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Title: Size-selective fishing of *Palaemon serratus* (Decapoda, Palaemonidae) in Wales, UK: implications of sexual dimorphism and reproductive biology for fisheries management and conservation.

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Abstract

The common prawn (*Palaemon serratus*) supports a small-scale but economically important seasonal static-gear fishery in Cardigan Bay, Wales (UK). Due to a lack of statutory obligation and scientific evidence, the fishery has operated to date without any harvest-control-rules that afford protection from overfishing. In response to fluctuations in landings and in pursuit of increased economic returns for their catch, some members of the fishing industry have adopted a size-selective harvesting regime, which we evaluate here using baseline data. Monthly samples were obtained from fishers operating out of five ports between October 2013 and May 2015 (n = 4,233). All prawn were sexed, weighed and measured, whilst the fecundity of females was estimated for 273 (44%) individuals. Peak spawning occurred during the spring and females were estimated to undergo a 'puberty moult' at a carapace length (CL) of 7.7 mm, whilst functional maturity was estimated at a CL of 9.9 mm. The sampled population exhibited sexual dimorphism, with females attaining a greater size than males. The current harvesting regime results in a sex bias in landings as even large mature males remained under the recruitment size to the fishery, unlike the large mature females. The temporal trend in sex-ratio indicates a continual decrease in the catchability of female prawn through the fishing season; however, whether this is caused by depletion via fishing mortality or migratory behaviour is yet to be resolved. Here, we provide a comprehensive baseline evaluation of population biology and discuss the implications of our findings for fisheries management.
INTRODUCTION

The fishery for the common prawn, *Palaemon serratus* (Pennant 1777; Neal 2008), is relatively small compared to other European “prawn” fisheries (*Nephrops norvegicus* and *Pandalus borealis*); however, in the UK it has significant regional economic importance. In Cardigan Bay (Wales), the fishery accounts for ~76% of total UK landings (estimate from 2013; MMO, 2015). Commercial exploitation of prawns in Cardigan Bay is exclusively an inshore static-gear pot fishery, with most vessels working within six nautical miles of the coast. The fishery begins to target prawn in early autumn and continues through to the following spring (Cardigan Bay Fishermen’s Association (CBFA) pers comm.). The fishing season is dictated by the reproductive migrations of *P. serratus*, which are thought to move inshore to release larvae during the summer and then move offshore in winter. Similar seasonal migrations are reported in a range of palaemonid species, including for *P. serratus* in other regions (Guerao & Ribera, 2000; González-Ortega *et al.*, 2006).

The seasonal migration of prawns inshore in the summer decreases static gear catches to levels that are no longer economically viable (CBFA pers comm.). Nonetheless, the Cardigan Bay prawn resource is integral in maintaining the economic viability of many fishing businesses as it provides income during a time of the year when the catchability of other target species, such as European lobster (*Homarus gammarus*), is low. The fishery is therefore an important element in a necessarily diverse static-gear sector.

Commercial demand for a larger-sized prawn has resulted in the introduction of voluntary size-grading of catch by fishers. Since 2008, many Cardigan Bay fishers have used a 10 mm bar-spacing riddle (CBFA pers comm.). Prawns that fall through the bars and into the sorting box are discarded overboard, whilst prawns retained by the riddle are stored onboard, usually within small viver systems.

