Effects of oxygen depletion on the ecology, blood physiology and fishery of the Norway lobster *Nephrops norvegicus*

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ABSTRACT: Biomass, population structure, food selection and blood (haemolymph) physiology of the Norway lobster *Nephrops norvegicus* (L.) were investigated in SE Kattegat, an area where low oxygen concentrations (< 2 ml l⁻¹, 30 % O₂-saturation) have occurred in the bottom water for 1 to 3 mo periods in most years in the 1980’s. During the study period (October 1984 to September 1989) lobster biomass decreased in the area from 10.8 kg h⁻¹ (catch per unit effort) to zero (estimated during the last 12 mo of the investigation). Males contributed on average 78 % of the population density, except in September 1988 during severe hypoxia when a reversed sex ratio was found and females (even berried) dominated (75 % of density). The food of *N. norvegicus* belonged to 4 major groups; crustaceans, echinoderms, molluscs and polychaetes. The dominant species eaten within these groups were also found to be dominants in the benthic infauna. This suggests that *N. norvegicus* are not feeding selectively but taking available organisms indiscriminately. In the field and in laboratory experiments *N. norvegicus* increased blood pigment (haemocyanin, Hcy) concentration in moderate hypoxia (20 to 40 % O₂-saturation), and reduced it in severe hypoxia (10 to 20 % O₂-saturation). At O₂-saturations below 15 % *N. norvegicus* ceased feeding and had empty stomachs. Thus in low oxygen concentrations the lobsters suffer from hypoxia-induced starvation rather than lack of food. Survival of *N. norvegicus* exposed to 15 and 10 % O₂-saturation was 4 wk and 2 to 4 d, respectively. After return to normoxia recovery of blood Hcy concentration was slow, probably due to lack of copper in the diet, which is essential for Hcy synthesis. We consider blood Hcy concentration to be a promising ‘in situ’ biomarker with ecological relevance.

INTRODUCTION

The Kattegat is shallow (mean depth 23 m) and has a strong halocline at 15 to 20 m depth during summer and autumn, which makes the area below the halocline susceptible to oxygen deficiency due to eutrophication. Between 1980 and 1989 oxygen deficiency has occurred in the bottom water during autumn in most years in the southern part of the Kattegat and has increased progressively in space and time causing adverse effects on the benthic macro- and mega-fauna and on demersal fish (Rosenberg 1985, Hagerman & Baden 1986, Rosenberg & Loo 1988, Pihl 1989). Dead and dying Norway lobsters *Nephrops norvegicus* (L.) were first reported by fishermen in the southern part of the Kattegat in October 1985. In 1988 the oxygen deficiency was more serious (O₂ < 1 ml l⁻¹, 15 % O₂-saturation), lasted for a longer time and extended to an area of about 4000 km². Few fish and no lobsters were caught in this area for at least 3 mo during that year.

Investigations on the effects of eutrophication on *Nephrops norvegicus* were carried out between 1984 and 1989. The aims of these investigations were to (1) make field observations on changes and interactions in lobster biomass, population structure and feeding, (2) study changes in blood (haemolymph) pigment concentration in response to hypoxia (O₂ < 20 % saturation) in the SE Kattegat and, (3) evaluate the use of blood haemocyanin concentration of lobsters as an ‘in situ’ biomarker of hypoxia in the Kattegat and Skagerrak and in experimental studies.

AREA INVESTIGATED

A 3000 km² area in the SE Kattegat at 25 to 50 m depth and with a soft sediment bottom was investigated (Fig. 1). The sediment structure in the area is silty-mud with a content of organic matter (ignition loss) of about 13 to 15 % (Rosenberg & Loo 1988). The
Fig. 1. The sampling subareas (N: north, C: center and S: south) in the SE Kattegat in which research trawling for *Nephrops norvegicus* has been carried out at 8 to 12 stations from 1984 to 1989. *N. norvegicus* from Stations 1 to 13 in the northern Kattegat and in the Skagerrak were collected by commercial trawlers for blood sampling, except for Station 9 near the Marine Research Station at Kristineberg, where monthly samples through a year were obtained with lobster creels. The areas 4257 and 4356 are 2 subareas from which commercial catch statistics are collected by ICES (International Council for Exploration of the Seas).

Water mass is characterized by a strong vertical stratification with the halocline normally situated at 15 to 20 m depth. Surface salinity varies between 15 and 25 %o and below the halocline between 32 and 34 %o. During summer stratification is even stronger as a thermocline also develops. Surface water temperature varies from 10 to 20 °C in summer and 0 to 10 °C in winter. Bottom water temperature is 4 to 12 °C, with highest values in late summer and autumn. Tidal amplitude is about 0.1 m (Svansson 1975).

In the Skagerrak *Nephrops norvegicus* were obtained from muddy coastal stations at depths of 33 to 60 m. Bottom water salinities and temperatures were similar to those found in the Kattegat.

**MATERIALS AND METHODS**

**Sampling.** *Nephrops norvegicus* in the SE Kattegat were sampled in subareas N, C, S (Fig. 1) by research vessels using a Glommen lobster trawl (ground rope 86 m, head rope 76 m, sweep 55 m and otter board 2.2 m) with a mesh size of 60 mm in the cod end. Samples were taken in spring (April) and in autumn (September to December) during each year from 1984 to 1989. In general 8 to 12 trawl samples (duration 30 min; speed 3 knots) were taken on each occasion during daytime. Lobsters were counted, weighed, measured (tip of rostrum to telson) and sexed on board.

In the NE Kattegat and the Skagerrak *N. norvegicus* were sampled (Stns 1 to 8, 10 to 13; Fig. 1) by lobster trawl from commercial boats and for experimental work they were creel-caught near the research station (Stn 9, Fig. 1).

