



Royal Netherlands Institute for Sea Research

This is a postprint of:

Witbaard, R, Duineveld, G.C.A, Bergman, M.J.N., Witte, H.IJ., Groot, L. & Rozemeijer, M.J.C, (2015). The growth and dynamics of *Ensis directus* in the near-shore Dutch coastal zone of the North Sea. *Journal of Sea Research*, 95, 95–105

Published version: dx.doi.org/10.1016/j.seares.2014.09.008

Link NIOZ Repository: www.vliz.be/nl/imis?module=ref&refid=244826

[Article begins on next page]

The NIOZ Repository gives free access to the digital collection of the work of the Royal Netherlands Institute for Sea Research. This archive is managed according to the principles of the [Open Access Movement](#), and the [Open Archive Initiative](#). Each publication should be cited to its original source - please use the reference as presented.

When using parts of, or whole publications in your own work, permission from the author(s) or copyright holder(s) is always needed.

1 The growth and dynamics of *Ensis directus* in the
2 near-shore Dutch coastal zone of the North Sea.

3

4 Rob Witbaard^a, Gerard C.A. Duineveld^a, Magda J.N. Bergman^a, Hans IJ. Witte^a, Lennart
5 Groot^a & Marcel J.C. Rozemeijer^b.

6

7 a: Royal Netherlands Institute for Sea Research (NIOZ), Postbus 59, 1790 AB Den Burg, the
8 Netherlands.

9 b: IMARES: Postbus 68, 1970 AB IJmuiden, the Netherlands.

10

11

12 Corresponding author. Tel: +31222369537.

13 E-mail addresses: Rob.Witbaard@nioz.nl

14 Gerard.Duineveld@nioz.nl

15 Magda.Bergman@nioz.nl

16 Hans.Witte@nioz.nl

17 Lennartgroot@gmail.com

18 Marcel.Rozemeijer@wur.nl

19

20

21 **ABSTRACT**

22

23 Here we present data on the wax and wane of a subtidal *Ensis directus* population which
24 settled in 2009 off the coast of Egmond (North Holland Coast in the Netherlands). Initial
25 densities decreased from a maximum of 700 m⁻² in early 2010 to about 50 m⁻² in June 2013.
26 In this period the average length increased from ~ 4 cm to ~12 cm. In 2011-2012 the
27 population was sampled at 3 to 6 week intervals and near bottom environmental conditions
28 were monitored continuously. Samples of animals that were collected were used to follow the
29 change in gonadal mass, tissue glycogen content, tissue weight and shell length. On basis of
30 these data well defined seasonal cycles were observed. The data indicate that the maturation
31 of gonadal tissue already starts early in the year, initially at the expense of somatic tissue.
32 Main spawning takes place in May. After spawning net somatic tissue growth starts after
33 compensation of losses due to spawning. Somatic growth precedes shell growth which starts
34 at water temperatures exceeding 12-14 degrees. Mortality, growth and production are
35 comparable to those found for populations in close-by intertidal areas. As such there is no
36 indication that this offshore population significantly suffers from nearby beach nourishments
37 along the Dutch Coast.

38

39

40

41

42 **Key words;**

43 North Sea

44 *Ensis directus*

45 Growth

46 Environment

47 Population dynamics

48 Beach nourishments

49

50 1. INTRODUCTION

51

52 The most abundant bivalve species occurring in dense aggregations along the Dutch coast is
53 the invasive American razor shell, *Ensis directus*. The species was first observed in the
54 German Bight in 1979 (Von Cosel *et al.*, 1982). Since then it has spread rapidly in north- and
55 southward direction (Armonies, 2001; Severijns, 2002) and has reached the Iberian coast
56 around 2011 (Arias & Anadón, 2012). In Dutch coastal waters it dominates the biomass of the
57 macrobenthic invertebrate community (Goudswaard *et al.*, 2013; Verduin *et al.*, 2011; Verduin *et*
58 *al.*, 2012). Interestingly the presently known standing stock of *E. directus* is substantially
59 higher than previously reported estimates of total bivalve biomass for that area. There is no
60 evidence that this newcomer outcompeted native species (Dannheim & Rumohr, 2012). Most
61 likely because its preferred (Dekker & Beukema, 2012) habitat of mobile sands with high
62 currents speeds has never been occupied by native *Ensis* spp. nor by other local bivalves.

63 *E. directus* can live in these dynamic sedimentary conditions because it can rapidly retract
64 itself deep in the sediment (Drew 1907; Trueman, 1967). Hence, accurate density estimates
65 are often hampered by sampling difficulties. Moreover a significant part of *E. directus*
66 population lives in areas like the very shallow shoreface at depths which are difficult to access
67 with ships large enough to operate sampling gear which is suitable for collecting this species
68 quantitatively. Population size in Dutch coastal waters is therefore, likely even higher than
69 reported by for instance Goudswaard *et al.* (2013). Nowadays, densities are that high that a
70 commercial fisheries for this species has developed. As part of the regulation of this fisheries,
71 annual stock estimates are being made which gives good insight in the long term population
72 development (Goudswaard *et al.*, 2013) along the Dutch coast. Where many researchers
73 thought that the population of this invader would collapse after its first years of successful
74 settlement, it managed to maintain high population levels.

75 Various studies illustrate that since *Ensis directus* invaded European waters it has become an
76 ecologically important species in coastal waters (Tulp *et al.*, 2010). Fish and birds have started
77 feeding on *E. directus*. Armonies and Reise (1999) speculated that the presence of dense *E.*
78 *directus* beds might indirectly lead to a higher diversity of associated fauna, especially since
79 *E. directus* tends to occupy a habitat which formerly was underutilized by native fauna. The
80 dense beds are also likely to facilitate ecological processes such as sedimentation and burial
81 of the finest sediment fraction. In terms of processes the high population biomass also may
82 act as means by which nutrients are retained which dampens the effects of eutrophication
83 and seasonal river nutrient run off (Norrko *et al.*, 2001; Vaughn & Hakenkamp 2001).

84 Large parts of the near-shore Dutch coastal zone especially those where highest densities of
85 *Ensis directus* are found, are nowadays protected as habitat H1110 under the EU Habitat
86 directive. At the same time this coastal area is prone to erosion and needs continuous
87 maintenance by means of beach nourishments and shoreface replenishment of sand
88 (Ellerbroek *et al.*, 2008; Rozemeijer *et al.*, 2013). The direct effects of such mass dumps of

89 sand on the fauna will be evident but with the sand also a small percentage of silt is released.
90 Tides transport this material and thereby potentially cause an effect over a larger area than
91 the dumping plot (van Duin e.a., 2007). High concentrations of suspended fine sediments are
92 known to interact negatively with bivalve filter feeding (Gremare *et al*, 1998, Roper & Kickey,
93 1995; Szostek *et al*, 2013). Concern about the possible impact of beach nourishments on the
94 production of the local *Ensis directus* population led to a project initiated by Rijkswaterstaat
95 (RWS-Ministry I&M) with the ultimate objective to develop a Dynamic Energy Budget (DEB)
96 model (Kooijman, 2010) for *E. directus* allowing assessment of the effects of sand mining and
97 beach nourishments on its secondary production. A first version of a DEB model for *E.*
98 *directus* has been published (Wijsman, 2011), later adjusted and used by Schellekens &
99 Witbaard (2012) and Schellekens (2012).

100 In the present study which was a part of the RWS project we focus on growth and
101 development in a field population of *E. directus* living in a shallow coastal site where beach
102 nourishments were taking place. The stock lives 1 km offshore of Egmond at ~10 m depth on
103 the steep edge of the shoreface. In the summer of 2009 mass settlement of *E. directus*
104 occurred in this area and this settlement was subsequently monitored from the end of 2009 till
105 June 2013.

106

107 **2. MATERIAL AND METHODS**

108 **2.1 Study site and research outline**

109 The project started in September 2009 when a research partner (MEDUSA Explorations;
110 <http://www.medusa-online.com>) commenced with measurements of the sediment silt
111 composition. They sampled sediments along transects extending up to 6 km out of the coast
112 off Egmond. Their methodology also included sampling of sediments with a boxcorer (Vries &
113 Koomans, 2010). We continued their work in spring and autumn 2010 by sampling sediments
114 and *Ensis directus* with a boxcorer along their southern transect and by deploying a
115 measurement platform (lander) to collect semi continuous data on near-bottom environmental
116 conditions. The platform (lander) carried various instruments (section 2.2) to monitor biotic
117 and a-biotic conditions at heights between 30 cm and 200 cm above the bottom. The
118 deployment site was located about one kilometre off the coast of Egmond at a depth of ~10
119 meter (Fig. 1). In February 2011 the platform was redeployed at this location and continuously
120 kept in operation until the end of November 2012. At approximately monthly intervals the
121 deployment site was visited for maintenance of the platform and for sampling of the *E.*
122 *directus* population. In June 2013 the site was visited a final time to sample the *Ensis*
123 population.

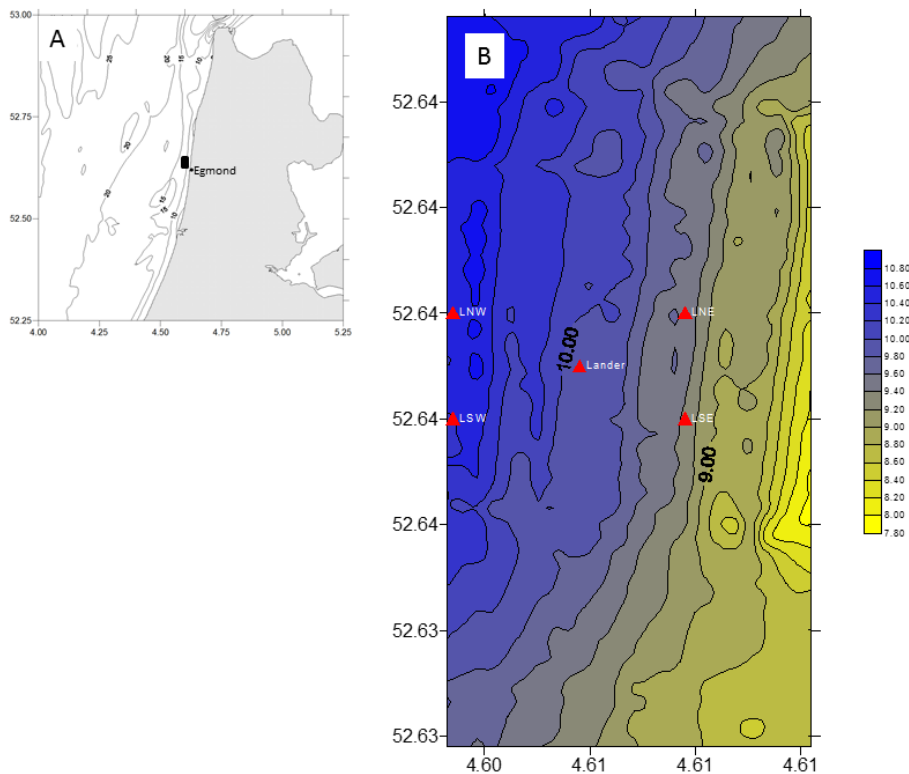
124 In June 2011 a supplementary sampling survey was carried out to determine the distribution
125 and densities of *Ensis directus* at a wider scale covering a large part of the coast of the
126 province of North Holland. During this survey boxcore samples were collected along 8
127 transects perpendicularly oriented to the coast. Each transect extended up to 6 km from the
128 coast. The aim of this sampling survey was to determine if the distribution and densities of
129 *Ensis directus* around the "lander" location represented distribution patterns over a wider
130 area.