As with many small-scale artisanal fisheries, the Welsh prawn fishery is considered data-poor, with little information pertaining to the fisheries biology of the species. Combined with limited management and the lack of a formal stock-assessment, there is considerable uncertainty about the future sustainability of the fishery. Indeed, fluctuations in inter-annual landings in the Irish fishery (Fahy & Gleeson, 1996; Kelly *et al.*, 2009) suggest a variable biomass that may be vulnerable to periodic overfishing or recruitment failure in the absence of management. Understanding the interaction of fishing activities with the species biology is necessary to inform future evidence-based management of the fishery and more generally, understanding the reproductive biology of a fished species is critical information when considering “supply-side” ecology of benthic populations with economic value (Underwood & Fairweather, 1989; Anger, 2006).
The common prawn is patchily distributed throughout European inshore waters (Kelly et al., 2009) and occurs between the Mediterranean Sea in the south and the temperate coastal waters of the United Kingdom and Ireland in the north (Forster, 1951). Although the longevity of the species has been speculated to be up to five years (Cole, 1958; Forster, 1959), *P. serratus* are more likely to have a relatively short life span, with individuals persisting for between two to three years (Forster, 1951; Fahy & Gleeson, 1996). Similar to other palaemonids, *P. serratus* is sexually dimorphic, with adult females attaining significantly larger sizes (Forster, 1951; Berglund, 1981). Sexual dimorphism may influence mortality rates between the sexes, from both size-selective commercial exploitation and natural mortality through predation (Berglund & Rosenqvist, 1986). For female palaemonids, a larger body size also allows for increased fecundity (Guerao et al., 1994).

Compared to other similar species, *P. serratus* broods contain larger eggs with high nutritional values (Morais et al., 2002), which are thought to reflect environmental conditions and increase successful recruitment through the larval phase (Parker & Begon, 1986). The planktonic larval phase is characterised by temperature dependent periods of incremental growth and metamorphosis (Reeve, 1969a; Kelly et al., 2012), while salinity has been shown to influence mortality rates during the early life stages (Kelly et al., 2012).

The aim of this research was to fill the knowledge gaps for this data-poor fishery by presenting baseline catch and population biology characteristics (length frequency, sex ratio, size at maturity) during the adult stage of the species life-history and to highlight several potential implications of a mandatory technical conservation measure of riddling catches at 10 mm.

**MATERIALS AND METHODS**

In August 2013, six commercial fishermen operating from five different ports in Cardigan Bay, Wales, were each given three standard prawn traps (referred to hereafter as “science pots”). The cylindrical pots were fitted with 8 mm mesh on all sides with 35 mm circular entrance at both ends. Once a month, when possible, each fisher recorded the date and GPS location of a haul and the entire contents of each science pot were kept separate and stored frozen. Samples were retained for scientific analysis during two fishing seasons (2013-2014 and 2014-2015), ending in May 2015.

Scientific pot samples were assessed in the laboratory using a dissecting microscope. All animals caught in the science pots were identified, weighed and measured. Palaemon species were identified according to the
illustrated key published by González-Ortegón & Cuesta (2006). Sex was recorded; male prawns were identified by the presence of an appendix masculina on the second pleopod pair. All morphometric measurements were recorded to the nearest 0.1 mm and included the carapace length (CL; the distance between the posterior of the eye-orbit to the posterior of the cephalothorax carapace segment), carapace width (CW; the widest point of the cephalothorax carapace) and pleura width (PLW; the widest section of the second abdominal pleura). The reproductive state (ovigerous or not) was also noted for female prawns and the fecundity of ovigerous females were calculated from a subsample of 10% of the entire egg mass (wet weight). The fecundity was estimated using the following formula (1); where \( \Upsilon \) indicates the subsample calculated as a proportion of the total eggs mass (T), which was then used to calculate fecundity (F).

\[
Y_{\text{approx}=0.1} = \frac{\text{Weight}_{\text{subsample}}}{T_{\text{weight}}}
\]

\[
F = \frac{\text{Count}_{\text{subsample}}}{\Upsilon}
\]

(1)

All statistical analyses were run in “R” (R Core Team, 2014). Prior to statistical modelling data were tested for normality using the Kolmogorov-Smirnov test and inspected visually using a Q-Q plot. Heteroskedasticity was tested using Levene’s test and a Cook’s distance plot was used to check for outliers. A Hartigan’s dip test was used on length distribution data for non-unimodality. The likelihood of the sample having a 1:1 sex-ratio was tested using a G-test. Since we were not able determine size-at-age for the sampled population, age cohorts are inferred from the observed length distribution. A mixed population approach was used to determine statistical differences between sexes and cohorts within a mixed bi-modal dataset. Using the R packages “MIXTOOLS v1.0.3” (Young et al., 2015) and “MIXDIST v0.5-4” (Macdonald & Du, 2011), the mean and standard deviation of the two modes in aggregated male and female length distribution data is presented alongside a goodness-of-fit Chi-square test. We use the results to evaluate the length distributions of immature and mature populations as well as sexual dimorphism within a single mixed-population cohort.

