

Desktop study on autecology and productivity of European lobster (*Homarus gammarus, L*) in offshore wind farms

Author(s): M.J.C. Rozemeijer, K.E. van de Wolfshaar

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Summary

This desk study describes the biology of the European lobster *H. gammarus*. Using the obtained data a model was developed to describe the growth of the European lobster under assumed conditions on the anti-scouring of monopiles in Dutch OWFs. One of the main questions to answer was, if, theoretically, local productivity supports the continuous harvesting of lobsters with passive fishery methods.

Main findings

Lobster ecology

The first finding is that lobster is characterised as a large top end predator. It exhibits a k-strategy life style, able to reach very old ages (e.g. an estimated 72 years). The planktonic and early benthic stages are very vulnerable and susceptible to predation. It grows slowly, taking at least four years to reach maturity in favourable conditions although in general maturity is thought to start from five years old depending on ambient conditions. Reproduction occurs in a two to three year cycle. The eggs are held on the pleopods for approximately a year until hatching. In itself the reproduction is sufficient to maintain populations at constant levels (given the high age). However, global lobster populations have suffered considerable fishing pressure and stock collapse, only recovering slowly now at some regions partly due to appropriate management (the MCRS and stock enhancement programmes, (Wickins & Lee, 2002, Prodöhl et al., 2006, Phillips, 2013)).

In principle European lobsters have a high enough mobility to colonize the entire North Sea using wrecks, hard substrates etcetera as stepping stones (Krone and Schröder, 2011). Yet, unexpected differentiated genetic clusters occur which implies that at a North Sea scale realized mobility is more limited than potential mobility. Apparently the exchange is less than based on theoretical estimates of mobility.

The limited exchange implies that when exploiting lobsters at OWFs an exploitation management plan should be established based on the local OWF population dynamics and the presence of lobsters nearby on the soft sediments. Also stock enhancement using larvae or juveniles from local populations might be an additional option. The seafloor at the OWF could be an issue in choosing at which size to release juvenile lobsters since the earliest benthic stages can dig their holes. When the seafloor is more sandy than solid mud, early stages are entirely depending on the crevices between stones and cobbles. In addition they actively harvest the walls of their residence. The more productive the seafloor bottom is (in terms of worms, amphipods etc.) the less frequent these vulnerable stages have to expose themselves to predation and currents (Lawton & Lavalli, 1995, Jensen et al., 2000, Wickins & Lee, 2002). In this sense older and larger stages (> an estimated 15 mm CL) are less dependent on bottom type (and more on availability of crevices).

European lobster prefer hard substrate with crevices to hide, except for the pelagic larval stage. At younger stages they are entirely dependent on shelter in order to evade predation. At later stages they become less dependent on shelter but keep a preference for hard substrate. Crevices should have the size to accommodate the complete body and should be available for all sizes. Younger animals of European and American lobsters move to larger crevices while growing (Lawton & Lavalli, 1995, Jensen et al., 2000, Dunnington et al., 2005, Phillips, 2013)

Taking the expected crevice size into account for OWF Prinses Amalia and OWF Luchterduinen (maximally 8 x 8 x8 cm, Table 7), theoretically there is a lack of suitable crevices for larger sized lobster in the OWFs preselected for TKI WinWind. Adding extra hard structures with crevices suitable for lobster might stimulate the presence of legal catchable lobster (at 85 mm CL, an estimated 240 mm total length, Prodhöl et al., 2006). Monitoring is needed to establish which size-classes of *H. gammarus* are present in the OWFs. A potential new habitat should be carefully designed accommodating several sizes classes of lobster, having sufficient water and oxygen refreshment and also enabling stocks of the typical food like mussels and crabs.

Lobster densities at suitable substrates of a fished population are reported to be 1 lobster per $\sim 150 \text{m}^2$. American lobsters can have much higher natural densities (1 lobster every 6m^2) but one should keep in mind that this species is migratory (Dunnington et al., 2005, Geraldi et al., 2009, Phillips, 2013), which might explain that a given substrate does not have to provide food year round. In addition, they have a higher mobility so likely have a greater foraging area.

DEB modelling and lobster production

The DEB model predicts lobster growth in line with growth reported in literature, although length at age from field data shows a large variability. Based on the current model parameter setting resource production is limiting lobster growth and abundance while ensuring that lobsters reach marketable sizes. The current model setting predicts that 1 lobster can reach marketable size after 3 years, provided a stocking length at 50 mm CL. Model resource productivity was based on mussel density estimated at a limited survey of monopiles in a Dutch OWF. However, resource production might differ between monopiles, parks and temperature regimes, and higher productivity will support more lobsters and/or have a positive effect on individual growth rates.

The population model does not take recruitment into account, the reason being that a single monopile surface area is being studied and that little is known on factors determining recruitment success in wild populations, nor the geographical extent of such populations in the North Sea. In addition, no lobsters were observed at a studied wind park 4 years after operation started (Bouma and Lengkeek, 2012). This suggests that at least the chance of natural recruitment on monopiles in wind parks in the North Sea is low. However, when the expectancy is 1 lobster per monopile, changes of encounter during a scientific dive are not that high. Note that energy loss through egg production is accounted for in the model.

The modelled European lobster productivity is in line with low densities found in the field, using the OWF hard substrate and anti-scouring data from Bouma & Lengkeek (2012). One lobster could be supported by one monopile given its anti-scouring surface area (1 per 364 m²).

Enhancement strategies

The current literature survey on lobster ecology linked observations to selected enhancement strategies for local populations (habitat, stock and food). All strategies seem to have a potential added value for European lobster production but measurements and monitoring in OWFs is the first step to describe the actual situation in OWFs. DEB modelling suggests that food availability is the first limiting factor. Therefore food enhancement (by e.g. improving quality by seeding with favoured food or improving availability by depositing discards of fisheries) can help to increase productivity. Habitat enhancement by means of extra hard substrate with extra crevices can serve to increase the amount of crevices available in OWFs, especially for larger lobsters. In addition, extra hard substrate can serve as fundament for extra biota and thereby provide food. Stock enhancement can help to surpass the vulnerable pelagic and early settlement stages (up to ~15 mm CL or even larger). It can also safeguard supply of animals since natural recruitment is low.

1 Introduction

The multi-use of offshore wind farms (OWFs) on the Dutch National Continental Flat (NCP) with harvesting natural stocks and or aquaculture is becoming more and more realistic and accepted. Amongst others the passive low-impact fisheries of European lobster (*Homarus gammarus*) and brown crab (*Cancer pagurus*) has been proposed as an excellent opportunity. To that extent a TKI tender has been awarded and the project TKI Win-Wind on enabling this type of fisheries is in progress.

As a preparation for this project knowledge on the ecology of both species is necessary. This desk study will focus on the European lobster and make an overview of relevant features.

To be able to grow additional biomass¹ on locations like OWFs, food availability to enable metabolism and growth of such additional biomass is essential. However, the production potential of the NCP for European lobsters and brown crabs is not yet well known, and hardly any spatial planning based on ecosystem productivity is currently considered. In general is valid for the benthos of the North Sea that quantity and quality of the primary production (PP) are important for the amount and quality of the biomass of the benthos. Also hydrodynamics are important determining availability. A benthos community dominated by suspension feeders like the Dutch Coastal zone or the German Bight is thought to represent a food saturated situation (Kröncke, 2006). Smaal et al. (2017) use food concentration (in μ g Chlorophyll-a (Chl-a)/l) as a factor in habitat modelling of the European oyster (Ostrea edulis) in order to select the best suitable OWFs for pilot reef reconstruction experiments. They do not calculate maximum productivity potential of the European oyster based on available PP since there are hardly any recent measurements on PP in the Dutch NCP. The tools to support adequate spatial planning, and ecosystem based productivity (a driver for site selection and food harvest potential) are not yet available for utilisation. This information should be available to choose and optimize harvest and locations (in time and space) and also to be able to evaluate management measures in order to increase local productivity (e.g. evaluation of habitat enhancement, stock enhancement or food enhancement).

Growth and production models could yield such insights to assess the site dependent potential of ecosystem productivity and production species occurrence (where species can grow, and what production can be achieved). In this study a first step is taken to model productivity of European lobsters in OWFs. In this first inventorying study a model is selected best suited for our questions on productivity potential of lobster on the anti-scouring of one monopile in an OWF. An important question for the TKI Win-Wind project is whether it is possible and effective to enhance the populations of European lobsters in OWFs in order to increase the potential profits of harvesting. Therefore a first limited exploratory review will be performed on potential population enhancement strategies. **Table 1** gives a definition of the population enhancement strategies that are commonly used for European lobsters (Jenner et al., 2000) that will be studied in this study.

1.1 Problem definition

It is currently unknown what the local maximum production capacity is for aquaculture in general and lobster in particular in especially OWFs and what drives or limits aquaculture production.

The added presence of hard substrates in de form of anti-scouring and monopiles in an OWF should lead to increased benthic biomass and diversity in a soft substrate dominated seafloor (van Moorsel et al., 1991, Leewis et al., 1997, van Moorsel & Waardenburg, 2001, Bouma & Lengkeek, 2012). There is little knowledge on the offshore carbon fluxes in general and how the carbon fluxes have changed

¹ Either passively by just offering constructions to enhance settlement of desired species or actively by stimulating biomass grow by e.g. additional food.

within the food web after addition of hard substrates. The potential maximum carrying capacity for aquaculture stocks is consequently also undefined. Information is needed to assess maximum aquaculture production and driving forces thereof in time and space in order to make substantiated management choices in allocation of OWFs for multi-use purposes such as aquaculture. These insights can also aid in selecting the appropriate tools for the potential improvement of local populations of European lobsters (e.g. enhancement of habitat, stock or food). Since assessing production capacity from a holistic point of view is an extensive research question, a stepwise approach is necessary. This desk study represents a first step in this approach (a simplified production capacity estimation model).

1.1.1 Target group and knowledge

<u>Target groups</u>: are governments as area managers, aquaculture companies, OWF operators and fishermen being potential users of the knowledge to optimise their business case. <u>Knowledge</u>: the project will lead to knowledge and tools to manage and optimise multi-use of OWFs aiming at aquaculture in general and lobster and brown crab in particular.

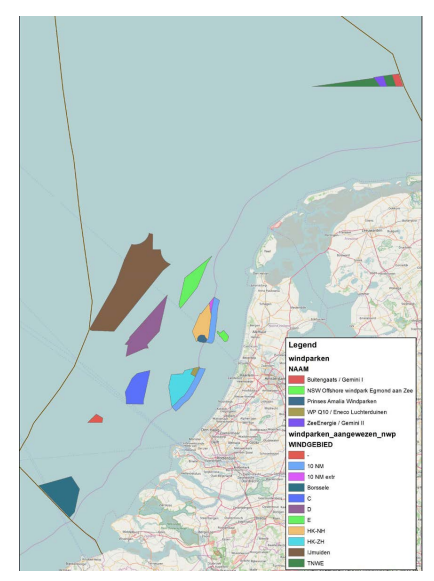


Figure 1. Map of the locations of existing and planned offshore wind farms (OWFs) in the Dutch section of the North Sea (DSC: Dutch Continental Shelf). The existing OWFs are the first names and colours given in the legend. HK-NH: Hollandse Kust Noord-Holland. HK-ZH: Hollandse Kust Zuid-Holland (fig. from Smaal et al., 2017).

1.2 Objectives of the project

The objective of the project is to generate management information on the potential of aquaculture in wind parks in the North Sea:

- 1. This study will describe the general ecology and life history of the European lobster (*H. gammarus*) from which suggestions will be derived if and which population enhancement strategies can be potentially successful to promote production.
- 2. A first, limited exploratory review will be performed on potential population enhancement strategies (Table 1).
- 3. The literature study obtains the necessary data to support the (already existing but adapted) production capacity estimation model.
- 4. A already existing DEB model will be adapted and developed within a population dynamical framework to study the production capacity of lobster at a monopile.
- 5. The productivity of lobsters will be determined with the developed model fed with resource levels based on encountered biomass data on antiscouring of OWF Prinses Amalia (Bouma and Lengkeek, 2012).
- 6. The stock enhancement strategy will have a first exploratory test with the production capacity estimation model.

Table 1. Definitions of habitat enhancement, stock enhancement and food enhancement.

- Habitat enhancement: E.g. pre-adult and adult lobsters are reef-obligate and select sites that supply sufficient food and oxygen and shelter from currents and predation. The brown crab exploits a broad-range of environments, ranging from soft muds into which it can hide or dig for food to hard, rocky substrata where it exploits, and seeks shelter, in crevices. (Linley et al., 2008). Next to shelter, additional habitat will yield extra food possibilities. Reef type habitat can be improved by improving the texture of the anti-scouring construction (Lengkeek et al., 2017) or by adding artificial reef structures outside the anti-scouring zone (with specific structures and features, e.g. Jensen et al., 2000, Linley et al., 2008, Buck et al., 2017). Both options have their disadvantages: anti scouring protection has specific design criteria thereby limiting the possibilities for optimising design to meet species preferences. Adding new and extra structures outside anti-scouring locations obliges both Water-act permits and the removal of the structures on decommissioning the park (Lengkeek et al., 2017). A first exploratory literature review will be given on possibilities and applicability for the specific Dutch case in order to support management decisions on the next pilot phase.
- 2. Stock enhancement: is an option to increase the natural occurring number of brown crabs or preferably the more profitable lobster. Stock enhancement beholds the release of juvenile lobsters (either larvae, post larva or older stages) that have been reared in hatcheries thereby increasing survival of the vulnerable stages. Release of reared individuals enables tagging and thereby following individual growth in time and other quantitative evaluation of lobster release programmes (Jensen et al., 2000, Prodöhl et al., 2006, Cornwall National Lobster Hatchery website²). For the Oosterschelde ideas exist to initiate a local hatchery. Risks exist in the sense that e.g. the Oosterschelde populations represent an isolated distinct genetic group. Furthermore the released recruits i) represent also a specific (favoured) genetic pre-selection ii) that has not been selected (survived earlier stages) by local circumstances (Prodöhl et al., 2006). The literature review can be considered exploratory.
- Food enhancement: lobsters and brown crabs are opportunistic in their food preferences. An option to improve populations might be to increase the amount of available food. Potentially low value fish catches that used to be defined as by catch, can serve as additional food supplies. In Sub-WP-E3: Monitoring and habitat modelling, insight will be generated on the food availability in relation to stocks.

² https://www.nationallobsterhatchery.co.uk/whats-it-all-about/ d.d. 24-05-18, 09-12-18.

Literature review on lobster

This section gives the results of the literature review on European lobster. In its intention it does not aim at being complete on the aspects mentioned. Because the American lobster has a similar size range and general ecology and more literature on this species is available, information on this species is added to the review. Data on *H. americanus* are often used as a model or representative for *H. gammarus*. General feeling is that the species are very much alike and that the data of *H. americanus* can be used as such (Phillips, 2013). Though it should be noted that *H. americanus* is bigger, has migration and occurs to -700m, much deeper than -165m of *H. gammarus* (Phillips, 2013). This suggests at least some differences in physiology. When *H. americanus* data are used, it will clearly mentioned.

2.1 Description

2

The scientific name of the European lobster is *Homarus gammarus*. They possess two large claws, one blunt and designed for crushing and the other sharper and smaller normally used for slicing. Males have larger claws than females, females have wider abdomens (Agnalt et al., 2009, Sørdalen et al., 2018). Their exoskeleton is blue/navy on the dorsal side but slightly lighter underneath with an orange tinge. The body length up to 60 centimetres and weighing up to 5–6 kilograms, although specimen of more than one meter are encountered too. Commercial catches in Cornwall (UK) showed large males of > 150mm CL and females of 140-150mm (Hepper, 1978, Figure 6). In Norway up to 140 CL for both males and females. The largest specimen on record measured 1.26m and weighed 9.3kg, caught in 1931 in Fowey, England².

2.1.1 Age

They can reach considerable ages. Sheehy et al. (1999) estimated³ males to average 31 years, maximum 42 \pm 5 years and females to average 54 years, maximum 72 \pm 9 years (Figure 2). Weights were maximally 4.09 kg for an eleven year old male (166 mm Orbital Carapace Length, OCL, from the back of the eye socket (the orbital), see Appendix 1) and 2.09 kg for the 72 year old female (170mm OCL). These were all from a commercially bought group in the UK preselected on large size (>130mm OCL, bought at Bridlington and Scarborough considered to have originated primarily from the offshore fishery, 8–20 km from the coast). The sizes of the largest males and females ranged between 150–170 mm OCL. Appendages and limbs lost through injury can be regenerated, but this usually has a negative effect on both the intermoult period and the growth increment (Jensen et al., 2000, Phillips, 2013).

