regressing corpus luteum in the left ovary. The animal was nulliparous, judging from the condition of her reproductive tract. Three other nulliparous females with small, relatively regressed corpora lutea were killed in August and September and small corpora lutea, with less vascularization and connective tissue, were found in four specimens taken during the period of August to October. The differences in macroscopic and histological appearance are shown in Figures 16 (J, K & E) and 17 (E & F), and their smaller sizes and associated follicular patterns are outlined on Table X. Unless there is an inexplicably great difference in the rates of regression of corpora lutea of pseudopregnancy, for several of these seals were taken long after the embryo should have been apparent, it is difficult to believe that all these corpora lutea resulted from breeding season ovulations. A ninth-year seal, which had borne young in the past, was found not pregnant in early September. The corpus luteum, which had presumably resulted from a breeding season ovulation, was regressing. This may have been an example of missed impregnation. It is suggested, in the belief that corpora lutea of pseudopregnancy are likely to regress

after a relatively short time, perhaps after the normal lapse in time of preimplantation, that the March specimen represents an ovulation which occurred in the winter season. The regressing corpus luteum taken in August may have resulted from ovulation shortly before the breeding season. The corpora lutea of pseudopregnancy taken from September 30 and October 9 seals are younger in appearance and may have resulted from ovulations after the normal breeding season. More direct evidence for such unseasonable ovulation is shown in a fifth-year female secured on October 4. The left ovary of this specimen contained a large follicle, which appeared all but mature (Table X, Fig. 16A). It will be noted that the phenomenon of pseudopregnancy seems almost confined to the younger, nulliparous seals (see also the section on virginity in the young females, page 37); only one seal of the nine in which pseudopregnancy was known or suspected had borne young previously. It may be that the follicular cycle is fixed by the advent of full maturity, or by the first pregnancy, or by both.

#### Implantation and Development of the Embryo

It has been shown that the female is impregnated shortly following the birth of the young. The first pregnant specimen was taken on August 6, 1954, but unfortunately the thickened uterine tract was eaten by dogs. Twenty-eight embryos, in various stages of development, were removed from females in August to early October. Another 7 embryos from later dates were available in the collections of the National Museum of Canada. Measurements of a foetus killed on January 9, 1954, in northern Baffin Island, complete the series available for study. The smallest embryos were removed shortly after mid-August, in southwest Baffin Island; their average size at this time suggested that implantation had occurred one to two weeks earlier. If we consider that impregnation occurs in mid-April, a delay in the development of the foetus of about  $3\frac{1}{2}$ months is implied. This is somewhat longer than the delays found in most other pinnipeds.

Attempts were made, without success, to locate unimplanted blastocysts in fresh and preserved material. Since the centers of the uterine swellings of three tracts in August were 28 mm., 35 mm., and 24 mm. from the junctures of the uterine cornua, this gives some indication of the length of tract which would have to be sectioned with a view to discovering pre-implantation embryos in serial section. Also, anomalies do occur, such as the trans-uterine migration of the blastocyst, described above. Thus it was felt that the size and state of the corpus luteum gave sufficient evidence of hidden pregnancy.

Figure 18 illustrates the size of these embryos and their dates of removal. A straight line of least squares fitted to these points (not shown on the Figure) indicated an average date of implantation on August 27 and a birth size of 657 mm. on March 1. A spread of implantation times from August 3 to September 17 (45 days) was derived from lines parallel to the line of least squares running through the two extreme deviations on either side of this line. Actual birth size approximates 65 cm. (from three full-term foetuses and two newly born white-coats in northern and southwestern Baffin Island and Foxe Basin). If the

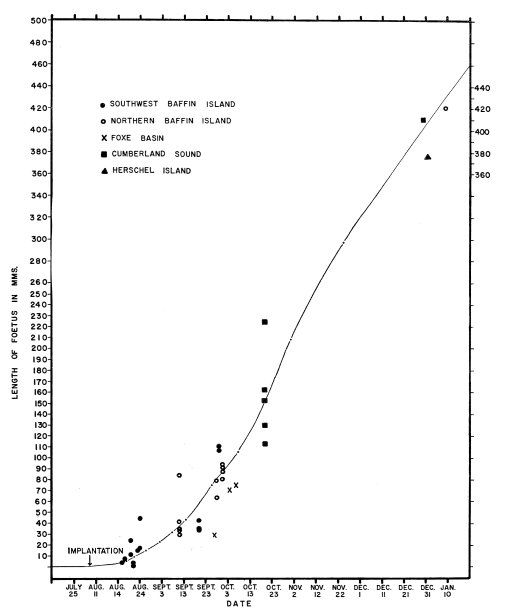
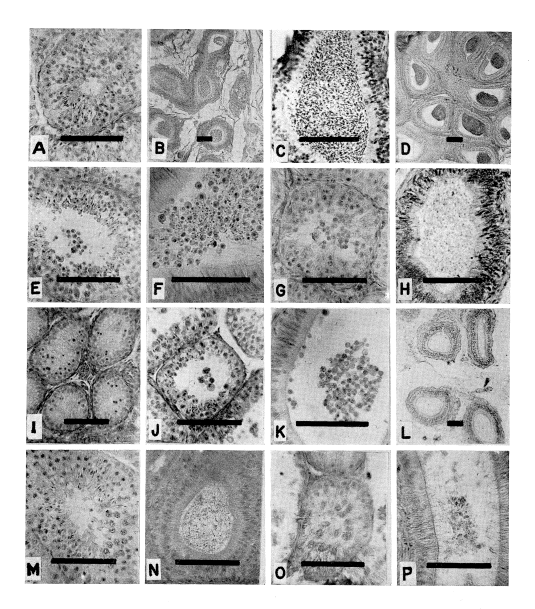


FIGURE 18. Early growth of the ringed seal foetus. The curve is derived from human embryonic growth data, as described in the text.

FIGURE 15. Photomicrographs of testis and epididymis sections from adult *Phoca hispida*. All were sectioned at  $7\mu$  and stained with Delafield's haematoxylin, and eosin. The black bars represent  $75\mu$ .

- A. SD-20, April 1, 1954. 25-plus years old. Testis. Highly active, with all stages up to spermia present.
- B. SD-18, March 23, 1954. Almost 8 years old. Epididymis. Almost all tubules are packed with spermia, although the mass of cells is somewhat shrunken away from the wall in most tubules.
- C. SD-18, March 23, 1954. Almost 8 years old. Epididymis. A single epididymis tubule showing packed spermia and high, active lining epithelium.
- D. A-8, May 15, 1955. 9-plus years old. Epididymis. Tubules packed with spermia. Testis of this specimen still highly active.
- E. A-6, May 15, 1955. 10-plus years old. Testis. Still quite active, but note the spermatogenetic cells (chiefly spermatids) in the lumen.
- F. A-6, May 15, 1955. 10-plus years old. Epididymis. A few spermia are present, but the bulk of the content is made up of sloughed spermatogenetic cells and detritus.
- G. SD-70, May 24, 1954. 15-plus years old. Testis. No spermia are present, but spermatocytes and spermatids, some of which are shown sloughed in the lumen, are common.
- H. SD-70, May 24, 1954. 15-plus years old. Epididymis. Degenerate cells and colloid fill the lumen, although a few spermia were found in other tubules.
- I. A-16, June 7, 1955. 10-plus years old. Testis. A relatively inactive testis with a few dark-staining spermatids, some detached in the lumen, but with no spermia.
- J. A-20, June 7, 1955. 9-plus years old. Testis. Quite active, with sloughed spermatocytes and spermatids in lumen.
- K. A-20, June 7, 1955. 9-plus years old. Epididymis. Detail of spermatogenetic cells in lumen. No spermia were present in any of the tubules, although these were still being produced by the testis (see J).
- L. A-35, June 24, 1955. 10-plus years old. Epididymis. Empty but for colloidal material. Epithelium somewhat less columnar than those earlier in season. This condition is typical of summer-killed seals.
- M. S-55-50, October 10, 1955. 20-plus years old. Testis. Highly active testis. No other specimens showed such activity at this season.
- N. S-55-50, October 10, 1955. 20-plus years old. Epididymis. Spermia-packed tubule, one of several such in the section.
- O. S-55-13, September 24, 1955. 10 years and six months old. Testis. More typical testis tubule of autumn season. The large cells are probably spermatocytes.
- P. S-55-12, September 24, 1955. 16 years and six months old. Epididymis. More typical contents of autumn specimens, consisting largely of degenerate cell products.



51238 - 4 - 4a

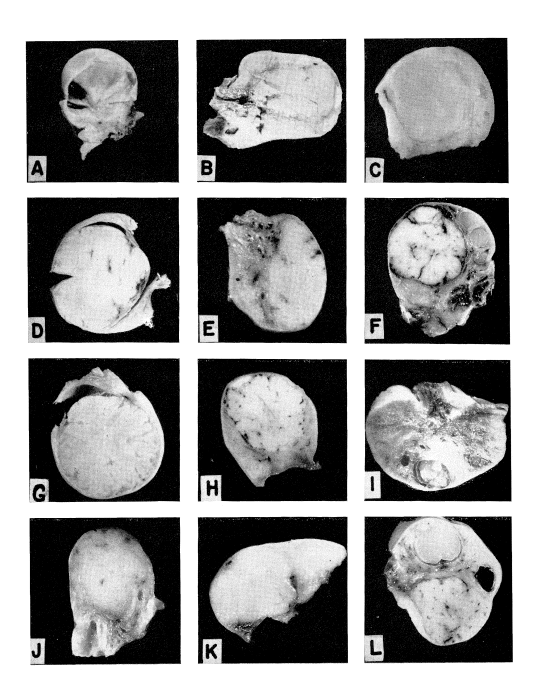


FIGURE 16. Macroscopic appearance of ovaries in section.

- A. S-55-28. Enlarged follicle of fifth-year ovary in section. Follicle 8.5 mm. in maximum diameter and probably close to ovulation size.
- B. SD-443. New corpus luteum of eighth-year post-partum female in mid-April. Very little vascularization or connective tissue in gland, which is somewhat crumbled due to loose packing of cells. The gland is 8.5 mm. by 11 mm. in section.
- C. A-33. A fairly recent corpus luteum from a female of nine-plus years taken on June 23. Lack of vascularization typical of corpora lutea at this stage of pre-implantation of the embryo. The gland is 13 mm. by 11 mm. in section.
- D. # 50. Corpus luteum of a fourteenth-year female killed on August 11. Tract was very slightly swollen, but field examination revealed no embryo. The gland is large (14 mm. by 13 mm.) and some vascularization (dark areas) is evident.
- E. SD-461. Corpus luteum of fourteenth-year female on August 20, supporting an 11 mm. embryo. The gland is 14.5 mm. by 8 mm. and slight vascularization is evident.
- F. S-36. Corpus luteum of eighth-year female on August 24, with a 44 mm. embryo. The gland is not very large (11 mm. by 11 mm.) but much vascularization is shown.
- G. S-55-39. Corpus luteum of a female in her tenth year on October 7, with a 74 mm. embryo. Very little change from F, although the vascularization is less dark due to differences in preservation. The gland is 12 mm. by 11 mm. in section,
- H. SD-527. Corpus luteum of a ninth-year female on September 29. The seal was carrying a 110 mm. foetus and the gland, which was 11 mm. by 11 mm. in section, shows little change from G and F in amount of vascularization.
- I. SD-58. The corpus albicans of a female fifteen or more years in age on May 17. The scar was probably more than one year old: the scar of the more recent corpus luteum, not shown in this section, was more conspicuous in this same ovary.
- J. SD-15. The corpus luteum of a nulliparous female, about five years old, on March 25. The ovaries were frozen and poorly preserved for histological study, but the gland is small (8.5 mm. by 8 mm.) and shows vascularization too extensive for recent formation,
- K. SD-291. A rather young corpus luteum from a nulliparous seventh-year female on August 6. The gland was only 8.5 mm. by 8.5 mm. in section, pale yellow and unvascularized, and was probably not supporting an unimplanted blastocyst.
- L. SD-533. The corpus luteum of an eighth-year female which was not pregnant on September 30, long after the normal implantation time. The gland is small (9 mm. by 9 mm.) and quite heavily vascularized.

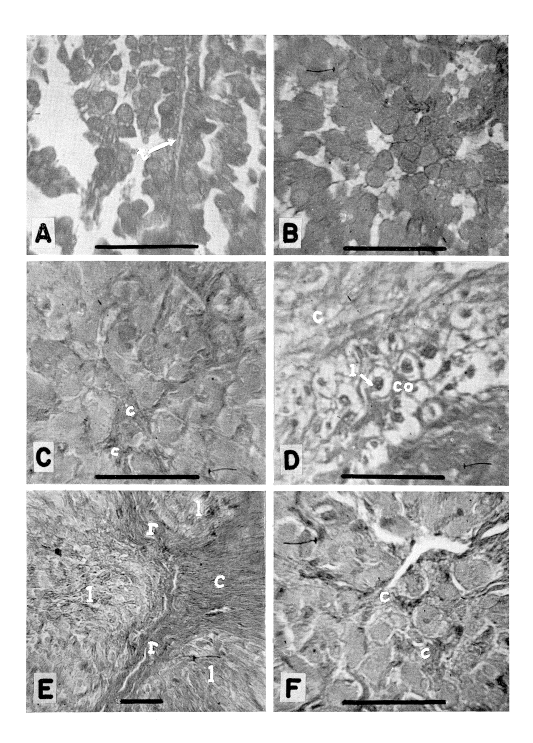


FIGURE 17. Photomicrographs of sections of corpora lutea from *Phoca hispida*. All were sectioned at  $10\mu$  and stained with azocarmine, phloxine, and fast-green. The lines represent  $100\mu$ .