Oxygen concentration in the Kattegat was determined on most sampling occasions during 1985 to 1989. Measurements were made every 1 m below, and every 5 m above the halocline. At the bottom the measurement was made less than 0.5 m above the sediment surface. An air and Winkler calibrated Yellow Springs Instruments model YSI-58 oxygen meter was used. On 6 occasions additional Winkler titrations were made simultaneously on samples taken close (10 cm) to the bottom. Deviations between YSI-meter measurements and Winkler titrations were in general < 0.1 ml l⁻¹. In the Skagerrak oxygen was measured only at Stn 9 by Winkler titration (Lindahl pers. comm.).

**Blood, stomach and gill analysis.** *Nephrops norvegicus* caught in SE Kattegat during the period October 1986 to September 1988 were analysed for blood pigment (haemocyanin, Hcy) concentration (only lobsters in the intermoult stage) and stomach content. Twenty specimens (if possible) from each trawl station were analysed. In the Skagerrak blood samples were also taken at 2 stations in the period November 1987 to November 1988. In addition, blood samples were taken from Skagerrak lobsters found dying in creels or when emerging from the sediment and caught in an anchor-seine net. Blood samples were obtained from *N. norvegicus* caught in areas not subjected to low oxygen conditions in the Moray Firth, eastern Scotland, January 1987 (Hagerman & Baden 1988) and from Loch Torridon, western Scotland in September 1988. The latter population of lobsters had never been commercially fished.

Samples of prebranchial blood were taken with hypodermic syringes (0.3 to 0.5 ml) from the arthrodial membrane at the base of the fifth leg as soon as possible after capture. The blood was diluted to double volume with distilled water to prevent clotting, frozen and returned to the laboratory. The influence of freezing for a period of 1 wk was tested by Hagerman & Baden (1988) and shown to give < 5 % reduction of the Hcy concentration. When thawed a 200 µl sample was diluted with 1 ml distilled water in a 10 mm quartz cuvette and the absorbance measured at 339 nm, using a Shimadzu spectrophotometer UV 2100. An extinction coefficient of $E_{339}^{\text{cm}} = 17.26$ was used. This $E_{339}^{\text{cm}}$ value
was calculated from $E_L^{\infty} = 2.83$ given by Nickerson & van Holde (1971) and Antonini & Brunori (1974) on the basis of a functional subunit of haemocyanin with a molecular weight of $74 \times 10^3$.

The weight of the stomach contents (frozen) was recorded and its constituents analysed to species or larger unit. In 1988 the occurrence of dark and black gills was recorded in *Nephrops norvegicus* from both the Kattegat and the Skagerrak.

Experimental analysis of changes in haemocyanin levels related to food and oxygen.

The rate of changes in Hcy in *Nephrops norvegicus* exposed to different food conditions and oxygen tensions were studied in tanks of base area 1 m$^2$, each containing 300 to 400 l of seawater. To prevent turbidity when catching the *N. norvegicus* for the blood sampling, sand instead of mud was used as a substrate and the lobsters were provided with artificial plastic burrows. To mimic the natural light conditions at 40 m depth, the tanks were illuminated by green light using a 9:15 h day/night cycle.

**Food:** Experiments were carried out at different times of the year, and at different regimes. Expt 1 lasted from December 1987 to January 1988 (40 d) with temperatures between 7 and 9.5 °C. Expt 2 lasted from February 1988 to April 1988 (81 d) with temperatures of 4 to 6 °C. As the water was pumped from 40 m depth in a flow-through system the temperature varied as in nature. Each experiment comprised 3 basins containing 4 to 7 *Nephrops norvegicus* of differing size and sex. In one of the basins *N. norvegicus* were starved, while in the second and third basins they were offered shrimps *Pandalus borealis* L. and brittle stars *Amphiura/Ophiura* spp. respectively, daily and in constant and generous amounts (2 g ww per *N. norvegicus*). Surplus non-eaten food was recorded and removed. According to stomach content analysis of field-captured lobsters, polychaetes and molluscs are also eaten by *N. norvegicus*.

**Oxygen:** Two experiments with *Nephrops norvegicus* in hypoxic conditions were carried out in January to April 1989. Four tanks of base area 1 m$^2$ were used, two with 100 % O$_2$-saturation (controls) and two with reduced oxygen concentrations (10 and 20 % in Expt 3; 12 and 15 % saturation in Expt 4). The control series of *N. norvegicus* showed the same percentage change of Hcy in the 2 experiments (these are combined in Fig. 7). Nine intermoult stage *N. norvegicus* were held in each tank. Nitrogen was used to reduce oxygen and air bubbling to provide full oxygen. Gaseous control occurred in 2 m high and 0.3 m wide cylinders separate from the experimental tanks. The water was recirculated, and to reduce accumulation of ammonium, 10 % of the volume was changed every other day. The increase of the oxygen concentration caused by change of water was small and returned to the former value within 1 h. Stable oxygen tensions in the hypoxic tanks were obtained using an oxygen regulator (E. Larsen, University of Århus), connected to an YSI-oxygen meter. The oxygen meter was connected to the outflow of a submersible water pump circulating the water in the tank. A piece of copper wire near the electrode prevented microbial growth on the membrane. Winkler titration was carried out every third day to calibrate the YSI-oxygen meter. The temperature during the experiments was 8 to 10 °C.

Blood samples were taken every 3rd to 14th day. Surplus non-eaten food was recorded and removed. According to stomach content analysis of field-captured lobsters, polychaetes and molluscs are also eaten by *N. norvegicus*. In a pilot test, these prey groups [mainly *Mytilus edulis* (L.) and *Glycera alba* (L.)] were offered both live and dead but not accepted.