131

132 **2.2 Long term environmental measurements**

133 The measurement platform (lander) consists of an triangular aluminium frame (heightxwidth:
134 2 x 2 m) with a series of ballast weights (total 500 kg) fixed onto the lower support that stands
135 on the seafloor. The measurement platform was equipped with a series of sensors measuring
136 current, temperature, salinity, turbidity and fluorescence. Current speed and direction (3D)
137 were measured every 10 minutes at 140 cm above the bottom with a NORTEK Aquadopp
138 Doppler current meter. This instrument also yielded a record of the acoustic backscatter.
139 Temperature and salinity were measured every 10 minutes with a pumped version of the
140 Seabird SM37 CTD system (<http://www.seabird.com/>). In addition to the NORTEK Aquadopp
141 current meter a NORTEK Vektor current meter (<http://www.nortek-as.com>) was mounted at
142 the lander at a height of 30 cm above the bottom. Every 10 minutes this instrument made
143 burst measurements during 2 minutes with a frequency of 1 sec⁻¹.

144 Simultaneous measurements of optical backscatter (OBS) and fluorescence were made at
145 four heights above the bottom, i.e. 30, 80, 140 and 200 cm, using ALEC Compact-CLW's
146 (<http://ocean.jfe-advantech.co.jp>). In the following sections we refer to the material being
147 measured by OBS as SPM (Suspended Particulate Matter). The fluorescent fraction is
148 referred to as Chlorophyll or Chl-a. The ratio of Chlorophyll to total SPM is used to express
149 the amount of chlorophyll per unit suspended matter and can be seen as an expression of
150 food quality. All ALEC sensors were calibrated in the lab over a range of known Chlorophyll
151 and SPM concentrations.
152 The data collected with this measurement platform were used as environmental
153 characterisation of the research area. Detailed information on these long term time series of
154 measurements on chlorophyll, suspended matter and hydrographical parameters has been
155 reported by Witbaard et al, 2013.
156



157
158 Figure 1. A) Study area offshore the coast of Egmond (the Netherlands; Province of North Holland) with (B)
159 bathymetric map with locations of the measurement platform (lander) and the four surrounding sample stations
160 (LNW, LNE, LSW, LSE). Colorscale indicates waterdepth in meter.
161

162 **2.3 Measurements on the *Ensis directus* population near the lander**

163 **2.3.1 Sampling the stock**

164 During the years 2011-2012 the site was visited every 3 to 6 weeks to service the platform
165 (Fig. 1) and to collect *Ensis directus* from four locations (LNW,LNE,LSW,LSE) around the
166 platform (Table 1 & Fig. 1). Hereto two boxcore samples were taken at each of the four corner
167 locations around the measurement platform. If the boxcore sample was of sufficient quality,
168 i.e. > 10 cm deep and with an undisturbed surface layer, a small subcore of the top 5 cm was
169 preserved for sediment grain size analyses. The remainder of the boxcore was sieved over a
170 1 mm screen and the live *E. directus* were collected and stored for density estimates, size
171 measurements and ash free dry weight (AFDW) determination. In this way a time series of the
172 growth and population development of the local *E. directus* stock could be obtained on basis
173 of population averages. For the analyses of shell and tissue growth also the boxcore samples
174 taken in 2010 were used. In 2010 the site was visited 6 times. In 2011 and 2012 the site was
175 visited 19 times. In 2013 the site was revisited once in June.

176

177 Table 1. Positions of the measurement platform ("Lander") and the four stations around it which were sampled 6
178 times in 2010 and 19 times in the period between 2011-2012.

179

Station	Latitude (N)	Longitude (E)
LNE (Lander North East)	52° 38.280'	4° 36.356'
LSE (Lander South East)	52° 38.216'	4° 36.380'
LSW (Lander South West)	52° 38.220'	4° 36.220'
LNW (Lander North West)	52° 38.281'	4° 36.220'
Lander	52° 38.249'	4° 36.294'

180

181 In the 2011 and 2012 surveys the 8 boxcore samples around the measurement platform were
182 supplemented with additional boxcores to collect about 70 *E. directus* specimens for the
183 assessment of seasonal trends in condition related parameters and gonadal development.
184 Samples for this purpose were not confined to any of the four stations, but were still taken in
185 the vicinity of the measurement platform and its four surrounding stations. From all collected
186 specimens a sub-sample of 50 *Ensis* was used for determination of their seasonal change in
187 gonado-somatic index. Another 10 specimens were collected for analysis of the tissue
188 glycogen content. On board all samples were stored refrigerated. In the lab the samples for
189 glycogen content were stored at -80°C. Statistical analyses and description of the data was
190 done in R (R-coreteam, 2012).

191 **2.3.2 Size and weight measurements**

192 For the determination of the seasonality of shell growth and tissue growth the living animals
193 were measured with digital callipers to the nearest 0.1 mm. Three measurements were made:
194 length, width and thickness. For incompletely sampled or broken animals the total length was
195 estimated from shell width. In addition to these size measurements the ash free dry weight

196 (AFDW) was determined. For this, the soft tissue was removed from the shell, dried at 60°C
197 until constant weight and then incinerated at 540°C during 4 hours. The weight difference of
198 dry weight and ash-weight is the ash free dry weight (AFDW). These data were used to
199 calculate the body mass index (BMI: Dame, 1996) or condition by dividing the AFDW by shell
200 volume. Here we calculated shell volume as (length×height×width) .
201

202 **2.3.3 Gonado somatic index.**

203 For every survey date a selection of 50 individuals of different size classes was used to
204 measure the gonadal development. Hereto the gonads were dissected from the somatic
205 tissue under a stereomicroscope. Both tissue types were kept separate, dried and incinerated
206 to obtain their AFDW (see above). These weights were used to calculate the contribution of
207 the gonadal mass in the total AFDW of the animals. Based on these determinations and the
208 collection of data over the two years, the seasonal change in gonadal mass over the years
209 2011 and 2012 could be determined.
210

211 **2.3.4 Glycogen content.**

212 The glycogen content of 114 specimens was determined according to the enzymatic-
213 colourographic method of Keppler & Decker originally published in 1970 and adjusted in 1974
214 (Keppler & Decker, 1970; 1974). Average sample weight for each determination was
215 approximately 40 mg. Glycogen present in the sample is first hydrolyzed to glucose. This
216 glucose is enzymatically transformed (with glucose-6-P-dehydrogenase) into 6-
217 fosfogluconolacton. In this reaction NADPH is formed which is spectrophotometrically
218 determined at a wavelength of 340 nm. A calibration line was used to back calculate the
219 observed extinction values into concentrations. The calibration line was determined on basis
220 of pure mussel tissue glycogen (Sigma G1508-5G).

221 **2.4 Sampling the *Ensis directus* population in the wider coastal zone**

222 The aim of the sampling survey in June 2011 was to determine if the distribution and densities
223 near the "lander" location represented distribution patterns over a wider coastal (along shore)
224 area. During this survey the coastal zone between IJmuiden and Petten was covered. In total
225 8 transects perpendicularly to the coast with 12 boxcore stations each were sampled. Each
226 transect extended up to 6 km from the coast. The three stations closest to the beach were
227 approx. 100 m apart, the stations furthest away from the beach were 1000 m apart and the 4
228 stations in between were 300 meter apart. One boxcore was taken at each station. From each
229 boxcore a small-sized 5 cm deep subsample was taken for the determination of sediment
230 grainsize and mud content (% <63 µm). The remaining boxcore contents were sieved over a
231 1 mm screen. From the residue, living *Ensis directus* were collected, counted and shell sizes
232 measured. Because of the draft of the ship, we could only sample in waters deeper than
233 approximately 10 meter.

234

235 **2.5 Data analyses**

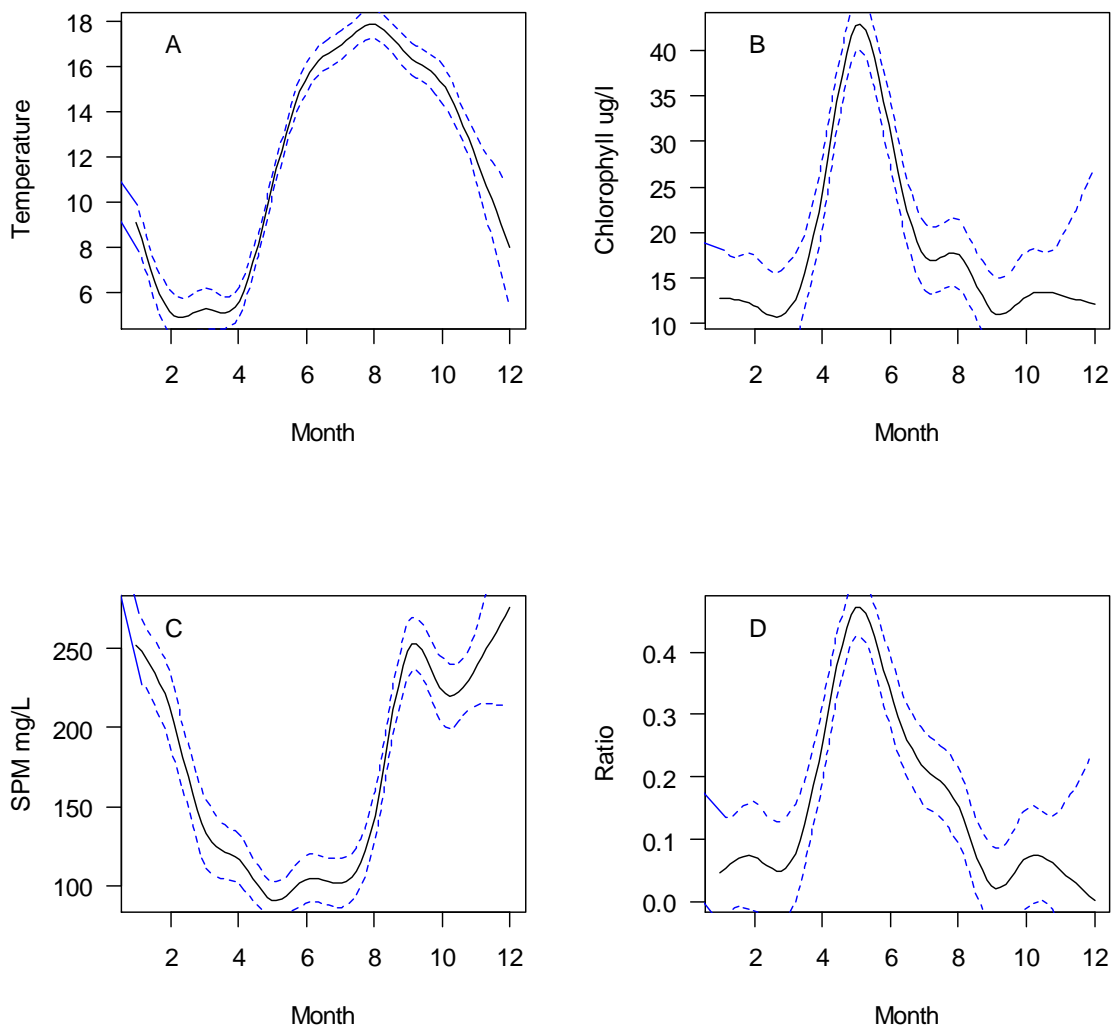
236 Generalized additive modelling (GAM) (Wood, 2006) was used to describe the average
237 seasonal trends in shell growth, tissue growth and condition parameters. GAMs were also
238 used to describe the average seasonal trends in environmental parameters. In this method
239 the determination of the seasonal trends is based on the calculation of a smoothed regression
240 spline which is penalized by the number of variables used. This modelling was done with the
241 package “mgcv” (Wood, 2006) within R (R Core Team, 2012).