The size of functional maturity was estimated by relating growth parameters (CL) and ovigerous status (binary variable, where 0 = no eggs and 1 = gravid) of females using a logistic regression model (Roa et al., 1999) reformulated by Walker (2005) to give:
where $P_i$ is the proportion of the female population gravid at a given CL. Model parameters were estimated using generalized linear model with logit link function and a binomial error structure. Confidence intervals were added by bootstrapping the generalized linear model (1000 runs). The base R code was constructed by Harry (2013) and is available online.

To describe morphometric maturity and determine at what CL positive allometry occurs, an iterative search procedure was used whereby PLW is modelled against CL for male and female populations separately using piecewise linear regression. The analysis examines the linear morphological relationship (CL:PLW) and searches for significant deviations between male and female growth patterns, indicating sex-specific morphological changes in preparation for sexual reproduction described as a “puberty moult” (Hartnoll, 2001). The method searches each potential “breakpoint” or “inflection” ($c$) within a predetermined range until the model has found the point at which the total residual mean standard error is minimised (Crawley, 2007). The model simulation then produces a value (CL) at which the linear models above and below the breakpoint $c$ show the statistically strongest inflection. The model applied to both male and female datasets is described mathematically using the equation 3.

$$y_i = \begin{cases} 
\beta_0 + \beta_1 CW_i, & CW_i < c \\
\beta_2 + \beta_3 CW_i, & CW_i \geq c 
\end{cases}$$

(3)

where $y_i$ is the CL of individual $i$, $c$ is a breakpoint (inflection) between linear relationships applying above and below the value of carapace length equal to $c$, and the $\beta$ parameters are the intercepts and slopes of the two linear relationships.

In order to relate the morphological estimate of population characteristics, fisheries catches (CL) results are converted to CW using the following equations (4) produced by linear regression ($p < 0.05$):

$$CW_{\text{Male}} = 0.563 CL_{\text{Male}} + 0.643$$

$$CW_{\text{Female}} = 0.6389 CL_{\text{Female}} - 0.297$$

(4)

Individuals with a CW < 10 mm are assumed to be discarded through the use of a 10 mm spaced riddle.

RESULTS
Severe weather conditions during the 2013 and 2014 fishing seasons limited the fishing opportunities and the number of individual prawns that could be sampled within that season (n = 765). In total, fishers returned 82 pot-samples and 4,233 *P. serratus* underwent laboratory analysis (table 1).

Sexual dimorphism and riddling

Sexual dimorphism was evident in the length distributions of all samples. Moreover, prawn populations showed bimodal distributions when data was aggregated by fishing season and location (Hartigan’s dip-test; $D_{\text{Male}} = 0.95$, $D_{\text{Female}} = 0.04$, p-value < 0.001). The majority of male prawns and the smaller sized cohort of female prawns caught in the small mesh science pots were of a size that would be discarded using the 10 mm riddle employed by Cardigan Bay fishers (Fig 2).

Carapace width varied significantly between sexes and two cohorts were identified using a mixed population cohort analysis (1+ and 2+; summary statistics and ANOVA results in Table 2a). Table 2b compares dimorphism highlighted by Forster (1951) and the present study. A higher proportion of the males (78.3 %) caught were smaller than 10 mm CW compared to the females (39.7 %) in catches.

The maximum size observed in the sampled population showed females grew to a size considerably greater than males, whilst the length distribution of catches show that the average male prawn within the 2+ cohort does not reach a size at which it recruits into the Cardigan Bay prawn fishery.