- A. SD-443. New corpus luteum of mid-April. Note the loose-packed cells, denser along the trabeculum. t = trabeculum.
- B. SD-59. Corpus luteum on May 20, probably about one month old. The luteal cells are loose-packed and there is little connective tissue between them.
- C. SD-528. Corpus luteum of September 29, supporting a 106 mm. foetus. Note the development of connective tissue between the more closely packed luteal cells. c = connective tissue.
- D. SD-443. Old, post-partum corpus luteum in mid-April. Note the large connective tissue masses and the network of collagenous fibres constricting the luteal elements.
   c = connective tissue mass.
   co = collagenous network.
   l = luteal cell.
- E. S-39. Regressing corpus luteum of nulliparous sixth-year female on August 24. Bodies of luteal tissue are being cut off by connective tissue instrusions.
   c = large connective tissue mass.
   r = radiating arm of this mass.
   l = luteal tissue.
- F. SD-533. Corpus luteum of a nulliparous eighth-year animal on September 30. No embryo was present, but the structure of the gland is similar to that of pregnant seals (as in C, above). c = connective tissue.

spread of implantation dates is related to the length of the breeding season, this should be about 45 days (equivalent to April 1 to May 15). Thus the line of least squares for these data gives a birth size and implantation spread in keeping with the expected figures. But the line does not fit well the sizes of the earliest embryos, taken from August 16 to August 24 in southwest Baffin Island; these are larger than the line would imply. Arey (1954) gives figures of human embryonic growth. Table XI converts these data into the hypothetical growth curve of the ringed seal embryo. No measurement exactly comparable with the crown-rump length of the human could be applied to the ringed seal foetuses, which were measured from tip of nose to tip of tail, along curvatures and flexures, but the proportionate growth is considered to be of the same form. The growth period is here considered to be 240 days (for the sake of a round figure, from August 1 to March 29), and the birth size to be 650 mm. The growth curve derived from these assumptions fits the points of Figure 18 somewhat better, particularly the earlier growth stages.

Hu	MAN	Ringed Seal				
Age	Crown-rump length	Age	Nose-tail length			
days	mm.	days	mm.			
$7 \\ 14 \\ 17.5 \\ 21 \\ 24.5 \\ 28 \\ 35 \\ 42 \\ 49$	$\begin{array}{c} 0.1 \\ 0.2 \\ 1.5 \\ 2.0 \\ 2.5 \\ 5.0 \\ 8.0 \\ 12.0 \\ 17.0 \end{array}$	$ \begin{array}{r} 6 \\ 12 \\ 15 \\ 18 \\ 21 \\ 24 \\ 30 \\ 36 \\ 42 \\ \end{array} $	$ \begin{array}{c} 0.2\\ 0.4\\ 2.8\\ 3.7\\ 4.6\\ 9.3\\ 14.9\\ 22.3\\ 31.6 \end{array} $			
56 70 84 112 140 280	17.0 23.0 40.0 56.0 112.0 160.0 350.0	42 48 60 72 96 120 240	$\begin{array}{c} 31.0\\ 42.7\\ 74.4\\ 104.0\\ 208.0\\ 297.0\\ 650.0 \end{array}$			

TABLE XI. Conversion of rate of human embryonic growth data into ringed seal embryonic growth.

Figure 18 reveals a quite large spread in implantation time. This is perhaps confirmed by the occurrence of one embryo of 24 mm. on August 20 and another of 28 mm. on September 27, suggesting that the range of implantation time is, at least, just over one month. The straight line growth curve, when applied to the extremes of the range, suggests an implantation period from August 3 to September 17 (45 days). The hypothetical curve gives an implantation period from July 9 to August 22 (46 days) and is more in keeping with the sizes of early-August embryos. Since the period of rut of the male and post-pregnancy availability of the female is also about 45 days (early April to mid-May), the spread in implantation dates may reflect the period of impregnation time.

It has been suggested (page 37) that nulliparous females might be impregnated earlier in the breeding season. No statistical difference could be shown in the comparison of deviations from the hypothetical line of nulliparous and parous females; nor could differences be demonstrated in the means of implantation times of embryos from the four localities, southwest Baffin Island, northern Baffin Island, Foxe Basin and Cumberland Sound, although the pupping season may be somewhat later at higher latitudes. The sample is small for statistical treatment and it may be that any effect of impregnation date on implantation time is masked by inherent variations of implantation times among females.

#### Birth and Care of the Young

THE BIRTH-LAIR. The young seal is always, as far as is known, born on land-fast ice. From March 25 to April 15, 1954, no pregnant or post-parturient female was taken in many days of hunting at the tide-rips within the fast ice of southwest Baffin Island. Kumlien (1879) suggests that the relative scarcity of females in the tide-rips of Cumberland Sound, as early as the beginning of March, indicates that the females have chosen localities for producing pups by this time.

The young seal is born on the ice, generally in an excavation hollowed out by the female in the overlying snow (Fig. 19, 20A). The cavity is usually elongate, 10 or more feet long, with the entrance from the water situated at one end. The height of the roof above the ice depends on the depth of snow. The

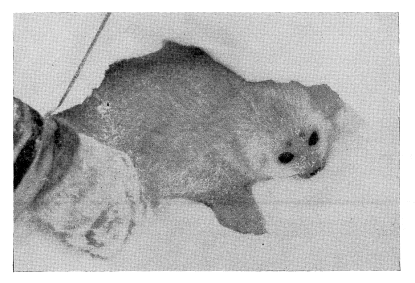


FIGURE 19. Ringed seal pup, April 8, 1954.

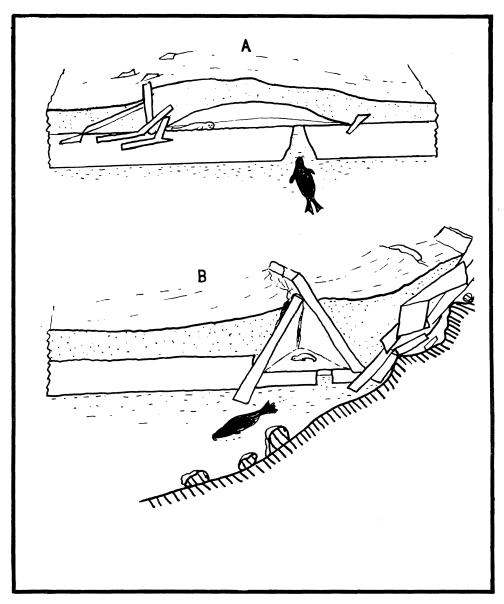


FIGURE 20. The two types of pupping-lair. (A) is the form hollowed out by the female in drifted snow. Note the exposed broken ice, causing drift. (B) shows a pup in a natural pressure-ridge lair.

extremities of the lair may be very low, suggesting that it is constructed by scraping and ploughing with the extended foreflippers. Radiation and breathing of the seal increase the stability of the lair, through icing of the inner surface. Many of those lairs observed in the spring of 1954 could be broken into only by vigorous and repeated jumping on the roof.

A sufficient depth of snow is required for the construction of the birth-lair. It is interesting to note that most of the lairs observed in 1954 were built in the drifted snow which tends to develop in areas of rough ice, but it is not known if suitable snow depth is chosen directly. Within the areas where fast ice forms in the autumn, the seals may tend to congregate in patches of open water, some of which result from the break-up of newly formed ice. This would enable the seals, for a short time before refreezing occurred, to maintain a simpler existence, without the necessity for breathing holes. Indirectly, a sufficient final snow cover would result on the rough ice formed at the temporary break-up. The following observations, extracted from the author's field-notes, may indicate an active choice, or 'prediction', of suitable snow depth.

On April 2, 1954, 16 breathing holes, 100 or more feet apart, and four pupping-lairs, from one of which a white-coat was removed, were discovered by the dogs in a long, narrow band of heavily drifted broken ice. By its general appearance, this band was thought to have represented a boundary between ice and open water, at some time during the season of freezing. Floating ice, trapped and frozen at this boundary, or the grinding action of newly formed ice sheets against the older sheet of ice, could have resulted in the rough ice pattern as found in April. Several traverses of smooth, poorly covered ice in the area failed to reveal any birth lairs, although breathing holes, including those of several malodorous males, were found in such ice.

This distribution suggests that the females moved to the edge of the older sheet of ice before further freezing took place, or that they moved into the rough ice at the boundary through recently congealed surroundings. Such areas of broken ice may be encompassed by stable, early-formed ice, so that ice stability and snow cover, both desirable for pupping success (see page 60) need not in this case be antagonistic. Observations in northern Baffin Island (Ellis, 1957) show that seals may often appear in surprisingly large numbers at newly formed tidecracks in the fast ice in mid-winter. The seals may be taking advantage of easier conditions for keeping open the fresh breathing holes in the new ice which forms in these cracks. Drifted snow may also form along such cracks, thus making them more suitable for birth-lairs. Most seals give birth in deep bays and close to shore, where the basic snow cover and ice stability are optimal. Figure 20A depicts a lair of the broken-ice type.

Some seals produce pups in the pressure ridges which often form just outside the ice which lies on the shore at low tide (Fig. 20B). The chief advantage of this sort of lair is that its canopy of solid ice makes it less assailable by foxes and humans, but its instability probably more than outweighs this. Ice movement may take a heavy toll of pups born in these pressure-ridge lairs.

THE YOUNG SEAL. By all accounts, the great majority of white-coats are born from mid-March until mid-April, though a few may be produced outside these dates. Freuchen (1935) states that in the Canadian eastern arctic and north of Hudson Bay, "in the years of the [Fifth Thule] Expedition . . . most of the seals had their young in the beginning of April". The Eskimos of southwest Baffin Island were of the opinion that a similar average birth date applies in that area. Ellis (1957) considers that the pupping period in northern Baffin Island extends from about April 9 to at least April 16 and it seems likely that the season is delayed somewhat in higher latitudes as Freuchen also suggests.

In southwest Baffin Island, a full-term foetus, taken by the natives near the beginning of March, weighted 9.5 lb. (4.3 kg.); and a recently-born female of 10.5 lb. (4.8 kg.), with a raw umbilical cord, was tagged on April 2. Another fully developed foetus and a new-born white-coat in northern Baffin Island weighed 10 and 11.5 lb. respectively (4.5 and 5.2 kg.). Kumlien (1879) gives a range of weight of 4 to 6.5 lb (1.8–2.9 kg.) for new-born pups in Cumberland Sound, weights considerably smaller than those found in the above localities. An older male white-coat tagged on April 8, weighed 24 lb. (10.9 kg.), indicating a rapid gain in weight. As stated earlier, the length at birth appears to be about 65 cm.  $(25\frac{1}{2} \text{ inches})$ .

The young begin to shed the foetal coat after two weeks (Pedersen, 1930) or at least before 20 or 30 days (Kumlien, 1879). A white-coat on Southampton Island exhibited slight darkening on the face and flippers on May 2 (A. W. Mansfield, field-notes, 1955), but another pup examined by the present author on southwest Baffin Island had only small amounts of foetal hair clinging to the middorsal area on May 14. On the other hand, a young of the year in Foxe Basin was still almost all white on June 10 (*Calanus* field-notes, 1956) and patches of white hair were still present on another killed on June 18 in northern Baffin Island (D. V. Ellis, personal communication). The latter two localities are more northerly and the later retention of the foetal coat may be the result of later birth dates.

The stomach of the pup taken on May 14 contained nothing but curdled milk. In late May several young were observed sunning themselves in the depressions formed after the birth-lairs had caved in. A large seal, probably the mother, was observed lying beside one of these young on May 26. Milk was found in the incised mammae of three females killed on May 17, 19, and 20. No other post-parturient female was taken before June 26, when a single nonvirgin female showed no signs of lactation. According to native information, some females lactate until at least late June and the stomachs of young seals in mid-June may contain milk, a mixture of milk and sea-food, or sea-food alone. The statement of Freuchen (1935) that "milk is to be found in the stomachs as late as in September or October" seems erroneous; no first-year animal taken by the author in summer or fall was milk-fed.

This long period of parental care, potentially from April until the break-up of winter ice, is of great significance in the determination of movements and distributions of this species and will be enlarged upon in the last section.

#### The Sex Ratio

Too few new-born pups were taken for an estimate of the sex ratio at birth. During the study, 259 males and 229 females were taken, these figures excluding all samples on which sexual selection might have been practised by Eskimos or the author. There is, however, some sexual segregation of the adults in certain seasons. During the pupping season, only adult males and immature seals of both sexes were killed in the open water of tide-rips within the fast ice; the adult females presumably were occupied with their pups at this time. The adults are about equally represented on the fast ice, following the breeding season. Four adult females and 11 adult males were taken in Foxe Basin, some distance from the favoured breeding areas, in the autumn of 1955. Most of the adult females would be pregnant at this time, and they may have been attracted to the more suitable areas in anticipation of the requirements for pupping in the following spring. The immature seals are probably unaffected by the sexual urges and requirements which cause segregation of the adults. Of seals younger than 6 years old, 190 males and 165 females were killed, a ratio of 53.5 males to 46.5 females. Sleptsov (1943) found a sex ratio in this species of 40 males to 60 females, but concluded that the preponderance of females was due to sampling error and that the true ratio is close to 50:50.