242

243 **3 RESULTS**

244 **3.1 Environmental setting lander site**

245 In 2011-2012 the lander study site has a marked seasonal cycle in bottom water temperature,
246 with highest temperatures of about 18°C at the end of summer (Fig. 2). Salinity varies
247 between 26 PSU in winter-spring and 32 PSU in summer. At 140 cm above the seafloor the
248 alongshore semi-diurnal tidal current has maximum speeds which vary over the neap-spring
249 cycle between 70 and 120 cm s⁻¹. The average pressure difference measured over 10 minute
250 long intervals during the deployment corresponds to an average wave height of 1.3 m. The
251 maximum measured pressure amplitude was equal to a water column height difference of 6 m
252 (in December 2011), which is about half of the local water depth. Hence the area can be
253 characterized as an energetic environment.



254 Figure 2. GAM fits describing the averaged (2011-2012) seasonal cycles in (A) temperature, (B) Chlorophyll, (C)
255 SPM and (D) Ratio of Chl-a/SPM as measured over the period February 2011-November 2012. Curves are GAM fits
256 with 95% confidence limits around it. GAMs were constructed with the R package "mgcv" (Wood, 2006).
257
258

259 Chlorophyll concentrations peaked in May with an average near-bottom concentration of 50
260 $\mu\text{g l}^{-1}$. Total SPM concentrations 30 cm above the bottom are at minimum ($\sim 100 \text{ mg l}^{-1}$)
261 between April and August. In autumn and winter the average SPM concentration is here
262 approximately 250 mg l^{-1} . During storms, SPM peak concentrations surpassing 3000 mg l^{-1}
263 have been measured.

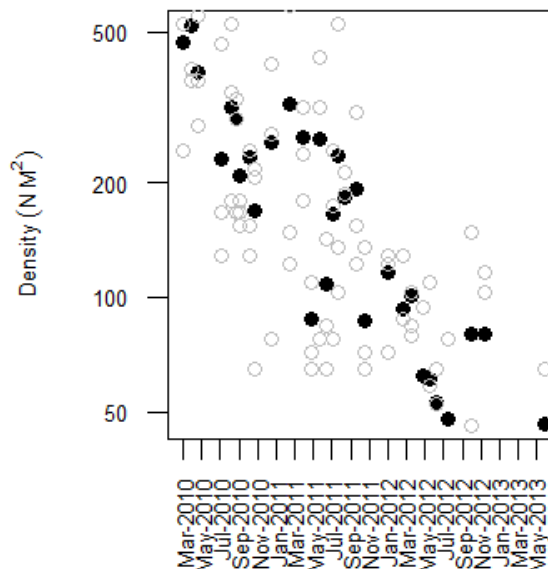
264

265 3.2 Measurements on the *Ensis directus* population.

266 3.2.1 Population density

267 In 2009 a new cohort of *Ensis directus* had settled in the near coastal area off Egmond and in
268 early March 2010 we found initial densities of $\sim 2300 \text{ ind m}^{-2}$ close to but not at the permanent
269 sampling stations (Lander, LNW, LNE, LSW, LSE). At that time a maximum density of ~ 700
270 ind m^{-2} was found at these four stations (Fig. 3). Initially the average densities at the southern
271 two stations (LSW and LSE in Fig. 1) were higher than at the northern stations (LNW and
272 LNE). Later this difference disappeared. At all stations, densities gradually decreased over
273 time. At the end of November 2012 the average density was 80 ind.m^{-2} and half a year later
274 on June 6th 2013 the average density had decreased to 46 ind m^{-2} . On basis of this observed
275 decrease and over an annual time base, the finite survival rate was estimated to be 49% (Fig.
276 3) which is very similar to rates reported by Armonies & Reise (1999) and for the Wadden
277 Sea by Dekker & Beukema (2012).

278



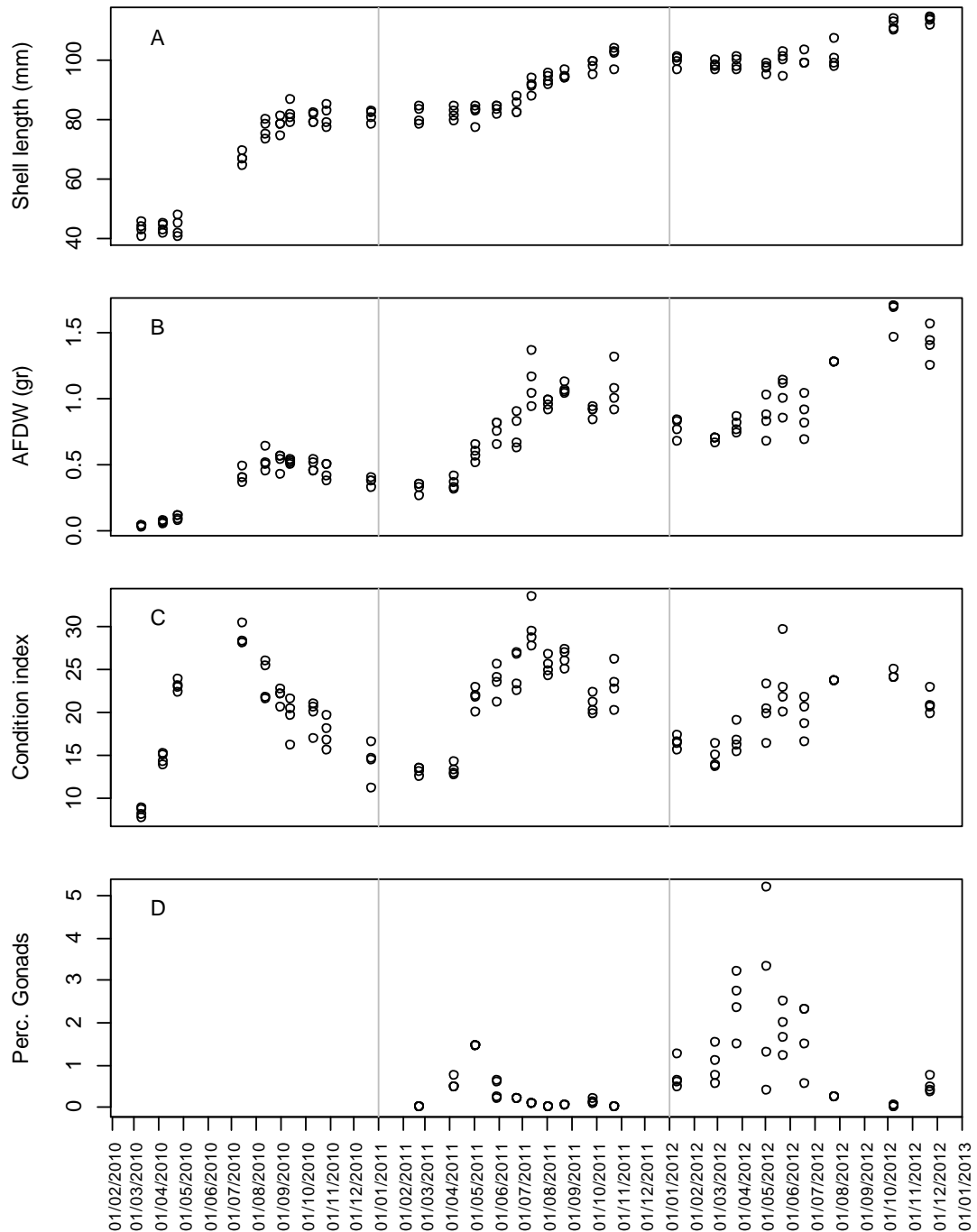
279

280

281 Figure 3. Densities of *Ensis directus* over time sampled at the locations around the measurement platform (LSE,
282 LSW, LNW, LNE) between spring 2010 and June 2013. Grey open circles the density estimates for each of the four
283 stations. Filled dots give the average densities. Note the logarithmic y-axis.

284

285 Morisita's index of dispersion (Young & Young, 1998) was calculated on basis of numbers
 286 caught in all separate boxcore samples during each sampling occasion. This yielded values
 287 which were smaller than those expected in case *Ensis directus* had a clumped distribution
 288 suggesting that at this location and spatial scale of sampling *Ensis directus* is randomly to
 289 evenly distributed.



290
 291 Figure 4. Temporal development of (A) shell length, (B) AFDW, (C) condition index (AFDW shellvolume⁻¹) and (D)
 292 the amount of gonad tissue as percentage of total bodymass between February 2010 and November 2012 measured
 293 for the 2009 cohort of *E. directus*.

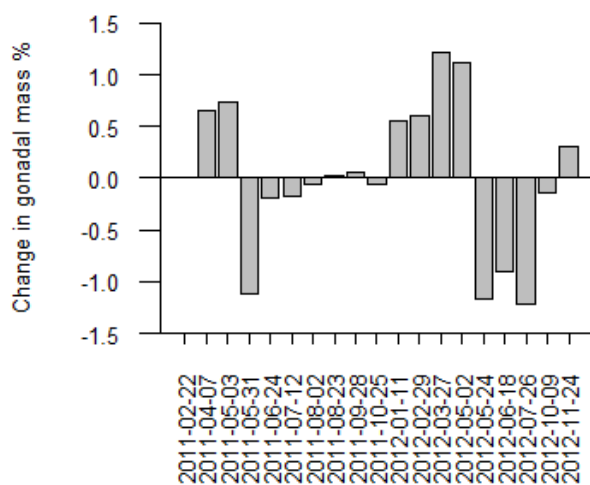
294 **3.2.1 Shell length, AFDW, condition, gonads**

295 The temporal variation of shell length, AFDW and condition index of individuals which had
 296 settled in 2009 and were collected at the four locations around the measurement platform
 297 (LNW, LNE, LSW, LSE) during the study period are shown in Fig. 4. Determination of age
 298 (cohort 2009) was made on basis of externally visible growth lines (Cardoso *et al.*, 2013). The
 299 patterns in Fig. 4A, B illustrate that tissue growth (AFDW) starts before length growth. The
 300 total AFDW increases in spring and early summer. In autumn and winter AFDW decreases
 301 again.