2.1.2 Physical characteristics

Scientifically published data on weight, CL, age, total body length (TL) for European lobster are given by Jensen et al. (2000), Kristiansen et al. (2004), Van Stralen and Smeur (2008) Schmalenbach (2009) Schmalenbach et al. (2011), Tangelder et al., 2015) In this study the calculated weight at CL relationships were gathered and given in Table 2 and Figure 3. Observing the graphs the impression is raised that there are differences between locations and differences between male and female lobsters from Helgoland. This could reflect differences in life history (temperature, food conditions, habitat) and sexes. Figure 2 and Figure 4 confirm that males are heavier than females. On the other hand, one should keep in mind that the steep exponential part of the relationship is determined by the larger and heavier animals which are rarely caught. Data points of these larger animals make the graph much

³ It is difficult to estimate age in lobsters. It has to be done indirectly, usually based on the accumulation of lipofuscin. Lipofuscin, is a by-product of cellular metabolism. It accumulates in all cells at a rate determined by metabolic rate of the individual and is quite stable after formation and therefore has met with some success in crustacean age estimation.

steeper than the equation based on data leaving out these older animals. So differences between these relations can be a mathematical difference rather than an ecological difference.

In Table 2 several measured carapace length at total length (TL) relationships were gathered. This ratio seems rather stable, the slopes are alike. For the relation between TL and CL, We calculated a slope of 0.37 ($R^2 = 0.9963$) from the data of Schmalenbach (2009). Kristiansen et al. (2004) calculated a slope of 0.35 ($R^2 = 0.9519$) for reared juvenile European lobsters from Norway, Mercer et al (2001) a slope of 0.36 for reared lobster from Ireland and Scotland.

Table 2. Relations between different physical aspects like carapace length (CL) and weight of different authors and total length (TL) and CL or vice versa for different authors. For Schmalenbach 2009 given year class averages were used to calculate the relations by ourselves

Origin of animal	Relation	R ²	Country	Reference
CL vs Weight			country	
Reared juveniles	W=0.0003*CL ^{3.1522}	0.9851	Norway	Kristiansen et al. (2004)
Larger wild lobster caught in the sea	W=0.0002*CL ^{3,2953}	0.9219	Norway	Kristiansen et al. (2004)
Larger wild lobster caught in the Oosterschelde	W=0.000328*CL ^{3,1708}	0.9779	Netherlands	Van Stralen and Smeur (2008)
Larger wild lobster caught in the Oosterschelde	$W = 0.0016 * CL^{2.803}$	0.9840	Netherlands	Tangelder et al. (2015)
Larger wild female lobster caught at Helgoland	$W = 0.001 * CL^{2.9211}$	0.9996	Germany	Schmalenbach (2009)
Larger wild male lobster caught at Helgoland	$W = 0.0005^* CL^{3.1046}$	0.9982	Germany	Schmalenbach (2009)
TL vs CL				
Wild male and female lobster caught at Helgoland	CL = 0.3727*TL - 5.4775	0.9963	Germany	Schmalenbach (2009)
Reared juveniles	CL = 0.3512*TL	0.9519	Norway	Kristiansen et al. (2004)
EBP Juveniles reared in either Shellfish Research Laboratory, Carna, or the Orkney Fisheries Association in Orkney, Scotland.	CL = 0.3552*TL + 0.170		Ireland, UK	Mercer et al. (2001)

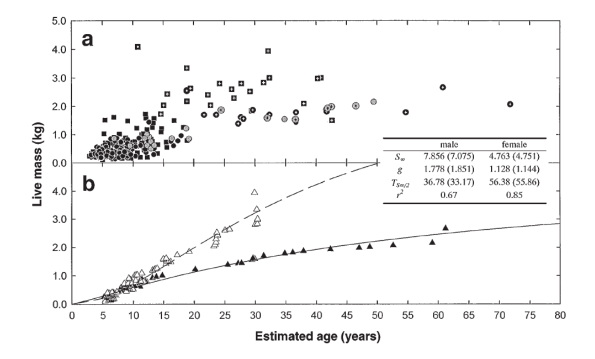


Figure 2. Relationship between live mass and estimated age for sampled wild Yorkshire *H. gammarus*. (a) Raw estimated age – live mass data: squares, males; circles females; solid black symbols, Yorkshire lobsters collected in 1996 and originating primarily from inshore fishing grounds with gravid females shown as solid grey circles; black symbols with cross, Yorkshire lobsters collected in 1997 and originating primarily from offshore fishing grounds with gravid females shown as grey circles with cross; black symbols with dot, Cornish lobsters. Live mass data were not available for the reference lobsters.

(b) LOWESS-smoothed data from Figure 2a, excluding Cornish lobsters: open triangles, males; black triangles, females; dashed and solid lines, best-fitting logistic models for males and females, respectively. The regression coefficients for the models are shown in the inset. Values in parentheses and r² coefficients refer to the same model when fitted directly to the raw data in Figure 2a. Figure taken from Sheehy et al. (1999).

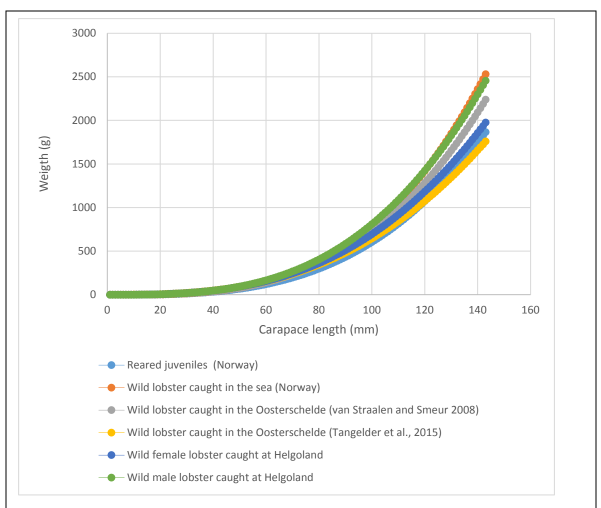


Figure 3. Calculated weight at carapace length relationships obtained from Kristiansen et al., 2002, van Stralen and Smeur, 2008, Schmalenbach (2009), Tangelder et al. (2015). For the weight at carapace length relationship of Schmalenbach (2009) the average data for each year class were used.

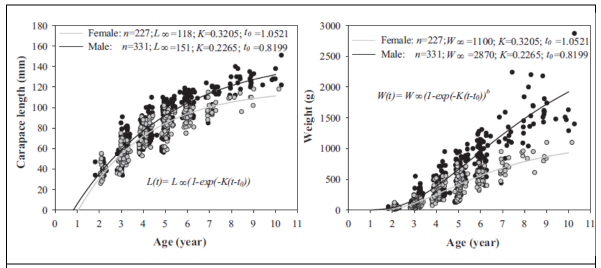


Figure 4. Growth curves from length-at-age and weight-at-age data for female and male European lobsters (*Homarus gammarus*) from Helgoland (Germany) using the von Bertalanffy growth function (Schmallenbach et al., 2011)

2.2 Distribution

The European lobster is distributed throughout coastal areas of the northeast Atlantic from mid-Norway to the North Sea, south to the north African coast and into the eastern Mediterranean (Figure 7, Phillips , 2013). However, their 'commercial range' is currently limited to northern France, Britain and Ireland. The northernmost populations are found in the Norwegian fjords Tysfjorden and Nordfolda, inside the Arctic Circle (Agnalt et al., 2009). In Norway and Shetland stocks have declined significantly or collapsed (Figure 5). In the Mediterranean and off the Iberian peninsula, north to the Bay of Biscay, lobsters are rare and generally well below commercial levels. Their stronghold, therefore, is a relatively small proportion of their previous commercial distribution (Holthuis 1991, Prodöhl et al., 2006, Phillips, 2013).

2.3 Landings

2.3.1 Regulations and management on landings

European regulations on European lobster state (European Commission, 2006): "The catching, keeping on board, transhipping, landing, storing, selling and displaying or offering for sale of **berried female** crawfish (*Palinuridae* spp.) and berried female lobster (*Homarus gammarus*) shall be prohibited. Berried female crawfish and berried female lobster shall be returned to sea immediately after accidental catching or may be used for direct restocking and transplantation within management plans established pursuant to either Articles 18 or 19 of this Regulation."

Dutch regulations state that with all gears egg-bearing lobsters, freshly caught lobsters and lobsters caught between July 15 and the last Thursday of March are put back immediately after the catch⁴.

The minimum conservation reference size (MCRS) in Dutch Belgium and German waters 85 mm CL⁵ (Schmalenbach, 2009); in British waters is 87mm⁶; in Norwegian waters 88mm (Agnalt et al., 2009). In 2019, parts of the West and North coasts of Scotland and south west England (Cornwall) will have a MCRS of 90 mm CL and the East coast remains 87mm^{2, 7}.The American lobster *Homarus americanus* has a MCRS of 83 mm CL in Maine (Dunnington et al., 2010).

Also V-noticing is applied on female tails after which they are returned ensuring at least one possibility to bear eggs.

2.3.2 Landings

In the past 70 years, the total annual European landings varied between 1,600 and 5,600 tonnes (Figure 5, Prodöhl et al., 2006, Phillips, 2013). At the beginning of the sixties an annual catch of 3,000 -3,500 tons was not uncommon, but the annual catch dropped in the 1970s to less than 2,000 tons at the beginning of the eighties. Since then, a slow increase to 3,200 tons has been observed and after 2006 to 2010 a revival to about 5,000 tons (Phillips, 2013). Lobster catches vary considerably from country to country: between 1950 and 1975, Scotland accounted for 26% of total landings; Norway for 18%, followed by England, Wales and France with 16%, Ireland with 9% and Sweden, Denmark and Spain with less than 5%. Before the 1960s, Norway recorded annual catches of 600 to 1,000 tonnes, but during the following two decades a collapse of fishing was observed and annual catches are now less than 60 tonnes (Figure 5). Within the Mediterranean countries, the annually reported landings have never reached the same levels as those in the north-western distribution area (Prodöhl et al., 2006). In the year 2016 the Netherlands landed 64 tonnes (1.4%); the UK 3281 (71%); France

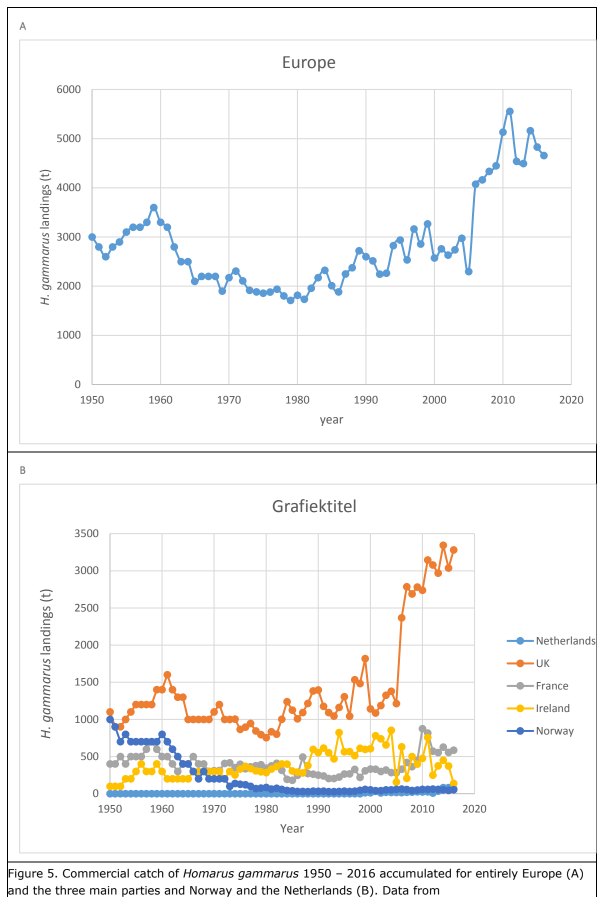
⁴ https://wetten.overheid.nl/BWBR0024539/2017-10-05 d.d. 20-12-18

⁵ http://www.zeevissers.com/minimummatenzeevis.html D.d. 07-12-18.

⁶ https://www.gov.uk/government/publications/minimum-conservation-reference-sizes-mcrs/minimum-conservation-reference-sizes-mcrs-in-uk-waters#crustaceans d.d. 07-12-18,

⁷ https://www2.gov.scot/Topics/marine/Sea-Fisheries/InshoreFisheries/crab-lobster-landingcontrols?fbclid=lwAR2Qrkl9mUois5g9wyKiO8V5SYRpUaWHTqFpNKLigFWhW6e2we0LcL9Lf78

587 tonnes (13%); Ireland 138 tonnes (3%) and Norway 54 tonnes (1.2%), summing with other countries altogether to 4656 tonnes landed in 2016.



http://www.fao.org/fishery/statistics/global-capture-production/query/en (d.d. 21-12-18)

2.3.3 Sizes caught

Giving some examples without trying to be extensive, the size distributions of male and female European lobsters caught on the Northumberland coast (North sea, UK) were around ~80-82 mm CL (Skerrit at al., 2012). Average sizes estimated were e.g. 83 to 88 mm CL in Cornwall and 77 mm CL in Yorkshire (Hepper, 1978), in Westermost Rough Offshore Windfarm Park (near the Humber, North sea, UK) most abundant class of lobsters after a period of closure for fisheries was ~87 mm CL class whereas the control site had the most abundant size class at ~80 mm CL (Roach et al., 2018). In Norway average sizes were encountered between 74 mm (Stefjord, a special genetic subpopulation) and 108 mm CL (Stavfjord) in Norway (Agnalt et al., 2009). The lobsters caught in lobster pots are usually 23–38 cm long and weigh 0.7–2.2 kg. Hepper (1978) showed 75-90 mm CL range being the most frequent commercially caught lengths in Cornwal (Figure 6). Schmalenbach et al. (2011) caught larger European lobsters at Helgoland with the main fraction between 90 to 130 mm CL.

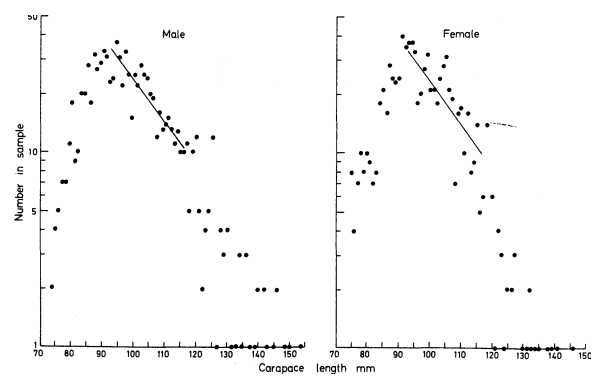


Figure 6. Size frequency distribution of lobsters measured from the Cornwal lobster fishery plotted on a logarithmic scale (the catch curve) (Hepper, 1978).

2.4 Biology and habitat

European lobsters are usually located at lower than mean low water neaps (sublittoral fringe) to depths of 165m; usually not deeper than 50 m (Table 3). The LD 50 for adult *H. gammarus* was at a salinity of 24‰ according to Mercer at el. (2001, temperature ranging from 9 to 11°C) and according Wolf and Sandee (1971) lobsters from the Oosterschelde and Veerse Meer can survive down to 9‰ during wintertime (~3°C). EBP stage European lobsters had a LD50 of 16‰ (Mercer et al., 2001). Low salinities will lead to reduced moulding and below 11‰ no more moulding will occur (Schuiling and Smaal, 1998). Older and larger lobster are encountered in deeper regions, although this could be an impact of catchment rather than habitat preference. They are vulnerable to current speeds of 0.6 m/s and higher and wave energy. Then they get carried away (Howard, 1988, Linely et al., 2008, van den Boogaard et al., 2019).

They are usually found on hard substrates: rock or hard mud, but also soft substrates are used for food and nursery grounds. The animals are nocturnal and territorial, living in holes or crevices (Holthuis, 1991, Lawton & Lavalli, 1995, Jensen et al., 2000, Turner et al., 2009, Phillips, 2013, Seitz et al., 2014, Skerrit et al., 2015). Holes and crevices are especially important for the younger stages that are vulnerable to predation and cannibalism (Table 4, Jensen et al., 2000, Mercer et al., 2001, Phillips, 2013). As lobsters grow they gain a size-refuge from predators, and in turn their association with shelter tends to relax. Therefore, smaller lobsters are more common in the middle of cobble patches, whereas large lobsters are more common on edges (Skerrit, 2014). In England, areas with habitats that include less structure and fewer large-scale outcrops for adults produce lobsters of smaller size than other areas, indicating the importance of the habitat for growth (Jensen et al., 1994). Crevices and ledges remain still important (see e.g. Howard, 1988, Jensen et al., 2000, Galparsoro et al., 2009). Table 4 (taken from Lawton & Lavalli, 1995) gives an excellent overview of the different life history phases of H. americanus with associated behaviour and ecological processes, which might be quiet valid for H. gammarus as well (Jensen et al., 2000, Phillips, 2013). Note that the sizes of the different stages are slightly different. And the column on realms occupied emphasis the seasonal movement of *H. americanus*, which is not applicable to *H. gammarus*, which remain mostly at one location once having encountered a suitable crevice.

Like other crustaceans, lobsters have a hard exoskeleton which they must shed in order to grow, in a process called ecdysis (moulting). This may occur several times a year for young lobsters, but decreases to once every 1–2 years for larger animals until becoming an annual part of the mating, spawning and egg hatching cycle (Hepper, 1978, Holthuis, 1991, Phillips, 2013).