Sex ratio

The sex-ratio of catches varied significantly from the expected 1:1 ratio, with both male and female directed skews being observed throughout the sample period (Fig 3a)
For all locations sex-ratios were female skewed in autumn and winter samples, with a higher proportion of males caught in spring. Where an extended time-series was available from a single location, data exhibited strong temporal trends in the sex-ratio and declining abundance of females as the fishing season progressed in New Quay (3b); however, the data trend was less clear in the samples from Aberystwyth (Fig 3c).

**Size at maturity (SOM)**

Using an iterative search procedure, an inflection point was detected in the linear relationship between CL and PLW in the female dataset. The data suggests that for pleura morphometrics, males display an isometric growth pattern and females an allometric growth pattern. For females, the CL:PLW inflection point was detected at 12.5 mm CL (Fig 4).

Size at maturity (L50)

Maturity is expressed as L50, which is the size (CL) at which 50% of the females were observed to be gravid (carrying eggs). The maximum likelihood estimate of L50 estimated by the generalised linear model with a binomial distribution was 15.9 mm CL (upper and lower confidence intervals = 16.4 mm and 15.4 mm CL respectively; Fig 5).

Fecundity

Of the 616 gravid prawn that were captured by scientific pots, 273 (44%) were analysed for fecundity using the equation described (1). Prawn ranged in size from 14.2 mm to 25 mm (CL) and produced fecundity estimates of between 221 and 5,121 eggs per animal.

A Spearman’s correlation was run to assess the relationship between CL and fecundity. There was a strong positive correlation, which was statistically significant ($r_s = 0.48$, $p <0.001$) and is explained by the power relationship below (figure6; equation 5). The fecundity data exhibits a high degree of variability with CL.
explaining just 22.2% of the variation in fecundity. Data points shown as triangles represent available fecundity data from Forster (1951).

Fecundity $= 92.546 e^{0.1465 \cdot CL}$

DISCUSSION

Sexual dimorphism

Our results confirm that *P. serratus* in Cardigan Bay are sexually dimorphic, with females occupying a broader length-distribution than males in the sampled populations. These results mirror the sexual dimorphism that has been reported elsewhere for *Palaemon serratus* (Guerao & Ribera, 2000) and many other *Palaemon* spp, with typically slower growth rates and smaller sizes in males. (e.g. Berglund, 1981; Ito et al., 1991; Bilgin et al., 2009; Al Maslamani et al., 2013).

The evolutionary cause for dimorphism in this species is likely to have resulted from selection based on the differing reproductive roles of the sexes (Shine, 1989). The current size-selective exploitation and resulting pressure on mature females could potentially result in evolutionary responses that change growth and reproductive patterns at a genetic level (e.g Conover & Munch, 2002; Walsh et al., 2006; Swain et al., 2007).

Given the short life span of *P. serratus*, fishery-induced responses such as decreasing size-at-maturity and size-at age may occur over a timescale of years or decades (Reznick et al., 1997; Thompson, 1998; Koskinen et al., 2002; Stockwell et al., 2003), a phenomenon that has been demonstrated in a number of other exploited populations (e.g Grift et al., 2003; Olsen et al., 2004; Barot et al., 2004). Indeed, the selection pressure towards large females and potential decrease in growth rates may have a negative effect on the value of the species in the long term, which runs contrary to the larger prawns desired by the market. Hence it is important to continue monitoring these life history characteristics, in order to determine any long term changes, particularly in females. It would also be valuable to compare populations with varying degrees of commercial exploitation.