# LOCAL MOVEMENTS, DISTRIBUTIONS, AND FACTORS OF ABUNDANCE

### Reproductive Ecology

The formation of the birth-cave has been described on page 55. As far as is known, this lair is never found outside the fast ice. The Eskimos believe that if a section of this ice is broken off around the time for pupping, any pregnant seals will desert the area for another patch of fast ice.

The young may not be completely independent until they are more than two months old, this long period of parental care probably being a reflection of the habit of pupping in the fast ice. Bearded seals and harp seals, both of which bear their young in moving and unstable pack ice, wean the young after a much shorter period of parental care. Obviously, on fast ice, the mechanics of attending the young are simplified and the dangers of separation of mother and young are lessened. These circumstances probably enable the female to feed normally in the vicinity of the young seal until the June season of intensive fasting (page 21). The female harp seal, on the other hand, loses much weight through lactation and starvation, during the shorter, more intense suckling period.

Of course, any effect of pupping habits will apply equally to the distributions and movements of adult males and females. The proximity of pupping and impregnation times indicates that the adult males must, in general, follow the patterns set by the adult females.

THE AMOUNT OF FAST ICE. Due to the habit of pupping in the fast ice, the total ringed seal production in a region is likely to be in some way proportional to the amount of fast ice available for the birth-lairs. In southwest Baffin Island, as in most areas of the eastern Canadian arctic, the extent of fast ice in winter is more or less fixed by the nature of the coastline and is much the same from year to year. Figure 21 shows the outer limit of fast ice. Obviously different sections of this coast differ markedly in the amount of fast ice which forms. Areas of fast ice, in ten-mile strips extending inshore from the ice edge, have been determined from the topographic map of the region. During the pupping season, coastal sections A, B, C, D, and E (Fig. 21) have approximately 43, 48 165, 193, and 181 square miles of stationary ice respectively. Thus a strip from the ice edge in the relatively complex areas of C, D, and E delimits about four times the amount of fast ice of an equivalent strip in the areas A and B.

ICE STABILITY AND SNOW COVER. The total amount of fast ice is only one of the factors determining the suitability of an area for pup production. In regard to the formation of fast ice, several generalizations may be made. Ice forms from the shore outwards. The deepest inlets and most sheltered, islandfilled regions are frozen earliest and most securely. Islands act as anchor-points in the formation of stable ice in offshore waters. The most recently formed ice, usually nearest the ice edge, is the least stable, and is more subject to the destructive effects of winds and tides. These factors indicate the greater suitability of areas of coastal complexity for stable ice formation. If the distribution of pups is considered to be randon throughout the fast ice, and if a strip of ice, one mile in width, is broken off at the ice edge from A to E (Fig. 21) carrying all its pups to death through separation and starvation, the resulting losses in the year's pup production would be about 22% in areas A and B and only about 5.5% in areas C, D, and E. This example, involving pup production as a function of ice stability, is considerably oversimplified. Nevertheless, there are several facts which suggest that ice instability has its effect on the structure of seal populations.

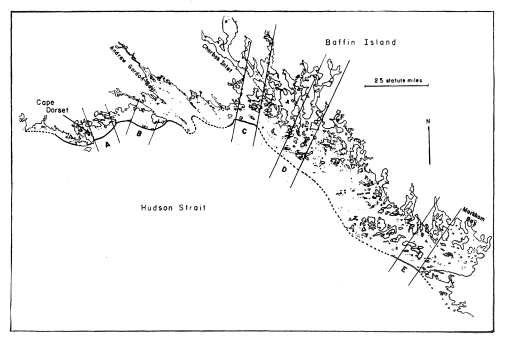


FIGURE 21. The limits of fast ice in southwest Baffin Island. Observed segments of the ice margin are indicated by a solid line; the dotted portions joining these are estimated approximate positions.

Not infrequently, young seals showing obvious signs of malnutrition are killed by the Eskimos of southwest Baffin Island. Two seals of this sort were killed by the natives and observed by the author. One weighing 19.5 lb. (8.8 kg.) was killed at the ice edge on July 1, 1954 (Fig. 22). The other, which weighed 20 lb. (9.1 kg.), was killed well offshore on August 6. Five other seals of the year killed during the summer season in southwest Baffin Island ranged from 28.5 to 38 lb., mean 33.2 lb. The Eskimos call these small seals *sierkolik*, the term referring to the prominence of the knee joint, and presume them to be starvelings which had been abandoned by the mothers before being properly weaned. The

Eskimos have found the *sierkolik* on large floating ice pans off the ice edge in the spring. Whether the young seal has been born, perhaps abortively, on moving ice, or whether the ice has been separated from the fast ice-sheet after the birth of the young, is not known. The latter possibility seems to be more likely. The majority of starvelings are taken by natives of southern Andrew Gordon Bay and from camps to the west of Cape Dorset (Fig. 21). In June, when the bearded seal is hunted from boats launched at the ice edge, the *sierkolik* is taken in the open water. In this same season, young ringed seals of the year, in the peak of condition, are being taken by Eskimos hunting on the fast ice.



FIGURE 22. A starveling, July 1, 1954, on southwest Baffin Island.

There is evidence of an average size difference between seals produced in the peripheral ice and those taken in areas where the ice is stable until late in the spring. The skins of first-year seals, which are known as "silver jars" in the fur trade, are of some commercial value. In this respect, the skins taken by the natives of camps situated deep within the archipelago to the east of Cape Dorset are considered to be the best. Figure 23 shows one of these larger, more valuable skins, taken from near the camp marked F on Figure 21, compared with a skin taken in the same season (early August) near Cape Dorset. The smaller skin was the intermediate in size of 3 skins from the Cape Dorset area, and the larger was close to the mean in size of 7 skins from its area. The largest of the 3 skins from the Cape Dorset region was about equal in size to the 2 smallest skins from the camp to the east. A possible cause of such a size difference of pups lies in the different periods of parental care which are expected in these two regions. In northwest Foxe Basin, the same differences of white-coat and "silver jar" size were suggested by a few specimens and confirmed by statements from the natives. In the areas around Igloolik Island, close to the winter ice edge, the young are small. Around Fury and Hecla Strait, Richards Bay and Murray Maxwell Bay, all areas of stable winter ice and far from the ice edge, the young are larger (H. N. Andersen & E. H. Grainger, personal communications).

Premature separation of mother and young is not the only danger in areas of unstable ice. Occasionally the natives find pups in pressure-ridge lairs (page 57) apparently crushed to death by shifting ice. High winds and tides, and the resulting ice pressure and movement, would have their greatest effect at the periphery of the fast ice.

A sufficient depth of snow is required for the construction of the birth-lair. Some mechanisms whereby the female may "predict" the snow depth of a localized area have been discussed on page 57. A good depth of snow is probably required for its insulatory effect, for pups born too early in the spring season run the risk of death by freezing. A small (59 cm. 23.2 in.) white-coat was found dead in its lair by E. H. Grainger (*Calanus* field-notes, 1956) on March 14, and the natives of Foxe Basin informed him that this was a common fate of young born in March. The amount of snow is directly related to the time of formation of the fast ice; it is also governed by the amount of shelter and windbreak. It is easily seen that snow depth is a function of the same forces that control ice stability.

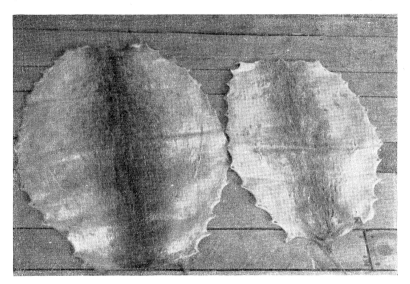


FIGURE 23. Comparison of a large first-year skin of *Phoca his/pida* from the complex coast, and a small first-year skin from the simple coast of southwest Baffin Island.

In Greenland, many young lose their lives in bad weather, according to Hansen (no date), and bad weather may also sometimes cause the mother to abandon her young. Clearly a pup born on an unmoving substrate, under a thick blanket of snow, is little affected by the vagaries of wind and weather.

These features of fast ice—stability and snow cover —result in a much higher production potential in the coastal complex between Chorkbak Inlet and Markham Bay than on the relatively simple coasts of the region near Cape Dorset. Thus, the ratio 4:1 between amounts of fast ice in areas C and A (Fig. 21) will be much smaller than the ratio in seal production in these two regions. ICE CONDITIONS AND THE AGES AND SIZES OF SEALS. As a general rule, the more suitable the ice of an area for pupping purposes, the greater the average age of adult seals making use of this ice. Young adult and newly matured seals are found in the regions of coastal simplicity and in the peripheral ice of more complex coastlines. Older seals are proportionately more common in the innermost bays and lagoons of ramified coastlines. This pattern suggests that penetration of the most suitable breeding areas is a function of experience with age. Kumlien (1879) observed six instances of "young" females pupping on the ice with no snow cover whatsoever. "The young exposed in this manner almost always fall prey to foxes and ravens before they are old enough to take care of themselves." Another factor which may affect this age-cline is population pressure, where the age of the seal might determine the success of competition for the most suitable areas.

Residents of the Canadian arctic, both white and Eskimo, are quite familiar with these differences in seal populations. The names "bay-ice seal" and "seaice seal" are used to differentiate the larger and smaller types of seal. Several authors, of whom Freuchen (1935) is the most specific, have commented on these matters. Freuchen found great differences in the size of adult seals "in the Upernavik district, where on the outer coast, for instance at the colony of Upernavik itself, or at the trading post of Taiussaq, one finds full-grown seals that are no longer than about one metre, whereas in near the glaciers, or up in Melville Bay, there are seals of two metres or more." In this he makes "due allowances for the fact that the seal . . . requires about five years to become fully developed"; but since he did not have available an accurate method for age determination, it is probable that these size differences may be, in fact, largely based on age.

In southwest Baffin Island the Eskimos are convinced, through observation, that "bay-ice seals" average larger regardless of age. The largest seal measured by the author in this area was a male, 25 or more years old, 140 cm. (55 inches) long, and 192 lb. (87 kg.) in weight. The innermost bays and lagoons at the north end of the area D of Figure 21 are said to harbour many such large seals. Unfortunately, time did not permit a trip to this area in the breeding season of 1954. In early September many very big seals were seen there, but these were too wary and none was killed. It was felt that if the seals of the innermost bays were larger, the same phenomenon might be reflected in differences between seals of simple and complex coastlines. Unfortunately, measurements of seals from the complex coast, the area between Markham Bay and Chorkbak Inlet, are too few to demonstrate possible differences. However, Figure 24 does suggest that the average sizes within the same age-class do differ in the two regions.

Perhaps the most satisfactory explanation of this situation revolves around the ice ecology and care of the young, which have been elaborated above. Summer starvelings are more common on open coasts, and there may be an average size difference in summer yearlings of simple and complex coasts. The ringed seal is not truly migratory and it is probable that a given section of coastline is replenished to a large extent by its own residents. If small young are the rule, the entire population may bear this stamp. Actually, as will be pointed out in a later section, dispersal is probably directed from complex to simple coasts. Thus, if the size at weaning is of significance, a wider range of seal size might be expected on simple coasts. There is a suggestion of this wider range in Figure 24.

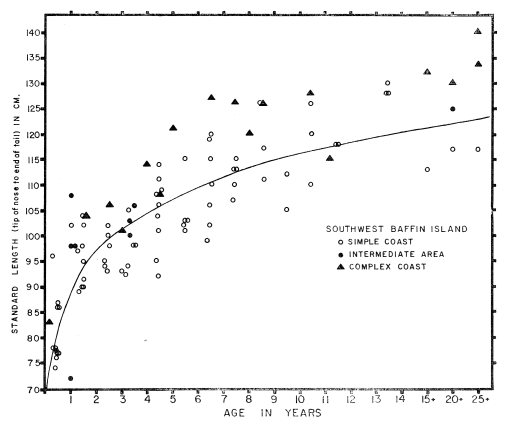


FIGURE 24. Size and age of *Phoca hispida* of southwest Baffin Island. The curve is drawn only through the simple coast specimens, with the aid of several straight lines of least squares.

Other factors may enter in: first, if there is competition for the best breeding sites, the success of the competitor may depend more on size than on age; secondly, it is not known if the age and size of the mother have an effect on the size of the young at birth. As regards the latter possibility, Pedersen (1930) suggests that old female ringed seals give birth about two weeks later than younger females, on the average. Direct correlation between the age-structure of a population and the size of the young produced would be valid only where all the young sampled were still under parental care. Many "silver jar" skins taken in these circumstances were shipped out of Cape Dorset before the problem presented itself. After weaning, nutritional factors, which may have an individual effect, are not likely to act at the population level, especially as the young, fast-growing seals from both the simple and complex coasts of southwest Baffin Island are almost all feeding in the same offshore waters, on the planktonic *Themisto* (see page 17).

Over its wider range, in the Canadian eastern arctic, the ringed seal shows the same variations in size. Latitude, as it governs the formation of fast ice, is superimposed on the effect of coastal complexity. Thus, the common belief among inhabitants of the north that seals in the high arctic areas are larger than those of the south has some basis in fact. Soper (1944) considers that a cline of seal size is to be found from southwest to northeast Baffin Island. An examination of his original field-notes (*idem*, 1924-25 & 1928-29) indicates that this cline may be partly a result of sampling variability; the coastlines on which his observations were made show a corresponding increase of complexity, which would effect the age structure of the populations.