Pre-adult and adult lobsters are reef-obligate and select sites that supply sufficient food and oxygen and shelter from currents and predation (Linley et al., 2008, Table 4). A number of criteria, including oxygen supply, length, entrance size, presence of multi-openings (escape routes), internal aspect ratio (manoeuvring space) are all aspects that may influence the suitability of cracks and crevices for lobsters (Howard, 1988, Lawton & Lavalli, 1995, Jensen et al., 1994, 2000, Linley et al., 2008). As lobsters grow (in steps following ecdysis) they need to move to increasingly larger crevices. As a given cohort of lobsters grows, it suffers mortality that is dependent on a number of factors. These include predation, competition and diseases.

Table 3 gives rather broad ranges for the tolerance of European and American lobsters towards abiotic factors⁸. European Lobsters have a large range of distribution (Figure 7) expressing both this tolerance and the suitability of local habitat. In general it is thought that suboptimal circumstances (in salinity, oxygen etc.) will lead to higher metabolic costs and reduced growth (Jensen et al., 2000, Linley et al., 2008). There estuarine environments, with altering salinities and particularly those associated with periodic reductions in temperature, these conditions are stressful these conditions are stressful and likely to be predominantly unsuitable environments

A first wreck suitability map for the Dutch NCP is made by Van den Boogaart et al. (2019, Figure 8) based on divers observations on wrecks. Suitable and less suitable spots appear next to each other whereas abiotic circumstances seem not that differentiating (Table 7, Smaal et al., 2017). More research is needed in order to determine habitat suitability and differentiating factors on the Dutch NCP.

⁸ Nb data on *H. americanus* are often used as a model or representative for *H. gammarus*. General feeling is that the species are very much alike and that the data *of H. americanus* can be used as such (Phillips, 2013). Though it should be noted that *H. americanus* is bigger, has migration and occurs to -700m, much deeper than -165m of *H. gammarus* (Phillips, 2013). This suggests at least some differences in physiology.

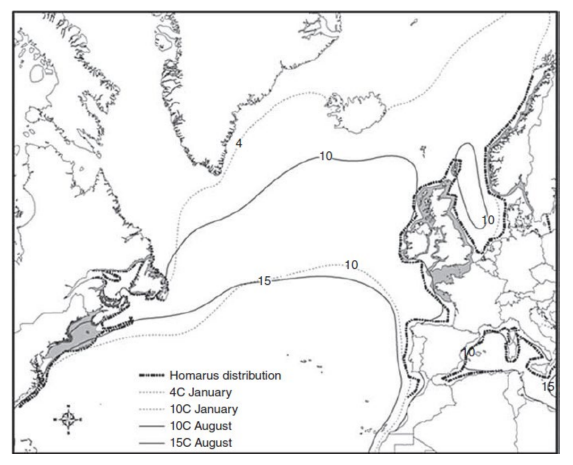


Figure 7 Geographic distribution of *H. americanus* and *H. gammarus* in the North Atlantic relative to seasonal isotherms. Dotted boundary is species geographic range; areas in grey are prime commercial fishing grounds (from Phillips, 2013).

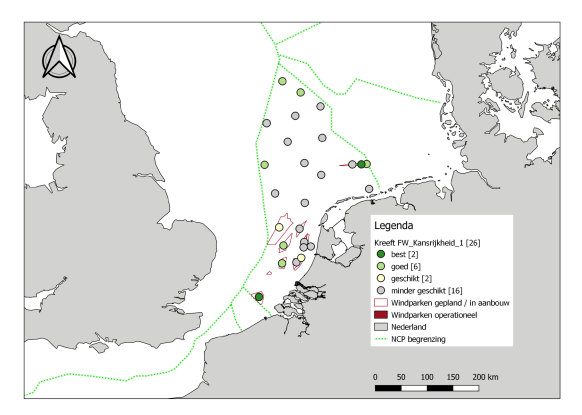


Figure 8 Opportunity map for catching North Sea lobster *H. gammarus* at (potential) offshore wind energy locations for the Dutch NCP (van den Bogaart et al., 2019).

Table 3. Ranges of relevant abiotic aspects under which *H. gammarus* is encountered.

Desirable levels for key v al. 1980 (<i>H. americanus</i>) compiled by Kristiansen Wolf and Sandee (1971) is claimed to be the optim) and Wickins & et al. (2004). Da and Schuiling a	Natural ranges in which <i>H. gammarus</i> is encountered (https://eol.org/pages/46505673_d.d_10-12-18)			
Parameter	Optimal condition	Natural range		Encountered range	
Temperature (°C)	18 - 22	1 – 25	<0, > 31	Temperature range (°C)	8.214 - 12.243
Salinity (‰)	28 - 35	28 - 35	<8, >45	Salinity (‰)	33.5 - 35.6
Salinity (‰) lobsters Oosterschelde	21- 27 -32	9-35			
O2 (mg l⁻¹)	6.4	4 -8.2	<1, >saturation	Oxygen (mg l ⁻¹)	5.4 - 6.7
pН	8	7.8 -8.2	<5, >9		
Ammonia (mg l ⁻¹)	<0.14	0 - 0.3	>1.4		
				Depth range (m)	0 - 165
				Nitrate (µmol l ⁻¹)	2.7 - 8.5
				Phosphate (µmol l ⁻¹)	0.32 - 0.63
				Silicate (µmol l ⁻¹)	1.9 - 4.5

Table 4. Life history phases of Homarus americanus with associated behaviour and ecological processes (from Lawton & Lavalli, 1995). Sizes at transition between all phases may vary geographically. For adolescent and adult lobsters, the sizes specified are the minimum carapace lengths (CLs) for entry into these life history phases. Size at maturity varies geographically, such that functional maturity may not be reached until ~100-mm CL in some areas.

Phase	Size (mm CL)	Activity pattern	Foraging mode	No. of shelters occupied	Realms (maximum abundance)	Ecological processes
Larval (stages I- III)	~2-4	Vertical migrator; poor swimmer; passive (?) drifter	Raptorial feeder	NIA	Pelagic (offshore to inshore depending on area)	Dispersal; development
Postlarval (stage IV)	~4-5	Strong surface swimmer; benthic settler	Raptorial/ suspension feeder	Selects preexisting shelter or excavates new one	Pelagic(inshore) to benthic	Settlement; predation while settling or shortly thereafter
Shelter- restricted juvenile	~4-14	Recent settler; remains under cover; subterranean movements within spatially complex habitats	Suspension feeder; browser within shelter; ambusher at shelter entrance	Usually one, sometimes several within a contiguous shelter space	Benthic (shallow)	Recruitment; predation
Emergent juvenile	~15-25	Mostly shelter confined; limited movements outside of shelter, but remains in close vicinity	Browser; ambusher	One to several	Benthic (shallow)	Predation
Vagile juvenile	~25- size of physiological maturity (~40)	Shelter user, but more extensive movements out of shelter for food	Ambusher; pursuer; searcher	One to several	Benthic(shallow)	Competition
Adolescent	Physiological maturity, but not functional maturity (~50)	Active, mostly nocturnal; may participate in seasonal movements with reproductive animals	Pursuer; searcher	One to many, depending on seasonal movement	Benthic (shallow to deep)	Competition; indirect fishing mortality due to retention in traps
Adult	Functional maturity (>50)	Active, mostly nocturnal; seasonal, reproductively mediated movements	Pursuer; searcher	One to many, depending on seasonal movement	Benthic (shallow to deep)	Mate selection; reproductive success; direct fishing mortality

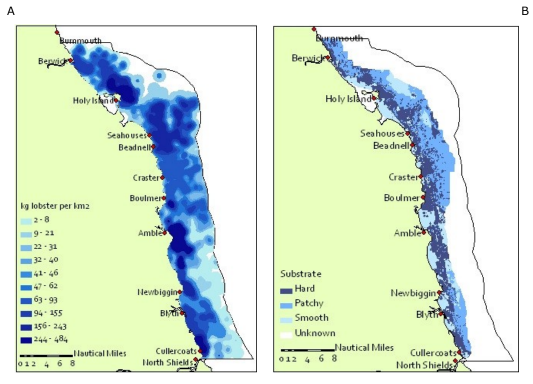


Figure 9. data from Northumberland, UK (from Turner et al., 2009).

A. Estimated distribution of lobster landings by port (mean annual totals 2004-2007) B.Larger scale substrate characterisation near Blyth

2.5 Densities

In general, density of the European lobster is irregular and seems to be regulated by the presence of hard substrate however not always (Figure 9, Figure 10, Figure 13, Turner et al., 2009, Skerrit et al., 2012, 2015, Wallace, 2015, Wright, 2018). In this sense it is important to define what is considered the seize of the area of reference and the resulting density.

An important finding is that there is little density data in literature on *H. gammarus*. There is a clear omission in this sense. We will give an overview of what was found on *H. gammarus* and also provide some densities of *H. americanus* (see also Table 5). Agnalt et al. (2009) estimated average density of lobsters in km⁻¹ shoreline, not in km⁻²: 169 (\pm 79) lobsters km⁻¹ shoreline in Stefjord, 189 (\pm 122) lobsters km⁻¹ in Mannfjord, and 99 (\pm 39) lobsters km⁻¹ in Mørsvikfjord. Turner et al. (2009) estimated landings of kg lobsters on Northumberland, UK, giving some indications for densities (Figure 9, Figure 10). They estimated landings per unit area from a minimum of 2 kg km⁻² yr⁻¹, to a maximum of 484 kg km⁻² yr⁻¹ (at an average weight of 0.5 kg per lobster is ~0.001 lobster caught m⁻²). The highest estimates of landings per unit area were at Holy Island, with maximum values at all other ports below 300 kg km⁻² yr⁻¹ (~0.0006 lobster m⁻² yr⁻¹).

In literature densities are encountered ranging from 0.00037 calculated with the entire surface of the Oosterschelde or (van Stralen and Smeur, 2008), 0.002 (Skerrit, 2014, Skerrit et al., 2012) to 0.27 lobsters m⁻² (Jensen et al., 1994) (see Table 5). The density given by Jensen et al., (1994) is high. It represents a situation with an artificial reef with high densities of crevices especially for lobsters. In Blyth a summer/winter difference was encountered in densities (Skerrit, 2014, Skerrit et al., 2012, Table 5, Figure 10). In winter a density of 0.002 lobsters m⁻² and in summer 0.006 to 0.007 lobsters m⁻². Jensen et al. (2000) and Howard (1988) give overviews on densities in which they cited personal communications ranging from around 0.03 lobsters m⁻² (unspecified British sites; R.J. Handford cited in Howard, 1988) similar to observations in Lough Hyne of 0.03 lobsters m⁻² (Robinson et al., in preparation, in Jensen et al., 2000, based on observed nearest neighbour distances ranged from 0.75 Wageningen Marine Research report C109/18 21 of 64

m to 35 m). In a natural, horizontal rock crack in Scottish waters, densities were encountered from 0.4 to 5.0 m⁻² (mean 1.5 m⁻² Comelyand Ansell, personal communication, cited by Jensen et al., 2000). Van Stralen and Smeur (2008) estimated a total of 130000 lobsters in the Oosterschelde. We calculated that to a roughly estimated 0.00037 lobsters m⁻² when using the entire surface of the Oosterschelde (350 km²)⁹. Using an estimated dyke surface (240 km total length according to Brummelhuis et al. (2013)) and an average of 9 m depth¹⁰, the roughly estimated densities are much higher: 0.060 lobster /m². It should be noted average depth takes also the shallow parts (like schoals and mud flats) of the Oosterschelde into account whereas most dykes are next to the deep tidal channels, reducing the density due to a larger average depth. In addition in the approach the dyke is a vertical wall whereas in fact it is an slowly sloping surface, also increasing the surface of the dykes thereby reducing the average. A tenth of the estimated density seems more appropriate (triple average depth and triple length of the slope). The densities on Dutch NCP soft sediments were roughly estimated at 0.000002 larger lobsters m⁻² (2 lobsters km⁻², P. Molenaar, WMR, pers. comm.).

For *H. americanus* densities were encountered up to 0.340 lobsters m^{-2} (Table 5). Given the migration behaviour of *H. americanus* densities vary in time and also depend on the substrate (Dunnington et al., 2005, Geraldi et al., 2009, Phillips, 2013).

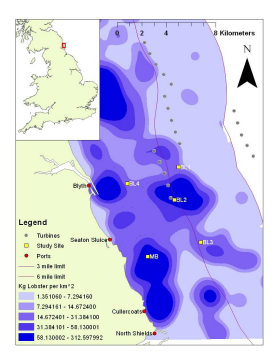


Figure 10. Lobster distribution near Blyth, Northumberland, UK based on landings, vessel sightings and average vessel home-range (from Skerrit et al., 2012). The spot B4 is where Skerrit et al (2012, 2015,) and Skerrit (2014) have performed their research. Figure 13 gives more details of location B4.

⁹ https://nl.wikipedia.org/wiki/Nationaal_Park_Oosterschelde d.d. 26-03-19.

¹⁰ https://duikeninbeeld.tv/scubapedia-oosterschelde/ d.d. 26-03-19

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are used.												
Species	Average density (ind/m2)	Average m2 per individual (m2/ind)	Size class average CL (mm)	Size range Low (mm)	Size range High (mm)	Remark	Substrate	Depth (m)	Location	country	sea	Reference
EL	0.267	3.75					Artificial reef: blocks made from stabilized coal-fired power station waste materials (stabilized Pulverized Fuel Ash (PFA))	-10	Poole Bay	United Kingdom	The English Chanal	Jensen et al. (1994)
EL	0.006	166.67	79	65	114	Summer	extensive areas of rock and cobble and soft substarte as well	-27.2 (-16.7- 31.8)	Northumberland coast, Blyth	UK	North Sea	Skerritt et al. (2012)
EL	0.002	500.00	79	65	114	Winter	soft/mixed habitat	-27.2 (-16.7- 31.8)	Northumberland coast, Blyth	UK	North Sea	Skerritt et al. (2012)
EL	0.007	142.86	82	64	114	Summer	extensive areas of rock and cobble and soft substrate as well	-27.2 (1-6.7- 31.8)	Northumberland coast, Blyth	UK	North Sea	Skerritt (2014)
EL	0.00037	2702.70	~80	<47	~150	Using total water surface of Oosterschelde (350 km ²)	dyke protection of bolders, cobble.	0 to-48,80	Oosterschelde	Netherlands	Oosterschelde	Van Stralen and Smeur (2008)
EL	0.060	16.62	~80	<47	~150	Using total dyke length (240 km) and an average depth of 9 m	dyke protection of bolders, cobble.	0 to-48,80	Oosterschelde	Netherlands	Oosterschelde	Van Stralen and Smeur (2008)
AL	0.155	6.45		11	83		Artificial reef: rocks, 5-100 an in diameter up to 15 an thiok	-18	Northumberland Strait	Canada	Gulf of St. Lawrence	Scarratt (1973)
AL	0.178	5.62				Natural lobster grounds	glacial till: coarse sand and gravel		Northumberland Strait	Canada	Gulf of St. Lawrence	Scarratt (1973)
AL	0.34	2.94		<75	>150		Artificial reef: pumice concrete 1 and 3-chambers	-6.4	Point Judith, Rhode Island	Canada	Northumberland Strait, Gulf of St. Lawrence	Sheehy (1976)
AL	0.227	4.41					Breakwater wall of the Harbor of Refuge with strong tidal currents, no significant wave action, coarse sand 300 µ	-5.6	Point Judith, Rhode Island	Canada	Northumberland Strait, Gulf of St. Lawrence	Sheehy (1976)
AL	0.065	15.38	≥50	50		mid-summer peak density	cobble/boulder, ledge, soft sediment	-30	Vinalhaven Island, at the mouth of PenobscotBay	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.09	11.11	66	50	<83	end of august	Cobble	-30	Carvers Harbor, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.065	15.38	66	50	<83	end of august	Ledge	-30	Carvers Harbor, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.04	25.00	66	50	<83	end of august	Sediment	-30	Carvers Harbor, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.026	38.46	≥83	83		end of august	Cobble	-30	Carvers Harbor, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.002	500.00	≥83	83		end of august	Ledge	-30	Carvers Harbor, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.005	200.00	≥83	83		end of august	Sediment	-30	Carvers Harbor, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.18	5.56	66	50	<83	Mid-july	Cobble	-30	Sheep Ledges, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)

Table 5. Different densities at which *H. gammarus* (EL) and *H. americanus* (AL) are encountered. Only traceable densities published in journals or grey literature are used.

