Immune functions in beluga whales (Delphinapterus leucas): evaluation of natural killer cell activity

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Received in revised form 18 February 1997; accepted 4 March 1997

Abstract

Natural killer (NK) activity, an important non-specific defense mechanism against viral infections and tumors, was demonstrated in beluga whales using two different methods: 51Cr release and flow cytometry. Using the 51Cr release assay, NK activity in belugas was shown to be higher against K-562 than against YAC-1 cell lines. Moreover, it was enhanced by the addition of human recombinant interleukin-2 with both cell lines. NK activity evaluated by flow cytometry in the peripheral blood of eight belugas increased when the effector:target cell (E:T) ratio increased, and averaged 13.9% ± 3.8% (range 9.9% to 17.8%) at an E:T ratio of 100:1. While NK activity could be readily detected using both methods, the lack of radio-isotopes and related laboratory room make the flow cytometric method a viable and safe alternative. The evaluation of this function in cetaceans could lead to a better understanding of the early events that lead to viral epizootics in populations of marine mammals in different parts of the world, as well as to the high prevalence of neoplasms in St. Lawrence beluga whales. © 1997 Elsevier Science B.V.

Keywords: NK activity; Beluga whale; Cetacean; Delphinapterus leucas

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PH S0165-2427(97)00035-4
1. Introduction

Natural killer (NK) cells represent a heterogeneous population of CD3 negative, T-cell receptor negative, large granular lymphocytes that commonly express surface markers such as CD16 and CD56 in humans (O'Shea and Ortaldo, 1992). NK activity, which is mainly directed against tumor cells (O'Shea and Ortaldo, 1992) and virus-infected cells (Biron and Welsh, 1982, Welsh et al., 1991, O'Shea and Ortaldo, 1992, Cook and Splitter, 1989), is not restricted to the major histocompatibility complex (MHC), does not need previous sensitization, and is readily enhanced by interleukin-2 (IL-2) (O'Shea and Ortaldo, 1992, Henney et al., 1981). While NK cells remain difficult to define owing to the heterogeneous nature of their subpopulations, they have been characterized on the basis of their activity for several species including man (Hiseroth et al., 1982), dog (Knapp et al., 1993), cat (Tompkins et al., 1992), horse (Chong et al., 1992), cow (Cook and Splitter, 1989), swine (Salmon et al., 1989), mouse (Welsh et al., 1991), rat (de Jong et al., 1980, Lindsay and Allardyce, 1982), seal (Ross et al., 1996), hamster (Datta et al., 1979), guinea pig (Altman and Rapp, 1978), poultry (Sieminski-Brodzina and Mashaly, 1991, Keller et al., 1992), and fish (Greenlee et al., 1991). NK activity has, until now, not been demonstrated in cetaceans.

Relatively high concentrations of environmental contaminants including PCBs and DDT have been measured in the tissues of many species of marine mammals from different parts of the world (Tanabe et al., 1983). The effects of these ubiquitous environmental contaminants on the health of these animals are still not well known. Numerous epizootics have recently been reported in marine mammals (Geraci et al., 1982, Domingo et al., 1990, Osterhaus et al., 1988), and questions have arisen about the effects of the contaminants on the initiation and spread of these epizootics, possibly through immunosuppression. The importance of NK activity during the initial phase of viral infection (Welsh et al., 1991) as well as the demonstration of impaired NK activity in seals fed with contaminated fish (Ross et al., 1996) make this function particularly interesting. The possibility of deficient NK activity has also been suggested in St. Lawrence beluga whales to explain the large number of tumors observed in this isolated and highly polluted population (De Guise et al., 1994).

In the course of our studies to investigate the effects of environmental contaminants on health of marine mammals, we developed a series of assays to evaluate immune functions in cetaceans. In the present study, we describe quantitative assays to measure NK activity in beluga whales (Delphinapterus leucas) using two different methods: the $^{51}$Cr assay and flow cytometry.

2. Materials and methods

2.1. Animals

Fresh blood from a captive beluga held at the Shedd Aquarium in Chicago, IL, USA, was used for a preliminary study on the technique, which was then applied to cryopreserved leucocytes from wild ranging belugas captured in August 1993 in the
Churchill River estuary, Western Hudson Bay, Canada. The animal from Chicago was a male (Naluark) that also came from Churchill River. The wild animals were 5 males (2 adults, 3 immatures) and 3 females (2 adults, 1 immature). Pairs of belugas in which there were calves (young animals estimated to be younger than 2 yrs old) were not captured for sampling.