Seals of known length and determined age are available from a number of localities, through the courtesy of several field workers, for comparison with the southwest Baffin Island material. The growth curve of Figure 24 is fitted by eve to the standard lengths of the simple-coast seals of southwest Baffin Island. This curve is used as a standard for comparison with seals from other areas in Figures 25, 26, and 27. The Southampton Island sample, composed chiefly of seals from the South Bay area, is from about the same latitude as southwest Baffin Island. Ice conditions here are comparable with conditions on the simple coasts of southwest Baffin Island and the seals exhibit approximately the same average size with respect to age. The two seals from Churchill, Manitoba, give little evidence of the smaller size of seals which would be expected, due to the unsuitability of ice conditions along the smooth, open coastlines of southwest Hudson Bay. Both these seals were adult females, which were not pregnant in February, another indication of poor conditions for breeding. Ice conditions in northwest Foxe Basin are excellent for care of the young, and the fast ice may not be completely gone from the harbour at Igloolik until the end of July or later, whereas at Cape Dorset Harbour on southwest Baffin Island, and Coral Harbour on Southampton Island, the ice has gone by early July. The seals from northwestern Foxe Basin are correspondingly larger. In northern Baffin Island and Ellesmere Island, where the complex fiord-coasts and lower mean temperatures result in conditions of stable, long-lasting ice, the seals are considerably larger. Only 3 of the 29 seals from these regions are smaller than the mean sizes determined for the simple-coast seals of southwest Baffin Island.

It cannot, of course, be stated definitely that the size differences between northern and southern populations as shown in Figures 24 to 27 are entirely environmentally imposed. However, the facts suggest that the control of size through the mechanism of ice conditions is the overriding cause. Genetic differences can hardly be postulated to explain the size differences in the continuously distributed population inhabiting the 135 mile coastline of southwest Baffin Island, and it is reasonable to suppose that genetic differences need not be invoked to explain a similar north to south cline in seal size. The well-known

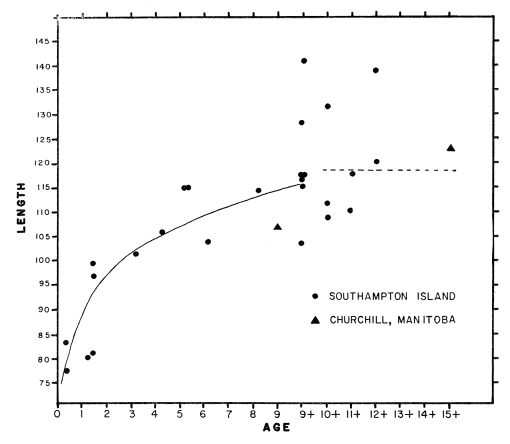


FIGURE 25. Size and age of *Phoca hispida* of Hudson Bay. The curve is for southwest Baffin Island specimens, as in Figure 24. As the older seals from Southampton Island are all claw-aged, the curve is not extended beyond 10 years; the dotted line is drawn through the mean size of 10 to 25 + years seals from Figure 24.

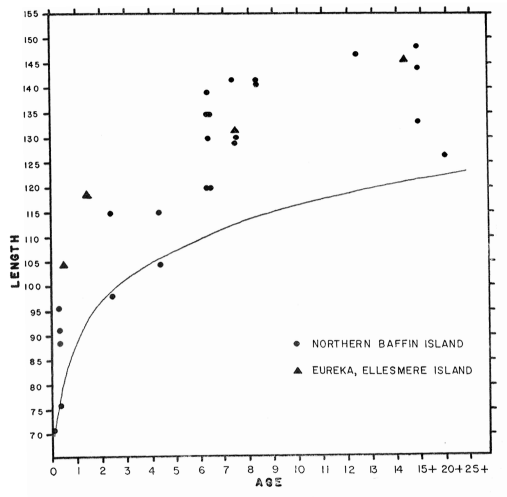


FIGURE 26. Size and age of *Phoca hispida* of two high arctic localities. The curve is for southwest Baffin Island specimens, as in Figure 24.

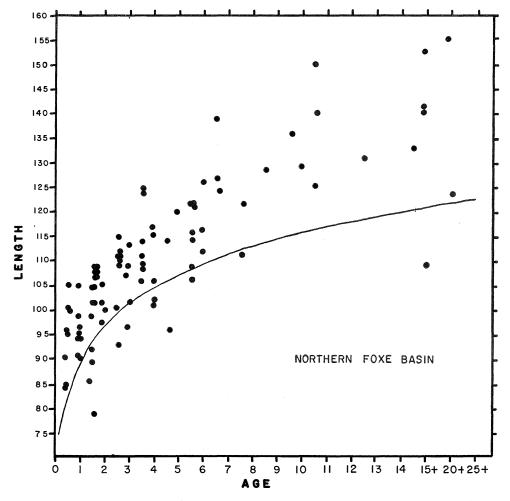


FIGURE 27. Size and age of *Phoca hispida* of northern Foxe Basin. The curve is for southwest Baffin Island specimens, as in Figure 24.

advantage of larger body size in homeothermic animals living in cold latitudes (Bergmann's rule) would not be very effective among ringed seals which live in relatively uniform temperatures throughout their range.

#### MOVEMENTS

The sedentary habits of the adults. There is a general tendency for the mature seals to become year-round residents within the areas delimited by the winter fast ice. This sedentary habit is apparently a reaction to the selection imposed by ice quality on reproductive success. Thus, even in the summer season adults are concentrated in bays, fords and among islands, while the immatures live a more offshore existence. Johansen (1910) and Pedersen (1926, 1930 and 1942) found this phenomenon very marked on the ford coasts of East Greenland. Here, in the offshore waters, "it is the small seals (the young 'troll' seals and a little older animals) which are met with . . .; the full-grown animals, which are so common the whole year through in the fjords, are not seen at all out here" (Johansen, 1910). Pedersen (1930) considers the young seals to be very rare within the breeding areas, and he even suggests that the young are dependent on the breathing holes of the adults when they find themselves in the bays at the time of winter ice.

THE PATTERNS OF ANNUAL MOVEMENTS. In general, the adult ringed seals are inhabitants of the inshore waters; the immatures prevail offshore. The seasonal and geographical variations in this distribution seem to constitute an overall pattern of small-scale annual migration. Figure 28 illustrates this pattern on two simulated coastlines, drawn respectively for simplicity and complexity of form.

Figure 28A shows the picture at the height of mid-winter. Note the preponderance of old adults in the inner bays of the complex coast, the presence of younger adults in the peripheral ice, and the scarcity of young animals in the fast ice.

Figure 28B shows the pattern of distribution at the time of break-up, in late June, on southwest Baffin Island. The majority of immature seals have moved from the open water to the periphery of the fast ice. Johansen (1910) states that young seals in the offshore East Greenland pack ice are never seen lying out on floating ice pans. In fact, the author has only rarely seen this species on small ice pans in the Canadian eastern arctic, although on July 14, 1954, 27 animals were counted on a very large ice pan, measuring about one mile by three-quarters of a mile, which had been recently torn from the ice edge. It would appear that the ringed seal prefers to haul out on fast ice. The movement of young seals into this ice was most marked in southern Andrew Gordon Bay (see Fig. 21) after mid-June of 1954. On June 12, and for several days afterwards in this area, the weather was chilly and occasionally sunny; the ice remained relatively solid. А few seals were seen on the ice at this time, and a number, chiefly young adults, were taken. From June 17 to 20, days of warm sunshine and warm rain, the

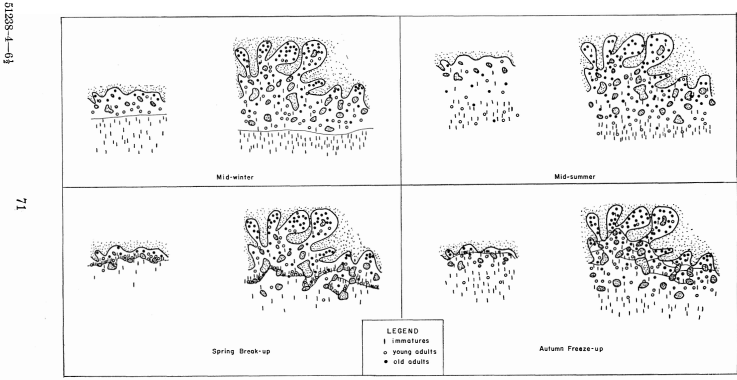


FIGURE 28. The patterns of annual movement of Phoca hispida. Explanation in the text.

ice underwent considerable change. The extensive and deep pools of melt-water dispersed, and the ice became rotten with sink-holes and fissures. At this time large numbers of young seals began to haul out. Most were close to the ice edge, though some had penetrated considerable distances through leads in the ice. Several offshore whaleboat trips in late June and early July produced very few ringed seals, and few of the species were seen. This movement of the immature seals into the fast ice in spring was also observed in Foxe Basin in the spring of 1955 (E. H. Grainger, personal communication) and in northern Baffin Island in 1954 (Ellis, 1957).

Figure 28C shows a return to the "normal" distribution in mid-summer. The young seals have moved offshore, apparently following the natural tendency which had been overcome by a desire to haul out on the fast ice in the spring. In the simple coast area of Cape Dorset, seals of all ages were rather scarce in the inshore waters in summer; the most profitable hunting was carried out in the waters a few miles to the southeast of the Cape. This was not so in the more complex coast to the east. In early September large numbers of seals of all sizes were seen in the waters around the camp marked F in Figure 21. Two seal counts in this area on September 7 and 9 resulted in 57 seals being seen in approximately 28 square miles of water. These figures are minimal, since great care was taken not to recount individual seals. Several immature seals and young adults were killed in these waters, but none of the warier older seals was taken. No very large seal was seen on the return trip to Cape Dorset, outside the archipelago.

Figure 28D shows the population status at the time of freeze-up. The author was unable to observe the pattern at this season in southwest Baffin Island and the figure is based on native information. There is a movement of young and old seals into the coast in the fall, but the movement of young seals is not as pronounced as it is in the spring. According to the natives, the fall movement of young seals is analogous to the spring movement, where the desire to haul out on the fast ice seems to be the chief cause. As the nights lengthen and the days become cold, fewer seals are seen sunning on the ice. Most young seals move out with the progressing ice edge, leaving the adults in their winter residence under the fast ice. Thus the winter pattern is established and the distribution remains relatively static until spring. It has been suggested by the natives living in the complex coast area that some seals may return from year to year to the same winter residence. Pupping-lairs may be found at the same point each year in this area, along well-established dog-team routes, and seals wounded by bullets of known peculiarities have been killed later in the same locality.

There are two main differences between the patterns of movement on simple and complex coasts. First, a higher concentration of immature seals is found, both summer and winter, in the inshore waters of complex coasts; secondly, seals of all ages, including adults, are less common in the inshore waters of simple coasts in summer. No further basic assumptions are required to explain these differences. It seems reasonable to assume that coastal complexity itself has a modifying, damping effect on seal movement and dispersal. Thus, for example,

the coastal area where large numbers of immature seals were seen (near F, Fig. 21) was actually a maze of bays, lagoons and islands; in fact, every entrance to this area is probably cut off by bars and overfalls at low tide. It is thus probable that many young seals find themselves trapped in such circumstances, even when the instinct to escape to the open sea is strong. The fast ice in such complex areas forms rapidly and extensively, cutting off large expanses of open water in its progression. Little chance of avoiding this ice would be offered to young seals living in the area at freeze-up. Immature seals living in the inshore waters of simple coasts in autumn would find it easier to avoid the fast ice as it crept more slowly outward. The same reasoning may apply to the summertime abandonment of inshore waters by adult seals of simple coasts, keeping in mind the fact that adults are inclined to be sedentary. Random movements of dispersal, even on a small scale, would disrupt any tendency for the adults to be confined to bays in the summer. In effect, once a seal has left a bay in the Cape Dorset area, it is all but offshore; when a seal has left its particular breeding-bay in the complex coast, it may be still perhaps 30 miles from the open sea. Another possibility is that the suitability of an area for pupping and the age and experience of the adult seal have effects on the tendency of a given seal to remain sedentary in the immediate area of its past choice, and it has been shown that the adult population of the simple coasts is younger in make-up.

In East Greenland the lack of mixing between adult and immature populations is very pronounced (Pedersen, 1942; Johansen, 1910). The coastal complex of southwest Baffin Island is not a ford coast; ford coasts are deeply indented and may harbour a great deal of stable fast ice, but they do not approach southwest Baffin Island in complexity of configuration. Normal seal movements and dispersal may not be damped to the same extent on true ford coasts. For example, if the young seal, in avoiding inshore waters, were sensitive to directional tidal-flow, this mechanism would be of less value in the coastal maze Dunbar (1941) describes a summertime departure of southwest Baffin Island. from the coasts of the Lake Harbour area. This area is, in distance from the more complex coast and in degree of complexity, similar to the Cape Dorset area, where the same seal movements are found. In northwest Foxe Basin the patterns of distribution and movements are in accord with those found in southwest Baffin Island, according to observations made by E. H. Grainger and H. N. Andersen in 1955 and 1956 (personal communications). In fact, the general pattern of movements seems the same throughout the eastern Canadian arctic, with modifications imposed by the local geography.