**RESULTS**

**Oxygen concentration**

The seasonal variation in the mean oxygen concentration of the bottom water in the study area is shown for the period August 1986 to September 1989 in Fig. 2A. Low oxygen values (O$_2 < 2$ ml l$^{-1}$) were recorded for 1 to 3 mo during autumn each year. The oxygen concentrations generally decreased with increasing depth on all sampling occasions, and the depth distribution is given for subarea C in the period August to December 1988 (Fig. 2B). The study area in the SE Kattegat was divided into 3 sub-areas (Fig. 1) based on the frequency of oxygen deficiencies in the bottom water since 1981 (Table 1). The northern subarea has been affected by low oxygen (O$_2 < 2$ ml l$^{-1}$) during autumn in 6 out of 8 yr, the southern subarea in 5 out of 8 yr and the central subarea only in the last 3 yr. The oxygen concentrations at Stns 1 to 13 are not known, except at Stn 9.
Table 1. Minimum oxygen concentrations (ml l⁻¹) during autumns of 1981 to 1988 in 3 subareas of the SE Kattegat (see Fig. 1). Recordings from 1981 to 1984 by Sundberg & Rydberg (1986) and from 1985 to 1988 are own measurements.

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<th>Center</th>
<th>South</th>
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<td>&gt;2</td>
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<td>&gt;2</td>
<td>1.5-2.0</td>
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<tr>
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<tr>
<td>1987</td>
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<td>1.4-2.1</td>
<td>1.4-2.1</td>
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<tr>
<td>1988</td>
<td>0.7-0.7</td>
<td>0.7-1.4</td>
<td>0.7-1.4</td>
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Biomass

The mean biomass of Nephrops norvegicus found was estimated between 0 and 10.8 kg (wet weight) per hour trawling in the area investigated (Fig. 3). During spring low biomasses were recorded each year. The highest biomass was recorded in November 1984. In 1985 and 1986 the biomass had decreased to between 0.8 and 2.3 kg h⁻¹. During the autumn of these years large quantities of dead lobster (up to 50% of the total catch) were caught by Danish and Swedish fishermen in the area investigated. In the autumn of 1987 a higher biomass (3.3 to 5.0 kg h⁻¹) was observed compared to the 2 previous years. In September 1988 the mean biomass at the 12 stations was 3.5 kg h⁻¹, but in the following 3 mo and in April and September 1989 no lobsters were caught at these stations.

Population structure

The length frequency distribution and the sex ratio of the Nephrops norvegicus population is given for the 3 sub-areas in Fig. 4. The length distribution and the mean total length were similar in the 3 sub-areas on all sampling occasions and there was no clear change in distribution during the investigation. The sex ratio was about the same in the sub-areas during the period 1984 to 1987 where the percentage of males varied from 62 to 93 (mean = 78, SD = 8.2, n = 20). Females in the catch were mainly small and non-berried. However, in September 1988 a reverse in the sex ratio was observed.
in the central and southern sub-areas with males only contributing between 23 and 26% of the total. During this sampling, low (<1 ml l⁻¹, 15% O₂-saturation) oxygen concentrations were recorded, which might have forced the females (even berried individuals) to leave their burrows. In the northern sub-areas only dead _N. norvegicus_ were found on this sampling occasion.

**Catch statistics**

Annual mean of reported catches [catch per unit effort (CPUE), kg h⁻¹] of _Nephrops norvegicus_ in the Swedish fishery was assessed during the period 1978 to 1988 in 2 ICES squares (4356 and 4257) in the E Kattegat (Figs. 1 and 5). ICES square 4356 covers the NE Kattegat and 4257 the SE Kattegat, including most of the area studied in this investigation. During the period 1978 to 1982 an increase in CPUE was observed in both areas, especially 4257, followed by a decrease. The year-to-year variation in CPUE (expressed as coefficient to variation) was calculated to be 47% in 4257, which is about 2½ times higher than in 4356. In 1988 the overall CPUE was dominated by the catch in September, when a high mean CPUE of 10 kg h⁻¹ was recorded, coinciding with the beginning of the hypoxic period when the _N. norvegicus_ are emerging from the sediment. Very low CPUE values (<2 kg h⁻¹) were recorded later in the autumn of that year.

**Lobster diet**

No differences in diet of _Nephrops norvegicus_ were found between the sub-areas, so the data for the SE Kattegat were pooled. All together 35 different food items were found in the stomachs of _N. norvegicus_ (Table 2). This is a minimum number because the food was masticated and difficult to identify. The identified species belong to 4 major groups; crustaceans, echinoderms, molluscs and polychaetes.

The 4 groups were all represented in an average of 46 to 66% of the stomachs (Table 2). _Amphiura_ spp. and _Ophiura_ spp. could not always be separated and were therefore pooled. They occurred together with _Abra_ spp. in a high percentage of the stomachs. Other species which commonly occurred were _Diastylis_ spp., _Nucula_ spp., _Glycera_ spp. and _Pectinaria_ spp. The method of using percentage occurrence only gives an indication of the most important food items. Species such as _Pagurus bernhardus_ (L.) and _N. norvegicus_ and rapidly digested polychaetes will be underestimated. When present in the stomachs _P. bernhardus_ and _N. norvegicus_ represented most of the food taken, whereas small animals such as _Diastylis_ spp. made up only a small proportion. Sometimes black deposits, presumably due to melanosis, were seen on remnants of _P. bernhardus_ and _N. norvegicus_ indicating that they could have been dead when eaten. _N. norvegicus_ were mainly eaten in October 1986 and 1987 after periods of low oxygen in the bottom water.

The variation in frequency of occurrence of the 4 groups during the 2 yr study period (October 1986 to September 1988) was approximately the same with a maximum in April 1988 and a minimum in September 1988 (Table 2). _Amphiura/Ophiura, Abra_ spp. and ostracods occurred in highest percentages in the stomachs during spring 1988. The low figures in September 1988 arose because most of the stomachs were empty at that time. If not empty, the stomachs of many berried females contained eggs from their pleopods and spines from _Brissopsis lyrifera_ (Forbes). These spines were also found among the eggs on the pleopods. The mean ratio between the weight of the stomach contents and the overall body weight ranged from 0.8 to 1.2% during the period October 1986 to April 1988 and showed no seasonal pattern (Table 2). However, in September 1988 the ratio was very small (only 0.1%).