302 Because AFDW is linked to shell size the seasonal change in condition index being AFDW
 303 divided by shell volume (Fig. 4C) might give a better estimate of the temporal evolution of
 304 tissue weights as it is independent of shell size. This shows that the increase in condition is
 305 largest between April and June and parallels the development of the spring bloom (Fig.2B).
 306 After the bloom period, the condition of *Ensis directus* almost immediately starts to decrease
 307 and reaches minimum values in winter and early spring of the following year. This leads to a
 308 cyclic seasonal evolution of the condition index.

309 The proportion of gonad tissue is largest in the beginning of May. (Fig. 4D). The gonadal
 310 mass increases from about 0.5% to a maximum of 3.5 % of the AFDW. Around the time of
 311 spawning 90 % of the dissected specimens had developed gonads and could be sexed. The
 312 difference in the average percentage of gonadal mass between subsequent sampling dates
 313 was used to detect the main spawning season (Fig. 5). This illustrates that in 2011 the largest
 314 loss of gonadal mass was at the end of May. In 2012 the loss of gonadal tissue was spread
 315 over a longer period and lasted from the end of May until the end of July. This evidences
 316 inter-annual variability in the length of the spawning period. In 2012 the one year older and
 317 thus larger animals have a larger proportion of gonadal mass which might imply that a
 318 population with larger and older animals may spawn over a prolonged period.

319



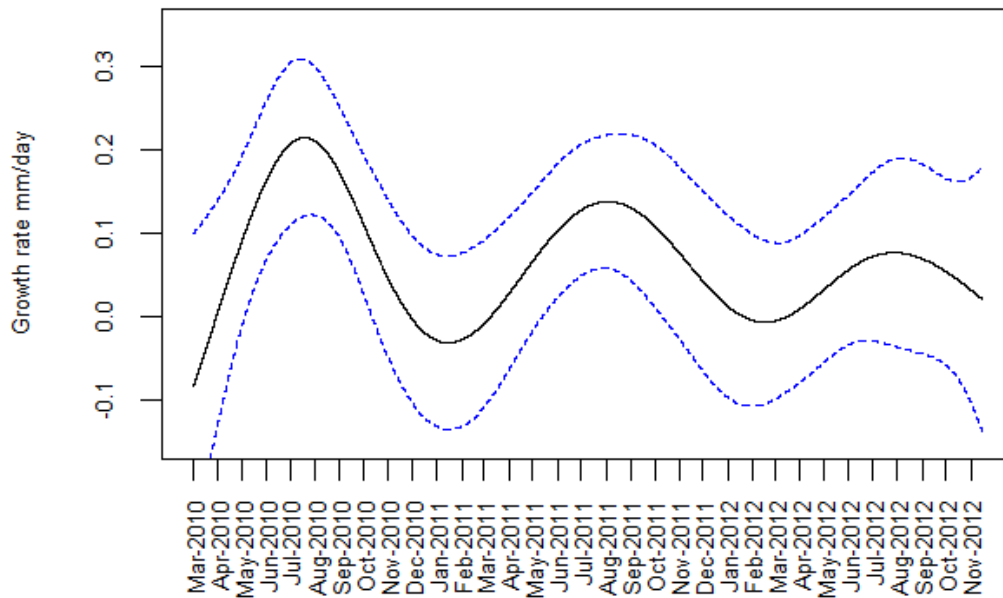
320

321

322 Figure 5. Change in the percentage of gonad tissue between February 2011 and November 2012

323
324
325
326
327
328
329
330
331
332

On basis of the average shell sizes per sample, location and date, shell growth rates could be estimated from the change in average length divided by the time (days) passed since the previous measurement. For this calculation only shells which belonged to the 2009 cohort were used. Figure 6 shows that maximum growth rates decrease with time and illustrates that the older and larger animals have lower absolute shell growth rates. Maximum shell growth rates decreased from about 0.25 mm day⁻¹ in the first year to approximately 0.05 mm day⁻¹ in their third year (Fig 6). The maximum rates were achieved in the summer months only.



333
334
335
336
337
338

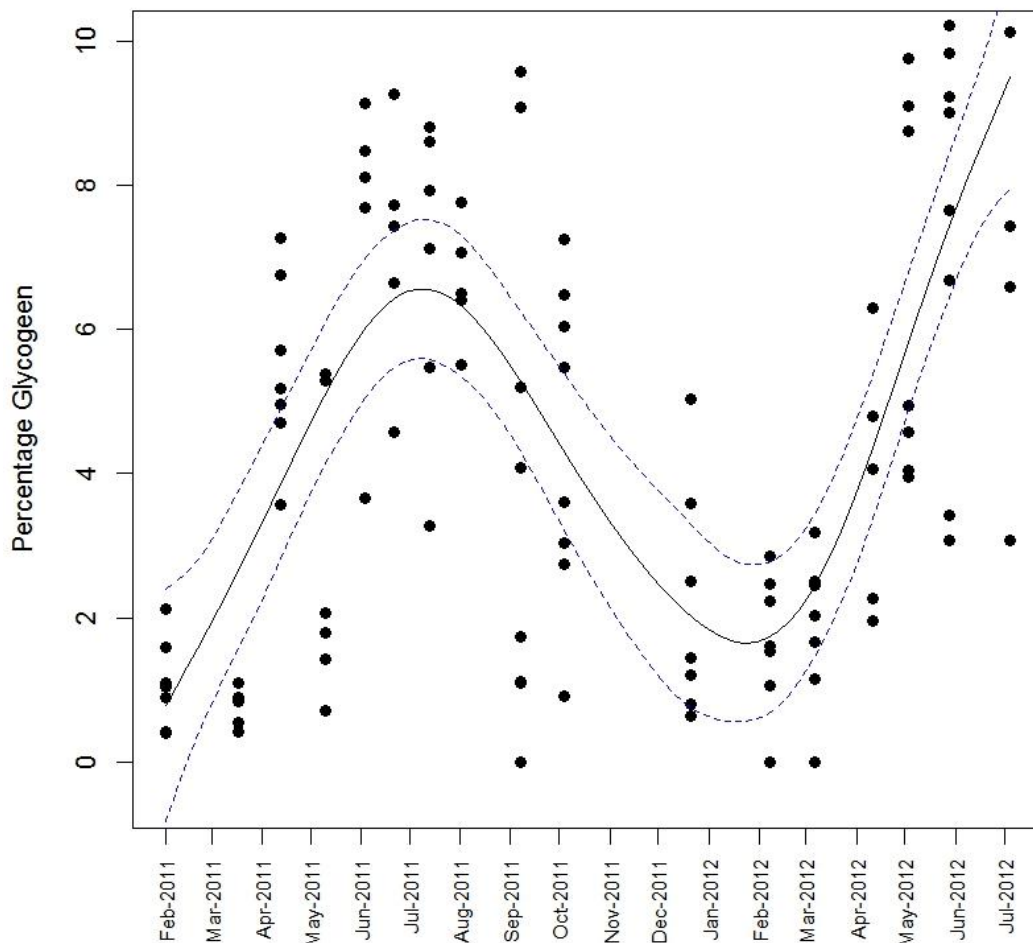
Figure 6. Variation in estimated shell growth rate over time for the 2009 cohort. the black line is a fitted GAM model (Wood 2006) illustrating the change in absolute growth rates over time with its 2.5 % confidence limits around it (dotted lines). The fitted function is highly significant $p < 0.01$.

339 3.2.2 Glycogen values

340 Likewise as the condition index, the energy content of the soft tissue mass also reflects a
341 change in condition as relative amounts of fat, proteins and carbohydrates vary.

342 One of the carbohydrates which is stored as reserve is glycogen. The analyses of the
343 glycogen content showed that the maximum glycogen tissue percentage (by weight) is about
344 10% (Fig. 7). In autumn the percentage glycogen rapidly decreases and between November
345 and March a minimum of less than 1% was found. The observed seasonal cycle in glycogen
346 content matches the observed seasonal cycle in overall condition index (Fig. 4C). The

347 condition index (bodymass shellvolume⁻¹) appeared to be a good predictor of the period
 348 averaged glycogen content. The regression between both is highly significant and the
 349 condition explains 72% of the variance in glycogen content ($p < 0.05$). This strong relationship
 350 shows that the classical determination of the condition index based on AFDW and shell size is
 351 a good predictor of the energy reserves stored within the animal. Sample size per collection
 352 date was too small to examine a potential relationship between glycogen content and body
 353 (shell) size.

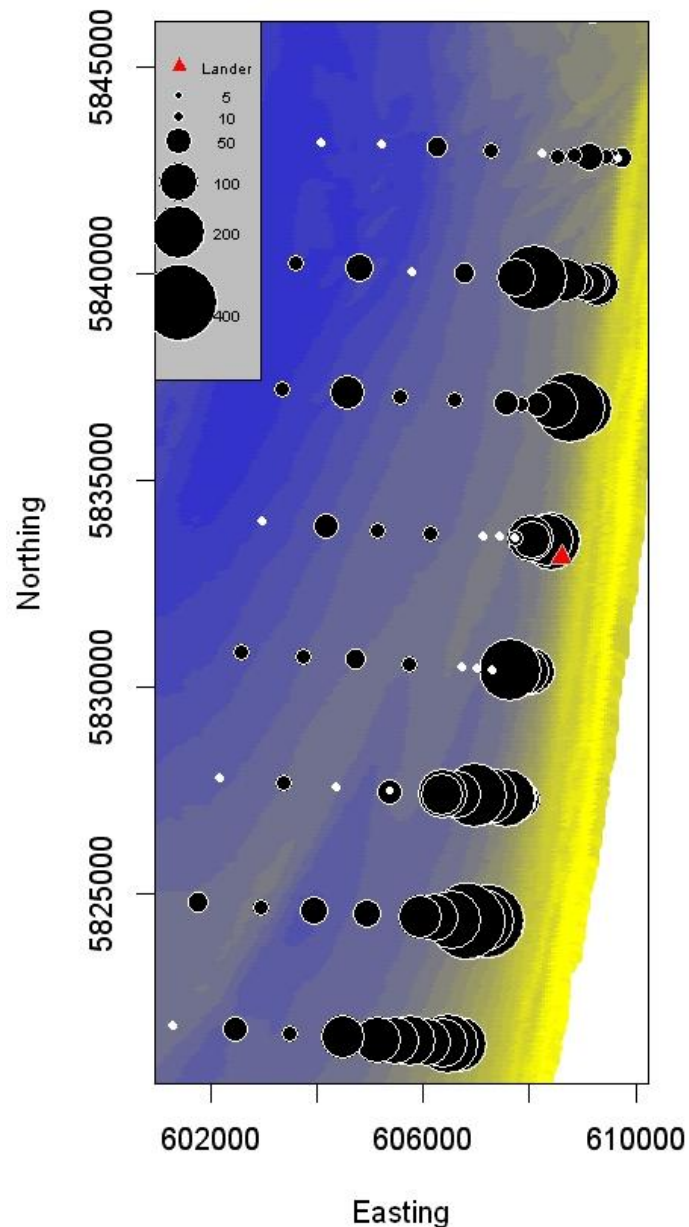


354
 355 Figure 7. Gam fit (Wood, 2006) of the percentage glycogen in homogenized freeze dried *Ensis directus* tissue in the
 356 period 2011-2012. Fitted line is significant at $p < 0.001$.
 357