DISPERSAL. The ringed seal is not truly migratory, and its movements, other than the small-scale patterns described above, seem to represent dispersal from productive centers, the extent and nature of which varies with the area concerned. Dispersal may be directed offshore or along coasts; both types are discussed here.

The largest summer concentration of seals in the Cape Dorset area is to be found between about 3 and 10 miles offshore. Hunting becomes rapidly less productive when the outer limit is exceeded. Farther out in Hudson Strait ringed seals are decidedly scarce, although harp seals may be seen in large numbers at certain seasons. Seal hunting is very poor 50 miles across the strait on the northern Quebec coast. Johansen (1910) found no ringed seals farther than about 80 miles from land; they occur "only rarely in the outer parts of the drift ice, but in the middle and inner parts of this [ice] . . . everywhere on the coasts of East Greenland". Thus the ringed seal is essentially a coastal species.

There is evidence that the populations of areas of poor breeding potential may be maintained, to a large extent, by areas of high seal production along the same coast. It is believed that the highly successful hunting at Cape Dorset and at Lake Harbour to the east is mainly due to the large reservoir of breeding seals in the complex area between these two trading posts. Seals rapidly become less common around the west coast of Foxe Peninsula. This may indicate the limits of dispersal from the east. If many of the seals found around Cape Dorset are actually produced in the production center to the east, then we might expect that the take of seals of the year will show an increase in the Dorset area throughout the period following the pupping season and after the departure of the fast ice. This seems to be the case. Table XII shows the change in the catch near Cape Dorset, west of Andrew Gordon Bay (Fig. 21), through the period of July to the end of September. The September increase is believed to be real, although a  $\chi^2$  analysis of the material in Table XII (contingency table, with Yate's correction for continuity) reveals a 9% probability of such difference occurring by chance. September and October always produce more of the valued "silver jars" in the Cape Dorset area, according to Eskimo information. After mid-July, 17 first-year and 22 second-year animals were taken from the offshore waters near Cape Dorset. Assuming that all the young age-classes are well represented offshore in summer, first-year seals should have outnumbered the second-year animals in this sample, if all the second-year animals were born in the area. The figures therefore suggest that local pup production is supplemented from the more suitable breeding areas to the east. In northwest Foxe Basin, samples from around Igloolik and Skeoch Bay, some distance from the main productive centers around the entrance to Fury and Hecla Straits, produced 15 second-year animals and 9 first-year animals in September and October. The same factors are probably responsible for this unbalanced proportion.

	July	August	September
First-year seals	1 <sup>a</sup>	8	11
Older seals	6	73	38
Percentage first-year seals	14	10	22.4

TABLE XII. Change in the proportion of first-year seals in the catch near Cape Dorset, southwest Baffin Island.

<sup>a</sup> This seal was the starveling shown on Figure 22.

#### POPULATION SAMPLES

Large collections of teeth for ageing from known localities were taken in southwest Baffin Island in 1953 and 1954. In developing the ecological patterns of the ringed seal populations of this area, little reference has been made to these collections; if the reasoning is sound, the teeth should reflect differences in the age-structure of populations inhabiting various localities at different times of the year.

Six populations have been distinguished, chronologically or geographically, and their make-up is depicted in Figures 29 to 31. The few seals taken before April 1 (the birthday of seals for the purpose of these figures) are put into the age-class of the following seal-year. Only seals for which there is definite evidence of source and date of collection are included in the histograms. In the discussion which follows, all seals of the seventh year or older are considered to be adult.

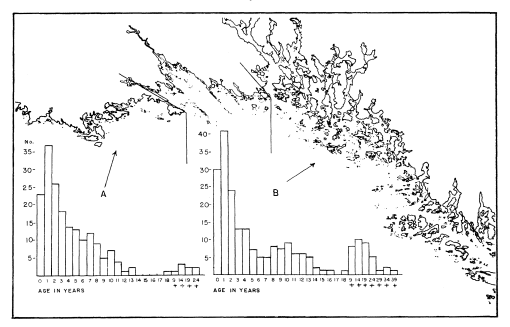


FIGURE 29. Age composition of the total samples of Phoca hispida from southwest Baffin Island.

Figure 29 shows the total samples for 1953 and 1954 from the simple coast near Cape Dorset (A) and from the more complex coast to the east (B). The difference between the adult populations of these coasts is quite apparent. (A) probably gives a fair picture of the structure of population of this coast, as it represents seals taken in all seasons, under all conditions. (B) does not represent a random sample of the immature seals of this coast, as large numbers of the teeth for ageing were taken in the late winter, when most immatures would be excluded from the catch. This, however, does not affect the difference between the adult sections of the histogram; the complex coast sample is considerably older in make-up. Figure 30 compares populations living in the fast ice. The adult population dwelling in the less suitable ice of simple coasts and near the outer periphery of the ice of more deeply indented coastlines is younger in age-structure. The complex coast sample was taken before mid-June, before the large numbers of young seals had invaded the area during and following spring break-up. Thus it is believed that all, or almost all, the young seals shown on this histogram were winter residents. Most of the immature seals shown on the simple coast sample were taken during late June, when these had entered the fast ice from the offshore waters.

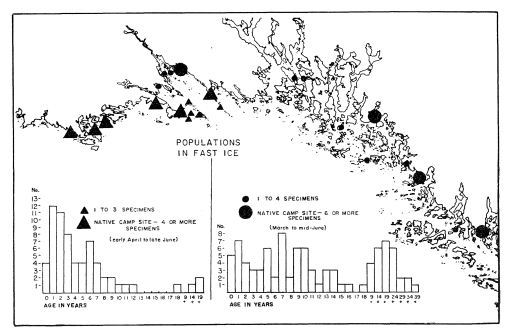


FIGURE 30. Age composition of populations of *Phoca hispida* inhabiting the fast ice of simple and complex coasts of southwest Baffin Island.

Figure 31 depicts the populations living offshore at all fast-ice edges in the winter and off Cape Dorset in summer. The winter sample is exclusively immature. The rapid falling off of adult seals in the older age-classes and the scarcity of first-year animals are both apparent in the summer offshore samples. A sample of open water seals from Foxe Basin in the period of January 31 to April 18 was similarly young in make-up. There were 10 one-year-olds, 9 two-year-olds, 7 three-year-olds, 5 four-year-olds, 2 five-year-olds, and 3 six-year-olds. The only older animal from the fast-ice edge was a sexually active male, 15 or more years old, taken on February 27. Residence in the fast ice, away from the open water, is a response to breeding needs. Neither of two adult females, of 9 and more than 15 years, taken in the open water off Churchill, Manitoba, on February 22, 1954, was pregnant. This unusual condition probably accounts for their presence there. An adult male jaw secured several miles off shore in the

pack ice off Cape Dorset (SD-36, 1954) was in a badly diseased condition. The canines, which were very worn, were decayed through to the pulp cavity. Rings in the cementum, which were not affected by the decay, suggested an age of about 10 years. The presence of this animal outside the fast ice was probably related to this abnormality.

It can be seen from the diversity of these seasonal and geographical samples that it is difficult to obtain a true picture of the total population inhabiting any stretch of coast. For this reason, such features as mortality rates and effect of hunting pressure may be impossible to determine accurately, regardless of the size of the population samples.

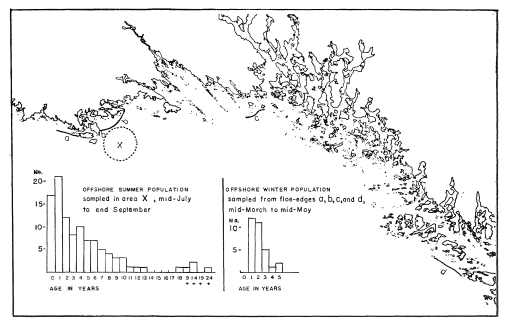


FIGURE 31. Age composition of offshore populations of *Phoca hispida* in southwest Baffin Island in summer and in winter.

DISTRIBUTION FACTORS IN THE EASTERN CANADIAN ARCTIC

The economy of many Eskimo groups is based on the availability of the ringed seal. This animal is scarce, regionally and seasonally, in many parts of its range and the economy of Eskimos living in such localities is often depressed. The previous few sections have dealt with the importance of quantity and quality of fast ice in determining local seal production, particularly in southwest Baffin Island. Ice conditions may well be the key to ringed seal distribution throughout its range.

Temperature, as this affects the formation of fast ice, certainly delimits the southern range of this species. This is undoubtedly the limiting factor in southwest Greenland, where little ice forms south of latitude 67° N and the ringed seal is correspondingly scarce. In Canada, the animal becomes progressively scarcer

south along the Labrador coast, although a few may pup as far south as northern Newfoundland (Fisher, 1950). Even in northern Labrador conditions may not be optimal. Late freezing and light snow cover is found in the fiords of this area and seals are scarce on the bay ice until the late spring thaw, although they may be more common on the ice outside the fiords earlier in the spring (Wheeler, 1953). At Hopedale, at about 55°N on the Labrador coast, an Eskimo who was familiar with conditions on southwest Baffin Island informed the author that few seals give birth on the bay ice of this area. More seals pup on the fast ice of the outer skerries, which, due to the influence of the ice-laden Labrador Current, is formed early and develops a sufficient snow cover.

In the far north the ice may be extremely heavy, and this too may be a limiting factor. Immature seals live offshore in southwest Baffin Island, apparently to avoid the fast ice, but little open water is available in winter to offshore seals in some of the channels of the northern arctic archipelago and in the Polar Sea. Stefansson (1944) found seals in the Polar Sea "in numbers in winter only under such ice as represents water of the preceding season." Pedersen (1942) speculates that the greater abundance of adult seals in active-glacier fords of East Greenland is determined by the presence of leads, fissures and relatively thin ice, which result from the activities of glaciers and icebergs in such fiords.

Local wind conditions may affect the stability of the fast-ice, and springtime river outflow certainly aids in clearing ice from bays and fiords. The latter influence may be of greatest importance on continental coasts. The shore ice may be cleared from the mainland of the western Canadian arctic long before the departure of the heavy offshore pack (Jenness, 1953), and the same is true of Ungava Bay and southwestern Hudson Bay, both areas of relative seal scarcity. The river outflow from the arctic islands is much smaller, and in southwest Baffin Island the offshore waters were navigable before the departure of much ice from the bays in 1954.

Coastal configuration is probably of greatest importance in influencing seal production, for it is the primary control on fast ice in southwest Baffin Island. The east coast of Foxe Basin (see Manning, 1943), the Hudson Strait coast of Quebec, and most of Hudson Bay are relatively unindented and are poor in ringed seals. Frobisher Bay, Cumberland Sound, the Igloolik area, the fiord coasts of eastern and northern Baffin Island, and most of the high arctic islands are all complex in coastline and plentifully supplied with seals. The coasts of Ungava Bay and northwestern Hudson Bay, intermediate in complexity, are intermediate in hunting productivity. Although no exact correlation may be drawn between seal abundance and coastal topography, nevertheless, a connection seems to be established on the basis of known seal distribution.

It appears, then, that population make-up, seal productivity, and abundance are related chiefly to natural geographical circumstances, as these affect the formation of stable fast ice for the purposes of reproduction.

### SUMMARY

1. The material consists of large collections of stomach contents, jaws and claws for ageing, reproductive organs, and body measurements from *Phoca hispida* of southwest Baffin Island and northern Foxe Basin, with smaller collections from other areas of the eastern Canadian arctic.

2. Over 750 seals were aged by the dentinal annuli of the canine teeth. Incompletely calcified dentine, which may be thin and transparent or vacuolar and reflective, is laid down in the spring, generally during the fasting season. It may be that Vitamin D, which has been found to be an active agent in dentine deposition, is required in large quantities for the formation of the densest dentine, although seal fatness may be maintained by smaller amounts of food. The Eskimo method of ageing by pelage is of limited reliability. The claws are marked by alternating light and dark bands, the light band being formed in the spring and early summer; these are useful for ageing seals up to about the tenth year, beyond which wearing makes them unreliable.

3. The nose-tail length increases from about 65 cm. at birth to about 135 cm. in adults, the females being slightly smaller. An extreme adult size of about 5.5 feet (168 cm.) and 250 pounds (113 kg.) is suggested. There is little anteroposterior growth gradient, but the flippers grow less rapidly than the animal as a whole.

4. At least 72 food organisms were detected in stomachs from seven localities. The crustaceans *Themisto libellula* and *Mysis oculata*, and the polar cod (*Boreogadus saida*) were the most common food elements. The seal may dive to about 50 fathoms or 100 metres, but generally turns from inshore feeing on *Mysis*, decapods, or fish to strictly planktonic feeding when in deeper offshore waters, and in southwest Baffin Island its offshore food is almost exclusively *Themisto*. The patterns of feeding suggest that food is not a limiting factor in the distribution and abundance of the ringed seal.

5. There is a general relaxation of feeding in the early spring, but intense fasting does not take place until June and early July, when most seals are basking on the fast ice. Feeding is resumed and blubber is restored after departure of the ice. Not much blubber loss occurs before the basking season, and immediately after the departure of the fast ice, seals killed in the open water are more inclined to sink. There is some evidence that seals living inshore in deeply indented coastlines, where the late departure of the ice may facilitate a more prolonged basking and fasting season, show a greater lag in the return to floating condition in the summer. Annual variations in the departure time of the ice may also affect the proportion of seals which sinks in the summer.