2.2. Blood sampling and isolation of mononuclear cells

Blood samples were drawn from the tail fluke (Chicago) or caudal peduncle (Churchill) into heparinized tubes (Vacutainer, Becton Dickinson, Rutherford, NJ, USA). The blood was kept cool until it was used: it was either shipped from Chicago to Bilthoven (National Institute of Public Health and Environmental Protection, The Netherlands) within 24 h, or transported from the field study site on Hudson Bay to the laboratory at Churchill, where it was processed within three hours of collection. A complete blood count was performed for each animal to detect any individual abnormalities. For the evaluation of NK activity, blood was diluted in an equal volume of RPMI 1640 containing 10% fetal calf serum, 100 U/ml penicillin, and 100 μg/ml streptomycin (complete RPMI), and the peripheral blood mononuclear cells (PBMC) were isolated by density gradient centrifugation on Ficoll (specific gravity 1.077, Pharmacia Biotech, Montréal, Què.) for 35 min at 900 × g. PBMC were resuspended in complete RPMI, washed three times, and viable cells enumerated using Trypan blue and light microscope or acridine orange and propidium iodide and fluorescence microscope. The cells were then used immediately or cryopreserved for further use (wild belugas).

2.3. Cryopreservation

For cryopreservation, cells were suspended at a concentration of 1 × 10^7/ml in fetal calf serum with 10% DMSO, and aliquoted in cryogenic vials (Nalgene, Rochester, NY). The cell suspensions were frozen and stored in liquid nitrogen. For subsequent cell culture, the cells were thawed at 37°C, and washed once and resuspended in complete RPMI.

2.4. Target cell lines

YAC-1, a murine lymphoma cell line, and K-562, a human erythroleukemic cell line, both generally recognized as NK sensitive, were used in the evaluation of NK activity. These cells were obtained from ATCC (Rockville, MD) and were cultured in complete RPMI at 37°C with 5% CO₂. The cells were passed at least once, 24 h prior to the assay.

2.5. Cr release assay

Both YAC-1 and K-562 cell lines were washed once in complete RPMI, and adjusted to 1 × 10^7/ml. For labelling, 100 μl of each target cell suspension (1 × 10^6 cells) was incubated for 45 min at 37°C with 100 μCi of ^51^Cr (Amershaw International, Little Chalfont, Buckinghamshire, England). The cells were then washed 5 times with 10 ml of
complete RPMI, and resuspended in complete RPMI at a concentration of $1 \times 10^5$/ml. Beluga whale PBMC were adjusted to $1 \times 10^7$/ml for an effector:target ratio of 100:1, $5 \times 10^6$/ml for an effector:target ratio of 50:1, and $2.5 \times 10^6$/ml for an effector:target ratio of 25:1. For each effector cell concentration, 100 μl was plated in triplicate in round-bottomed 96-well plates for each target cell line (YAC-1 and K-562), and for each time of incubation (6 and 18 h). For each target cell line (YAC-1 and K-562), and for each time of incubation (6 and 18 h), 100 μl of the $1 \times 10^7$/ml effector cell concentration was also plated in triplicate with 200 UI/ml of recombinant human interleukin-2 (rh-IL-2; EuroCetus, Amsterdam, The Netherlands). 100 μl of either $^{51}$Cr-labelled target cell line was added to the effector cells. For each time of incubation (6 and 18 h), 100 μl of either $^{51}$Cr-labelled target cell line was also added to 6 wells with 100 μl of complete RPMI to assess spontaneous release, as well as to 6 wells with Triton 1% (Merck-Schuchard, München, Germany) to assess maximum release. The plates were then centrifuged at $200 \times g$ for 1 min, and cultured for 6 and 18 h respectively at $37^\circ$C 5% CO$_2$. The plates were then centrifuged at $200 \times g$ for 10 min, and the supernatant from each well was transferred into counting vials, and counted 1 min in a gamma-counter. The triplicates (and the 6 samples for the spontaneous and maximal release) were averaged, and the specific release was calculated as follows:

$$\frac{\text{experimental release} - \text{spontaneous release}}{\text{maximal release} - \text{spontaneous release}} \times 100\%$$