358 3.3 - Distribution *Ensis directus* in the wider coastal zone

359 The distribution of *Ensis directus* in the deeper parts of the (>10m) coastal zone is plotted in
 360 Fig. 8 which shows that the highest densities are found in a rather narrow band closest to the
 361 coast. The maximum observed density during this inventory in 2011 was 397 individuals m⁻²
 362 with a mean density of 90 individuals m⁻². At the stations nearest to the shore, the average
 363 size of the animals was considerably smaller (88 mm) than at the stations lying further

364 offshore (120 mm). This was mainly due to the difference in age of the specimens collected
365 from near shore and off shore locations. Over the entire area the average size was ~99 mm.
366 The analyses of the sediment samples taken along the transects showed that median grain
367 size increased with distance from the coast. The stations furthest away from the coast had
368 slightly lower mud concentrations when compared to the near coastal stations. Over the
369 sampled area the median grain size decreases from north to south, but only the most
370 northerly transect had a significantly different median grain size (Tukey HSD, $p < 0.05$). The
371 mud content did not differ between transects.
372 The map in Fig. 8 furthermore illustrates that densities of *Ensis directus* at the sampling sites
373 near the platform (lander) and hence the data on growth and production of the *E. directus*
374 population, are representative for a larger part of the near-shore coastal zone of North
375 Holland.
376



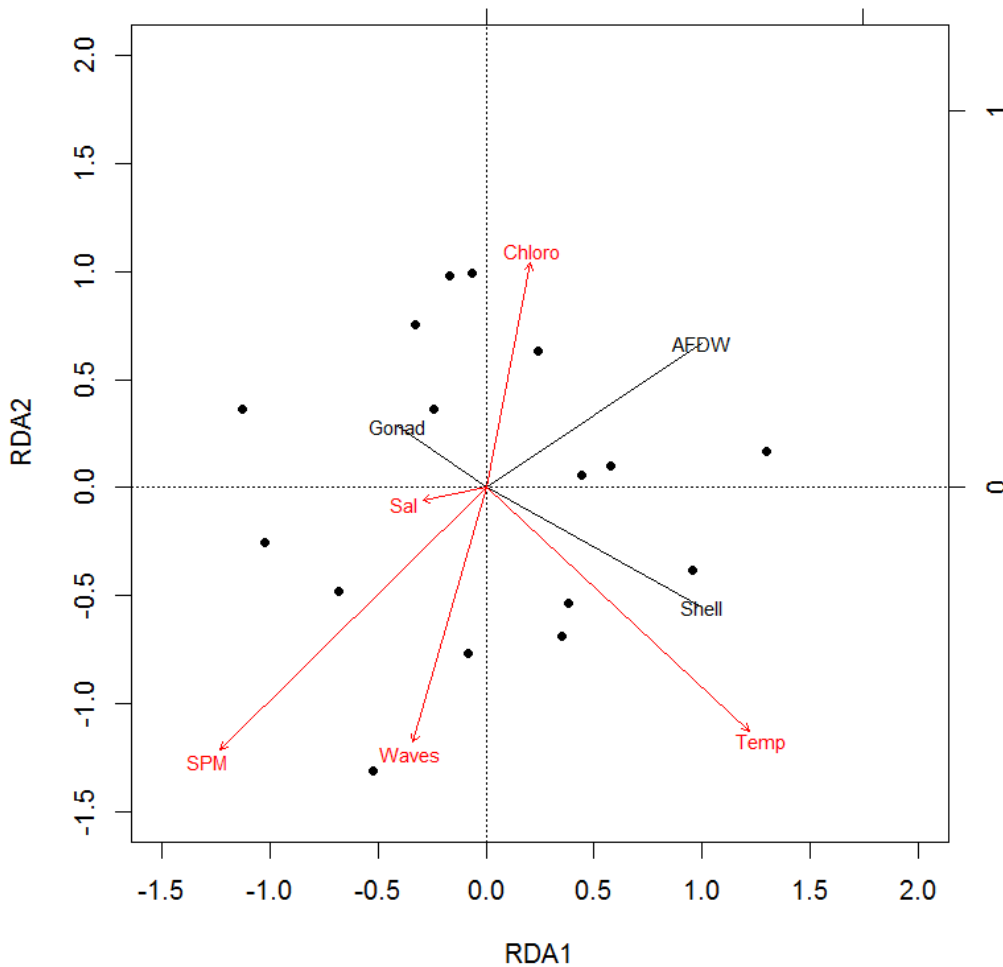
377
 378
 379
 380
 381
 382
 383

Figure 8. Distribution and density of *Ensis directus* (all year classes) along the coast of North Holland in June 2011. The shallow area (<10m) could not be sampled. Hence, no information on densities in this area are available. Black filled circles of different size indicate density. White circles indicate absence from sample. The triangle indicates the deployment location of the measurement platform (lander).

384 **3.4. Growth and condition of *Ensis directus* in relation to environmental conditions**

385 We used a redundancy analyses (RDA, R-package: vegan, Oksanen *et al*, 2013) to obtain
 386 insight in the relation between shell and tissue growth of *Ensis directus* and seasonal
 387 variations in environmental factors. For this we used the data collected with and around the
 388 permanent measurement platform (Fig. 1). During the deployment of the platform we
 389 continuously measured several variables, i.e. temperature, salinity, pressure (waves), SPM
 390 and fluorescence, all of which can potentially influence the growth of *E. directus*. For this RDA
 391 analyses we used the station averaged data, i.e. the means of growth and condition of the

392 four stations (LSE, LSW, LNE, LNW) during each of the measurement periods, together with
 393 the period averaged environmental conditions simultaneously recorded by the lander at the
 394 central location (Fig. 1). Selection of explanatory variables and covariates used in this
 395 analyses was made on basis of their contribution to variance inflation ($VIF < 3$). Therefore
 396 condition and glycogen were omitted from the analyses. The result of the RDA analysis is
 397 given in Fig. 9.



398
 399 Figure 9. RDA correlation triplot showing the relationships between abiotic variables (averaged over periods) and the
 400 *E. directus* growth parameters determined at the end of each of these periods. Measurements cover the years
 401 2011-2012. Shell= shell growth, AFDW= Ash free dry weight growth, Gonad,=change in relative gonadal mass.,
 402 Chloro=average chlorophyll concentration at 30cm above the seafloor, Sal=average Salinity, SPM = average
 403 Suspended matter concentration, Waves = average waveheight and Temp= average temperature.
 404
 405

406 The explanatory variables explain about 49% of the observed variance in growth and
 407 condition parameters of *Ensis directus*. The (correlation) triplot made on basis of the first two
 408 RDA axis shows that wave height and SPM are strongly linked and are inversely related to
 409 tissue growth. Tissue growth (AFDW in Fig. 9) is positively related to chlorophyll (Chloro).
 410 Strikingly, shell growth (Shell) appears to be unrelated to tissue growth (AFDW). Instead shell
 411 growth is positively correlated to water temperature. The change in gonadal mass (Gonad) is

412 unrelated to SPM or chlorophyll. The plot in Fig. 9 suggests an inverse relationship between
 413 gonadal mass and temperature and independency from tissue growth or condition.

414 A permutation test was used to determine the order of importance of separate environmental
 415 factors in explaining the observed growth responses. This showed that both temperature and
 416 SPM contribute significantly in explaining the observed shell and tissue growth rates and
 417 condition changes (Table 2).

418

419 Table 2. Overview of the test statistics of the permutation test to determine the order of importance of the
 420 various environmental factors in explaining growth and condition parameters (Shell growth, AFDW growth,
 421 change in Gonadal mass and average caloric content as measured in *E. directus* in 2011 and 2012.
 422

	Df	AIC	F	N.Perm	Pr(>F)
SPM	1	16.549	3.9725	9998	0.007301 **
Temp	1	13.891	4.3925	9998	0.007401 **
Chloro	1	14.115	1.4090	9998	0.253925
Sal	1	15.721	0.2743	9998	0.852485
Waves	1	17.633	0.0549	9998	0.981798

423

424 The calculated variance inflation factors suggest that SPM and Chlorophyll could be used
 425 simultaneously in the RDA (correlation <0.7). Nevertheless there is an inverse relationship
 426 between both as they have strong opposite seasonal trends ($r=-0.61$). This not necessarily
 427 indicates a causal relationship but complicates the interpretation of Fig. 9. Therefore the RDA
 428 analyses was repeated with the ratio (Chloro:SPM) instead of the separate factors (Chloro
 429 and SPM). The ratio expresses qualitative aspects (μg Chlorophyll per mg SPM) and
 430 circumvents the potential dependency of Chlorophyll and SPM in the RDA. It describes the
 431 seasonal effect in the relative amounts of SPM and Chlorophyll as one quantity. Incorporating
 432 the ratio in the RDA analysis showed that the total variance explained by the explanatory
 433 variables becomes 38%. This analysis suggests that tissue growth is positively correlated to
 434 the quality of suspended matter (Chloro:SPM =Ratio). Shell growth remains indifferent to the
 435 suspended matter quality. A permutation test showed that both temperature and the Ratio
 436 (Chloro:SPM) are highly significant ($p<0.05$). The RDA analyses suggests relationships
 437 between growth and environmental controls with SPM concentration being most important. It
 438 is however not entirely clear whether the absolute amounts of SPM or the quality of the SPM
 439 (ratio Chloro:SPM) is the key factor.

440

441

442 4 DISCUSSION

443 4.1 Growth of the *E. directus* population

444 Concern about the possible impact of beach nourishments on the production of the local
445 *Ensis directus* population initiated the present project. There are various studies which review
446 the potential effects of beach nourishments (Essink, 1999; Greene, 2002; Speybroeck et al,
447 2006). Increased turbidity is one of the potentially important factors identified by above
448 studies. However, actual measurements on turbidity close to the bottom where bivalves live,
449 its natural seasonal variations and how they relate to maintenance activities are lacking.

450 Therefore we focussed on growth and energy allocation in *E. directus* in relation to variation in
451 environmental conditions. These data were anticipated to give insight into factors that are
452 important for shell and tissue growth and their seasonal timing.

453 All our measurements on tissue composition and growth of *E. directus* demonstrated strong
454 seasonal cycles most of which closely followed the spring bloom peak in chlorophyll
455 concentrations except for gonadal development which started before the spring bloom. In our
456 study the earliest signs of a seasonal increase of the gonadal mass were visible in December
457 2012 (Fig. 4D). But also in the previous winter of 2011-2012 the gonadal mass had already
458 started to increase in February-March. This early start of gonadal development implies that
459 most of the energy uptake in winter is allocated to gonad growth as by that time no shell or
460 significant tissue growth was observed. In fact our data suggests that this early in the year,
461 total AFDW still decreases thus gonad weights apparently increase at the expense of the
462 somatic tissue. In early May, the gonadal mass is maximal and is followed by spawning.
463 Gonad development peaks during the phytoplankton spring bloom and as a consequence of
464 this timing the planktonic larvae appear in a period in which they optimally profit from rich food
465 conditions in the water column. For *Ensis arcuatus* Darriba et al. (2005) observed similar
466 timing with an early gonadal development before the phytoplankton blooms in their study area
467 had started. This shows that the gonads in *E. arcuatus* also developed at the expense of
468 stored energy but that the larvae of *E. arcuatus* are released in a period of maximal food
469 availability.