6. The moulting season extends from mid-May to mid-July, with a peak in June. Some abnormal individuals may moult much later.

7. Studies of the testes and bacula reveal that the male does not mature, on the average, until 7 years old. Most females ovulate for the first time, usually in the left ovary, when about 6 years old; 12% were mature in the fifth year, 22% in the sixth year, and 79% in the seventh year. Early ovulations are not as likely to result in pregnancy.

8. The males are in rut from sometime in March until mid-May at the latest, and the females normally ovulate shortly after parturition, while still lactating; the peak of mating activity probably occurs in mid-April.

9. The control of folliculation by the corpus luteum imparts an alternation of ovulation between right and left ovaries. Transmigration of the ovum or blastocyst to the uterine cornu opposite its side of origin may be one of the factors upsetting this alternation.

10. The corpus luteum increases in size and develops more connective tissue and vascularization at the time of implantation of the blastocyst. It regresses rapidly after parturition, freeing the opposite ovary for ovulation.

11. First ovulations do not always result in pregnancy, and it appears that the newly matured female may ovulate outside the normal breeding season.

12. The blastocyst implants, on the average, in early August, after about  $3\frac{1}{2}$  months, delay in development. Embryos may implant over a period of about 45 days, corresponding to the length of the breeding season. The curve of human embryonic growth fits the data on ringed seal embryonic growth very well, when adjusted for birth-size and gestation period.

13. The young seal is born on the fast ice in a lair hollowed out by the female in the overlying snow or in a natural hollow in a pressure ridge near shore. Seals may move into areas of broken ice in autumn or appear at tide-cracks in winter; in both situations drifting results in suitable depths of snow for construction of pupping-lairs.

14. The peak of pup production occurs near the beginning of April, or perhaps somewhat later at higher latitudes. The new-born seal weighs about 10 pounds (4.5 kg.) and measures about  $25\frac{1}{2}$  inches (65 cm.) in standard length. The foetal hair is completely shed in May or June. The females may attend the young seals until the break-up of winter ice, although weaning may occur sooner in most cases.

15. The sex ratio of immatures is about 53.5 males to 46.5 females.

16. The habit of pupping on the fast ice makes ice quantity and quality of primary importance in the reproductive ecology of this species. Coastal configuration governs the amount of fast ice and also the depth of snow cover which is available for construction of the pupping-lair. Ice stability is affected too, and this is important, for pups born on the unstable ice of simple coasts are in danger of premature separation from the females. Starvelings are most common in such areas, and there may be a difference in the average size of first-year seals from complex and from simple coasts. The most suitable ice for pupping

purposes is inhabited by the older adults, and younger adults are proportionately more common in less stable ice. This may result from experience with age, or from competition for more suitable areas. There is a size difference, without respect to age, between seals of simple and complex coasts. This probably results from the larger average size of pups produced on the stable-ice complex coasts. Seals of higher latitudes are considerably larger, probably through the same effect of better ice conditions in more northerly localities.

17. The adults tend to be sedentary in the areas delimited by the fast ice of winter, while the immatures are generally offshore residents. There is a movement of immature seals into the fast ice at break-up, and a less pronounced movement in the same direction at freeze-up. Coastal complexity may exert a damping effect on seal movements, and this may be the factor behind the greater abundance of immature seals inshore on complex coastlines and the smaller numbers of inshore summering adults on simple coastlines. Ringed seals do not disperse very far offshore, and dispersal in southwest Baffin Island seems to be directed from the more productive complex coasts to the simple coasts of lesser production potential.

18. The factors affecting ice formation appear to determine the distribution and abundance of the ringed seal, throughout its wide range in the eastern Canadian arctic.

#### ACKNOWLEDGMENTS

During this study the author has been aided by many individuals and organizations, and he is especially indebted to the following:

1. Dr. M. J. Dunbar of McGill University for directing the problem initially, for offering much useful advice, and for identifying some of the stomach-content Amphipoda.

2. Personnel of the *Calanus* expeditions of 1953, 1955, and 1956 for their aid in collecting material, particularly Dr. E. H. Grainger and Capt. H. N. Andersen, who made valuable collections in the winter of 1955-1956.

3. Several workers who have collected ringed seal material incidental to their own field-problems in the arctic; particularly A. W. Mansfield, who also suggested many improvements after reading the first draft of this paper, S. D. MacDonald, D. V. Ellis, and A. H. Macpherson.

4. Mrs. G. Lubinsky for identifying stomach-content Mollusca from Foxe Basin, and H. J. Squires for identifying some of the decapods from Ungava Bay.

5. Residents of Cape Dorset, N.W.T., especially Mr. and Mrs. N. Ross and Mr. and Mrs. R. Peyton of the Hudson's Bay Company, who provided living quarters during the author's stay at Cape Dorset, and Father F. Trenel of the Roman Catholic Mission, who accompanied the author on several trips and whose services were always available for Eskimo-English translation.

6. The many Eskimos who aided the author in the field, particularly Pootoogok, Paulassie, and Johnniebou of southwest Baffin Island.

#### REFERENCES

- ALLEN, J. A. 1880. History of North American pinnipeds. U.S. Geol. and Geog. Surv. of the Territories, Misc. Pub. No. 12. Washington, 748 pp.
- ANDERSON, R. M. 1942. Two new hair seals from arctic Canada with key to the Canadian forms of hair seals (Family Phocidae). Ann. Rept. Provancher Soc., Quebec, 1942, pp. 23-47.
- AREY, L. B. 1954. Developmental anatomy. W. B. Saunders Co., Philadelphia. 6th Ed., 680 pp.
- BERTRAM, G. C. L. 1940. The biology of the Weddell and crabeater seals, with a study of the comparative behaviour of the Pinnipedia. Sci. Rept. British Graham Land Exped. 1934-37, 1(1):1-139.
- CAMPBELL, N. J., and A. E. COLLIN. MS, 1956. A preliminary report on some of the oceanographic features of Foxe Basin. Fish. Res. Bd. Canada, MS. Rept. Biol. Sta., No. 613, 41 pp.
- CHAPSKY, K. K. 1940. [The ringed seal of the western seas of the Soviet arctic.] Vsesoiuznyi Arkticheskii Institut, Trudy, 145:1-72. Lenningrad.
- DOUTT, J. K. 1942. A review of the genus Phoca. Ann. Carnagie Mus., 29:61-125.
- DUNBAR, M. J. 1941. On the food of seals in the Canadian eastern arctic. Canadian J. Research, D. 19:150-155.
  - 1949. The Pinnipedia of the arctic and subarctic. Bull. Fish. Res. Bd. Canada, No. 85, 22 pp.
  - 1951. Eastern arctic waters. Ibid., No. 88, 131 pp.
  - 1953. Arctic and subarctic marine ecology: immediate problems. Arctic, 6(2):75-90.
- ELLIS, D. V. 1957. Some observations on mammals in the area between Coppermine and Pond Inlet, N.W.T., during 1954 and 1955. *Canadian Field-Nat.*, 71(1):1-5.
- FISHER, H. D. 1950. Seals of the Canadian east coast. Fish. Res. Bd. Canada, St. Andrews Station Circulars, Gen. Ser., No. 18, 4 pp.
  - MS, 1954a. Studies on reproduction in the harp seal, *Phoca groenlandica* Erxleben, in the northwest Atlantic. *Fish. Res. Bd. Canada, MS. Rept. Biol. Sta.*, No. 588, 109 pp.
  - 1954b. Delayed implantation in the harbour seal, *Phoca vitulina* L. Nature, 173:879-880. London.
- FISHER, H. D., and B. A. MACKENZIE. 1954. Rapid preparation of tooth sections for age determination. J. Wildlife Management, 18(4):535-537.
- FREUCHEN P. 1935. Mammals, Part II: Field notes and biological observations. Rept. Fifth Thule Exped., 1921-24, 2(5):68-278.
- HAHN. 1884. De la presence de galets dans l'estomac des Otaries (Otaria jubata). C. R. Soc. Biol. Paris, 8(1).4.
- HAMILTON, J. E. 1939. A second report on the southern sea lion, Otaria byronia (De Blainville). "Discovery" Rept., 19:121-164.
- HANSEN, P. No date. In: Vejdledning i behandling af Grønlandske Produkter. Greenland Administration, 64 pp. (Mimeographed).

- HARRISON, R. J. 1948. On the development and fate of the corpus luteum in the vertebrate series. Biol. Reviews, 23:296-331.
  - L. H. MATTHEWS and J. M. ROBERTS, 1952. Reproduction in some Pinnipedia. Trans. Zool. Soc. London, 109, B:481-512.
- JENNESS, J. L. 1953. The physical geography of the waters of the western Canadian arctic. Geog. Bull., 4:33-64. Ottawa.
- JOHANSEN, F. 1910. Observations on seals and whales made on the Danmark-Expedition, 1906-1908. Medd. om Grønland, 45(2):201-224.
- KUMLIEN, L. 1879. Contributions to the natural history of arctic America made in connection with the Howgate Polar Expedition, 1877-78. Bull. U.S. Natl. Mus., No. 15, 179 pp.
- LAWS, R. M. 1953a. A new method of age determination for mammals with special reference to the elephant seal (*Mirounga leonina Linn.*). Falkland Islands Dependencies Survey, Sci. Rept., No. 2, 11 pp.
  - 1953b. The elephant seal (*Mirounga leonina* Linn.). I. Growth and age. *Ibid.*, No. 8, 62 pp.
  - 1956a. Growth and sexual maturity in aquatic mammals. Nature, 178: 193-194. London.
  - 1956b. The elephant seal (Mirounga leonina Linn.). III. The physiology of reproduction. Falkland Islands Dependencies Survey, Sci. Rept., No. 15, 66 pp.
- MANNING, T. H. 1943. Notes on the mammals of south and central west Baffin Island. J. Mammal., 24(1):47-58.
  - 1944. Hunting implements and methods of the present-day Eskimos of northwest Hudson Bay, Melville Peninsula and southwest Baffin Island. *Geogr. J.*, **103**:137-152.
- MANSFIELD, A. W. (In press). The breeding behaviour and reproductive cycle of the Weddell seal (Leptonychotes weddelli Lesson). Falkland Islands Dependencies Survey, Sci. Rept.
- MELLANBY, M. 1928. The influence of diet on the structure of teeth. *Physiol. Reviews*, 8(4):545-577.
- PEDERSEN, A. 1926. Beiträge zur Kenntnis der Säugetier- und Vogelfauna der Ostküste Grönlands. Medd. om Grønland, 68(3):149-249.
  - 1930. Forgesetzte Beiträge zur Kenntnis der Säugetier- und Vogelfauna der Ostküste Grönlands. *Ibid.*, **77**(5):341-507.
  - 1942. Säugetiere und Vogel. Ibid., 128(2):1-119.
- PIKHAREV, G. A. 1947. [Food of the seal Phoca hispida.] Izvestiia Tikhookeanskovo N. i. Inst. Rybnovo Khoziaistva i Okeanografii, 22:259-261. Vladyvostok.
- PLEKHANOV, P. 1933. [Determination of the age of seals.] Sovetskii Sever, 4(1):111-114.
- RAND, R. W. 1954. Reproduction in the female Cape fur seal, Arctocephalus pusillus Schreber. Proc. Zool. Soc. London, 124(4):717-740.

SCHEFFER, V. B. 1946. How deep will a fur seal dive? Murrelet, 27(2):25.

- 1950. Growth of the testes and baculum in the fur seal, Callorhinus ursinus. J. Mammal., **31**(4):384-394.
- SCHEFFER, V. B., and F. WILKE. 1953. Relative growth in the northern fur seal. Growth, 17(3):129-145.
- SIVERTSEN, E. 1941. On the biology of the harp seal, *Phoca groenlandica* Erål. *Hvalradets* Skrifter, No. 26, 166 pp.
- SLEPTSOV, M. M. 1943. [On the biology of reproduction of Pinnipedia of the Far East.] Zool. Zhurnal, 22(2):109-128. Moscow.

- SOPER, J. D. MS, 1924-25. Report on the Baffin Island Expedition of 1924-1925. (Unpublished field-notes.)
  - 1928. A faunal investigation of southern Baffin Island. Bull. Natl. Mus. Canada, 53, Biol. Ser. No 15, 143 pp.
  - MS, 1928-1929. Explorations in southwestern Baffin Island, Northwest Territories, on the expedition of 1928-1929. (Unpublished field-notes.)
  - 1944. The mammals of southern Baffin Island, Northwest Territories, Canada. J. Mammal., 25(3):221-254.
- SQUIRES, H. J. 1957. Decapod Crustacea of the Calanus Expeditions in Ungava Bay, 1947 to 1950. Canadian J. Zool., 35:463-494.

STEFANSSON, V. 1944. The friendly arctic. Macmillan Co., N.Y. New Ed., 812 pp.

THIBERT, A. 1954. Eskimo-English and English-Eskimo dictionary. University of Ottawa, 174 pp.

VIBE, C. 1950. The marine mammals . . . Medd. om Grønland, 150(6):1-115.

WHEELER, E. P. 1953. Notes on Pinnipedia. J. Mammal., 34(2):253-255.

## APPENDIX

Tables giving body measurements of ringed seals, and details of the food found in their stomachs.