2.6. Flow cytometric assay

K-562 cells were used as target cells for a flow cytometric assay slightly modified from Chang et al. (1993). The target cells were washed once in complete RPMI, and adjusted to $1 \times 10^6$/ml. One ml of the suspension was forcefully added to 10 μl of a 3 mM solution of the lipophilic carbocyanine membrane dye 3,3'-dioctadecyloxycarbocyanine perchlorate (DiO, Molecular Probes, Eugene, OR) dissolved in dimethyl sulfoxide (DMSO, Sigma, St. Louis, MO). The cells were then incubated for 20 min at $37^\circ$C in 5% CO$_2$, followed by 2 washes in complete RPMI. The cells were then resuspended in complete RPMI at a concentration of $1 \times 10^5$/ml. Beluga whale PBMC were adjusted to $1 \times 10^6$/ml, and 1000, 500, 250, 125 μl of the cell suspension were put in triplicate in 12 × 75 mm polystyrene test tubes (Falcon, Mississauga, Ontario) for an effector:target ratio of 100:1, 50:1, 25:1, and 12.5:1 respectively. The volume in each tube was brought to 1 ml with complete RPMI. One hundred μl of the labelled target cell suspension were then added to each tube, as well as to six tubes with 1 ml of complete RPMI to assess spontaneous mortality. The tubes were centrifuged at $750 \times g$ for 30 s, and placed at $37^\circ$C in 5% CO$_2$ for 150 min. The tubes were centrifuged for 10 min at $250 \times g$, the supernatant was discarded, and the cells were resuspended in 300 μl of a 200 μg/ml solution of ethidium bromide. The tubes were placed on ice and analyzed by flow cytometry using a FACScan (Becton Dickinson). The beluga PBMC were distinguished from the K-562 target cells on the basis of their fluorescence at 530 nm (FL1), and were excluded from the acquisition using an electronic gate. The viability of the K-562 was assessed on the basis of their fluorescence at 630 nm (FL3). The
triplicates were averaged, and the net NK activity was calculated as the difference between the mortality of the K-562 incubated with effector cells and the spontaneous mortality of the same cells (cultured without effector cells).

2.7. Statistics

The results are expressed as mean ± standard deviation, and comparisons between groups were made using student t-test (p = 0.05).

3. Results

The $^{51}$Cr assay revealed slight NK activity against K-562 cells after 6 h (2.3%), but this activity was clearly enhanced in the presence of rh-IL-2 (16.8%; Fig. 1). Nevertheless, this activity was proportional to the effector:target ratio with higher E:T ratios having the highest specific release. NK activity against YAC-1 cells after 6 h was detected only with the presence of rh-IL-2, and was low (2.0%; Fig. 1). After 18 h, the NK activity against both cell lines was greatly increased, was proportional to the effector:target ratio against both cell lines, and was enhanced by the presence of rh-IL-2 (Fig. 2). The activity against K-562 was higher than that against YAC-1, except in the presence of rh-IL-2 after 18 h (Figs. 1 and 2). Spontaneous release after 6 h was relatively low (14.9% and 10.4% of the maximal release for YAC-1 and K-562 respectively), while it was respectively moderately and slightly increased after 18 h for YAC-1 and K-562 (39.3% and 16.7% of the maximal release respectively).

The results of the flow cytometry-based assay demonstrated that K-562 could easily be discriminated from PBMC on the basis of their fluorescence at 530 nm (FL1) (Fig. 3), and that the distinction between live and dead K-562 was evident (Fig. 4). Average

![Fig. 1. Specific $^{51}$Cr release in a beluga whale as an evaluation of NK activity after a 6-h incubation of PBMC with $^{51}$Cr labelled YAC-1 and K-562 cell lines at different effector:target cell ratio, with or without IL-2.](image-url)
Fig. 2. Specific $^{51}$Cr release in a beluga whale as an evaluation of NK activity after an 18-h incubation of PBMC with $^{51}$Cr labelled YAC-1 and K-562 cell lines at different effector:target cell ratio, with or without IL-2.