AL	0.053	18.87	66	50	<83	Mid-july	Ledge	-30	Sheep Ledges, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.004	250.00	66	50	<83	Mid-july	Sediment	-30	Sheep Ledges, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.028	35.71	≥83	83		Mid-july	Cobble	-30	Sheep Ledges, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.023	43.48	≥83	83		Mid-july	Ledge	-30	Sheep Ledges, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.005	200.00	≥83	83		Mid-july	Sediment	-30	Sheep Ledges, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0		66	50	<83	Mid-august	Cobble	-30	Browns Head, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.156	6.41	66	50	<83	Mid-august	Ledge	-30	Browns Head, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.016	62.50	66	50	<83	Mid-august	Sediment	-30	Browns Head, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0		≥83	83		Mid-august	Cobble	-30	Browns Head, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.017	58.82	≥83	83		Mid-august	Ledge	-30	Browns Head, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.006	166.67	≥83	83		Mid-august	Sediment	-30	Browns Head, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.09	11.11	66	50	<83	end of august	Cobble	-30	Dogfish Island, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.066	15.15	66	50	<83	end of august	Ledge	-30	Dogfish Island, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.037	27.03	66	50	<83	end of august	Sediment	-30	Dogfish Island, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.016	62.50	≥83	83		end of august	Cobble	-30	Dogfish Island, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0		≥83	83		end of august	Ledge	-30	Dogfish Island, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)
AL	0.007	142.86	≥83	83		end of august	Sediment	-30	Dogfish Island, Vinalhaven Island	Maine, USA	Atlantic Ocean	Dunnington et al. (2005)

2.6 Reproduction and juveniles

2.6.1 Mating

In most areas lobsters do not mature before 5–8 years and males reach sexual maturity earlier than females (depending on water temperature). Lobster are long-lived and iteroparous, capable of reproducing many times. Size at functional and physiological maturity of females also varies geographically presumably related to temperature. Females can start bearing eggs at 71 mm. The size range of mature females (bearing eggs) found in the wild ranged from ~71 (mostly North Sea) to 150 mm CL. The range of estimates of L₅₀ for functionally mature female lobster from different locations range from 95 to 140 mm (Figure 11, Contarini et al., 2008, Lizarraga – Cubedo et al., 2003, Laurans et al., 2009, Phillips, 2013, Wood, 2018).

The lobster mating system is polygynous whereby larger, competitively dominant males are more successful in obtaining mating shelters and in courting females. Also larger claws lead to higher mating success (Phillips, 2013, Sørdalen et al., 2018). Male *H. americanus* have sperm present in the vas deferens at 45 – 50 mm CL, but given their small size mating is not likely to occur yet. Data for *H. gammarus* are not available. Reproduction takes place during summer (around July) and is linked with the moulting cycle. Mating usually takes place shortly after the female molts; a spermatophore is deposited internally by the male in the seminal receptacle of the female. Sperm may be stored for several years. After extrusion, the eggs are held on the pleopods for approximately another year until hatching the following summer. In general, moulting cycles are two to three years from smaller to larger females. Large females (>120 mm carapace length) have been shown to moult and then undergo two successive spawns before moulting again, suggesting the capacity for sperm storage. Both *Homarus* species egg sizes are large, clutch sizes small, and brood period long relative to other marine decapods of similar size (e.g., spiny lobsters and crabs) (Prodöhl et al., 2006, Phillips, 2013). Clutch sizes range from a few thousand to 20,000 larvae, increasing with CL and age (Contarini et al., 2008, Figure 11).

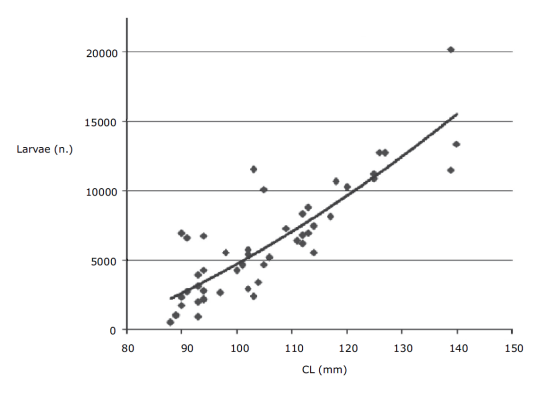


Figure 11. Number of hatched larvae (stage I) in function of maternal carapace length (CL) (taken from Contarini et al., 2008).

2.6.2 Pelagic larval stage

The first few post-hatching weeks are characterised by a pelagic phase usually lasting 10-18 days depending on water temperature (three larval instars and one post larval instar are pelagic)(Wickins & Lee, 2002, Skerrit, 2014, Figure 12). During this period, larvae undergo metamorphosis to a lobster-like body at the molt from Stage III to post larva. There is a distinct change in body-structure, with the claws and tail moving from their dangling positions to point out straight from the thorax and abdomen respectively. This alteration towards the post-larval form of an adult allows the post larva to swim forwards using the pleopods and seek a suitable substrate to settle (gravel or coarse sand).

Literature and transport models suggest that most larvae are generally transported between 6 to <100 km (Jensen et al., 1994, Krone & Schröder, 2011, Phillips, 2013). Local currents and residual currents seem to determine distribution. Local availability of substrate and benthic habitat determine settlement. For example, patterns of post larval concentration predicted for the Gulf of St. Lawrence bore little resemblance to spatial patterns in the harvest, suggesting other factors, such as the availability of benthic habitat, may ultimately limit adult abundance (Phillips, 2013).

Larval production and transport are critical components of the dynamics of lobster populations. It has been estimated that only 0.005% of the hatchling lobsters survive the planktonic phase to reach the benthic phase (Phillips, $2013)^2$.

2.6.3 Early settlement

The metamorphosis is accompanied by changes in behaviour and habitat preference. Two to four days after metamorphosis to post larva bottom-seeking behaviour appears, in order to find appropriate nursery habitat (gravel or coarse sand). Then the post larva makes the transition from pelagic to benthic environment and becomes a permanent resident of the sea bed. Based on indirect measurements and extrapolations, larval settlement strength of *H. gammarus* is associated with local sea temperatures (production and mortality) and onshore winds (transport and availability) as is described for *H. americanus* as well (Sheehy and Bannister, 2002, Phillips, 2013). Mercer et al. (2001) observed that predation on unprotected EBO lobsters was generally rapid. They suggested that time can be a critical factor during settlement, and high mortality can occurs where suitable shelter or substrate is unavailable.

The juvenile phase of the life cycle is characterized by an early shelter-restricted period in which the lobster emerges infrequently from its burrow, followed by an increase in range of movement and habitat choice. In the beginning young *H. gammarus* are cryptic and spend their initial year of life almost totally underground in burrows feeding on infauna and possibly bacteria (Table 4). Small lobsters are well adapted to a burrowing existence. However at a certain moment the food reserves within and close to the burrow become increasingly incapable of providing complete sustenance. Food quantity, availability and nutritional variety within the burrow are major factors influencing when the juvenile lobster emerges to forage. Lobsters then starts to forage further afield seeking shelter whenever necessary to avoid strong currents, competition and predators. At this stage they will eat marine worms and other post-larval animals like small crabs, urchins and gastropods, as well as retaining the ability to filter-feed on plankton. In general at a carapace length of about 15mm they leave their burrows for crevices in rocky substrate to begin life as an adult. (Howard, 1988, Jensen et al., 2000, Phillips, 2013,²).

Despite multiple and widespread investigations, no extensive information is currently available on the early benthic phase (EBP) of the European lobster from settlement at 5-7mm CL until 20mm⁺ and juveniles are hardly found up to 40-45mm CL. It is not clear whether they simply are uncommon, or rarely settle in densities high enough to be detected. Alternatively, they could be in unexplored habitats. While densities of American lobster (*H. americanus*) juveniles are commonly found and used in stock assessments, it is not feasible to use EBP or early juveniles to predict future recruitment in *H. gammarus*. (Mercer et al., 2001, Prodöhl et al., 2006, Phillips, 2013).

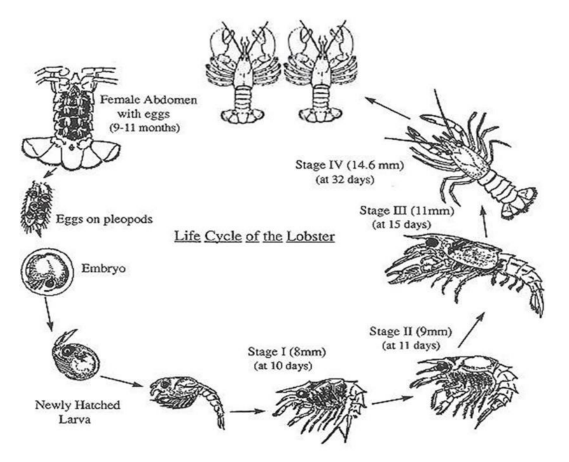


Figure 12. The life cycle of *Homarus* spp. Newly hatched larva are the beginning of the pelagic stage, and stage III represents the ontogenetic shift to a benthic stage (from Skerrit, 2014).

2.6.4 Refugia

Sheehy et al (1999) suggested that the lobster's natural longevity can be interpreted as a life history optimization strategy to enhance reproductive success. Larger (older) females do have higher number of eggs (Figure 11). This emphasizes the demographic importance of old lobsters as reproductive strongholds in closed areas like e.g. wrecks or selected OWFs more offshore. The meta-population analysis of *H. americanus* suggested that heavily fished inshore stocks could depend critically on larval input from offshore. Given the slow growth and reproduction these population 'refugia' could take many years to replace when fished out. This may also explain why some stocks have never recovered from heavy exploitation despite significant effort reduction. Refugia can also serve as safeguards for the genetic diversity (e.g. males and females can grow to larger sizes thereby safeguarding genes for larger growth, Sørdalen et al., 2018)

2.7 Diet

There is little detailed data on the diet of *H. gammarus*, while more information can be found for *H. americanus*. Benthic American lobsters are omnivorous, feeding on a great range of benthic organisms. The diet spectrum encompasses a broad range of plants and animals (even algae and zooplankton) but is dominated by molluscan, crustacean, fish, echinoderm, and polychaete remains (Table 12 in Annex 2). They are also cannibalistic. The range of prey species and amount consumed changes as a function of lobster size (age), temperature, prey availability, habitats, season and molt cycle (Holthuis, 1991, Elner & Campbell 1987, Sainte-Marie and Chabot, 2002, Hanson, 2009). At one location mussels (*Modiolus modiolus, Mytilus edulis*) appeared the most consistently important identified prey species, in terms of frequency-of-occurrence and estimated volume (points) indexes, while at another location crustaceans were the most important species (Hanson, 2009).

Lobsters eat until satiated hiding any left-over food for consumption at a later time. The growth and survival of juvenile lobsters is greater where there is a consistent and plentiful food supply, the latter occurring as it precludes the need for risky excursions in pursuit of food (Phillips, 2013).

There is some debate on whether *H. americanus* is a selective feeder or not. Elner and Campbell (1987) state that lobsters are selective feeders with a complex foraging behaviour and they are capable of maintaining gross diet composition despite strong variations in prey availability. Others think that lobsters are either simply opportunistic omnivores whose 'stomach contents reflect the relative abundance of prey species in the habitat' (Miller et al. 1971,) or 'scavengers' (Herrick, 1895, cited by Elner and Campbell, 1987) and 'most unspecialized feeders' consuming 'almost anything, regardless of its possible nutritional value.

For this study it is sufficient to conclude that bivalves are part of the lobster diet. Blue mussel is mentioned (Elner & Campbell 1987, Sainte-Marie and Chabot, 2002, Hanson, 2009).

2.8 Mobility

Mobility of European lobster can be divided in daily activities and more migratory movements. European lobsters are sedentary animals with migration ranges varying from 0 to 45 km. They prefer to remain at the locations where they have settled and are living. In general *H. gammarus* has restricted movements around its territory; <4 km for periods of up to a year (Figure 13, Jensen et al. 1994, Jensen et al., 2000, Smith et al., 2001, Skerrit et al., 2015). On a daily basis *H. gammarus* seems to make short random movements away from shelter with a maximum range of 125 m (Skerrit et al., 2015, Lees et al., 2018), which could be influenced by local competition for food, shelter and mates. They are capable of fast propulsion using their telson, but cannot maintain this momentum, relying on walking for sustained movements. E.g. *H. americanus* typically walks in five minute bouts; with a mean walking speed of 0.9 m min⁻¹, increasing to 2.5 m min⁻¹ (Skerrit, 2014). In term of long term movement and migration, Bannister et al. (1994) observed most recaptures of tagged lobsters within 6 km of known release positions. Only a small number of larger individuals have been observed to travel up to 15.7 km in a season (Jensen et al., 1994) or even up to 45 km (Smith et al., 2001), presumably finding new territories.

As an illustration the findings reported in Skerrit et al. (2015) are discussed in more detail. They observed seasonal patterns in activity: an utilisation distribution ranged from 244 to 7,722 m² during spring (mean \pm SE: 11,104 \pm 397 m²), and declined to 237–784 m² during autumn (mean \pm SE: 455 \pm 66 m²). They also found behavioural differences between sexes, with males using more space than females. Both sexes conduct the majority of their activities within similar areas, male lobsters had a greater propensity for travelling further from shelter and were more likely to have multiple cores of usage (Figure 13). Taking the diversity in habitat use and high individual variation is seems that populations are governed by individual 'personalities traits' in the form of variation in boldness, habitat-use, exploration and movement which in turn are likely governed by environmental opportunities and individual fitness. As demonstrated in Figure 13 *H. gammarus* travels over soft habitat. Studies of colonisation by lobsters of an artificial reef in the UK, positioned 3 km away from suitable lobster ground, also show that ability (Jensen et al., 1994).

On population level, both adults and larvae are considered to be able to use wrecks, OWFs and other hard substrates like oil and gas platforms as stepping stones to reach all parts of the North Sea (Krone & Schröder, 2011).

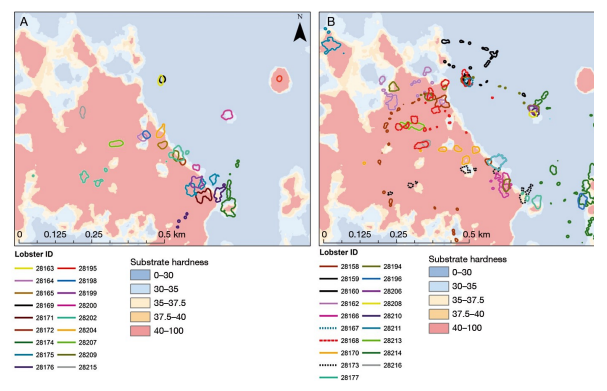


Figure 13. 95% utilisation distribution (95UD) home-ranges of (A) 18 female and (B) 19 male European lobsters during the spring study period (from Skerrit et al., 2015)

2.9 Population genetics

H. gammarus can be divided into six to seven genetically distinct clusters. Ellis et al. (2017) distinguished four genetic cluster (Figure 14): An Atlantic and Swedish cluster (with high connectivity) and in between those a transitional zone within the eastern North Sea, and an isolated Aegean cluster. The transitional zone between the Atlantic and Swedish cluster suggests an isolation by distance and suggests that direct gene exchange between these stocks is limited which fits a stepping-stone model. Prodöhl et al. (2006) and Agnalt et al. (2009) suggested two more distinct populations: one in the north of Norway (the "midnight-sun lobster") and another one in the Oosterschelde. Triantafyllidis et al. (2005) suggested even two different subpopulations in the north of Norway. Though the separation between the two clusters is only 142 km of shoreline, probably the hydrological characteristics of the two fjord systems prevent sufficient exchange of larvae. Taking into consideration the potential for migratory behaviour and larval dispersion, genetic exchange between European lobster populations seems unexpectedly restricted (see also Krone & Schröder, 2011).

Based on the low degree of genetic differentiation revealed in the European lobster and its limited capacity for dispersal, Triantafyllidis et al. (2005) suggested that all populations have been established from a common refuge after the end of the last Ice Age (< 15,000 years ago). The northern Norway, Netherlands and Aegean groups exhibit reduced gene diversity (Triantafyllidis et al., 2005, Prodöhl et al., 2006). Although the overall level of genetic differentiation among European lobster populations is low, still the genetic differences seem important adaptive aspects. It is extremely likely that lobsters living at the edges of environmental tolerance for the species are adapted to some degree to these differences in conditions (Triantafyllidis et al., 2005, Prodöhl et al., 2006, Agnalt et al., 2009).

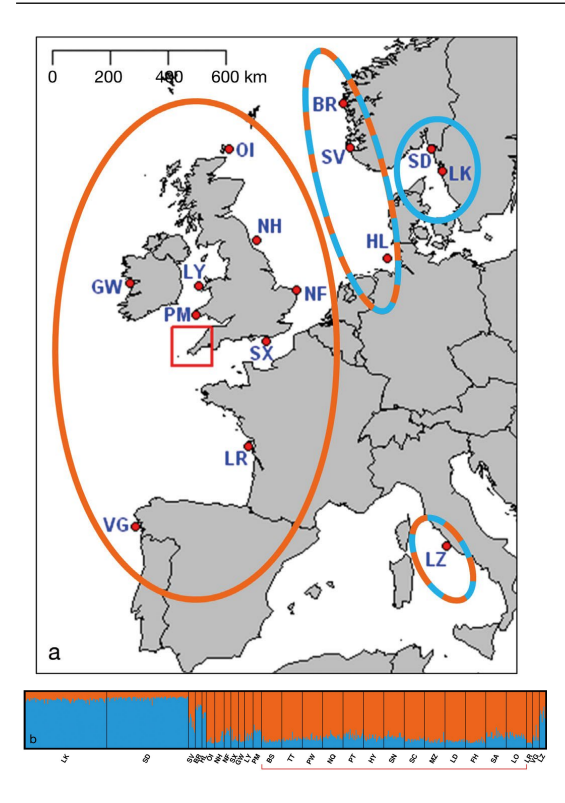


Figure 14. Assignment and map of population clusters. (a) Sample groupings as indicated by colour composition of individual samples (vertical bands) in (b). (b) Destruct plot of 5 converged iterations of K = 2 using a priori location data. The inset red area in (a) and red underlined section in (b) denote fine-scale samples from Cornwall, UK. (Ellis et al., 2017).