Table I.	Detailed body measurements of 102 Phoca hispida from southwest Baffin Island and	1
	northwest Foxe Basin. The measurements are defined in the text.	

ş	er				th	anus	sn			.			
Age in years	Field number		-	Maximum girth	Axillary girth	) a1	Penis to anus	Foreflipper length	Foreflipper width	Hindflipper length	Hindflipper width	ß	
u v	nu		1 arc	nu	ury	Navel to	Ę	lipi	Idil	diff	lip	les les	ht
е і.	pla		gtl	Ek.	illa	ve	nis	gtl	lth	gtl	lth	ckibt	Weight
Ag	Fie	Sex	Standard length	Bir	Ax	Za I	Pe	Fo: len	Wic	Hi	Hiu wid	Blubber thickness	We
<b>Curr</b>													
			cm.	cm.	cm.	cm.	cm.	cm.	cm.	cm.	cm.	mm.	lb.
Full-term													
foetus	SD-33	М	61		40	16	9	12	7.5	15	18	10	9.8
0.3	SD-267	F	78	72	72	22	n.a.	15	10.5	21	27	18	32
0.3	SD-282	F	74	54	55	21	n.a.	14	9	19	24	17	20
0.4	S-8	M	76	68	65	24			11		27	25	••
0.4	S-10 S-26	F	87 86	68 64	66	24	n.a.	••	11.5		· 28	25	
0.4	S-20 S-49	M F	80 77	65	64 64	23 21	11.5		11 9		30 24	25 28	33 29
0.4	S-49 S-50	г F	77	65 65	64 65		n.a.		11		24 27.5	28 25	29 29
0.4	S-50 S-51	F	86	72	70	19.5 24	n.a.	••	11		27.5	25 32	38
0.4 0.5	S-51 S-55-6	г М	101	86.5	86.5	24 28.5	n.a. 15	 15	11.5	 23.5	29	29	58 61
0.5	S-55-16	F	85	80.3 77	75.5	28.5 26	n.a.	13	11.5	23.5	29	29 27	45
0.5	S-55-17	M	95	82.5	81.5	20	n.a. 16	14.5	11.3	23	27.3	32	43 59
0.5	S-55-32	M	90.5	83	79	28	15	10	. 11	21	27.5	28	
1.0	SD-3	M	102	83 98	88	29.3 31	15			21		43	83
1.4	S-7	F	95	98 79	76	25.5	n.a.	•••	11.5		 29	32	
1.4	S-9	F	81	70	68	23.5	n.a.		11.5		28	25	••
1.4	S-12	M	104	83	83	29	14		14		35	25	63
1.4	S-29	M	98	77	71	27	13		11.5		29	22	48
1.4	S-31	F	92	73	70	25.5	n.a.		11.5		28.5	28	45
1.4	S-47	м	102	79	77	29	14.5		13		32	25	57
1.4	S-48	F	90	77	75	26	n.a.		11.5		30	25	48
1.5	S-55-29	м	99	94	88.5	29.5	17	18	12	19.5	27	37	88
1.5	S-55-33	м	102	83	83.5	30	15.5	16	12.5	23	30	31	
1.5	S-55-51	F	109	97	93	32	n.a.	16.5	14	24	34	43	
1.5	S-55-54	F	105	91	89.5	31	n.a.	15	13	23	32.5	31	81
1.5	S-55-57	M	109	88.5	87	31.5	16	17.5	12.5	25	32	27	
1.5	S-55-66	F	79	75	72.5	24	n.a.	13	10	19	26	32	••
2.4	S-11	M	102	85	83	30	15		14		35	32	••
2.4	S-22	М	93	84	81	28	14	••	11		28.5		••
2.4	SD-482	M	106	91	90	29	15	16.5	14	26	33	26	82
2.5	S-55-11	F	111	94	91	30.5	n.a.	15	11.5	21.5	31.5	30	••
2.5	S-55-53	М	93.5	74.5	72	23	13.5	13.5	11	20	24	30	49
2.5	S-55-55	M	115	96	92	34.5	18	16.5	13	24	31	38	92
3.0	SD-22	М	101	92	90	27	13	••				40	76
3.2	SD-119	F	105	85	83	29	n.a.	19	14	29	32	30	75
3.4	S-32	F	98	80	79	26	n.a.		12		30.5	28	57. <b>5</b>
3.4	SD-479	M	106	91	91	31	16.5	18	15	27	34	29	81
3.5	S-55-3	M	114	96.5	94	33.5	18	18	14.5	28	35.5	42	84
4.4	SD-215	M	95			29	14	16	13.5	24	33	33	63 67
4.4	SD-266 SD-275	F	104	80	81	29	n.a.	17	13 13.5	26 24	34.5 33	32 18	67 66
4.4		M F	108	80 99	78 95	31 28	16	15	13.5 14	24	33 37	18 38	00 90
4.4	S-14	Г	111	99	95	28	n.a.		14	••	51	30	90

TABLE I. Continued

													<u>.</u>
Age in years	Field Number	Sex	Standard length	Maximum girth	Axillary girth	Navel to anus	Penis to anus	Foreflipper length	Foreflipper width	Hindflipper length	Hindflipper width	Blubber thickness	Weight
<u></u>	-		cm.	cm.	cm.	cm.	cm.	cm.	cm.	cm.	cm.	mm.	lb.
$\begin{array}{c} 4.4\\ 4.4\\ 4.4\\ 4.5\\ 5.5\\ 5.4\\ 5.4\\ 5.4\\$	$\begin{array}{c} {\rm S-16} \\ {\rm S-19} \\ {\rm S-33} \\ {\rm SD-496} \\ {\rm S-55-28} \\ {\rm S-17} \\ {\rm S-34} \\ {\rm S-34} \\ {\rm S-39} \\ {\rm S-45} \\ {\rm SD-516} \\ {\rm S-55-15} \\ {\rm S-55-52} \\ {\rm S-55-52} \\ {\rm S-55-56} \\ {\rm SD-296} \\ {\rm S-3} \\ {\rm S-15} \\ {\rm S-21} \\ {\rm S-41} \\ {\rm SD-485} \\ {\rm SD-485} \\ {\rm SD-485} \\ {\rm SD-493} \\ {\rm S-55-9} \end{array}$	M M M F F M M F M M F M F F F F M M M M	<i>cm.</i> 101 106 114 109 114 102 101 103 101 115 103 122 106.5 116 114.5 122 109 121 119 102 115 106 120 115 102 103 122 106 114 115 122 106 114 115 122 106 114 115 122 106 114 115 122 106 114 115 122 106 114 115 122 106 114 115 102 116 114 119 102 115 106 120 115 106 120 115 106 120 115 106 120 127 115 106 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 139,55 130 130 130 130 130 130 130 130	cm. 83 86 88 84 113 80 84 83 76 93 94 102 81.5 99 96.5 103 98 99 99  92 90 89 97 94 107.5	<i>cm.</i> 76 81 88 83 105  82 81 75 89 91 99 81.5 97.5 94 102 96 99 95 86 91 85 86 95 91 103.5	cm.           29         30.5           33         36           35            30.5         28           30         33           33         33.5           31.5         33           30         35           31.5         33           30         35           31         36           37         32           30         30.5           33         34	cm. 15 16.5 18 n.a. 16 16.5 15 n.a. 17 19 n.a. 17 19 n.a. 20 n.a. n.a. 10 13 14 15.5 22	cm.  21 20   18 16 16 18.5 16.5 16 19   19.5 15 20.5	cm.         11.5         13         14         13         12         11.5         13.5         14.5         13.5         14.5         12.5         14         13.5         14.5         15.5	cm.  26 23.5  25 29 25.5 28.5 27.5 28   27 22 29	cm. 32 32 36 33 31 32 33 30 36 32 33 37  37  32.5 38 33 36 33 37  37  32 33 34 35 33 37 37 38 38 39 39 30 30 30 30 30 30 30 30 30 30	mm.  32 33 49  32 28 35 36 48 28 44 48 40 49 44 42 45 32  27 38 44 31	lb.                 1122              54.5              54.5              54.5              54.5              54.5              54.5              54.5              54.5              54.5              54.5 <tr td=""></tr>
6.5 7.3 7.4 7.4 7.5 8.4	S-55-14 SD-274 SD-276 S-36 S-46 S-55-49 S-28	M M F F F M	127 113 107 115 113 112 126	110 94 83 95 98 91 105	108.5 93 81 94 95 91 103	39.5 32 29.5 36 34 31 33	19 16.5 16 n.a. n.a. n.a. 19	18 16 17  15.5 	14.5 14 12.5 13.5 13.5 12 14	30.5 22 22.5  26.5 	 34 30 34 33 33.5 37	55 31 24 42 44 33 44	142 90  91 97.5 
8.4 8.5 8.5 9.4 9.4 9.5	SD-480 SD-523 SD-527 S-55-47 S-20 S-38 S-55-26 SD-672	M M F M F M	126 117 111 129 105 112 136.5	96 97 110 117.5 84 94 119	92 92 103 113 85 88 114.5	37 32 31 36 28 34 40	20 17 n.a. 21 n.a. 1.a. 21.5	18 17 17 17 	15 13 14 15 11 13.5	27 23 25 28  	36 36 32  30.5 32 	26 49 52 52  32 56	104 106 118  81 
$10.3 \\ 10.3 \\ 10.4 \\ 10.4 \\ 10.5 \\ 10.5 \\ 10.5 \\ 11.4 \\ 10.5 \\ 11.4 \\ 10.5 \\ 10.5 \\ 11.4 \\ 10.5 \\ 10.5 \\ 11.4 \\ 10.5 \\ $	SD-273 SD-279 SD-464 S-25 S-55-13 S-55-20 S-55-39 SD-301	F M M M F M	123 118	114.5 111 99	110 95	37 32 35 33 40.5 42.5 37.5 36	n.a. 18 19 17 21.5 24 n.a. 20	20 17 19  19 23 17.5 17	13	29 27 27  30.5 31.5 27 26	37 34 39 33 38.5 39.5 38 35 35	34 33 29 37 54 30 41 31	121 97  100 169 150 
11.4 13.4 13.4 13.4 13.5 15+	S-4 SD-461 S-30 S-40 S-55-18 S-27	F F M F F	118 130 127 127 120 113	 106 105 106 118.5 86	104 104 105 102 113.5 84	36 38 38 38 33.5 33	n.a. n.a. 20 20 n.a. n.a.	 18  18.5 	14 15 14 13.5 14.5 13	28	38 39 36 38 35.5 33	44 33 38 33 52 25	 123 121.5 153 73

 $51238 - 4 - 7\frac{1}{2}$ 

.

TABLE I. Concluded

Age in years	Field Number	Sex	Standard length	Maximum girth	Axillary girth	Navel to anus	Penis to anus	Foreflipper length	Foreflipper width	Hindflipper length	Hindflipper width	Blubber thickness	Weight
			cm.	cm.	cm.	cm.	cm.	cm.	cm.	cm.	cm.	mm.	lb.
16.5	S-55-12	м	153.5	129	127	48.5	25.5	26	16	33		63	214
ca. 20	S-55-31	М	141	122	119.5	45.5	23.5	23	18	30	43	47	
20+	SD-302	Μ	117	102	97	35	16.5	18	13	28	34	29	
20+	S-55-50	Μ	156	131	131	49	24	20.5	16	30	41	52	222
25+	S-37	Μ	117	95	94	38	20	• •	13	• •	34	35	94

TABLE II. Certain measurements of *Phoca hispida*, expressed as percentages of standard lengths. These are derived from regression lines for which standard errors of the estimates are given.

	Navel to anus	Penis to anus	Fore- flipper length	Fore- flipper width	Hind- flipper length	Hind- flipper width
Female:						
Percentage of standard length of 60 cm	28.46		19.41	13.04	26.65	34.86
Percentage of standard length of 160 cm	29.07		12.00	10.46	18.85	26.00
Standard error of estimate	1.79		1.44	0.74	1.68	1.45
MALE:						
Percentage of standard length of 60 cm	27.97	14.57	18.26	13.79	24.83	33.08
Percentage of standard length of 160 cm	30.07	17.05	13.70	10.17	19.42	27.43
Standard error of estimate	1.57	0.97	1.23	0.78	1.59	2.00

.

#### TABLE III. Food specimens taken from *Phoca hispida* of Ungava Bay and northern Labrador in 1947 to 1950.

Origins of stomachs:

- A. Near Inuksilik, approximately 58° 47' N. by 68° 23' W. (1 stomach).
- B. Mouth of Koksoak River, approximately 58° 32' N. by 68° 10' W. (11 stomachs).
- C. Mouth of Leaf Bay, approximately 59° 00' N. by 68° 40' W. (5 stomachs).
- D. Tunnusaksuk Fiord, approximately 60° 15' N. by 64° 30' W. (7 stomachs).
- E. Keglo Bay, approximately 59° 10' N. by 65° 45' W. (1 stomach).
- F. Port Burwell and Forbes Sound, approximately 60° 22' N. by 64° 50' W. (3 stomachs).
- G. Button Islands, approximately 60° 25' N. by 64° 50' W. (1 stomach).
- H. Adlorlik Fiord, approximately 59° 30' N. by 65° 25' W. (1 stomach).
- I. Near Imilik, approximately 60° 45′ N. by 69° 27′ W. (1 stomach).
- J. West of Cape Hope's Advance, approximately 61° 05' N. by 69° 37' W. (2 stomachs).