Stomach analysis of lobsters caught near Stns 3 and 4 in the NE Kattegat in 1988 again showed that the diet, as in the SE Kattegat, comprised the same 4 groups: crustaceans, echinoderms, molluscs and polychaetes (Table 2). However, in this area crustaceans were found more frequently in the stomachs compared to the SE Kattegat. The hermit crab _Pagurus bernhardus_ was
scarcer in the stomachs from the NE Kattegat compared to in the SE Kattegat, perhaps because the low oxygen concentration in the SE Kattegat had forced the crabs to leave their shells and thus increased their exposure to predation.

Haemocyanin concentration and discoloured gills: field observations

Kattegat

From October 1986 to December 1988 the blood Hcy concentrations of *Nephrops norvegicus* in the Kattegat were generally low (Fig. 6A). In October 1986 low <0.11 mM) Hcy concentrations were found in lobsters from all subareas. The level of Hcy under normoxic field conditions was found to be between 0.4 and 1.0 mM, e.g. in *N. norvegicus* in Scottish waters (Hagerman & Baden 1988), corresponding to an oxygen-carrying capacity of the blood of 1.0 to 3.0 vol. O₂ % for most decapods (Mangum 1983). In subarea S the Hcy concentration in October 1986 was significantly (p<0.001, NMC test) lower than in the other subareas. However, in December 1986 concentrations in lobsters from subarea S had increased significantly (p<0.001, U-test)
to 'normal' Hcy concentration (>0.6 mM). This was not the case in the northern and central areas, where no significant (p<0.05, U-tests) recovery took place.

In September, October and November 1987 the Hcy concentrations were <0.2 mM (Fig. 6A) and no significant (p>0.05, KW test) difference was found between the subareas in September and October when compared at the same month. In November, however, significant differences in blood pigment were observed between all subareas (p<0.005, NMC tests) with the lowest value in the northern subarea. During the autumn 1987 lobster Hcy concentration increased significantly (p<0.001, KW test) in the northern and central subarea, however not to 'normal' values, while no increase (p>0.05, KW test) occurred in the southern subareas. In the central subarea the increase was significant (p<0.001, NMC test) between September and October while in the northern subarea a significant increase in blood pigment took place between September and October (p<0.005, NMC test) and between October and November (p<0.001, NMC test).

From November 1987 to April 1988, only the Nephrops norvegicus from subarea S showed a slight but significant increase (p<0.005, U-test) in Hcy concentration after return to normoxic conditions in the bottom water (Fig. 6A). Between April and September 1988 the levels of haemocyanin in lobsters from subarea C increased significantly (p<0.001, U-test) to 'normal' level (mean = 0.5 mM), but this recovery did not occur in subarea S. In the northern subarea only dead N. norvegicus were caught in September 1988. Comparing the Hcy levels of N. norvegicus during the same months but from different years, significant differences were found in subareas C (p<0.001, U-test) and S (p<0.01, U-test) from October 1986 to October 1987, but not in subarea N, and from September 1987 to September 1988 in subarea C (p<0.001, U-test) and S (p<0.001, U-test).

**Skagerrak**

The temporal change in Hcy concentrations of Nephrops norvegicus during November 1987 to November 1988 at Stn 9 (45 m depth) in the Skagerrak is shown in Fig. 6B. Monthly mean levels of Hcy concentration were between 0.4 to 1.2 mM. In August and September 1987 oxygen concentrations in the bottom water were around 3.5 ml l⁻¹ (50 % O₂-saturation), decreasing to 2.9 ml l⁻¹ (43 % O₂-saturation) in October. After this decrease in oxygen concentration the mean Hcy concentration in November was 1.1 mM with individual measurements of 1.8 mM. With such high blood pigment concentrations the ventral side of the abdomen was blue and the blood clotted immediately and became jelly-like when exposed to air. The oxygen concentrations in the bottom water varied between 4.5 ml l⁻¹ (67 % O₂-saturation) and 6.7 ml l⁻¹ (94 % O₂-saturation) in the period November 1987 to February 1988. Between November 1987 and January 1988 blood Hcy-concentrations did not decrease significantly (p>0.05, KW test). Between January and February, however, a significant (p<0.01, U-test) decrease in Hcy took place.

After the spring plankton bloom in 1988 the oxygen concentration decreased and reached 3.1 ml l⁻¹ (43 % O₂-saturation) in June. In this spring period the Hcy concentrations of lobsters showed a peak in April, which was significantly (p<0.001, U-tests) higher than both the March and May Hcy concentrations. By the beginning of August the oxygen concentration in the bottom water was down to 2.8 ml l⁻¹ (38 % O₂-saturation). From June to September the Hcy synthesis in N. norvegicus resulted in a significant (p<0.005, U-test) increase in Hcy concentration, with a further significant (p<0.005, U-test) increase in October. After low oxygen in the bottom water (1.4 ml l⁻¹, 21 % O₂-saturation) in October 1988 a significant (p<0.005, U-test) decrease in blood Hcy-concentrations was recorded between October and November.
Table 2. *Nephrops norvegicus*. Percentage occurrence of food items in the stomachs of lobsters from 31 trawl samples in SE Kattegat from 1986 to 1988, and from 4 trawl samples in NE Kattegat in October 1988. Number of stomachs analysed and stomach weight/body weight ratio (%) are given for each sampling occasion.

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<td>4.4</td>
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<td>2.3</td>
<td>0.4</td>
<td>0.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No. stomachs</td>
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<td>19</td>
<td>112</td>
<td>127</td>
<td>9</td>
<td>45</td>
<td>107</td>
<td>77</td>
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<td>1.0</td>
<td>0.8</td>
<td>1.1</td>
<td>1.2</td>
<td>0.1</td>
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In 1987 and 1988 blood pigment analyses of *Nephrops norvegicus* were carried out at Stn 13 (65 m depth; Fig. 1). The mean concentrations were < 0.4 mM Hcy in the 1987 samples, but significantly (p < 0.01, U-test) lower (about 0.15 mM Hcy) in the 1988 samples (Table 3).
Baden et al.: Effects of oxygen depletion on lobster

Fig. 6. *Nephrops norvegicus*. Mean concentration (mM) and standard deviation of blood haemocyanin (bars), in relation to the actual oxygen concentration (lines; mL l\(^{-1}\)) at each sampling occasion. (A) Sampling in the northern (N), central (C) and southern (S) subareas in the SE Kattegat (Fig. 1) between 1986 and 1988. (B) Sampling in the Skagerrak 1987 and 1988 (Stn 9, Fig. 1). Samples with only dead lobsters indicated by (+).