2.10 Enhancement of stocks

Lobster production can be increased actively (aquaculture) or passively (enhancing local stocks). For aquaculture two forms can be distinguished: **product enhancement** (wild caught lobsters are maintained in pounds where they are fed to improve quality/size). And secondly there is **full grow-out** (close cycle culture): rearing lobsters from egg to marked size. Until recently, full grow-out

culture of lobsters was not considered economically viable given the logistical implications related to the need to keep individual lobsters in separate compartments due to their cannibalistic behaviour (Wickins & Lee, 2002) and the lack of automated procedures for feeding and maintenance. In optimal rearing conditions (e.g. at constant 20°C), it is possible to rear a portion size lobster (250-300g, total length 210mm, CL 75mm, Prodhöl et al., 2006 or 345-400g, Wickins & Lee, 2002) from hatching in 800-900 days.

For enhancing local stocks three strategies have been defined: **habitat enhancement**, **stock enhancement** or **food enhancement**, which will be described in more detail below.

2.10.1 Habitat enhancement

Habitat enhancement has several aspects: 1) increasing the volume and surface available for lobsters 2) increasing the volume and surface available for food, and 3) offering crevices and caves which are suitable for all sizes lobsters emphasising especially the larger lobsters (Howard, 1988, Jensen et al., 1994, 2000, Wickins & Lee, 2002). When lobsters are present in the neighbourhood they will colonize the suitable new substrate rapidly by immigration of adult specimen not depending on larval settlement (Scarratt, 1973, Sheehy, 1976, Jensen et al., 1994, 2000, Phillips, 2013). The antiscouring of OWF monopiles is considered to be good potential new artificial habitat which can support lobsters (Linley et al, 2008, Buck et al., 2017, Roach et al., 2018). However not always European lobsters are encountered at such locations, raising doubts on suitability of the location or reachability (Bouma & Lengkeek, 2012, Hooper & Austen, 2014). Since the Dutch NCP is rather scarcely occupied with hard substrates, adding extra substrates could improve the potential for populations of H. gammarus, using e.g. the designs as mentioned in Jensen et al. (2000) or Buck et al. (2017). Lobsters themselves can learn us on the requirements of the habitats, e.g. on a cohesive clay/mud substrate, juvenile H. americanus create hydro dynamically advantageous burrow systems that facilitate the exchange of oxygenated water (Lawton and Lavalli, 1995). Optimum habitats for H. gammarus will therefore include mature, heterogeneous cobble-boulder layers overlaying penetrable substrate offering crevices for all sizes of lobsters (Howard, 1988, Jensen, 2000, Wickins & Lee, 2002).

The designs and materials used provide lobster habitat worldwide range from specially designed, new concrete (Reef Ball type), artificial reefs designed and built by engineers from non-waste materials, or unspecific casually deposited large, constructed from low-cost 'materials of opportunity' (like rock rubble or stabilised, pulverised fuel ash) (Wickins & Lee, 2002, Rozemeijer et al., 2017). More research is needed to get more insights of the required spatial needs (hydrodynamic conditions, tolerable nearest neighbour distances, foraging behaviour) and habitat needs (crevice size and shape preferences, food availability) and thus carrying capacity of a reef structure for lobsters of different sizes to survive and grow within a defined area (Sheehy, 1976, Wickins & Lee, 2002). To illustrate demands on habitats and their positioning, habitat enhancement constructs in Canada having their entrances towards the main hydrological forces were tumbled over after storms whereas otherwise oriented constructs remained unaffected (Sheehy, 1976, Wickins & Lee, 2002).

Monitoring is needed within OWFs to determine the local populations of European lobsters to determine whether this form of enhancement can be of service to enhance local productivity.

2.10.2 Stock enhancement

As is mentioned in sections 2.6.2 and 2.6.3 the pelagic stages and early benthic stages seem especially vulnerable. Therefore frequent hatchery programmes have been started in order to increase local populations both for *H. gammarus* as well as *H. americanus*, in order to improve local fisheries and restore stocks (Bannister and Addison, 1998, Jensen et al., 2000, Wickins & Lee, 2002, Phillips, 2013). These lobster hatcheries can be aiming at hatching eggs, and releasing stage I or stage IV larvae to supplement wild stocks (Prodhöl et al., 2006) or release more full grown animals at a size of e.g. 50 mm TL (Buck et al., 2017). Currently, programmes are in development to train the juvenile lobsters to enhance survival when being transferred from hatchery with low competition for food and shelter to the natural environment with food shortage, predators etc. (e.g. Agnalt et al., 2017). Despite the mobility of larvae and adults, (re)colonisation can be reduced, therefore stock

enhancement could be an option (Hooper & Austen, 2014, Buck et al., 2017). It seems important to use local lobsters in order not to have genetic pollution or less adapted animals to local circumstances (Ellis et al., 2017).

An important issue to tackle, before starting stock enhancement, is the cost–benefit analyses which should encompass aspects like:

- objectives of the release programme;
- scale of the operation;
- ownership rights;
- policy on population genetics and ecological impact of the released species;
- size and number of hatcheries;
- size at which the crustaceans are to be released;
- social benefits and externalities.

It can be worthwhile to do stock enhancement. A female lobster can have up to 20,000 eggs (Figure 11), however only one of these is expected to survive in the wild. Stock enhancement techniques are thought to improve this survival rate by about 1000 times. However a large investment in numbers released seems needed to improve stocks at the scale of areas. Bannister and Addison (1998) suggested a release of 10 000 lobsters per year on area level for 5 years, to demonstrate a significant measurable result. They were quit critical on claiming potential success. Others do see potential improvement of local populations (Wickins & Lee, 2002, Duffill-Telsnig, 2014, Buck et al., 2017). The National Lobster Hatchery Cornwall claims a >30% increase in lobster catchments whereas catchments on other crustaceans are declining due to their efforts².

Monitoring on focussing on juvenile lobsters is needed within OWFs to determine the local populations of European lobsters to determine whether this form of enhancement can be of service to enhance local productivity.

2.10.3 Food enhancement

In principle *H. gammarus* has a broad diet, yet food availability might be limiting. Given the large size and energy density of the flesh, an ample supply is necessary. In the TKI Win-Wind project food enhancement is one of the enhancement strategies to be investigated. Food enhancement can be achieved more passively: 1 increase of hard substrate and habitat; or by actively introducing 2. favoured food; or 3. extra food in the local environment. Fishery discards that are now to be landed without direct purpose could be a cheap source.

Seeding with e.g. the favoured mussels could promote food stock present. Disadvantage is e.g. that mussels can attract other predators too like fish and starfish (Wickins & Lee, 2002,). In addition seeding actively could destabilize a local system that is developing a certain direction due the prevailing (a)biotic circumstances at that moment.

Adding extra mussels to enhance local populations of favoured species or e.g. fisheries discards nearby could reduce the risks for lobsters running out of food. In addition feeding time could be enhanced when having food nearby due to providing discards nearby. Lobsters in particular cannot forage in high current speeds (Howard, 1988). Close supply of e.g. discards could reduce the need to forage away from the burrow, save foraging time and walking, and thereby extend the period of time over which they can safely feed before tide currents are too strong (> 0.6 m s⁻¹, speeds that can occur on the Dutch NCP, Howard, 1988, Wickins & Lee, 2002).

This short chapter gives the relevant data needed for the modelling of lobster on monopiles. The characteristics of the OWF Princess Amalia are treated in addition to a an example of habitat enhancement: Reef Balls.

3.1 OWF Princess Amalia

The aim is to model the growth of lobster in OWFs at monopile antiscouring. To this end information from the existing OWF Princes Amalia is used, although information of other parks is also available (Figure 1). OWF Princess Amalia was selected for a number of reasons, namely:

- 1. In this OWF a concrete test on lobster and brown crab fishing will start (TKI project WinWind).
- 2. Already data are available on the biodiversity and biomasses on the anti-scouring and other hard substrate are available.
- 3. In 2019 once again the anti-scouring and other hard substrate will be measured.
- 4. It is integral part of the planned OWF Hollandse Kust Noord-Holland (HK-NH) (Figure 1).
- 5. Being located at 19-24 m depth, it is in the transition zone between near shore and offshore.
- I is also reasonably close to the measuring station (bottom lander, Witbaard et al., 2014) from which data are used to derive as natural as possible annual developments of ChIA, SPM and T.

Relevant aspect for OWF Prinses Amalia are given in Table 6 and Table 7. The most relevant aspect is that the total anti-scouring surface in the park is 0.149% (0.073% for Luchterduinen) a minimal amount so the local consumption will hardly influence the primary production.

Table 6. Relevant aspects of OWF Prinses Amalia and OWF Luchterduinen (data obtained from Eneco). The data of OWF Luchterduinen are added because for the TKI project WinWind OWF Prinses Amalia is choosen and OWF Luchterduinen is the alternative.

Aspect	Unit	Prinses Amalia OWF	Luchterduinen
Number of windmills	N	60 (All 61 have scour protection included substation)	43 (ONLY 42 have SCOUR protection included OHVS)
Average distance between the monopoles	m	550-600	650
Capacity of the OWF	MW	120	129
Depth	m	19-24	19-26
Distance to the shore	km	23	23
Diameter scouring	m	21	18
Surface area antiscouring per pole	m ²	346.4	260.2
Characterisation of anti- scouring		one layer of filter with an armour layer of rocks on top of it. The median rock diameter is around 0.5 m.	0,3 m layer of filter layer with an armour layer of 1.2 meter. The rock layer has boulders of a median diameter of 40 cm
Average size of crevices (width*height*depth)	cm	8 x 8 x8 assume perfect circle	6 x 6 x 6 assuming perfect circle
Range width of crevices	cm	8-9	6-8
Range height of crevices	cm	8-9	6-8
Range depth of crevices	cm	8-9	6-8

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Aspect	Unit	Prinses Amalia OWF	Luchterduinen
Total Surface area Anti- scouring OWF	m ²	21128	10927
Surface OWF (km2)	km ²	14,218	15,035
Surface OWF (m2)	m ²	14218000	15035000
% anti-scouring vs total are OWF	%	0.149%	0.073%

Table 7. Abiotic aspects of some selected existing and planned OWFs: sea bed shear stress, suspended particles, water temperature (winter minimum and summer maximum), sea bed structure and motion, sediment composition (from Smaal et al., 2017).											
OWF	sea b	ed shear stro	ess	water		sea bed	sediment				
	(N/m	2)		matter	(mg/l)	tempe	rature (°C)	motion	composition		
	max	max wave action	avg	max	avg	min	max	cm/day			
Borssele	2	2	0,6	25	10	4	20	0 (local)	coarse to fine sand		
HK-ZH	4	5	0,6	35	10	4	18	?	fine sand		
OWEZ	8	8	0,8	50	20	3	20	?	fine sand		
HK-NH	5	6	0,6	35	10	4	18	?	fine sand		
Luchterduinen	5	6	0,6	35	10	3	20	?	fine sand		
Prinses Amalia	5	7	0,6	35	10	3	18	?	fine sand		
Buitengaats	4	7	0,4	40	10	3	18	?	fine sand		
Zee-energie	4	6	0,3	40	10	3	18	?	silty sand		

3.2 Habitat enhancement through Reef Balls

In order to enhance the habitat several options are possible that can serve as a lobster enhancing habitat (Jensen et al, 2000). The most important aspects are that several size (age) classes of lobster can be accommodated. Especially the younger stages need the protection of the crevice. Also the shape of the crevice is preferably elongated (Jensen et a., 2000, Linley et al., 2007). However, enhancing habitat for a large lobster is not easily defined given the harsh conditions near the bottom of the NCP. E.g. the monopiles need location specific anti-scouring protection with strict design rules (Lengkeek et al., 2017). Also placing constructions on the anti-scouring or directly on the sandy bottom needs a careful approach (Raaijmakers, Deltares, pers. comm.).

In the Netherlands extensive research is performed using e.g. Reef Balls as an example of general habitat improvement. Given the fact that we are currently in the preliminary stages of constructing an ecological habitat for lobster, Reef Balls might serve as an example and the characteristics can be included in our calculations. Natural reefs are variable in size, shape and hole density. Artificial reefs function better when they mimic nature. Reef Balls can be created with varying hole sizes to host differently sized lobsters. Reef Balls are made by pouring concrete into a fiberglass mold containing a central Polyform buoy surrounded by various sized inflatable balls to make holes (Figure 15, Table 8).

In general Reef Balls are designed such that over half of the weight is in the bottom of the ball and near the sea floor. All sizes of Reef Balls have withstood, without movement, heavy tropical storms in as little as 20 feet of water without anchors. Reef Balls are stable because the opening in the top of the unit breaks up the lifting force of the hydrofoil effect common to dome shapes. In essence they are chimneys and an under-pressure is generated by the passing currents thereby promoting stability. Side holes are wider near the centre of the walls and narrow near the units surface. This feature creates miniature vortexes which further reduce lifting forces and bring rich nutrients to organisms living on the reef.

Table 8. Dimensions of several types of reef balls. Data from: http://www.reefball.org/brochure.htm D.d.: 26-09-18.

D.a.: 26-09	9-18.						
Style	Width (m)	Height (m)	Weight (kg)	Concrete Volume (m3)	Surface Area	# Holes	Mold prices
Goliath							
Booster	2	. 1	1,818-2,727	1.19	16.7	15-25	
Ring							
Goliath	1.83	1.52	1,818-2,727	1.19	21.4	25-40	\$12,000
Super Ball	1.83	1.37	1,818-2,727	1.19	17.6	22-34	\$10,000
Ultra Ball	1.68	1.31	1,591-2,045	0.76	13.9	22-34	\$8,280
Reef Ball	1.83	1.16	1364-1,909	0.57	' 12.1	22-34	~\$8,280
Pallet Ball	1.22	0.88	682-1,000	0.25	7.0	17-24	\$7000
Bay Ball	0.91	0.61	170-341	0.08	2.8	11-16	\$3630
Mini-Bay	0.70		60.01	less than 4	ŀ	0.10	\$2300
Ball	0.76	0.53	68-91	50 lb. bags	5	8-12	
	0.61	0.40		less than 2		C 10	\$1222
Lo-Pro	0.61	. 0.46	36-59	50 lb. bags	5	6-10	
	0.40	0.20	14.20	less than 1		6.0	\$898
Oyster	0.46	0.30	14-20	50 lb. bag	I	6-8	

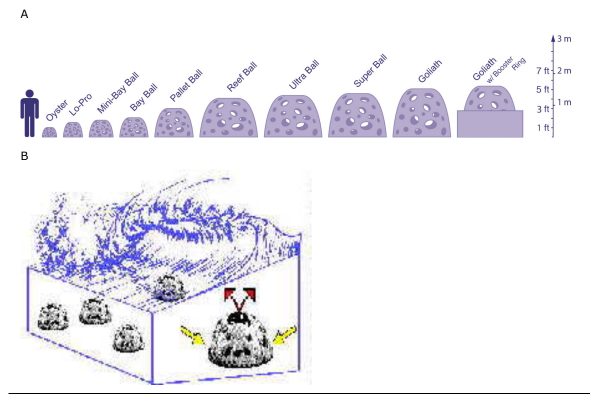


Figure 15. A. Different types of Reef Balls as described in Table 8. B. Chimney functioning of the shape yielding under pressure in the dome of the Reef Ball.

Modelling lobster production

An energetic population model was developed in order to assess lobster growth on a monopile given seasonal variation in temperature and its effects on lobster rates and resource productivity. Such a model allows for the necessary size based approached since fisheries is a size based activity. In order to incorporate effects of temperature and resource availability, as well as size, a DEB approach was used. Such a model can be used to model individual lobster growth under different circumstances, such as different temperature regimes to mimic different locations. A benefit of this model type is that modelled growth trajectories can be compared to growth trajectories in the field, thereby allowing model validation. In order to predict lobster density at a monopile food competition needs to be considered. To that end, the lobster growth model needs to be incorporated into a population dynamical shell and a dynamically modelled resource needs to be added. The population dynamical shell also allows to study the effects of stocking and differences in resource availability. Habitat enhancement, for example with Reef Balls, could also be incorporated in a later stage. Alternative modelling approaches are possible, but only a fully size-structured approach, such as DEB, results in growth curves that can be compared to field data. If there is a need to incorporate more food web components a different approach, such as stage-structure, will complexity but will also result in a loss of detail. This is a trade-off that is common to make to ensure feasibility (Lipscius et al., Under Review).

4.1 Method and parameters

4

The production of lobster is modelled for a single monopile. Model results will show lobsters growth and the number of lobsters that can grow given resource availability and temperature regime. Resource and lobster growth are modelled as a function of temperature.