470 In *Ensis directus* we observed that shortly after the observed peak in gonadal mass and
471 spawning, the somatic tissue starts to increase but at this time shell growth is still not yet
472 evident. We observed that significant shell growth did not take place before the water
473 temperature was about 12-14 °C. Latter observation is confirmed by the carbonate stable
474 oxygen isotope data as reported by Cardoso *et al.* (2013) and similar uncoupling between
475 tissue growth and shell growth has also been observed for other species (Hilbish, 1986; Lewis
476 & Cerrato, 1997).

477 The sequence of changes in the growth of gonads, somatic tissue, and shell as described
478 here was seen in both 2011 and 2012. In 2010 this sequence of tissue and shell growth could
479 not be determined. Various other studies have reported on the timing of reproduction of *Ensis*
480 *directus*. Mühlenhardt-Siegel et al (1983) estimated that a population living in the German

481 Bight spawns in March and April. This seems earlier in the year than what we found. But the
482 data collected by Pulfrich (1997) in the German Wadden Sea as well as the observations by
483 Cardoso *et al* (2009) are in line with our observations for the Egmond population in 2011 and
484 2012. While most studies report the main spawning in the spring, Cardoso *et al.* (2009) found
485 a second (small) peak in gonadal mass in July and August. Pulfrich (1997) and Philippart *et al*
486 (2014) found a similar second peak in larval abundance in the German and Dutch Wadden
487 sea respectively. It is thus likely that a second spawning event might exist in late summer.
488 This also has been suggested by the findings of Armonies & Reise (1999) and Strasser &
489 Günther (2001) and multiple spawning peaks have been observed for closely related species
490 as well (Darriba *et al*, 2004, Barón *et al*, 2004).

491 In our Egmond population we could identify a small peak in the number of animals which
492 could be sexed in September 2011 but it is questionable whether the percentage of gonadal
493 mass (0.1%) is high enough to represent a secondary spawning peak. The data however
494 show that in 2012 the period over which ripe animals were present was much longer than in
495 2011. This suggests that spawning may take place over an extended period or in several
496 pulses. Partial release of gametes may be possible in large animals whose energy reserve is
497 sufficiently large to maintain or build such a large gonadal mass. We observed that in the
498 primary spawning period (Feb-June) indeed a relation exists between the maximum
499 percentage of gonadal tissue and animal size. The data also show that the percentage of
500 gonadal mass in 2012 in comparison to 2011 was significantly larger (ANOVA, $P < 0.01$).
501 These observations thus suggest that larger and older animals in a population not only have a
502 relatively larger contribution to the total spawning mass but also might be able to generate
503 multiple spawning peaks.

504 Unpublished NIOZ-data collected in 2007 and derived from monthly water samples along a
505 east-west transect across the same coastal area, shows that pelagic larvae of *Ensis directus*
506 are present almost year round. This can only be explained by spawning which takes place
507 over prolonged periods possibly in combination with lateral transport of larvae, originating
508 from other source populations. Despite the above observations of an almost continuous
509 presence of larvae and the likely existence of multiple spawning peaks, we never observed
510 significant recruitment of new spat between 2010 and 2012. Only by the end of 2012 when
511 the density of adult *E. directus* had dropped to 40-60 m⁻², low numbers of recruits were found.
512 This suggests that density dependent processes might regulate settling. Other observations
513 that point to density dependent recruitment processes come from a pilot settlement
514 experiment performed prior to the present study in 2010. In this experiment we observed
515 settling of high numbers of juveniles in tubes which were filled with clean defaunated sand
516 while in tubes filled with living or dead *Ensis* no spat was found.

517

518 4.2 Relating density and growth to environmental conditions

519 The distribution of *Ensis directus* along the Noord-Holland coast (Fig. 8) shows that within the
520 extent of our survey, the highest densities are found in the transitional zone where the
521 seafloor abruptly descends to a depth of 10 meter and where waves have a strong effect on
522 the seabed due to the shallow depth and topography. This is also the type of habitat where
523 this species is typically found along the east coast of the US and Canada (Kenchington, *et al.*,
524 1998). Also in the Wadden Sea, maximal densities are found in a transitional zone between
525 intertidal and sub-tidal area (Armonies & Reise, 1999; Dekker & Beukema, 2012;
526 Freudendahl *et al.*, 2010). These areas are characterized by strong tidal currents and mobile
527 sands with low silt contents (Dekker & Beukema, 2012). The lander observations in this study
528 corroborate the view that *Ensis directus* prefers a dynamic environment. However, our data
529 do not support the idea that high *E. directus* densities are necessarily found in the sediments
530 with lowest mud contents. At the outermost stations (Φ 222 μm , Mud=2.9%) of the survey grid
531 only low densities of large animals were found. High densities of *E. directus* were found close
532 to the coast at water depths of approximately 10 meter and in sediments with an average
533 percentage of mud of approximately 5% and a median grain size of 185 μm . Dannheim &
534 Rumohr (2012) also found the highest densities in fine silty sands at water depths between 10
535 and 13 meter. We cannot exclude that *E. directus* densities at our location may attain even
536 higher levels in shallower water i.e. between low water mark and 10 m depth where we could
537 not sample due to the draft of our vessel. Therefore the survey does not give a complete view
538 of the distribution of this species over the entire depth range in the near coastal zone.

539 In this study, we have correlated growth of *Ensis directus* with in-situ measured environmental
540 parameters. A different approach to determine effects of silt and algae on growth was
541 adopted by Kamermans *et al.* (2013) who tested the effects of increased levels of kaolin and
542 algae on clearance rate and growth of *Ensis directus* at $\sim 18^\circ\text{C}$. They observed that the
543 clearance rate was reduced at suspended silt concentrations of 300 mg l^{-1} , but found that this
544 condition stimulated growth for medium sized animals. Their study showed that at the same
545 silt concentrations, a higher algal concentration led to increased tissue growth. The final
546 AFDW appeared dependent on the Chl-a concentration, but not on silt concentration. This
547 corresponds with our field observations showing a positive relationship between tissue growth
548 and Chlorophyll and between tissue growth and the Chl-a:SPM ratio.

549 The average SPM background concentration at 30 cm above the seafloor and over the entire
550 measurement period was 161 mg l^{-1} . During sand nourishments in June 2011 close to our
551 Egmond location, SPM showed elevated levels to approximately 250 mg l^{-1} . This value
552 approaches the critical concentration of 300 mg l^{-1} where the clearance rate of *E. directus* is
553 affected (Kamermans *et al.*, 2013). In comparison to SPM concentrations measured during
554 wind events the SPM increase due to the local maintenance works were relatively small. We
555 observed that waves during storms generate peaks in SPM to over 3000 mg l^{-1} . At our
556 location the observed variability and short lasting increase in SPM due to maintenance works
557 evidently fell within the natural variability range of SPM close to the bottom. Our time series

558 furthermore showed that during the main growing season for *Ensis directus* (March-Aug) the
559 SPM concentrations surpassed the 300 mg l⁻¹ threshold during only 6 % of the time. On basis
560 of this limited amount of time, it seems unlikely that a significant effect of elevated SPM levels
561 on growth, mortality and production of the local *Ensis directus* population can be
562 demonstrated. Especially since Kamermans et al (2013) also observed a positive effect of
563 elevated SPM on growth of medium sized animals.

564 To corroborate this assumption we have put our data on density, mortality and production of *E.*
565 *directus* off Egmond in wider perspective by comparing them with data from studies on
566 populations living elsewhere. Although the various studies refer to different years, cohorts and
567 locations the results enable a comparison with our data of the Egmond population.

568 The densities of early juveniles (end 2009 early 2010) at the lander site near Egmond (700-
569 2300 ind m²) appeared to have the same order of magnitude as found for spat and juveniles
570 in the subtidal and transition zone of the Balgzand area in the Wadden Sea (Dekker &
571 Beukema, 2012). The spat densities at various locations in the German Bight (Dannheim &
572 Rumohr, 2012) are based on sampling in July. These estimates are therefore much higher
573 when compared to our first density estimates which are based on sampling in December 2009
574 after mortality during autumn has already reduced the number of settlers.

575 Densities of older animals around the measurement platform are comparable to densities
576 reported in subtidal areas in the Wadden Sea (Armonies & Reise, 1999; Dekker & Beukema,
577 2012) or for the German Bight (Dannheim & Rumohr, 2012).

578 Survival and mortality rates are strongly dependent on the age and the cohort which has been
579 studied. For the Egmond location the 2009 cohort had an average survival rate of 49% over
580 the entire study period. This value compares well with the rate (51-45%) which is found for *E.*
581 *directus* living in the subtidal and transitional zone of the western Wadden Sea (Dekker &
582 Beukema, 2012) and the value of 55% given by Armonies & Reise (1999) for the German
583 Bight. Survival of the 0-age class appears to be very variable and is often only a few percent
584 of the initial densities. For our study area we could not estimate this quantity as we have no
585 data on densities shortly after settlement in summer and autumn 2009.

586 Longevity seems to be area dependent. For the Wadden Sea maximum ages of 5 years have
587 been mentioned (Dekker and Beukema, 2012). Palmer (2004) as well as Armonies & Reise
588 (1999) report a maximum age of 7 years for the Wash and the German Bight, respectively.
589 Based on external ring counts (see Cardoso et al, 2013) we estimate a maximum age of 8 to
590 9 years for individuals from the Egmond population. The maximum shell length found at
591 Egmond was 18.2 cm which compares well to the maximum shell lengths reported by
592 Armonies & Reise (1999) for the population around the isle of Sylt (German Bight).

593 Indicative for the conditions of growth is the average size at age. At the end of the first
594 growing season after settlement, the Egmond specimens have obtained a shell length of ~ 4-
595 7 cm. This fits the range as reported in the summary table made for various populations by
596 Dannheim & Rumohr (2012). The population and cohort we studied however seems to have
597 retarded shell growth in their 3rd and 4th year. Maximum sizes found in these years were

598 respectively 10.6 and 11.4 cm. Beukema & Dekker (1995) reported shell lengths of 12.6 cm
599 and 14.4 cm for the same yearclasses. Also Mühlenhardt et al (1983) and Armonies & Reise
600 (1999) report larger sizes for the third and fourth year of growth. Thus while densities,
601 mortality, maximum age of the Egmond population vary within the ranges reported in
602 literature, shell size at later age appears to be relatively small. This can point to an effect of
603 the coastal maintenance works in 2011. Therefore we checked whether body condition
604 parameters showed a potential effect of coastal maintenance works.