Reports from fishermen

During autumn 1987 *Nephrops norvegicus* were reported in gill-nets at 8 m depth and even in eel pots at 2 m depth near Stn 10 in the Skagerrak. *N. norvegicus* are usually restricted to depths below 25 to 30 m. Blood samples were taken from *N. norvegicus* (n = 52) caught at Stns 10, 11 and 12 (Fig. 1) in October and November 1987. Hcy concentrations were low (< 0.2 mM) in all individuals (Table 3). At Stn 10 Hcy concentrations were just above 0.7 mM in March and April and thus within the 'normal' range. In the northern part of the Kattegat (Stn 3) fishermen reported dead and dying lobsters in creels in October 1987. During November 1987 mean Hcy concentration in lobsters still alive in this area was 0.16 mM. In the autumn 1988 *N. norvegicus* were caught in pelagic herring trawls in the Kattegat, and in the Skagerrak (Stn 8) *N. norvegicus* were caught in unusually high densities in anchor-seine nets. The mean Hcy concentration of *N. norvegicus* in this catch was also low (0.15 mM).

Discoloured gills

Dark or black gills of live *Nephrops norvegicus* were observed from all stations in the Kattegat and from Stns 3, 7 and 10 in the Skagerrak. The percentage of coloured gills in the Kattegat area varied between 10 and 20 % (n = 254) for both sexes, but 73 % (n = 22) of females caught in the subarea S in September 1988 had dark gills. During autumn 1988 some *N. norvegicus* that had black gills also had about 10 mm diameter corroded holes in the carapax (Stns 3 and 7). The dark and black layer observed on the gills from *N. norvegicus* in the Kattegat, according to Dando & Notte (unpubl.), is a deposition of mineral particles of oxidized manganese. These were imbedded in the mucus of the gills under hypoxic conditions. The manganese layer did not seem to disappear in normoxia as the frequency of coloured gills in the spring was about the same as in the previous autumn.

Haemocyanin concentrations: experiments

Changes related to food

The influence of starvation on Hcy metabolism in *Nephrops norvegicus* resulted in an average decrease in Hcy concentration of 0.6 to 0.8 % d\(^{-1}\), corresponding to a 50 % reduction within 66 to 89 d (Table 4A).

In the feeding experiments *Nephrops norvegicus* consumed almost equal amounts of shrimps, *Pandalus borealis* and brittle stars *Amphiura/Ophiura* spp., about 7 to 9 g d\(^{-1}\) per kg body weight (Table 4B). The influence of different food types on the Hcy concentration showed that when feeding entirely on brittle stars, the blood Hcy level decreased in all individuals, whereas when feeding on crustaceans the blood Hcy increased in 50 % of the individuals and decreased in the remaining 50 %. The rate of change in Hcy concentration had a wide variation.

Changes related to oxygen concentration

*Nephrops norvegicus* exposed to severe hypoxia (< 20 % O\(_2\) saturation) showed, after a period of exposure, a sudden and rapid decrease in blood Hcy con-
Table 3. *Nephrops norvegicus*. Blood haemocyanin concentrations (mean, SD) of lobsters from different stations (see Fig. 1) in the coastal NE Kattegat and the Skagerrak. n: numbers sampled from each station.

<table>
<thead>
<tr>
<th>Stn</th>
<th>Date</th>
<th>n</th>
<th>Depth (m)</th>
<th>Mean Hcy (mM)</th>
<th>SD</th>
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<tr>
<td>1</td>
<td>17 Dec 1987</td>
<td>20</td>
<td>36</td>
<td>0.42</td>
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<tr>
<td>2</td>
<td>17 Dec 1987</td>
<td>20</td>
<td>50</td>
<td>0.30</td>
<td>0.21</td>
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<tr>
<td>3</td>
<td>4 Nov 1987</td>
<td>13</td>
<td>35</td>
<td>0.35</td>
<td>0.29</td>
</tr>
<tr>
<td>4</td>
<td>4 Nov 1987</td>
<td>12</td>
<td>36</td>
<td>0.16</td>
<td>0.17</td>
</tr>
<tr>
<td>5</td>
<td>14 Sep 1987</td>
<td>18</td>
<td>45</td>
<td>0.14</td>
<td>0.08</td>
</tr>
<tr>
<td>6</td>
<td>14 Sep 1987</td>
<td>13</td>
<td>40</td>
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<td>0.04</td>
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<tr>
<td>7</td>
<td>24 Oct 1988</td>
<td>6</td>
<td>45</td>
<td>0.21</td>
<td>0.08</td>
</tr>
<tr>
<td>8</td>
<td>5 Oct 1988</td>
<td>13</td>
<td>33</td>
<td>0.15</td>
<td>0.04</td>
</tr>
<tr>
<td>9</td>
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<td>See Fig. 6B</td>
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<td>20</td>
<td>39</td>
<td>0.20</td>
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<td>39</td>
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<tr>
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<td>39</td>
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<td>11</td>
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<td>55</td>
<td>0.13</td>
<td>0.07</td>
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<tr>
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<td>0.06</td>
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<tr>
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<td>42</td>
<td>0.32</td>
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<td>0.13</td>
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<td>20</td>
<td>42</td>
<td>0.14</td>
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The length of time before the decline began depended on the oxygen concentration (Fig. 7). When exposed to 10% O₂-saturation an immediate reduction of the mean Hcy concentration by 65% of the initial value occurred within 4 d. After 4 d mortality was 90% in a sample size of 10. The lobsters in 12% O₂-saturation first increased their Hcy concentrations, relative to the 4 series of control lobsters in fully saturated water (n = 38), to a threshold (12%; 15 d, 15%; 36 d). After that a decrease in Hcy level occurred at a rate similar to that recorded for lobsters in 10% O₂ saturation, and the percentage mortality after 4 d was high (100% for 12% O₂, 40% for 15% O₂). Lobsters exposed to 20% O₂ saturation were transferred to 15% O₂ saturation (Fig. 7) after 3 wk of exposure. This treatment resulted in a relatively rapid decrease of 58% Hcy within a further 13 d. The decrease in Hcy concentration of 30% within 33 d (0.9% d⁻¹) for the control groups is in accordance with the decrease for starving lobsters in this temperature range, as described above (Table 4A).