Lobster growth is modelled using a dynamic energy budget (DEB; Kooijman (2009)), which models individual growth and development. Model and parameter values were taken from the Add-My-Pet website (Kooijman and Augustine, 2017) (hereafter referred to as AMP). In order to allow growth to be temperature dependent, the scaling with Arrhenius function was added to the model and accompanying parameter values were estimated. The DEB model does not explicitly include moult.

The individual growth model was embedded in the Escalator Boxcar Train concept (EBT; (de Roos et al., 1992)). With this concept the fate of individuals and their resource can be followed over time, making the population dynamics an emergent property. The EBT model allows for the modelling of a dynamic resource, so that individual energy uptake leads to resource depletion and potentially induces resource competition among individuals. A similar approach was adopted to model the dynamics of brown shrimp (*Crangon crangon*) along the Dutch coast (Steenbergen et al., 2015). The equations are listed in Table 9 and the parameter values in Table 10.

The resource is modelled following semi-chemostat dynamics, mimicking *Mytilus edulis*, which is the most abundant species in number, biomass and energy at the anti-scouring surface (Bouma and Lengkeek, 2012). The actual productivity of the resource will vary with temperature. The resource is expressed as caloric content per square meter (J/m²), which feeds into the lobster growth model. The total consumption rate by lobsters, summed over all individuals present, is then subtracted in the resource equation (Table 9).

We assume that lobsters are being placed at the monopile at a given body size and this body size is set to the size at settlement or, in case of stocking, to 50 mm CL. Fishing mortality, or harvesting, is instantaneous and occurs when lobsters reach the minimal landing size of 87 mm CL Reproduction is accounted for as an energy loss for mature lobsters but there is no natural recruitment occurring at the monopile. This is based on the assumption that the monopile is not a closed system and currents

will take all offspring elsewhere. Hence, there is no population dynamical feedback through reproduction. New lobsters can only enter the population through stocking. Population dynamics occur only via resource competition.

Table 9. Variables, equations and functions for the shrimp population, physiology and the resources. Index *i* denotes an individual lobster cohort.

Variables			
Number of cohorts	D		
Number of individuals in cohort i	$N_i, i \in \{1, D\}$		
Volume of individuals in cohort i	V_i	cm ³	
Energy of individuals in cohort i	E_i	J	
Reproductive energy of individuals in cohort <i>i</i>	$E_{r,i}$	J	
Resource density	R	Jm ⁻²	
Lobster dynamics			
Volume of individuals in cohort <i>i</i>	$\frac{dV_i}{dt} = \kappa P_E - M/[E_G]$	cm ³ d ⁻¹	1
Energy of individuals in cohort <i>i</i>	$\frac{dE_i}{dt} = \varepsilon I - P_E$	J d ⁻¹	2
Reproductive energy of individuals in cohort <i>i</i>	$\frac{dE_{r,i}}{dt} = (1-\kappa)P_E - P_R$	J d-1	3
Number of individuals in cohort i	$\frac{dN_i}{dt} = -(\mu_s + \mu_f)N_i$		4
Resource dynamics			
Resource density	$\frac{dR}{dt} = r((Rf_{\tau}K) - R) - R(\sum_{i=1}^{D} S_m S A_{\tau}/Area)$	Jm ⁻² d ⁻¹	5
Lobster functions			
Maximum intake	$I_m = \{\text{PXm}\} V_i^{2/3}$	Jd ⁻¹	6
Search rate	$S = 1/(1 + R_j (F_m V_i^{2/3})/I_m)$	m ² d ⁻¹	7
Ingestion	$I = A_{\tau} S_m R_j S$	Jd ⁻¹	8
Maintenance	$M = A_{\tau} [P_M] V_i$	Jd ⁻¹	9
Energy utilisation	$P_E = \frac{\frac{E_i}{V_i}}{\kappa^{E_i}/V_i + [E_G]} \cdot \left(\frac{\left(\frac{\left\{PXm\right\}^{E_i}}{V_i} \in V_i^{\frac{2}{3}}\right)}{[E_M]}\right) + M$	Jd-1	10
Development and maintenance of maturity	$P_{R} = \begin{cases} \frac{V_{i}(1-\kappa)}{\kappa} M \text{ if } V_{i} \leq V_{P} \\ \frac{V_{P}(1-\kappa)}{\kappa} M \text{ if } V_{i} > V_{P} \end{cases}$	Jd ⁻¹	11
Cohort addition	$C \rightarrow C + 1$		12
Starvation mortality	$\mu_{s} = \begin{cases} 0 \ if \frac{V_{i}}{V_{max}} \ge S_{t} \\ S_{r} \left(\frac{S_{t}}{(V_{i}/V_{max})} - 1\right) if \frac{V_{i}}{V_{max}} < S_{t} \end{cases}$	d-1	13

Instantaneous Fishing mortality
$$\mu_f = \begin{cases} 0 \text{ if } L_i < L_h \\ 1 \text{ if } L_i \ge L_h \end{cases} \qquad \qquad d^{-1} \qquad 14$$

Carapace length

$$L_i = V_i^{1/3} / \delta_m \qquad \qquad \text{cm} \qquad 15$$

$$A_{\tau} = e^{\frac{TA}{Tref} - \frac{TA}{T_{\tau}}} \cdot \frac{\left(1 + e^{\frac{TAL}{Tref} - \frac{TAL}{TL}} + e^{\frac{TAH}{TH} - \frac{TAH}{Tref}}\right)}{\left(1 + e^{\frac{TAL}{T_{\tau}} - \frac{TAL}{TL}} + e^{\frac{TAH}{TH} - \frac{TAH}{T_{\tau}}}\right)} - 16$$

Arrhenius scaling at day $\boldsymbol{\tau}$

<u>Temperature</u>

Ambient temperature in area	$T_{\tau} = amp \sin\left(\pi \frac{(\tau - center)}{width}\right) + offset$	°C	17
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Temperature values were taken from Steenbergen et al. (2015), for the coast. OWF's placed in deeper areas will imply lower temperatures and hence reduced growth rates.

Table 10. Model parameters, there symbol used, value, unit and reference. AMP refers to the Add-My-Pet entry (Kooijman and Augustine, 2017).

Parameter	Symbol	Value	Unit	Reference
Individuals				
Maximum search rate	Fm	0.65	m ² cm ⁻² d ⁻¹	AMP
Handling	Н		1	
Maximum ingestion	{P _{Xm} }		250J cm ⁻² d ⁻¹	this study
Volume specific	[P _M]	24	4.3J cm ⁻³ d ⁻¹	AMP
Maintenance				
Cost of growth	[E _G]		100J cm ⁻³	AMP
Maximum energy density	[E _M]	8279	.36J cm ⁻³	АМР
Energy egg	Eв	43	3.9J	Pandian (1970)
Carapace length at	Lj	0.52	253cm	AMP
settlement				
Carapace length at	Lh	:	8.7cm	this study
harvesting				
Volume at maturation	Vp	10	5.4cm ³	AMP
Shape coefficient	δ_{m}	0.5	573-	AMP
Карра	к	0.89	959-	AMP
Conversion efficiency	3		0.8-	AMP
Starvation mortality rate	e S _r		0.5d ⁻¹	Steenbergen et al. (2015)
Starvation threshold	St	0.	.75	Steenbergen et al. (2015)
Arrhenius temperature	TA	80	000K	AMP
Optimum temperature	Topt	2	293K	Kristiansen et al. (2004)
Lower boundary of	ΤL	2	275K	Kristiansen et al. (2004)
tolerance range				
Upper boundary of	Тн	2	295K	this study
tolerance range				
Rate of decrease at	TAL	1	le6K	this study
lower boundary				
Rate of decrease at	Тан	1	le5K	this study
upper boundary				
Resource				
Carrying capacity	К	1.8	8E4J m ⁻²	this study

Parameter	Symbol	Value	Unit	Reference
Regrowth rate	r		0.1d ⁻¹	Kooijman (2009)
Arrhenius temperature	T _A	7	022K	AMP
Optimum temperature	Tref	293	3.15K	AMP
Lower boundary of	ΤL		275K	AMP
tolerance range				
Upper boundary of	Тн		296K	AMP
tolerance range				
Rate of decrease at	TAL	45	430K	AMP
lower boundary				
Rate of decrease at	Тан	31	376K	AMP
upper boundary				
A-biotics				
Temperature amplitude	amp	8	3.07K	(Steenbergen et al., 2015)
Temperature width	width	18	32.4d	(Steenbergen et al., 2015)
Temperature center	center	125.36d		(Steenbergen et al., 2015)
Temperature offset	offset	283.78K		(Steenbergen et al., 2015)
Monopile surface area	Area		364m²	this study, (Table 6: OWF
				Prinses Amalia)

The parameterization of the lobster model is taken from AMP (Kooijman and Augustine, 2017), while the values for the Arrhenius function were fitted for this application, based on oxygen consumption rates from Whiteley et al. (1990). The unit of the maximum search rate was converted from I cm⁻² d⁻¹ to m² cm⁻² d⁻¹ to match the units of the rest of the model and assuming a height of 10 cm within reach. The value of the maximum ingestion rate was raised to 250 J cm⁻² d⁻¹ (instead of 217 J cm⁻² d⁻¹) to increase the growth rate such that maturation occurs in better correspondence with field data, with an average of 6 years (Jensen et al., 2000).

For the resource parameter values were based on those of *Mytilus edulis*, as this is based on biomass the dominant species on the monopiles (Bouma and Lengkeek, 2012). For the regrowth rate *r* we used 0.1, the growth rate of *Mytilus Edulis (Kooijman, 2009)*. As we model the resource in terms of energy density, the mean ash free dry weight (AFDW) measured on the monopiles from Bouma and Lengkeek (2012) was converted to J/m2, based on the conversion of 22 J/mg AFWD (Brey et al., 1988), resulting in a maximum energy density of 1.8E4 J/m2.

4.2 Results

4.2.1 DEB stand-alone

As a test the DEB model for lobster was run under stand-alone conditions, meaning that the growth of a single individual is modelled with ad libitum food and given an annual temperature regime. The carapace length development of a lobster over time follows the expected growth curve, where growth slows down at large size (Figure 16). The annual temperature fluctuations are visible with retarded growth rate in the winter period and increased growth rate in summer. The length at age curve presented is compared to field data (Jensen et al., 2000) based on a mark and recapture experiment at the English coast. The carapace length predicted by the model complies with field observations (Bannister et al., 1994; Sheehy et al., 1999; Jensen et al., 2000).

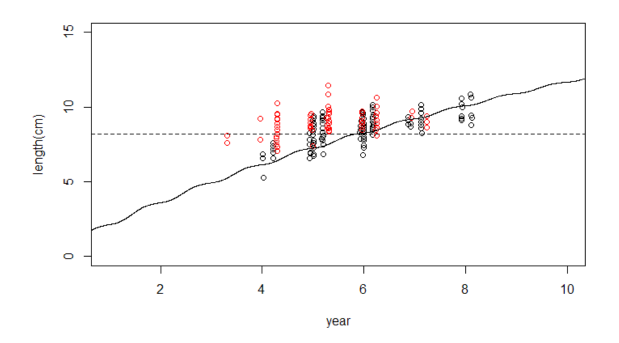


Figure 16. Carapace length (cm) in time for the DEB model (solid line) and length-at-age data taken from (Jensen et al., 2000) (red: 1983-1984; black: 1985-1988). The horizontal dashed lines denotes the size at maturity of 8.2 cm (AMP value).

From the basic DEB model and its parameters for lobster, one can calculate the time it takes for energy reserves to deplete from maximum energy to zero, for a given CL (Figure 17). This assumes no food intake and starting at the maximum energy density an individual can have given its length. Larger individuals have in total more energy to spend and energy depletion therefore takes longer. Energy depletion is affected by the ambient temperature. In cooler conditions energy depletion is slower than at higher temperatures (Figure 17).

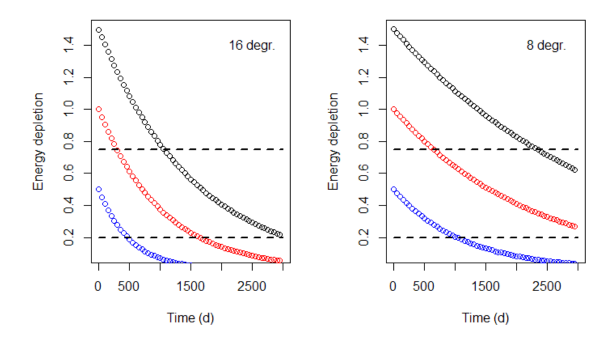


Figure 17. Energy depletion ratio relative to the maximum energy as function of time for different initial carapace lengths (black = 6 cm, red = 4 cm and blue = 2 cm). The left panel shows the depletion at 16 degrees and the right panel at 8 degrees. The horizontal dashed lines denote a depletion of 0.75 and 0.2.

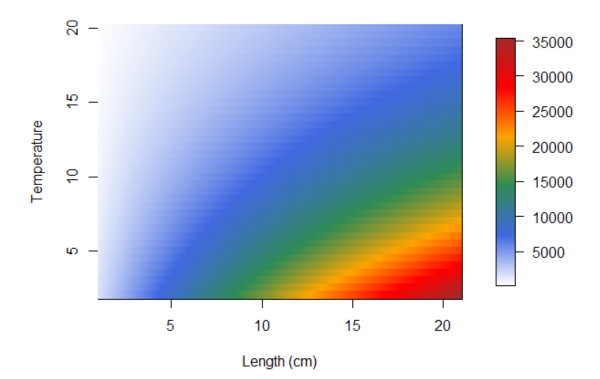
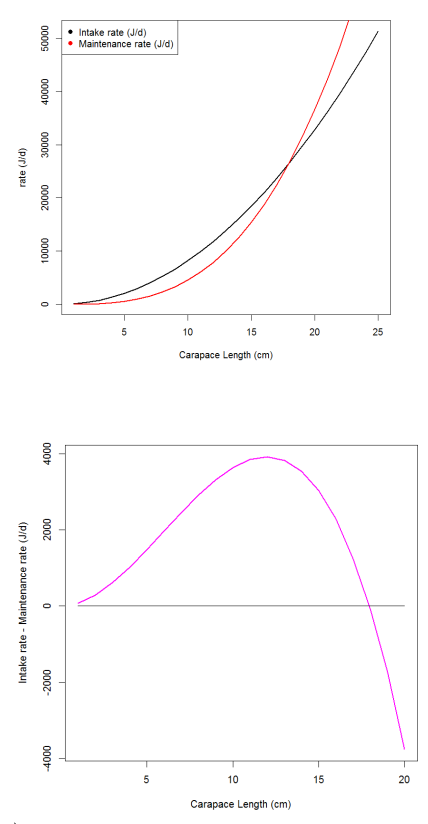
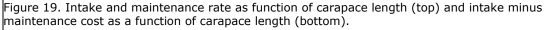


Figure 18. Time (in days, right axis) it takes in days to deplete energy to 20% of size specific maximum energy as function of carapace length and (constant) temperature (left axis).

From the energy depletion we can defer the time to deplete energy to 20% of the maximum as a function of length and water temperature (Figure 18). From this information it is clear that larger individuals can survive longer without food than smaller conspecifics. They can sustain lower food conditions due to the high energy content per unit of body weight relative to the maintenance costs per volume times the total amount of body weight available. This difference in survival time is largest at low temperatures. Warmer temperatures decrease the energy depletion time. Overall this result illustrates that lobsters can sustain periods with low resource conditions well.

When food is limited a size difference between individuals results in a competitive difference. Intake rate scales with surface area while maintenance scales with volume (Figure 19). Yet, the increase of both rates with lobster size differs in such manner that intake increases faster than maintenance rate for lobsters smaller than 12 cm. This implies that with constant but low food availability a larger lobster can realize a higher intake than a smaller conspecific, still being able to cover maintenance costs. This provides the larger lobster with a competitive advantage under low food conditions. However, the maintenance rate lobsters larger than 12 cm CL increases faster than intake, and therefore the competitive advantage is for the smaller lobsters. This difference in scaling of the rates and the breakpoint (at 12 cm CL) where smaller or larger individuals have a competitive advantage at low food conditions is illustrated by the humped shaped relationship of intake minus maintenance (Figure 19, bottom graph). When comparing the intake rate and the maintenance rate as a function of body size it is clear that intake exceeds maintenance for sizes smaller than 18 cm CL (Figure 19). This provides a natural limit where intake and maintenance are balanced. Note that these relationships are based on averages and that natural variability will provide variance around this theoretical value.





This result from the stand-alone DEB model already suggests that in a population dynamical setting with resource competition larger individuals may outcompete smaller conspecifics and that summer could be the most likely period for starvation mortality to occur.