605 Because we used a slightly different way to calculate the BMI a recalculation was done which
606 showed that the BMI (sensu Dekker & Beukema, 2012) for the Egmond population ranged
607 between 0.3 and 1.6 mg AFDW cm⁻³ which compares well to the range given in Cardoso et al
608 (2009). The minimum value we found (0.3) is slightly lower than the absolute minimum (0.37)
609 mentioned by Dekker & Beukema (2012).

610 Another measure for the condition of bivalves is the percentage of glycogen stored in their
611 tissue (Fernandez-Reiriz et al, 2007, Hummel et al, 1988). The seasonal trend in this
612 percentage shows that high values in *E. directus* follow periods of high food availability (Fig.
613 7, Fig 2B). It shows that glycogen is stored as energy reserve during such periods. In other
614 species a similar seasonal trend has been found (Hummel et al., 1988). The maximum
615 percentage glycogen in *E. directus* from Egmond was about 10% during the summer months.
616 This value is lower than the ranges of total carbohydrates (Glucose + Glycogen) in bivalves
617 as given by Beukema (1997) but is twice the maximum amount of glycogen which was found
618 in adductor muscles of the closely related *E. arcuatus* (Darriba et al, 2005). This suggests
619 that *E. directus* near the lander site had a good condition. A good condition is furthermore
620 supported by the strong correlation between the percentage glycogen and the condition index
621 ($r=0.88$, $p<0.001$, this study) especially since the condition indices compared well (see above)
622 with data published by Cardoso (2009) or Dekker and Beukema (2012). Also the maximum
623 amount of reproductive tissue which we found in *E. directus* from Egmond is very similar to
624 that found in specimens from the Wadden Sea (Cardoso et al, 2012). Thus the values of
625 various parameters indicative for the condition do not provide clear evidence that *E. directus*
626 from Egmond are performing poorly in the area of study.

627 From an ecosystem point of view, production by the *E. directus* population might be a good
628 measure of performance of the local population. Over the research period the biomass off
629 Egmond, varied between 50 and 322 gr AFDW m⁻². Total production of the *Ensis* population
630 was estimated to range between 200 gr and 50 gr AFDW m⁻² yr⁻¹ in 2011 and 2012,
631 respectively. These values are substantially higher than the average estimates (8.6 ± 3.4 , max
632 100gr AFDW m⁻² yr⁻¹) for the subtidal Wadden Sea population as given by Dekker & Beukema
633 (2012). Thus although length growth of individuals at Egmond in their 3rd and 4th year seems
634 to be less compared to other populations, annual production equals or even exceeds that of
635 the subtidal population in the western Wadden Sea.

636 In summary, there seems no evidence that population development and production of the
637 Egmond *Ensis* population deviates markedly from what could be expected on the basis of

638 data from other populations. Hence, we cannot prove that there is a negative effect of
639 elevated SPM levels on performance of *Ensis directus* in this area.

640

641 **4.3 Potential explanations for the dense near coastal distribution of *E. directus***

642 Within our survey the highest densities were found in a distinct zone at approximately 10
643 meter depth. This pattern was observed along all transects perpendicular to the coastline and
644 below we present some hypothetical explanations for this pattern.

645 A first hypothesis involves the temporal mismatch between temperatures high enough for
646 shell growth in *E. directus* and the presence of food with a sufficient high quality. Our data as
647 well as those by Cardoso *et al.* (2013) show that there is no or minimal shell growth at
648 temperatures below 14 °C during times that food quality peaks. The zone where we found
649 highest *E. directus* densities (Fig. 8) coincides with persistently high near-bottom
650 concentrations of SPM (van der Hout, 2014 submitted) but at the same time is so close to the
651 upper shoreface that the seasonal rise in water temperatures will closely follow the seasonal
652 rise in air temperature and thus minimize the mismatch between food availability and
653 occurrence of temperatures high enough for shell growth.

654 A second hypothesis explaining the peak in densities of *E. directus* so close to the coast is
655 that high numbers of larvae passively accumulate in this zone similarly as high concentrations
656 of SPM. The hydrography in this area (van Rijn, 1995; de Boer, 2009) triggers cross shore
657 currents with down and upwelling in case of salinity stratification due to fresh water discharge
658 by the river Rhine. The persistent presence of high SPM loads close to the bottom in this area
659 (van der Hout, 2014) lends support to the “larval accumulation” hypotheses. A comparable
660 mechanism has been proposed by Shanks & Brink (2005) to work on much larger spatial
661 scales. Such a mechanism would imply that the high *E. directus* densities in this nearshore
662 zone are merely caused passively and not based on larval selection.

663 An alternative hypothesis is that summer growth of microphytobenthos in the shallow
664 foreshore might be important for the nearby *E. directus* population. Our continuous
665 measurements of chlorophyll and SPM showed that chlorophyll concentrations peaked after
666 strong winds, even in periods outside the main bloom season. Microscopic analyses of macro
667 aggregates which were collected in June 2011 indeed showed the presence of large numbers
668 of benthic diatoms in this material. We speculate that the chlorophyll peaks, observed during
669 strong wind events, originate from the shallow part of the shoreface where enough light
670 reaches the bottom to generate primary production by benthic diatoms. During strong winds
671 these algae are detached from the substratum and become resuspended and concentrate
672 just below the steep slope of the shore face where SPM accumulates and high densities of
673 *Ensis* are found as well. Tidal mixing in this zone keeps these algae in resuspension and
674 available to the dense *Ensis* beds. This source is likely to provide them with fresh algal
675 material outside the (pelagic) phytoplankton bloom season. Due to depth limitations of the
676 research vessel we could not test this hypotheses but preliminary measurements of light

677 intensity at our 10 meter deep study site suggest that at that depth and during calm weather
678 sunlight reaches the sea floor. So for the more shallow shore face it is even more likely that
679 enough light reaches the seafloor to support primary production by an epi-benthic diatom
680 community.
681

682 **5. CONCLUSIONS**

683 Following a local *Ensis directus* population at a shallow subtidal coastal North Sea site
684 revealed marked seasonal cycles in gonadal development, soft tissue growth and shell
685 growth. Somatic tissue growth does not take place until spawning in May. Thereafter shell
686 growth begins when temperatures reaches 14°C or higher. The population characteristics
687 such as mortality, growth and production are comparable to population variables measured in
688 the Wadden Sea or German Bight. On basis of this observation in combination with the
689 observed range of variation in measured SPM concentrations we assume that the supposed
690 negative effects of elevated SPM concentration due to coastal maintenance works are
691 minimal.

692 An inventory of *Ensis directus* along the North Holland Coast shows that within the extent of
693 our survey the highest densities of *Ensis directus* are found closest to the coast at depths of
694 approximately 10 meter. To explain this distribution alternative hypothesis have been put
695 forward.

696 **ACKNOWLEDGEMENTS.**

697 This work has been made possible by financial support of the La Mer foundation (ref nr;LM-
698 006588), a grant of Ecoshape (projectnr NTW3.1 – NTW2.5) within the framework of BWN
699 and NIOZ itself. The practical work could not have been done without the enormous input of
700 the crews of RV "Terschelling", "Pelagia" and the "Navicula". We would like to thank Joost
701 van der Hoek, Evaline van Weerlee, Charlotte Saul, Job ten Horn and Carola van der Hout for
702 their help on board.

703

704

705

706 **REFERENCES**

707 Arias, A., Anadon, N. 2012. First record of *Mercenaria mercenaria* (Bivalvia: Veneridae) and
708 *Ensis directus* (Bivalvia: Pharidae) on Bay of Biscay, Iberian Peninsula. J. Shellfish. Res. 31
709 (1): 57-60.

710

711 Armonies, W., 2001. What an introduced species can tell us about the spatial extension of
712 benthic populations. Mar. Ecol. Prog. Ser. 209: 289-294.

713

714 Armonies, W., Reise, K., 1999. On the population development of the introduced razor clam
715 *Ensis americanus* near the island of Sylt (North Sea). Helgoländer Meeresun. 52: 291-300.

716

717 Beukema, J.J., 1997. Caloric values of marine invertebrates with an emphasis on the soft
718 parts of marine bivalves. Oceanography and Marine Biology; an annual review 35:387-414.

719

720 Beukema, J.J. Dekker, R. , 1995. Dynamics and growth of a recent invader into European
721 coastal waters: The american razor clam, *Ensis directus*. J. Mar. Biol. Ass. U.K., 75: 351-362.

722

723 Barón, P.J., L.E. Real, N.F. Ciocco & M.E. Re., 2004. Morphometry, growth and reproduction
724 of an Atlantic of an Atlantic population of the razor clam *Ensis macha* (Molina, 1782). Sci.
725 Mar. 68: 211-217.

726

727 Cardoso, J.F.M.F., Nieuwland, G., Witbaard, R., Veer, H.W., van der, Machado, J.P., 2013.
728 Growth increment periodicity in the shell of the razor clam *Ensis directus* using stable
729 isotopes as a method to validate age. Biogeosciences 10: 4741-4750.

730

731 Cardoso, J.F.M.F., Witte, J.I., Veer, H.W. van der, 2009. Reproductive investment of the
732 American razor clam *Ensis americanus* in the Dutch Wadden Sea. J. Sea Res. 62: 295-298.

733

734 Dame, C.F., 1996. Ecology of marine bivalves: An ecosystem approach. CRC Press, Boca
735 Raton. 272 pp.

736

737 Dannheim, J., Rumohr H., 2012. The fate of an immigrant: *Ensis directus* in the eastern
738 German Bight. Helgoland Mar. Res. 66: 307-317.

739

740 Darriba, S., San Juan, F., Guerra, A., 2004. Reproductive cycle of the rzaor clam *Ensis*
741 *arcuatus* (Jeffreys, 1865) in northwest Spain and its relation to environmental conditions. J.
742 Exp. Mar. Biol. Ecol. 311, 101-115.