**Behavioural observations**

When exposing *Nephrops norvegicus* to 20% O₂-saturation some of the lobsters stood high above the substrate with legs flexed ('tiptoeing') but were still eating when offered food at the beginning of the experiment. After 3 wk the lobsters stopped feeding. *N. norvegicus* exposed to <15% O₂-saturation showed no sign of feeding. When exposed to 12% O₂-saturation all lobsters were 'tiptoeing' and supported the elevation of the body from the substrate with the claws and telson. The lobsters remained elevated until they became moribund and sluggish, barely moving when touched. This stage lasted for 2 to 3 d after which they died.

**DISCUSSION**

**Biomass and population structure**

The emergent behaviour of *Nephrops norvegicus* in relation to their burrows in the sediment is known to vary due to factors such as temperature, salinity, oxygen, light, depth, sediment structure and reproductive status (Simpson 1965, Thomas & Figueiredo 1965, Bagge & Munch-Petersen 1979, Chapman 1980), and the catchability of the lobsters during trawling may well be partly dependent on these factors. As a consequence, trawling will not, in most cases, give a quan-
and then decreased (Fig. 5). Furthermore, a high year-
to-year variation in commercial catches was found in
this area, compared to the NE Kattegat (ICES area
4356). The southern part of Kattegat has been severely
exposed to low oxygen concentration during the
1980’s, whereas in the northern part the exposure has
been moderate. Low oxygen concentrations force lob-
sters to emerge from their burrows resulting in an
increased catchability (Bagge & Munch-Petersen 1979,
Baden et al. 1984). Variation in hypoxia between years
will therefore cause a variation in the catch of the
lobsters and this might explain the different patterns in
CPUE between the 2 areas.

The biomass (CPUE) of Nephrops norvegicus
showed a general decrease in the study area during the
investigated period and during the last 5 samplings no
lobsters were caught (Fig. 3). Increased exploitation
caused by moderate hypoxia and finally severe
hypoxia were most probably responsible for the decline
in biomass. The discrepancies between CPUE in Figs. 3
and 5 are because area 4257 covers not only most of the
study area, from which the CPUE in Fig. 3 derives, but
also areas less affected by hypoxic conditions, west of
the study area. When catches decrease in the study
area the fishermen can go to other areas within 4257.

The size distribution of the Nephrops norvegicus
population was similar throughout the period investi-
gated, and males were dominant (mean = 78% of
total) on most sampling occasions. These results are in
agreement with what has been found in other areas
where males were found to dominate in the catches
without any seasonal variation in size frequency distri-
bution (Thomas & Figueiredo 1965). Females stay
mainly in the burrows during the egg-carrying period
(September to May), as do the juvenile lobsters. During
low oxygen tensions, however, changes of the popula-
tion structure of N. norvegicus are
probably the main factors affecting the catchability of
N. norvegicus at least during autumn.

Since 1980 low oxygen concentrations have been
recorded in the bottom water of the SE Kattegat during
autumn of each year. Levels of oxygen concentration
and duration of the oxygen depletion varied between
years, mainly due to variations in nutrient load, inten-
sity in phytoplankton blooms and hydrographical fac-
tors. However, during the 1980’s a general decrease in
oxygen concentrations and an increase in duration of
time with low oxygen have been observed. In the
autumn of 1985, 1986 and 1988 low (< 1 ml l⁻¹) oxygen
concentrations were recorded in the bottom water for 2
to 4 wk, and during these periods dead lobsters were
captured (Petersen & Petersen 1986, this study). Catches
of Nephrops norvegicus from ICES area 4257 show that
catch per unit effort increased during the early 1980’s

Diet

The mean percentage occurrence of Crustacea,
Polychaeta, Echinodermata and Mollusca in the
stomachs of Nephrops norvegicus in SE Kattegat was
fairly similar, between 46 and 66 % (Table 2). This
suggests that they are not feeding selectively. It has
also been stated in other reports that N. norvegicus
seems to be a varied feeder, taking available organisms
indiscriminately (Thomas & Davidson 1962, Oakley
1979, Bailey et al. 1986). No detailed comparison has,
however, been made between food intake by N. nor-
The experiments also showed that even when exposed to hypoxic conditions in the SE Kattegat, about 80 to 90% of the population of *N. norvegicus* was well masticated and difficult to identify. Benthic data from 6 stations in the vicinity of the northern subarea, 7 in the center and 3 in the southern subarea were selected for a cursory comparison (depth range 24 to 57 m). Dominant organisms in terms of abundance and biomass in the northern and middle subareas were *Amphiura filiformis* (O. F. Müller) and *A. chiajei* (Forbes); other numerically dominant organisms were *Maldane sarsi* (Malme grades), *Myriochele oculata* (Zaks), *Philomedes globosus* (Lilljeborg), and in terms of biomass, *Brissopsis lyrifera*. In the southern subarea *P. globosus*, *Heteromastus filiformis* (Clearedre) and *M. sarsi* were dominant in abundance, and *Arctica islandica* (L.), *A. filiformis* and *B. lyrifera* were dominant in terms of biomass.