Fig. 3. The forward scatter (FSC) vs. side scatter (SSC) scattergram and corresponding fluorescence histogram at 530 nm (FL1) of unlabelled beluga PBMC (A, C) and of DiO-labelled K-562 (B, D) confirms the specificity of the staining for K-562 (larger cells) and not for PBMC.
net NK activity in eight beluga whales against K-562 cell line was proportional to the effector:target ratio, with relatively small variations among the animals (Fig. 5). The NK activity of beluga whale PBMC against K-562 cell line at an effector:target ratio of 100:1 was 13.9% ± 3.8% (range 9.9% to 17.8%). The spontaneous death of the target cells (in culture without effector cells) remained relatively low (11.6%). At an E:T ratio of 100:1, there was no significant difference (p > 0.05) between males and females, nor between adult and immature animals.

Fig. 5. Average NK activity in PBMC from eight beluga whales evaluated by flow cytometry as the percentage of dead K-562 for each effector:target cell ratio, from which spontaneous K-562 mortality was subtracted.
4. Discussion

Our results here represent the first demonstration of NK activity in PBMC isolated from a cetacean species. This natural cytotoxicity was demonstrated against two established NK-sensitive tumor cell lines, the murine YAC-1 lymphoma, and the human K-562 erythroleukemia, and increased in direct proportion to the E:T ratio in culture. Since the activity without IL-2 was higher against K-562 than against YAC-1, K-562 were chosen for further experiments.

NK activity in beluga whales was enhanced with the presence of rh-IL-2. This shows the cross-reactivity of human IL-2 in the beluga, and is consistent with the IL-2 responsiveness observed in other species (Henney et al., 1981), including the harbor seal (Ross et al., 1996). The pattern of increasing cytotoxicity with increasing E:T ratios, the relatively high cytotoxicity against both YAC-1 and K-562 cell lines, and the IL-2 responsiveness of the NK activity suggest that the effector cells responsible for the NK activity in beluga whales have similar properties to natural killer cells for other species.

While the $^{51}$Cr assay is a well-established test to assess NK activity in vitro, the problems associated with the manipulation of radioactive material might discourage its use in many circumstances. Numerous flow cytometric assays have been developed to assess NK activity without radioactive reagents. One method is based on the modification of the morphology of dead vs. live K-562 target cells (Vitale et al., 1989), while others used fluorochromes to evaluate either the mortality (Papa et al., 1988, Zarcone et al., 1986) or the viability (McGinnes et al., 1986, Shi et al., 1987, Chang et al., 1993) of target cells. Results have often been shown to parallel those of the classic $^{51}$Cr assay (McGinnes et al., 1986, Zarcone et al., 1986, Shi et al., 1987, Papa et al., 1988, Chang et al., 1993). The method we chose (Chang et al., 1993) offers the advantage of being easy and efficient, while leading to reliable discrimination of target and effector cells, as well as the further assessment of the mortality in only target cells. While NK activity could be readily detected using both methods, the lack of radio-isotopes and related laboratory room make the flow cytometric method a viable and safe alternative. Using flow cytometry, NK activity was demonstrated in eight belugas with relatively small variations between individuals. The differences in levels of cytotoxicity between the two methods might be due to the differences in incubation times, which differ because of the difference in the events measured by the different assays. While flow cytometry measures the loss of membrane integrity that allows a dye to penetrate a cell, the $^{51}$Cr assay measures the leak of a radioactive isotope from a cell after it has died and its membranes ruptured.

No significant differences were found between males and females, or between adult and immature animals in this study. This is in agreement with Lighthart et al. (1989) who found no difference in NK activity between young and aged humans on a per cell basis.

NK activity was demonstrated for the first time in a cetacean species, the beluga whale, in which it appeared similar to what is observed in other species. The evaluation of this function in cetaceans could lead to a better understanding of the early events that lead to viral epizootics in populations of marine mammals in different parts of the world, as well as the high prevalence of neoplasms in St. Lawrence beluga whales.
Acknowledgements

The authors wish to acknowledge Dr. Jeff Boehm from Shedd Aquarium at Chicago for providing blood samples, and Jack Batstone and his team for help in the capture of belugas in Churchill. This project was funded by US Environmental Protection Agency, Canadian Wildlife Toxicology Fund, Environment Canada, World Wildlife Fund, Washington, Biodôme de Montréal, Corporation professionnelle des Médecins vétérinaires du Québec. SD is supported by a MRC scholarship.

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