743

744 Darriba, S., San Juan, F., Guerra, A., 2005. Energy storage and utilization in relation to the
745 reproductive cycle in the razor clam *Ensis arcuatus* (Jeffreys, 1865). ICES J. Mar. Sci.
746 62:886-896.
747

748 De Boer, G.J. 2009. On the interaction between tides and stratification in the Rhine region of
749 the Fresh water influence. PhD thesis Delft University of technology. 210pp.
750

751 Dekker, R., Beukema, J.J., 2012. Long-term dynamics and productivity of a successful
752 invader: The first three decades of the bivalve *Ensis directus* in the western Wadden Sea. J.
753 Sea Res. 71: 31-40.
754

755 Drew, G.A., 1907. The habits and movements of the Razor shell clam, *Ensis directus* Con.
756 Biol. Bull. XII(3): 127-142.
757

758 Ellerbroek, G., Rozemeijer, M.J.C. Kok, J.M. de, Ronde, J. de, 2008. Monitoring and
759 Evaluation Programme Sand mining RWS LaMER, part B5 of the evaluation programme sand
760 mining. Ministerie van verkeer en waterstaat, Noord Holland.
761

762 Essink, K., 1999. Ecological effects of dumping of dredged sediments: options for
763 management. Journal of Coastal Conservation 5: 69-80.
764

765 Fernández-Reiriz, M.J., Pérez-Camacho, A., Delgado, M., Labarta, U., 2007. Dynamics of
766 biochemical components, lipid classes and energy values of gonadal development of *R.*
767 *philippinarum* associated with the temperature and ingestion rate. Comp. Biochem and
768 Physiol. Part A. 147:1053-1059.
769

770 Freudendahl, A.S.F., Nielsen, M.M., Jensen, T. Jensen, K.T., 2010. The introduced clam
771 *Ensis americanus* in the Wadden Sea: field experiment on impact of bird predation and tidal
772 level on survival and growth. Helgol. Mar. Res. 64:93-100.
773

774 Goudswaard, P.C., Perdon, K.J., Jol, J., vanAsch, M., Troost, K., 2013. Het bestand aan
775 commercieel belangrijke schelpdiersoorten in de Nederlandse kustwateren in 2013. Imares
776 rapport c133/13, 38pp.
777

778 Greene, K., 2002. Beach Nourishment: A Review of the Biological and Physical Impacts.
779 ASMFC Habitat Management Series # 7. Atlantic States Marine Fisheries Commission 1444
780 Eye Street NW, Sixth Floor Washington DC 20005: 174pp.
781

782 Gremare, A., Amouroux, J.M., Chaabeni, Y., Charles, F., 1998. Experimental study of the
783 effect of kaolinite on the ingestion and the absorption of monospecific suspensions of *pavlova*
784 *lutheri* by the filter-feeding bivalve *Venus verrucosa*. *Vie Milieu* 48(4): 295-307.
785

786 Hilbish, T.J., 1986. Growth trajectories of shell and soft tissue in bivalves: seasonal variation
787 in *Mytilus edulis*. *J. Exp. Mar. Biol. Ecol.* 96:103-113.
788

789 Hummel, de Wolf, H. L., Fortuin, A.W., 1988. The annual cycle of glycogen in estuarine
790 benthic animals. *Hydrobiological bulletin* 22:199-202.
791

792 Kamermans, P., Brummelhuis, E., Dedert, M., 2013. Effect of algae and silt concentration on
793 clearance- and growth rate of the razor clam *Ensis directus*, *Conrad. J. Exp Biol. Ecol.* 446:
794 102-109.
795

796 Kenchington, E., Duggan, R., Riddell, T., 1998. Early life history characteristics of the razor
797 clam (*Ensis directus*) and the moonsnails (*Euspira spp*) with application to fisheries and
798 aquaculture." *Can. Tech. Rep. Fish. Aquat. Sci* 2223: 32pp.
799

800 Keppler, D., Decker, K., 1970. Glykogen. Bestimmung mit amylogucosidase. *In*; Bergmeyer,
801 H.U., methoden der enzymatischen analyses, Verlag chemie Weinheim.- Berlin.
802

803 Keppler, D., Decker, K., 1974. Glycogen. Determination with Amylogulcosidase. *In*;
804 Bergmeyer H. U., Methods of enzymatic analysis., Verlag chemie Wennheim- Berlin.
805

806 Kooijman, S.A.L.M., 2010. Dynamic Energy Budget theory for metabolic organisation.
807 Cambridge University Press, Great Britain.
808

809 Lewis, D.E., Cerrato, R.M., 1997. Growth uncoupling and the relationship etween shell
810 growth and metabolism in the soft shell clam *Mya arenaria*. *Mar. Ecol. Prog. Ser.* 158:177-
811 189.
812

813 Mühlenhardt-Siegel, U., Dörjes, J., von Cosel, R., 1983. Die amerikanische Schwertmuschel
814 *Ensis directus* (Conrad) in der Deutschen Bucht. II Populationsdynamik. *Senck. Marit.* 15(4/6):
815 93-110.
816

817 Norkko, A., Hewitt, J.E., Trush, S.F., Funnell, G.A., 2001. Benthic-pelagic coupling and
818 suspension feeding bivalves: Linking site-specific sediment flux and biodeposition to benthic
819 community structure. *Limnol. Oceanogr.* 46(8): 2067-2072.
820

821 Oksanen, J., Blanchet, F.G., Kindt, R., Legendre, P., Minchin, P.R. , O'Hara, R.B., Simpson,
822 G.L., Solymos, P., Henry, M., Stevens, H., Wagner, H., 2013. vegan: Community Ecology
823 Package. R package version 2.0-9. <http://CRAN.R-project.org/package=vegan>
824

825 Palmer, D.W., 2004. Growth of the razor clam *Ensis directus*, an alien species in the Wash on
826 the east coast of England. J. Mar. Biol. Ass. U.K. 84: 1075-1076.
827

828 Philippart, C.J.M., vanBleijswijk, J.D.L., Kromkamp, J.C., Zuur, A.F., Herman, P.M.J., 2014.
829 Reproductive phenology of coastal marine bivalves in a seasonal environment. J. Plankton
830 Res. 0(0):1-16. doi:10.1093/plankt/fbu073.
831

832 Pulfrich, A., 1997. Seasonal variation in the occurrence of planktic bivalve larvae in the
833 Scheswig-Holstein Wadden Sea. Helgoländer Meeresun 51: 23-39.
834

835 R Core Team, 2012. R: A language and environment for statistical computing. R Foundation
836 for Statistical Computing, Vienna, Austria. ISBN 3-900051-07-0, URL [http://www.R-](http://www.R-project.org/)
837 [project.org/](http://www.R-project.org/).
838

839 Roper, D.S., Kickey, C.W., 1995. Effects of food and silt on filtration, respiration and condition
840 of the freshwater mussel *Hyridella menziesi* (Unionacea: Hyriidae): implications for
841 bioaccumulation. Hydrobiologia, 312: 17-25.
842

843 Rozemeijer M.J.C., de Kok, J., de Ronde J.G., Kabuta S., Marx S., van Berkel G., 2013. Het
844 Monitoring en Evaluatie Programma Zandwinning RWS LaMER 2007 en 2008-2012:
845 overzicht, resultaten en evaluatie. IMARES Wageningen UR Rapport C181/13, Deltares
846 Rapport 1207903-000-ZKS-004.
847

848 Schellekens, T., Witbaard, R., 2012. DEB ensis vs. data. Imares report C155/12. 30pp
849

850 Schellekens, T., 2012. Groei en conditie van zwaardschede (*Ensis directus*, Conrad) tijdens
851 en na geplande zandwinning in 2013-2017 berekeningen voor het RWS. IMARES rapport
852 C088/12. 33pp
853

854 Severijns, N., 2002. Verspreiding van de Amerikaanse zwaardschede *Ensis directus* (Conrad,
855 1843) in Europa 23 jaar na de introductie: opmerkelijke opmars van een immigrant. Gloria
856 Maris 40(4-5): 63-111,
857

858 Shanks, A.L., Brink, L., 2005. Upwelling, downwelling and cross-shelf transport of bivalve
859 larvae: test of a hypothesis. Mar. Ecol. Prog. Ser. 302: 1-12.
860

861 Speybroeck, J., Bonte, D., Courtens, W., Gheskiere, T., Grootaert, P., Maelfait, J.P., Mathys,
862 M., Provoost, S., Sabbe, K., Stienen, E.W.M., van Lancker, V., Vincx, M., deGraer, S.,
863 2006. Beach Nourishments: an ecologically sound coastal defence alternative? A review.
864 *Aquat conserv. Mar. Freshw. Ecosyst* 16: 419-435.

865

866 Strasser, M., Gunther, C.P., 2001. Larval supply of predator and prey: temporal mismatch
867 between crabs and bivalves after a severe winter in the Wadden Sea. *J. Sea Res.* 46: 57-67.

868

869 Szostek, C.L., Davies, A.J., Hinz, H., 2013. Effects of elevated levels of suspended particulate
870 matter and burial on juvenile king scallops *Pecten maximus*. *Mar. Ecol. Prog. Ser.* 474: 155-
871 165.

872

873 Trueman, E.R. 1967. The dynamics of burrowing in *Ensis* (Bivalvia). *Proc. R. Soc. Lond. B*
874 166: 459-476.

875

876 Tulp, I., Craeymeersch, J., Leopold, M.F., Damme, van, C., Fey, F., Verdaat, H., 2010. The
877 role of the invasive bivalve *Ensis directus* as food source for fish and birds in the Dutch
878 coastal zone. *Est. Coast. Shelf Sci.* 90: 116-128.

879

880 Van Duin C.F., Gotjé, W., Jaspers C.J., Kreft M., 2007. MER Winning suppletiezand
881 Noordzee 2008 t/m 2012. Grontmij Hoofdrapport, Definitief. 13/99080995/CD, revisie D1 242
882 pp, Grontmij, Houten.

883

884 Van der Hout, C.M., Gerkema, T., Nauw, J.J. Ridderinkhof, H., 2014. Observations of a
885 narrow zone of high suspended particulate matter (SPM) concentrations along the Dutch
886 coast. *Cont. Shelf Res.*, Submitted.

887

888 van Rijn, L.C., 1995. Dynamics of the closed coastal system of Holland. Delft hydraulics
889 report H2129. Delft hydraulics, Delft.

890

891 Vaughn, C.C. Hakenkamp, C.C., 2001. The functional role of burrowing bivalves in freshwater
892 ecosystems. *Freshwater Biol.* 46: 1431-1446.

893

894 Verduin E.C., Tempelman, D., Moorsel G.W.N.M. van, 2011. The Macro-benthic Fauna
895 Monitoring in the Dutch Sector of the North Sea, MWTL 2009 and a comparison with previous
896 data. Grontmij, report 290843. Amsterdam. 109 pp.

897

898 Verduin E.C., Tempelman, D., Moorsel G.W.N.M. van, 2012. The Macro-benthic Fauna
899 Monitoring in the Dutch Sector of the North Sea, MWTL 2010 and a comparison with previous
900 data. Grontmij and Ecosub, report 290843. Amsterdam. 143 pp.

901

902 Von Cosel, R., Dörjes, J., Mühlenhardt-Siegel, U., 1982. Die amerikanische Schwertmuschel
903 *Ensis directus* (Conrad) in der Deutschen Bucht. I. Zoogeographie und taxonomie im
904 vergleich mit den einheimischen Schwertmuschel-Arten. . Senck. Marit. 14: 147-173.

905

906 Vries, S. de, Koomans, R.L., 2010. Monitoring van slibgehalte en bodemligging voor de kust
907 van Petten/Egmond, Eind evaluatie. Medusa Explorations Groningen, 35pp.

908

909 Witbaard, R, G.C.A. Duineveld, M.J.N. Bergman, 2013. The final report on the growth and
910 dynamics of *Ensis directus* in the near coastal zone off Egmond, in relation of environmental
911 conditions in 2011-2012. NIOZ report 2013-2 78pp.

912

913 Wijsman, J.W.M., 2011. Dynamic Energy Budget. (DEB) parameters for *Ensis directus*.
914 IMARES Report C116/11; 39pp.

915

916 Wood, S.N. (2006) Generalized Additive Models: An Introduction with R. Chapman and
917 Hall/CRC.

918

919 Young, L. J. and J. Young (1998). Ecological Statistics, a population perspective., Kluwer
920 Academic publishers.