Most of these macrobenthic species were also frequently found in *Nephrops norvegicus* stomachs (Table 2), indicating that *N. norvegicus* tend to ingest what is readily available. *Abra* spp. were frequently recorded in the stomachs, but not among the benthic dominants. Both *Abra alba* (W. Wood) and *A. nitida* (Müller) can, temporarily, be locally abundant in the Kattegat (Rosenberg & Loo 1988). The highest occurrence in the stomachs was recorded for brittle stars (amphiprioids and ophiidioids). In the other studies cited above brittle stars were found less frequently, except in the Minch, off the Scottish west coast, where they formed 18 to 51% of the diet (Thomas & Davidson 1962). The high occurrence in the Kattegat was probably due to the dominance of brittle stars in this area, and it is not known how common this *animal* group was in other regions.

Feeding experiments in this investigation showed an average food intake of about 9 g d$^{-1}$ per kg *Nephrops norvegicus* (wet wt). According to Chapman (1980) and Smith (1988) the density of *N. norvegicus* is around 0.15 m$^{-2}$ in most areas but can vary from 0.1 to 3.8 m$^{-2}$ when estimated by underwater photography and by diving. The average total length of *N. norvegicus* in our samples from the SE Kattegat was around 14 cm (43 mm, carapace length) (Fig. 4) corresponding to a mean weight of 52 g (wet wt). If the *N. norvegicus* population is eliminated through low oxygen concentrations the result could be a reduction in predation pressure of ca 25 g (wet wt) benthic fauna m$^{-2}$. yr$^{-1}$, assuming an initial *N. norvegicus* density of 0.15 m$^{-2}$.

In addition to this, during the autumn with hypoxic conditions in the SE Kattegat, about 80 to 90% of the demersal fish fauna had disappeared (Pihl 1989) with a probable but still unknown decrease in the predation pressure on the infauna (Pihl unpubl.). Thus, apart from the likely direct effect of hypoxia, a considerable decrease in predation pressure might be an important influence for determining the community structure of the benthic infauna.

### Haemocyanin concentrations

#### Moderate hypoxia

Stress from hypoxia induces several parallel physiological changes to compensate for the effect of low O$_2$-saturation. In many crustaceans one way of compensating is to increase the capacity of O$_2$-uptake by increasing the blood Hcy concentration. In laboratory experiments moderate hypoxia of 20 to 40% O$_2$-saturation induced an increase in synthesis of Hcy in American lobster *Homarus americanus* (Milne Edwards) (Senkbiel & Wriston 1981), isopod *Mesidotea entomon* (L.) (Hagerman & Okama 1985), *Nephrops norvegicus* (Hagerman & Uglow 1985) and shrimp *Crangon crangon* (L.) (Hagerman 1986). In these investigations the rates of Hcy synthesis were found to be 6 to 9% of the initial value per day, provided enough food was available. Field results from the Skagerrak (Stn 9) suggest that *N. norvegicus* synthesized Hcy in moderate hypoxia (38 to 43% saturation). The O$_2$-concentration experienced by the lobster in the burrow was probably even lower than on the bottom (Atkinson & Taylor 1988, Rosenberg & Loo 1988). The rate of Hcy synthesis in the Skagerrak cannot be evaluated as the intervals between samples were too long. The maximum Hcy concentration of 1.8 mM recorded in *N. norvegicus* during moderate hypoxia was also found for *C. crangon* by Hagerman (1986).

During normoxia (January to June) a slow decrease to 'normal' blood Hcy concentrations occurred, while a faster decrease took place between October and November due to the onset of less than 20% saturation. In many crustaceans one way of compensating is to increase the capacity of O$_2$-uptake by increasing the blood Hcy concentration. In laboratory experiments moderate hypoxia of 20 to 40% O$_2$-saturation induced an increase in synthesis of Hcy in American lobster *Homarus americanus* (Milne Edwards) (Senkbiel & Wriston 1981), isopod *Mesidotea entomon* (L.) (Hagerman & Okama 1985), *Nephrops norvegicus* (Hagerman & Uglow 1985) and shrimp *Crangon crangon* (L.) (Hagerman 1986). In these investigations the rates of Hcy synthesis were found to be 6 to 9% of the initial value per day, provided enough food was available. Field results from the Skagerrak (Stn 9) suggest that *N. norvegicus* synthesized Hcy in moderate hypoxia (38 to 43% saturation). The O$_2$-concentration experienced by the lobster in the burrow was probably even lower than on the bottom (Atkinson & Taylor 1988, Rosenberg & Loo 1988). The rate of Hcy synthesis in the Skagerrak cannot be evaluated as the intervals between samples were too long. The maximum Hcy concentration of 1.8 mM recorded in *N. norvegicus* during moderate hypoxia was also found for *C. crangon* by Hagerman (1986).

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### Severe hypoxia

In severe hypoxia (< 20% O$_2$-saturation) *Nephrops norvegicus* in our experiments became less active and raised the body by extending their legs, a behaviour also found previously (Batterton & Cameron 1978, Senkbiel & Wriston 1981, Hagerman & Uglow 1985). The experiments also showed that even when exposed
to severe hypoxia, down to 12% $O_2$-saturation, the lobsters initially tried to compensate for the low oxygen concentration by increasing the Hcy concentration of the blood compared to controls. In this way $N.\ norvegicus$ survived about 4 wk in 15% $O_2$-saturation and about 2 wk in 12% $O_2$-saturation. Before starting to increase a slight decrease in the Hcy concentration occurred after 3 and 4 d for lobsters in 15 and 20% but not in 12% $O_2$-saturation. This is probably due to an initial increase of the haemocyanin $O_2$-affinity by raising the blood pH (Schmidt-Nielsen 1975). This compensation seemed to work for about a week, when exposing lobsters to 15 and 20% $O_2$-saturation, and the Hcy concentrations were the same as for lobsters in 100% $O_2$-saturation. However, to meet the stress from 12% $O_2$-saturation and prolonged exposure to 15 and 20%, the lobsters increased their Hcy concentration compared to lobsters in 100% $O_2$-saturation. The catabolism of haemocyanin during the hypoxic experiments was a rapid process lasting no longer than a week followed by mortality. In 10% $O_2$-saturation the lobsters were so stressed that no Hcy compensation to low oxygen levels occurred, and the lobsters survived for only 2 to 4 d. A rapid decrease in Hcy and high mortality was also found in 10% $O_2$-saturation in a similar experiment on $N.\ norvegicus$ by Hagerman et al. (1990).