Food species	Number of specimens	Number of times in stomachs	Number of times dominant	Areas of occurrence
Turritellidae (unidentified)	1	1		G
Limacina helicina (Phipps)	2	1		G
Cephalpoda (unidentified squid beaks).	6	5		J, H, F
Calanus hyperboreas Krøyer	1	1		B
Pareuchaeta norvegica (Boeck)	1	1		F
Mysis oculata (Fabricius)	8535	6		D, F, G, L
Mysis mixta Lilljeborg	2346	21	4	A, B, C, D, F, H
Gammaridea (unidentified)	5	5		B, C, F, H
Orchomenella pinguis (Boeck)	-	1		F
Ampelisca eschricti Krøyer	1	1		J
Haploops setosa Boeck	1	1		F
Ampeliscidae (unidentified)	1	1		F
Amphithopsis longicaudata Boeck	1	1		F
Atylus carinatus (J. C. Fabricius)	1	1		G
Rachotropis aculeata (Lepechin)		1		H H
Pontogeneia inermis (Krøyer)	1	1		B
Weyprechtia pinguis (Krøyer)	1	1		F
Gammarellus homari (J. C. Fabricius).	1	1		H
Gammarus oceanicus Segerstråle	7	4		A, B, I
Gammarus wilkitskii Birula	11	5		A, B, C, D
Ischyrocerus anguipes Krøyer	1	1		F
Themisto libellula (Mandt)		10		A, B, C, D, F, H
Euphausiacea (unidentified)	1	1		I I I I I I I I I I I I I I I I I I I
Thysanoessa inermis (Krøyer)		5	2	A, Č, D
Thysanoessa raschi (M. Sars)		9	4	A, B, C, E, F
Decapod larvae (unidentified)	1	1	···.	C
Natantia (unidentified)	1	2	1 <sup>a</sup>	F, I

\* Dominance as a group; specimens identified to species where possible.

TABLE III. Concluded.

Food species	Number of specimens	Number of times in stomachs	Number of times dominant	Areas of occurrence
Sergestes arcticus (Krøyer)	1	1		D
Pandalus montagui Leach	29	2	1	J, F
Hippolytidae (unidentified)	31	6	1ª	B, E, F, I, J
Lebbeus groenlandicus (J. C. Fabricius).	1	1		В
Lebbeus polaris (Sabine)	5	1		F
Eualus fabricii (Krøyer)	20	4		C, F
Eualus gaimardi (Milne-Edwards)	23	3		E, F
Eualus macilentus (Krøyer)	3	3	· · .	F, J
Spirontocaris spinus (Sowerby)	1	1		F
Argis dentata Rathbun	. 3	1		С
Fish (unidentified)	39	19	5ª	A, B, C, D, F
Boreogadus saida (Lepechin)	2	2	1	В
Reinhardtius hippoglossoides				
(Walbaum)	15	3	1	J, F
Ammodytes sp	55	1	1	В
Small sculpins (unidentified),	7	4	• • •	В, Ј
Triglops pingeli Reinhardt	9	1		В
<i>Triglops</i> sp	4	1		С
Lumpenus fabricii Reinhardt		1	•••	В
Lumpenus sp		2	• • •	В, С

<sup>a</sup> Dominance as a group; specimens identified to species where possible.

91

TABLE IV. Food specimens taken from *Phoca hispida* of southwest Baffin Island in 1953 and 1954. The contents of 153 stomachs, of which 30 were empty, are listed here. The geographical distribution of the dominant food forms is outlined on Figure 5. A plus sign following the number of specimens indicates that, due to examination of stomachs in the field or to subsampling for laboratory analysis, a full count cannot be given.

-			
Food species	Number of specimens	Number of times in stomachs	Number of times dominant
			A
Maldanidae (unidentified tubes)	2	1	· · · · ·
Pelecypoda (unidentified)	1	1	
Astarte elliptica Brown	1	1	
Astarte sp	2	1	
Cephalopoda (unidentified squid beaks and			
fragments)	8	3	
Balanus sp. (shell fragments)	1	1	
Mysis oculata (Fabricius)	6,365+	14	9
Mysis mixta Lilljeborg	7	6	
Mysis sp	350	* 1 ***	
Gammaridea (unidentified)	1	1	
Pseudalibrotus littoralis (Krøyer)	1	1	
Pseudalibrotus sp	4	2	
Ampeliscidae (unidentified)	300	1	1
Haploops setosa Boeck	106	5	
Pardalisca cuspidata Krøyer	1	1	
Calliopius laeviusculus (Krøyer)	3	1	
Gammarellus homari (Fabricius)	2	2	
Gammarus setosus Dementieva	269	4	1
Gammarus oceanicus Segerstråle	1	1	
Gammarus wilkitskii Birula	2	1	
Hyperia galba (Montague)	55	2	
Themisto libellula (Mandt)	18,645+	93	75
Euphausiacea (unidentified)	24+	5	
Meganyctiphanes norvegica (M. Sars)	34	10	
Thysanoessa inermis (Krøyer)	29	5	
Thysanoessa raschi (M. Sars)	328	25	
Thysanoessa sp	64	8	
Natantia (unidentified)	76+	2	1 **
Hippolytidae (unidentified)	13	7	12
Lebbeus polaris (Sabine)	50	8	3
Lebbeus groenlandicus (Fabricius)	15+	7	3
Spirontocaris phippsi (Sowerby)	5	3	
Spirontocaris spinus (Krøyer)	1	1	
Eualus gaimardi (Milne-Edwards)	6	3	
Eualus fabricii (Krøyer)	11	4	
Argis dentata Rathbun	81	3	1

• Dominance as a group; specimens identified to species where possible.

Food species	Number	Number of	Number of
	of	times in	times
	specimens	stomachs	dominant
Sabinea septemcarinata (Sabine).         Hyas coarctatus Leach.         Fish (unidentified).         Boreogadus saida (Lepechin).         Lycodes sp.?.         Small sculpins (unidentified).	$2 \\ 1 \\ 56+ \\ 94+ \\ 1 \\ 35+$	2 1 11 22 1 5	 7 19 1

TABLE IV. Concluded.

TABLE V.Food specimens taken from Phoca hispida of northern Foxe Basin in September and<br/>October, 1955. The contents of 51 stomachs, of which 6 were empty, are listed here. The<br/>geographical distributions of the dominant food forms are outlined on Figure 6. A plus sign<br/>following the number of specimens indicates that, due to examination of stomachs in the field or to<br/>subsampling for laboratory analysis, a full count cannot be given.

Food species	Number of specimens	Number of times in stomach	Number of times dominant
	2		
Polychaeta (unidentified setae)	;	1	
Maldanidae (unidentified tubes)	6	1	
Pelecypoda (unidentified)	1	1	
Nucula tenuis (Montague)	3	2	
Portlandia arctica (L.)	2	1	
Gastropoda (unidentified operculum)	1	1	• • • •
Margarita helicina (Phipps)	6	1	
<i>Fusus</i> sp	1	1	
Buccinum sp	2	1	
Cephalopoda (unidentified squid beak)	1	1	
Balanus sp. (unidentified shell fragment)	1	1	••••
Calanoidea (unidentified)	1	1	
Calanus hyperboreas Krøyer	1	1	
Mysis oculata (Fabricius)	8,933+	26	20
Gammaridea (unidentified)	1	1	
Pseudalibrotus glacialis G. O. Sars	147	9	
Pseudalibrotus nanseni G. O. Sars	70	6	
Stegocephalus inflatus Krøyer	9	2	
Acanthostephia malmgreni (Gøes)	2	2	
Apherusa glacialis (Hansen)	2,252	3	2
Atylus carinatus (Fabricius)	5	2	
Pontogeneia inermis (Krøyer)	12	1	
Gammarus wilkitzkii Birula	489+	15	3
Gammaracanthus loricatus (Sabine)	28+	10	• • • •
Themisto libellula (Mandt)	1,746	16	2
Hippolytidae (unidentified)	6	3	
Lebbeus polaris (Fabricius)	6	3	1
Spirontocaris spinus (Sowerby)	2	1	
Spirontocaris phippsi (Krøyer)	1	1	
Eualus gaimardi (Milne-Edwards)	12	4	
Eualus fabricii (Krøyer)	4	3	
Argis dentata Rathbun	2	2	
Chaetonymphon hirtripes (Bell)	1	1	
Fish (unidentified)	40+	14	5
Boreogadus saida (Lepechin)	66+	14	11
Cottidae (unidentified)	5	1	1
	_	-	_

TABLE VI. Food specimens taken from five *Phoca hispida* of Frobisher Bay in 1948, 1951 and 1952.

Origins of seals:

- A. Estuary of Sylvia Grinnell River, approximately 63° 45′ N. by 68° 36′ W. July 22, 1948.
- B. Ney Harbour, approximately 62° 51' N. by 67° 21' W. August 28, 1951.
- C. Head of Frobisher Bay. September 2, 1951.
- D. Same position as A. July 10, 1952.
- E. Jackman Sound, approximately 62° 15' N. by 66° 18' W. August 19, 1952.

Food species	Number of specimens	Number of times in stomachs	Number of times dominant	Areas of occurrence
Margarita helicina (Phipps)	19	1		С
Limacina helicina (Phipps)	1	1		D
Calanus finmarchicus (Gunnerus)	1	1		E
Mysis oculata (Fabricius)	701	4	2	A, B, D, E
Pseudalibrotus littoralis (Krøyer)	1	1		D
Gammarus sp	1	1		С
Themisto libellula (Mandt)	105	4		A, B, D, E
Thysanoessa inermis (Krøyer)	3	1		D
Thysanoessa raschi (M. Sars)	12	3		B, D, E
Thysanoessa sp.	4	1		D
Lebbeus microceros (Krøyer)	1	1		А
Fish (unidentified)	11	3	2	B, C, D
Boreogadus saida (Lepechin)	1	1		D
Small sculpins (unidentified)	1	2	1	D, E
Icelus sp		1		D

# TABLE VII. Food specimens taken from Phoca hispida of three high arctic localities in 1951,1952 and 1955.

Origins of stomachs:

A. Parr Inlet, approximately 82° 30' N. by 67° 30' W. August, 1951. (3 stomachs).
B. Slidre Fiord, approximately 79° 50' N. by 86° 10' W. August and September, 1955. (3 stomachs).

C. Mould Bay, approximately 76° 20' N. by 119° 30' W. August 23, 1952. (1 stomach).

Pelecypoda (unidentified shell frag.) Cephalopoda (unidentified beaks) Polychaeta (unidentified)	$\begin{array}{c c}2 & 2\\2 & 2\\\end{array}$		A
Cephalopoda (unidentified beaks)	2 2		A
	-   -	1	
Polychaeta (unidentified)			A, B
	1   1		В
Polynoidae (unidentified)	1 1		A
Mysis oculata (Fabricius) 11,	511 6	4	A, B, C
Pseudalibrotus glacialis G. O. Sars	5 2		В
Pseudalibrotus nanseni G. O. Sars	28 2		А, В
Anonyx nugax (Phipps)	480 1	1	A
Socarnes bidenticulatus (Bate)	127 1		A
Apherusa glacialis (H. J. Hansen)	100 1		A
Atylus carinatus (J. C. Fabricius)	49 2		A
Gammarus setosus Dementieva	50 1	1	А
Gammarus wilkitzkii Birula	21 2		А, В
Gammaracanthus loricatus (Sabine)	5 1		A
Themisto libellula (Mandt)	25 2		A, B
Natantia (unidentified)	1 1		В
Lebbeus polaris (Sabine)	1 1		В
Spirontocaris phippsi (Krøyer)	1 1		A
Sabinea septemcarinata (Sabine)	13 4		A, B, C
Small fish (unidentified fragments)	2 2		В
Boreogadus sp. (and larger Gadidae?).	5 1	1	А
Cottidae (unidentified)	1 1		А

TABLE VIII. Food specimens taken from Phoca hispida of Southampton Island in 1955.

Origins of stomachs:

- A. Off Native Point, 63° 41' N. by 82° 38' W. May 5. (1 stomach).
- B. Near Seahorse Point, approximately 63°47′ N. by 80° 10′ W. August 31. (2 stomachs).
  - C. South Bay, 64° 04' N. by 82° 24' W. September 1. (2 stomachs).

Food species	Number	Number	Number	Areas
	of	of times in	of times	of
	specimens	stomach	dominant	occurrence
Mysis oculata (Fabricius) Halirages nilssoni Ohlin Themisto libellula (Mandt) Thysanoessa raschi (M. Sars) Thysanoessa inermis (Krøyer) Lebbeus polaris (Sabine) Spirontocaris spinus (Krøyer) Fish (unidentified)	6603 1 980 122 1 4 3 4	3 1 3 1 1 1 2	1  2 1  1 	B, C, B B, C A, B, C C A A C

TABLE IX.Food specimens taken from two Phoca hispida near the mouth of the Churchill River,<br/>Manitoba, at 58° 54' N. by 94° 10' W.February 22, 1954.

Food species	Stomach 1	Stomach 2
Lysianassidae (unidentified). Hippolytidae (unidentified). Lebbeus polaris (Sabine). Lebbeus groenlandicus (Fabricius). Eualus gaimardi (Milne-Edwards). Eualus fabricii (Krøyer). Small fish (unidentified). Cottidae (unidentified).	1 4 3 47 11 12 2	1 2 1