4.2.2 Population dynamical model

In order to model a dynamic resource and the possibility of resource competition among lobster the DEB model was embedded within a population dynamical framework (de Roos et al., 1992). The growth of a single individual feeding on the benthic resource follows a similar pattern as the DEB stand-alone application (Figure 20). Lobster length increases with time, showing the effect of season with reduced growth in winter. Lobster growth levels off when the resource is depleted such that growth is no longer possible. The size at maturity (8.2 cm, AMP) is reached at the end of the fifth year and the size of harvesting (8.7 cm for the UK) is reached at the start of the sixth year. The resource follows the temperature driven sinusoidal pattern, yet showing an overall decrease in time due to lobster consumption. After about seven years the resource shows an additional pattern, a second depression in resource density occurring in early summer, which increases and then stabilises as lobster size becomes constant. This depression is caused as temporarily consumption exceeds resource regrowth rate, due to the differences in temperature scaling between lobster and its food (parameter values of the Arrhenius function); i.e. in spring consumption increases faster with increasing temperature than resource growth does. Halfway summer the resource regrowth rate can compensate consumption and resource density increases again. Resource levels are reduced towards an average of 3.5 KJ/m², in line with benthic productivity estimates in coastal areas (Steenbergen et al., 2015).

When assuming that a lobster is stocked at 5 cm CL the results growth curve and resource patterns are identical. With an initial size of 5 cm the harvesting length is attained after 3 years.

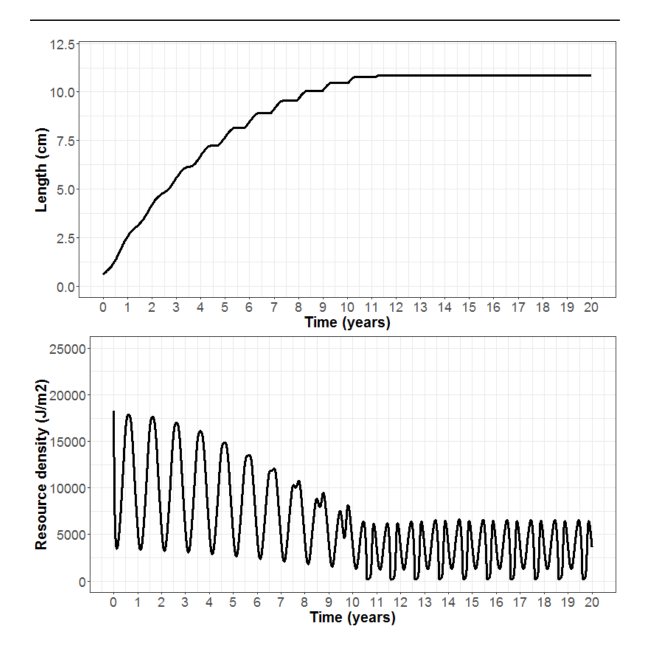


Figure 20. Individual lobster growth (CL top panel) and resource density (bottom panel) resulting from the population dynamics model, initiated with a single lobster.

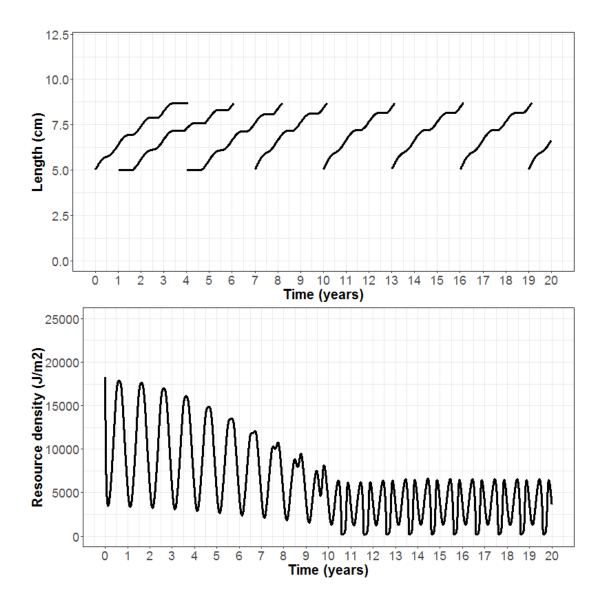
When more than one lobster is put into the system the attained length decreases due to severe resource depletion. Already with two individuals present the maximum attained size is 8.2 cm, the size of maturation. But this size is below the minimum landing size which prohibits harvesting. Interestingly, the lobsters do not suffer from starvation mortality, the low resource density suffices to cover the cost of being, but simply does not allow for further growth. If more lobsters are simultaneously introduced, the maximum attained length will be smaller as growth ceases at a smaller size.

Only increased resource productivity will result in increased growth, such that there can be more lobsters that reach harvestable sizes. There is a linear relationship between resource production and lobster growth to harvestable sizes.

4.3 Restocking

The model results show that with a single lobster in the system the length of harvesting is reached after 3 years when stocked with a length of 5 cm CL (Figure 20). We take therefore 3 years in between stocking events, and stock with individuals of 5 cm CL. Stocking with lobsters at a smaller

size will result in a longer period needed to reach the minimum landing size, which can be deducted from the growth curve in Figure 20. Similarly, stocking at a larger length will shorten the time needed at the monopile to reach the minimum harvestable length.



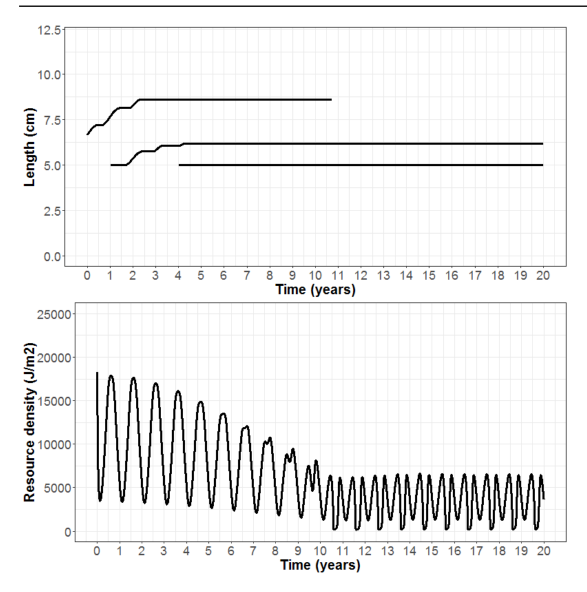


Figure 21. Individual lobster length and resource density in time. From year 1 onwards restocking occurs with lobsters of 5cm length. Top graphs: 1 lobster restocking, bottom graphs: 2 lobsters restocking. Lobsters of harvestable lengths (8.7 cm) are removed from the model.

Figure 21 shows the length of an individual lobster in time, with restocking every three years and removal of harvestable lobsters (8.7 cm CL). With one lobster added per stocking event lobsters reach harvestable sizes in three years (Figure 21, top graph). In such a system every three years one lobster can be harvested while one lobster is being placed at the monopile.

When under the same conditions two lobster are introduced at each stocking event, growth stops completely after the first event and the lobsters stocked at the second event do not start to grow (Figure 21, bottom graph). Resource levels are depressed to such levels that the smallest lobsters suffer from starvation mortality, meaning that latest additions starve to death in a few days. This starvation mortality is the result of resource competition between lobsters, which favours larger individuals. This is also demonstrated in Figure 18, where the reserve depletion as function of size and temperature is given. There is a linear relationship between resource productivity and the number of lobsters that reach the minimum landing size. A doubling of the value of carrying capacity doubles the number of lobsters reaching the minimum harvestable size. However, whether or not such increased carrying capacity values still relate to natural systems has to be established.

In case of the current model parameterization there can be one lobster growing from 5 cm CL to harvestable length in three years per monopile. This is a density of ca 0.002 individuals/ m^2 , which is compared to literature on the lower side, but within reported ranges (see Table 5). For American lobster densities in the order of 0.001-3.25 were reported (Howard, 1988; Dunnington et al., 2005).

Densities vary with temperature, season, productivity and lobster size, e.g. higher densities for smaller sized lobsters (Dunnington et al., 2005).

4.4 Model discussion

Any model is by definition a simplification. That said, a model allows for exploration of systems and circumstances that otherwise would be too time consuming, too costly or even impossible to pursue. In this study a DEB model was used both as stand-alone as well as embedded in a population dynamical framework. DEB is a model type with individual size and growth at its core, which allows for ontogenetic difference which in models ignoring size could lead to substantially different results. Above all, ignoring size and growth would mean ignoring one of the most obvious not to mention relatively easy to measure characteristic that defines an individual.

The DEB model shows length development which corroborates with field observations. Interestingly, the lobster energy content and metabolic rate as a function of size are such that larger individuals can sustain periods of resource limitation for a longer period than smaller individuals. Summer appears to be the period for which starvation mortality is most likely to occur if food conditions are poor. Differences in competitive dominance between size classes occurs due to asymmetry in the increase in maintenance costs vs intake rate as a function of size. When maintenance increases faster with size than intake small individuals are competitively dominant and large individuals win competition when the opposite holds, when intake rate increases faster with size than maintenance. The finding that small lobsters are inferior competitors to larger ones (till 12 cm) suggests that the size at stocking can play an important role in the survival probability of introduced individuals. Not only because larger individuals are less prone to predation but also because of their ability to sustain poorer resource conditions than small individuals.

The DEB stand-alone model with ad libitum food but with a seasonal temperature regime predicts that the harvestable length is reached at an age of 6. The population dynamical model predicts a similar age at which a harvestable size is reached. This is despite the fact that in the stand-alone version growth is limited only by the intake rate of lobsters while in the population dynamical model food availability plays a role. Not only is there a maximum in food density in the form of a given carrying capacity, there is also the seasonal effect of temperature causing a reduction in resource productivity at lower temperatures. The population dynamical model results suggest that this seasonality in resource productivity has little influence on growth. Stocking at a size of 5 cm CL would imply that this size is reached after 3 years. Stocking at smaller sizes will result in a longer period before harvestable sizes are reached, e.g. stocking at ~2 cm CL will result in a 5 year period. The seasonality in physiological rates due to changes in temperature are hence more limiting to growth than the seasonal reduction in resource productivity. Adding resources will only enhance lobster growth rate provided that resource productivity is limited and not the intake rate. Whether resource productivity is the most important limiting factor in field conditions will depend on ambient temperatures which in case are likely to show high inter annual variability and lobster size. Resource enhancement can aid growth rates when multiple individuals compete for resources and thereby reduce resource density to such levels that growth is hampered.

The surface area of a monopile provides food for a single lobster to reach the harvestable size based on the current parameterization. A lack of records on naturally occurring densities of (large) European lobster does not confirm nor reject this finding. Bouma and Lengkeek (2012) did not report the presence of lobsters in the wind park surveyed, 4 years after the park became operational. The reason for absence of lobster in unknown.

The current model does not take refuge holes into account. Such an addition might be useful if field observations demonstrate that lobster densities are indeed limited by shelter on the monopiles rather than food conditions. In such a case the population model can easily be adapted to include density regulation based on shelter availability.

Conclusions, discussion and recommendations

This desk study describes the biology of the European lobster *H. gammarus*. Using the derived data a model was developed to describe the growth of the European lobster under assumed conditions on the anti-scouring of monopiles in Dutch OWFs. One of the main questions to answer was if, theoretically, local productivity supports the continuous harvesting of lobsters with passive fishery methods.

5.1 Main findings

5.1.1 Lobster ecology

5

As a first finding lobster is characterised as a large top end predator. It exhibits a k-strategy life style, able to reach very old ages (e.g. an estimated 72 years). The planktonic and early benthic stages are very vulnerable and susceptible to predation. It grows slowly, taking at least four years to reach maturity in favourable conditions although in general maturity is thought to start from five years old depending on ambient conditions. Reproduction occurs in a two to three year cycle. The eggs are held on the pleopods for approximately a year until hatching. In itself the reproduction is sufficient to maintain populations at high levels (given the high age). However, global lobster populations have suffered considerable fishing pressure and stock collapse only recovering slowly now at some regions partly due to appropriate management (the MCRS and stock enhancement programmes, Wickins & Lee, 2002, Prodöhl et al., 2006, Phillips, 2013).

In principle European lobsters have a high enough mobility to colonize the entire North Sea as one genetic population using wrecks, hard substrates etcetera as stepping stones (Krone and Schröder, 2011). Still unexpected differentiated genetic clusters occur which should have been fully connected and exchanging based on this mobility. Apparently the exchange is less than based on theoretical estimates of mobility.

The limited exchange implies that when exploiting lobsters at OWFs an exploitation management plan should be established based on the local OWF population dynamics and the presence of lobsters nearby on the soft sediments. Also stock enhancement using larvae or juveniles from local populations might be an additional option. The seafloor at the OWF could be an issue in choosing at which size to release juvenile lobsters since the earliest benthic stages can dig their holes. When the seafloor is more sandy than solid mud, early stages are entirely depending on the crevices between stones and cobbles. In addition they actively harvest the walls of their residence. The more productive the seafloor bottom is (in terms of worms, amphipods etc.) the less frequent these vulnerable stages have to expose themselves to predation and currents (Lawton & Lavalli, 1995, Jensen et al., 2000, Wickins & Lee, 2002). In this sense older and larger stages (> an estimated 15 mm CL) are less dependent on bottom type (and more on availability of crevices).

All sizes European lobster prefer hard substrate with crevices to hide. At younger stages they are entirely dependent on shelter in order to evade predation. At later stages they become less dependent on shelter but keep a preference for hard substrate. Crevices should have the size to accommodate the complete body and for all sizes. Younger animals of European and American lobsters move to larger crevices while growing (Lawton & Lavalli, 1995, Jensen et al., 2000, Dunnington et al., 2005, Phillips, 2013)

Lobster densities at suitable substrates of a fished populations are 1 lobster per \geq 150m² (four observations). Two extremely high densities were found for European lobster at 1 lobster per ~4-6m². American lobsters can have on average much higher natural densities (1 lobster every 6m²) but one

should keep in mind that this species is migratory (Dunnington et al., 2005, Geraldi et al., 2009, Phillips, 2013) that might explain that a given substrate does not have to provide food year round. In addition, they have a higher mobility so likely have a greater foraging area .

5.1.2 DEB modelling and lobster production

The DEB model predicts lobster growth in line with growth reported in literature although length at age from field data shows a large variability. Based on the current model parameter setting resource production is limiting lobster growth and abundance while ensuring that lobsters reach marketable sizes. The current model setting predicts that 1 lobster can reach marketable size after 3 years, provided a stocking length at 50 mm CL. Model resource productivity was based on mussel density estimated at a limited survey of monopiles in a Dutch OWF. However, resource production might differ between monopiles, parks and temperature regimes, and higher productivity will support more lobsters and/or have a positive effect on individual growth rates.

The population model does not take recruitment into account, the reason being that a single monopile surface area is being studied and that little is known on factors determining recruitment success in wild populations, nor the geographical extent of such populations in the North Sea. Energy losses through egg production is taken into account. In addition, no lobsters were observed at a studied wind park 4 years after operation started (Bouma and Lengkeek, 2012). This suggests that at least the chance of natural recruitment on monopiles in wind parks in the North Sea is low. However, when the expectancy is 1 lobster per monopile, changes of encounter during a scientific dive are not that high.

The modelled European lobster productivity is in line with low densities found in the field, using the OWF hard substrate and anti-scouring data from Bouma & Lengkeek (2012). One lobster could be supported by one monopile given its anti-scouring surface (1 per 364 m²).

5.1.3 Enhancement strategies

The current literature survey on lobster ecology linked observations to the proposed enhancement strategies (habitat, stock and food). All strategies seem to have a potential added value to lobster production. As a basic requirement, monitoring is needed to establish how much and which size-classes of *H. gammarus* are present in the OWFs to think on any strategy at all.

<u>Habitat enhancement</u> can serve to increase the amount of crevices available. Taking the expected crevice size into account for OWF Prinses Amalia and OWF Luchterduinen (maximally 8 x 8 x8 cm, Table 7), theoretically there could be a lack of suitable crevices for larger sized lobster in the OWFs preselected for TKI Win-Wind. Adding extra habitat by means of hard substrate with crevices might stimulate the presence of legal catchable lobster (at 85 mm CL, an estimated 240 mm total length, Prodhöl et al., 2006). A potential new habitat should be carefully designed accommodating several sizes classes of lobster, having sufficient water and oxygen refreshment and also enabling stocks of the typical food like mussels and crabs. In addition, extra hard substrate can serve as fundament for extra biota and thereby food.

<u>Stock enhancement</u> has a potential to help to surpass the vulnerable pelagic and early settlement stages (up to ~15 mm CL or even larger). It can also safeguard supply of animals since natural recruitment is low. Not all OWFs have European lobster whereas one would expect presence. Also Figure 8 shows e.g. remarkable patterns of suitability of wrecks: wrecks nearby each other have marked differences in suitability. These phenomena can reflect differences in habitat suitability or supply of larvae.

Food seems the most obvious limiting factor. <u>Food enhancement</u> (by e.g. improving quality by seeding or improving availability by depositing discards or extra habitat) can help to increase productivity and populations.

5.2 Recommendations

5.2.1 Early stages

In literature major uncertainties exist on the larval and early settlement stages in terms of abundance, transport, settlement and survival, especially the early settlement stages (Phillips, 2013). One wonders whether to invest in research on early life stages in the case of commercially exploiting Dutch OWFs for lobsters. Especially given the low rates of colonisation (e.g. Krone and Schröder, 2011) and the vulnerability of the early life stages, difficulties in natural recruitment might be surpassed by active restocking (in the setting of exploitation).