Field data on lobster Hcy concentrations during hypoxia where the Hcy concentration decreased to less than 10% of what is regarded as the normal value (Hagerman & Baden 1988, this investigation) seem to be in accordance with our laboratory experiments. Thus, low haemocyanin concentrations found in $Nephrops norvegicus$ in the SE Kattegat were most probably due to the low $O_2$-saturations (less than 15%) over a 4 wk period.

In normoxia, catabolism of Hcy has been described as an effect of starvation, when the blood protein are used as an energy resource, for example in Carcinus maenas (L.) (Uglow 1969) and in Homarus gammarus (Hagerman 1983). Our experiments on the rate of catabolism when starving $Nephrops norvegicus$ under normoxic conditions correlate well with these results in showing nearly a 1% d$^{-1}$ decrement of the initial Hcy concentration.

The decrease of Norway lobster Hcy is unlikely to be explained by lack of food availability. As previously mentioned benthic infauna were abundantly available and during initial hypoxia perhaps even more accessible when both lobsters and some of the infauna emerged from the sediment (Dyer et al. 1983, Rosenberg & Loo 1988). In our experiments we found that when $O_2$-saturation was below 15% the lobsters stopped feeding and this was also the case in the field in September 1988, when all the lobsters had empty stomachs. Thus, in low oxygen concentrations, the lobsters suffer from hypoxia-induced starvation rather than direct lack of food.

The results from field samples of low Hcy concentrations in $Nephrops norvegicus$ from the Skagerrak indicate that severe oxygen depletion may occur even in some of the deeper coastal areas (Table 3).

Recovery

The recovery of blood pigment concentration to ‘normal’ concentrations was very slow after the return of normoxia in the bottom water. The only exception was from subarea S in autumn 1986, the first year when the effect of hypoxia on the lobsters was examined (Hagerman & Baden 1988). As indicated by our feeding experiments, where only lobsters fed on crustaceans increased their blood Hcy, and by Castell & Hudson (1974), Hagerman (1983) and Depledge & Bjerregård (1989), other crustaceans are an important part of the diet for crustaceans, supplying essential proteins, certain vitamins and copper for haemocyanin synthesis. The content of crustaceans in the diet of Norway lobsters in the southern Kattegat was only about 20% less than in the diet from areas in the northern Kattegat.

According to Depledge (1989) a translocation of part of the copper, mainly from haemocyanin to the hepatopancreas, occurs in crustaceans when starved and more than 50% of the copper may be excreted. Hypoxic conditions force not only the lobsters but also their food items into severe starvation. Under this condition more than 50% leakage of copper could be possible. As food and not seawater is the most important source of copper to crustaceans (Windom & Smith 1979, Bryan 1984, Depledge & Bjerregård 1989) the slow recovery of Hcy could more likely be due to copper deficiency than to the 20% lower abundance of crustaceans in the diet in the SE Kattegat compared to the NE Kattegat. How the layer of manganese on the gills might affect oxygen uptake in the ‘black gill’ individuals remains to be investigated. It is also possible that some elevation of the Hcy concentration took place during early summer, but the concentration may have decreased again due to low oxygen prior to the first sampling in September.

Haemocyanin concentration as a biomarker

In a review of physiological biomarkers, Mayer et al. (1990) conclude that it is important to develop ‘in situ’ biomarkers of ecological relevance.

The repeated occurrence of low blood Hcy levels in different localities in the Kattegat and the Skagerrak...
CONCLUSIONS

During the 1980's the SE Kattegat has suffered from severe and intensifying hypoxia which has been shown to affect benthic infauna (Rosenberg & Loo 1988) and demersal fish (Pihl 1986, 1989). In this study we have found that Nephrops norvegicus, due to its relatively sedentary life, is one of the most severely affected species.

Both ecological and physiological disturbance of Nephrops norvegicus has been shown to occur during hypoxia. In a 3000 km² area in the SE Kattegat these disturbances have been fatal, with no lobsters being caught at the sampling sites in the 12 mo since September 1988. During hypoxia a series of events make the lobsters unable to cope with the stressed environment. During moderate hypoxia lobsters emerge from the sediment, and are thereby overfished by trawlers. At even lower oxygen concentrations the lobsters are immobilized and unable to feed. Some lobsters had a black layer of mineralized manganese on the gills, and this layer did not disappear with return to normoxia. The layer might reduce oxygen uptake efficiency. At a later stage catabolism of the blood pigment protein haemocyanin helps the lobster to survive for a short period before death.

In the event of a much needed reduction of excessive nutrients levels, the underlying cause of hypoxia in the Kattegat, what are the chances of recovery of the Nephrops norvegicus stock in the area? As long as the hypoxia does not affect the NE Kattegat to the same degree as the southern part, the lobsters will most probably return, but after how long? The recruitment of N. norvegicus is not well documented. Juveniles are mainly found in small tunnels connected to the burrows of adult lobsters (Chapman 1980), but have also been observed to make their own burrows (Crnkovic 1968).

The highest survival potential may occur when juveniles are living in association with adults, perhaps by benefitting from food caught by adults. If this is the case, recovery would occur mainly by immigration from surrounding areas and this would be a very slow process, since migrations of adult Norway lobster are limited (Jensen 1965, Chapman 1980). If recovery can also occur by larval settlement, it could be faster. N. norvegicus reaches catchable size in about 4 to 6 yr, thus recovery of the population within a decade would be possible, provided all other factors such as food availability and sedimentary conditions are maintained.

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LITERATURE CITED


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