5.2.2 Presence, density and habitat

It is unclear whether European lobsters are present in OWFs. Bouma and Lengkeek (2012) did not encounter any during a limited diving survey (only four diving days, diving on neap tide, with two poor visibility days on three monopiles) . The TKI Energy project WinWind will pay attention to the presence, densities and population size of lobster in OWF Prinses Amalia and/ or OWF Luchterduinen. Also presence and densities of lobster on the surrounding substrates needs to be assessed in order to determine the potential of colonisation by larger lobsters.

In general densities are a rarely assessed aspect of European lobsters ecology (Skerrit, 2014). More data are needed to get more insight in the quantitative biology of *H. gammarus*. It is advisable to include length measurements, characterisation of the substrate, food availability and measurements of (a)biotic factors like temperature ranges, depth etc. (see Table 3).

In this sense Table 3 is striking due to its incongruence. It should be noted that the data compiled by Kristiansen et al. (2004) are from lobster aquacultures of both *H. gammarus* and *H. americanus*. The data of https://eol.org/pages/46505673/data are field data. This data base should be extended with measurements of Dutch circumstances especially in OWFs.

5.2.3 Position and functioning in the food web

More information is needed on the position and functioning of *H. gammarus* in the food web. What are the predators on European lobster in different life stages and what is *H. gammarus* preying upon in the Dutch NCP and OWFs specifically? Information is needed on stomach contents of both European lobsters of different sizes and their potential predators. Also the composition of the food web and trophic layers needs to be studied with and without European lobster in order to determine the potential (mutual) influences.

5.2.4 Physiology and DEB modelling

There is a lack of physiological data in literature. It is advisable to do some basic studies on the physiology of this species. A series of experiments could aid a parameterization of the DEB model for lobster specifically for North Sea conditions.

5.2.5 Primary production and benthic-pelagic coupling

Initially it was anticipated to start the productivity modelling at the trophic level of PP. However, both the PP and the benthic-pelagic coupling are scarcely investigated. The literature was not that detailed that trustworthy parameters and variables could be derived for a reliable productivity estimate at the tertiary production level of European lobster. This urges for more research on the benthic-pelagic and local processes driving secondary production in relation to PP and monopile characteristics.

5.2.6 Enhancement strategies

The literature survey on the enhancement strategies has underpinned the importance of enhancement to promote lobsters. Current results suggest that improvement of habitat in quantity and quality, in

order to have more and larger crevices and more food, are more useful to promote commercial exploitation. In the TKI Energy project Win-Wind more detailed information will be generated on the three strategies in order to get more insights in what enhancement strategy to adopt under which conditions. An important input for which strategy is most suitable will be the absence or presence of lobsters (number and size distribution) encountered. Information on food availability as well as crevice (size and numbers) needs to be gathered. The outcome of such surveys will determine the economic potential and also the urgency to adopt one or several enhancement strategies.

More research is needed in order to determine habitat suitability and differentiating factors on the Dutch NCP. Is that e.g. the suitability of the habitat (crevice numbers and characteristics), or is that food availability. It might even be supply of animals by recruitment by larvae or immigration by adults. Although the mobility of both should be enough and not be a limiting factor (see next sections).

6 Quality Assurance

Wageningen Marine Research utilises an ISO 9001:2015 certified quality management system. This certificate is valid until 15 December 2021. The organisation has been certified since 27 February 2001. The certification was issued by DNV GL.

Furthermore, the chemical laboratory at IJmuiden has NEN-EN-ISO/IEC 17025:2005 accreditation for test laboratories with number L097. This accreditation is valid until 1th of April 2021 and was first issued on 27 March 1997. Accreditation was granted by the Council for Accreditation. The chemical laboratory at IJmuiden has thus demonstrated its ability to provide valid results according a technically competent manner and to work according to the ISO 17025 standard. The scope (L097) of de accredited analytical methods can be found at the website of the Council for Accreditation (www.rva.nl).

On the basis of this accreditation, the quality characteristic Q is awarded to the results of those components which are incorporated in the scope, provided they comply with all quality requirements. The quality characteristic Q is stated in the tables with the results. If, the quality characteristic Q is not mentioned, the reason why is explained.

The quality of the test methods is ensured in various ways. The accuracy of the analysis is regularly assessed by participation in inter-laboratory performance studies including those organized by QUASIMEME. If no inter-laboratory study is available, a second-level control is performed. In addition, a first-level control is performed for each series of measurements.

In addition to the line controls the following general quality controls are carried out:

- Blank research.
- Recovery.
- Internal standard
- Injection standard.
- Sensitivity.

The above controls are described in Wageningen Marine Research working instruction ISW 2.10.2.105. If desired, information regarding the performance characteristics of the analytical methods is available at the chemical laboratory at IJmuiden.

If the quality cannot be guaranteed, appropriate measures are taken.

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Justification

Report C109/18 Project Number: 4318300086

The scientific quality of this report has been peer reviewed by a colleague scientist and a member of the Management Team of Wageningen Marine Research

Approved:

Dr.Ir. J.W.M. Wijsman Senior Researcher

Signature:

28 maart 2019

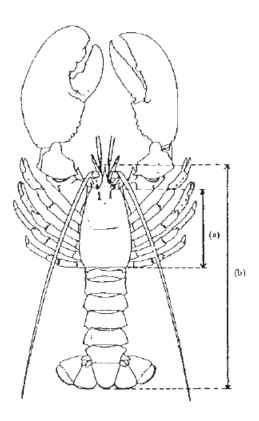
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Annex 1 Length definitions used for *Homarus spec.*

Carapace length

Based on European Community (2006). Council Regulation (EC) No 1967/2006 of 21 December 2006, concerning management measures for the sustainable exploitation of fishery resources in the Mediterranean Sea, amending Regulation (EEC) No 2847/93 and repealing Regulation (EC) No 1626/94 carapace length is measured from the rear of the eye socket to the rear of the carapace on a line parallel to the center line of the body shell. Make sure the gauge is at the extreme rear of the eye socket below the rostrum or horn. A common error is to measure from the horn located forward of the eye socket, which results in an improper measurement.



(Homarus) Lobster

Figure 22 Length determination of the lobster (*Homarus gammarus*) by measuring (a) the length of the Carapace or (b) the total length excluding antennae and scissors (European Community, 2006). Measured lobsters have a shell length (part a of the lobster) of 83 to 87 mm or larger depending on country.

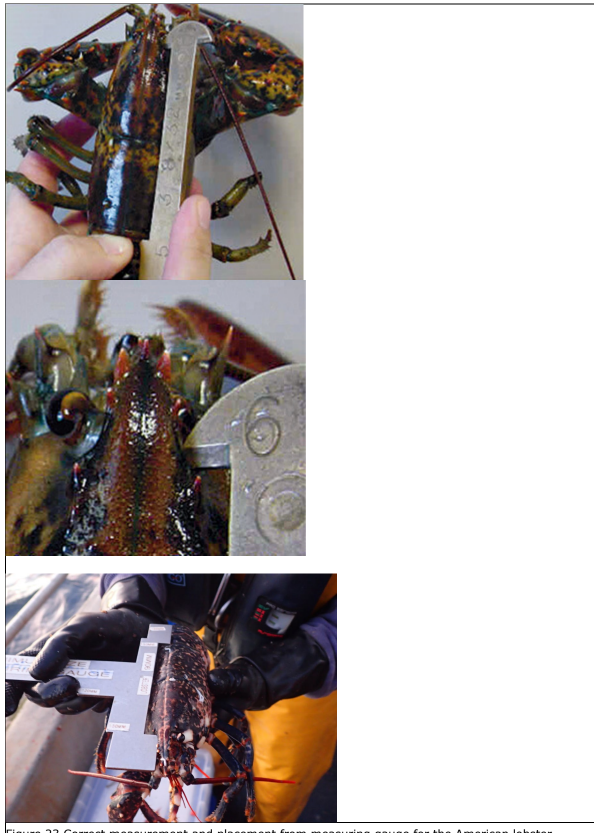


Figure 23 Correct measurement and placement from measuring gauge for the American lobster (*Homarus americanus*) (from http://www.eregulations.com/massachusetts/fishing/saltwater/how-to-measure-a-lobster/ d.d. 23-12-18) and European lobster (photo National Lobster Hatchery).

Orbital Carapace Length

As CL; length taken from the deepest point of the orbital (Figure 23).

Total length

The total length is measured, from the tip of the rostrum to the rear end of the telson, not including the setae (Figure 22).

Annex 2 Diet composition *Homarus americanus*

Table 11 Diet of *Homarus americanus*. Occurrence of prey taxa in stomachs by percentage frequency-of-occurrence (percentage point 3r the 4 area-date groups: A, macroalgal community August 1979 (n = 115); B, barrens (sandy sediments), Sep 1979 (n = 117); C, barrens (sandy sediments), July-August 1981 (n=400), barrens (sandy sediments), February-March 1982 (n = 400) (table derived from Elner and Campbell, 1987)

	Area-date groups							
Prey taxa	<u>A</u> <u>B</u>		<u>C</u>		D			
1. Animals	100	(94.2)	100	(92.8)	99	(97.8)	100	(97.3)
2. Protozoans/ foraminiferans	1	(0.02)	5	(0.21)	4.5	(1.5)	31	(1.6)
3. Poriferans	1	(0.02)	1	(0.19)	0.5	(0.07)		
4. Cnidarians	11	(0.31)	22	(0.64)	15	(039)	21	(1.2)
5. Bryozoans	1	(0.02)	-		1	(0.08)	-	
6. Molluscs	94	(18.8)	83	(21.2)	90	(19.6)	84	(16.5)
7. Chitons	9	(0.57)	34	(1.4)	21	(1.6)	9	(1.4)
8. Tonicella marmorea	-		2	(0.08)	9	(0.88)	3	(0.59)
9. Gastropods	78	(6 2)	44	(2.2)	10	(0.51)	4	(0.15)
10. Acmae testudinalis	2	(0.16)	1	(0.02)	2	(0.04)	-	
11. Lacuna vincta	26	(18)	6	(0.11)	1	(0.02)	-	
12. Skeneopsis striatum	2	(0.16)	-		1	(0.02)	-	
13. Crucibulum striatum	1	(0.04)	-		-		-	
14. Crepidula sp .	2	(0.05)	-		-		-	
15. Lunatia sp.	2	(0.16)	-		-		-	
16. Mitrella sp.	5	(0.26)	-		-		-	
17. Bivalves	63	(11.8)	71	(17.3)	87	(16.1)	78	(14.0)
18. Mussels (Mytilus edulis, Modiolus								
modiolus)	49	(9.6)	62	(16.2)	85	(15.0)	77	(13.8)
19. Cerastoderma pinnulatum	10	(0.57)	2	(0.03)	0.3	(0.01)	-	
20. Anomia sp .	4	(0.27)	4	(0.08)	-		-	
21. Hiatella sp .	-		4	(0.22)	1	(0.01)	-	
22. <i>Mya</i> sp .	-		1	(0.02)	-		-	
23. Musculus sp .	6	(0.26)	2	(0.03)	-		-	
24. Polychaetes	54	(4.0)	48	(2.8)	50	(5.7)	36	(5.6)
25. Nereis sp .	37	(1.37)	37	(0.96)	22	(2.4)	21	(3.0)
26. Polynoids	23	(1.1)	11	(1.4)	24	(2.1)	8	(1.5)
27. Spirorbis sp .	2	(0.05)	-		-		-	
28. Cistena sp.	3	(1.3)	2	(0.34)	0.5	(0.02)	-	
29. Crustaceans	86	(28.2)	79	(29.1)	41	(15.5)	31	(7.5)
30. Barnacles	3	(0.09)	2	(0.03)	-		-	
31. Isopods	11	(1.2)	1	(0.02)	0.3	(0.01)	-	
32. Idotea sp.	3	(0.57)	-		-		-	
33. Amphipods	23	(1.2)	3	(0 06)	2	(008)	0.3	(0.03)
34. Corophium	1	(0.02)	1	(0 02)	-		-	
35. Ampithoe rubricata	10	(0.73)	-		-		-	
36. Decapods	?6	(25)	74	(2A.1l	36	-15.2	28	(7.2)

38. Pagurus sp.			16	(6.1)	19	(11.9)	2	(0.51)
20 Caracar en	26	6.7)	11	(1.8)	-		-	
39. Cancer sp.	30	(9.5)	37	(14.9)	2	(0.53)	1	(0.49)
40. Hyas sp.	-		3	(1.5)	4	(0.76)	2	(0.78)
41. Carcinus maenas	2	(0.18)	-		-		-	
42. Caridion sp.	1	(0.02)	1	(0.02)	-		-	
43. Echinoderms	27	(6 2)	72	(19.4)	82	(27.4)	29	(14.3)
44. Holothurians	4	(2.0)	7	(2.2)	-		-	
45. Psolus sp.	-		1	(0.03)	-		I	
46. Cucumaria sp.	1	(0.29)	-		-		-	
47. Strongvlocentrotus droebachiensis	1	(0.04)	57	(2.6)	63	(8.1)	I	
48. Seastars	22	(4.1)	21	(5.2)	11	(6.6)	13	(2.5)
49. Ophiuroids	3	(0.07)	24	(9.1)	15	(8.7)	4	(3.7)
50. Ophiopholis aculeata	-		9	(5.4)	0.3	(0.13)	6	(5.8)
51. Amphiopholis sauamata	1	(0.02)	-		-		-	
52. Ascidians	28	(12.1)	12	(3.1)	0.5	(0.19)	0.5	(0.28)
53. Fish	21	(1.8)	9	(2.1)	0.5	(0.11)	1	(0.27)
54. Other animals ^a	5	(0.86)	3	(0.16)	-		I	
55. Unidentified animal tissue	79	(20.1)	69	(13.9)	95	(27.8)	100	(50.2)
56. Plants	83	4.2)	49	(4.2)	44	(2.0)	37	(2.6)
57. Algae	77	(3.9)	44	(3.3)	44	(1.9)	37	(2.6)
58. Brown algae	36	(1.1)	9	(2.2)	5	(0.12)	7	(0.64)
59. Green algae	14	(0.46)	1	(002)	3	(0.08)	4	(0.23)
50. Red algae	23	(0.06)	21	(0.53)	33	(1.2)	27	(1.3)
61. Tufted red algae ^b	-		14	(0.26)	29	(1.0)	26	(1.2)
62. CoralJina officianalis	36	(12)	15	(0.40)	9	(0.52)	2	(0.11)
63. Zostera marina	2	(0.04)	1	(0.03)	-		I	
64. Gravel	37	(1.5)	38	(1.5)	4	(0.16)	З	(0.10)
65. Foreign material ^c	2	(0.04)	7	(1.5)	-		I	
^a Eggs, nematodes, nemertines, turbellarians								
P Family: Ceramiaceae								
Plastic, rubber, wood								

Table 12 Major categories of food items, divided into specific food items when possible, and their overall volumetric contribution (total=100%) to stomach contents of all examined lobsters (<i>H. americanus</i>) from Baie de Plaisance, Magdalen Islands. Contributions more than 10% given in bald (Sainte-Marie & Chabot, 2002).						
	% volume contribution main groups	% volume contribution to specific group				
Formaniferans	0.31					
Macroalgae	3.94					
Coralline algae (Corallina officinalis)	1.78					
Hydrozoans	2.07					
Bivalves 16.57						

2.02

9.92

Mytilus edulis

Modiolus modiolus

Unidentified Pelecypoda	1	4.63
Gastropods	5.85	
Lacuna vincta		0.28
Unidentified Gastropoda		0.57
Polychaetes	5.97	
Nereidae		3.18
Polynoidae		2.71
Unidentified Polychaeta		0.08
Barnacles (Balanus sp .)	0.12	
Crustacean meiofauna	0.53	
Harpacticoida		0.03
Ostracoda		0.21
Unidentified minute Crustacea		0.29
Amphipods	0.54	
Corophium sp.		0.04
Gammarus sp.		0.03
Caprellidea		0.04
Gammaridae		0.16
Unidentified amphipods		0.27
Isopods	0.67	
Idotea sp.		0.13
Idoteidae		0.19
Unidentified valviferan isopods		0.34
Carideans	0.24	
Crangon septemspinosa	-	0.1
Unidentified carideans		0.13
Pagurids	4.16	
Pagurus acadianus		0.51
Paguridae		3.65
Rock crab (Cancer irroratus)	26.37	
American lobster (Homarus		
americanus)	0.76	
Echinoderms	2.22	
Strongylocentrotus droebachiensis		1.02
Ophiuroidea		0.12
Unidentified echinoderms		1.09
Fish	0.66	
Flesh	27.24	
Sum %s volume contribution	100	

Wageningen Marine Research T +31 (0)317 48 09 00 E: marine-research@wur.nl www.wur.eu/marine-research

Visitors' address

- Ankerpark 27 1781 AG Den Helder
- Korringaweg 7, 4401 NT Yerseke
- Haringkade 1, 1976 CP IJmuiden

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