Length frequency and commercial fishing

At present, the fishery is not subject to any statutory harvest-control-rules or technical-measures that aim to encourage sustainable exploitation of prawn populations in Welsh waters in addition to the requirement for
commercial fishers to hold a shellfish license. The voluntary riddling of catch using either a 10 mm riddle or > 10 mm pot mesh by some fishers ensures both a better market price and may return as many as 40% of females to sea including sexually immature individuals. Since the grading of prawns is an entirely voluntary practice, it is not possible to determine the relative proportion of common prawn landings in the UK that have been graded at sea, on the quayside or not at all. Whilst the mortality rate amongst prawns discarded at sea is still to be determined, personal observations indicate a very high level of mortality when prawns are graded on the quayside. The absence of information on discard mortality rates calls into question the real value of the ridding practice, particularly since the mortality rate is likely to be high. If the rate of mortality amongst discarded prawn is at a significant level, the sex-specific consequences of ridding may not be as severe as the data suggests. Nonetheless, there is a need to ensure that riddling is done at sea over fishing grounds and habitat from which the prawn were removed. Some fishermen argue that a larger mesh size on the fishing gear is a more appropriate conservation measure. We suggest that a gear comparison trial be conducted to determine the gear design that maintains catchability whilst promoting the escape of undersized prawn. Importantly the interaction between ridding and size-selectivity (i.e. that a riddle will retain prawn only of a size ≥ 10 mm) is an assumption in this study and not empirically validated. Future research needs to collect data on retention rates of a known size distribution of animals being graded in order to evaluate the real the real size and sex specific implications of the technical measure.

In 2008, when voluntary measures were adopted by some Cardigan Bay fishers, it was hoped that the discarding of small prawn at sea would provide additional ecological and economic value by improving market prices and releasing immature prawns to improve yield-per-recruit and spawner-per-recruit respectively (CBFA pers. comm.). Our results show that by applying a size-selective harvesting regime, the Cardigan Bay prawn fishery subjects the female population to a much higher level of removal relative to the male population. Indeed, the immediate consequence of the quasi-minimum-landing-size would have been the discarding of approximately 78% of male prawns caught in pots, compared to a female discard rate of approximately 40% on average throughout the fishing season. The bi-modal distribution of size-frequency data was present in all spatial and temporal combinations, representing a strong indication that two cohorts of prawns are present during fishing the season. With the assumption that commercial activities select prawn at a size 10 mm CW under the voluntary MLS, data shows females are recruiting into the fishery in their
second year at a mean size of 12.32 mm CW (SE ± 1.22). However, fewer 2+ males are recruited into the fishery, as the average male in their second year is 9.64 mm CW (SE ± 0.84).

Our study shows similar patterns in length-cohort distribution to previous studies. Forster (1951) reported female population attaining a greater modal size than males within the 2+ cohort (TL_{Mo} ≈ 92.5 mm; TL_{Mo} ≈ 77.5 mm). The above values are comparable to those reported in this study; however, the historical data indicates a smaller average size of prawn within the 1+ group than we observed in this study (see table 2b), although the difference is unlikely to be significant. The difference in 1+ size is likely to be as a consequence of differing sampling methods employed by the two studies; Forster (1951) using fishery independent trawl surveys in contrast to the present study, which used fishery-dependent ‘science pots’, which were fished alongside commercial gear and therefore targeted the larger prawns.

**Sex bias in the fishery**

A consequence to size-selective fishing and higher rates of removal of female prawn may be evident in the temporal trend of sex-ratios (Figure 3a), representing sex-overfishing on a regional scale. However, Fig 3b and 3c shows that decreasing catchability of female prawn is location specific, with samples from Aberystwyth showing a near 1:1 sex ratio late into the fishing season in comparison to fishing grounds to the south, although the proportion of females in spring is still lower than during winter. The decreasing abundance of females in catches marks the end of the prawn season as it is perceived by fishers as a weakening fishery that yields less marketable catch. Seasonal variation in sex-ratios have been observed in a range of palaemonid species (see Kim, 2005; Al-Maslamani et al., 2013) and has been attributed to differential migration patterns, seasonal habitat preferences and possibly mortality between males and females (Berglund, 1981). Female *P. serratus* are known to migrate between habitats to release larvae in Wales (Haig et al. *unpublished data*) and hence it is unsurprising that we observes temporal and spatial changes in sex-ratio as the fishing season progressed in Cardigan Bay as this may reflect localised differences in timing of migration or habitat availability.

On a regional scale, fishing behaviour follows an inter-annual pattern whereby fishers in the south experience the onset of the fishing season, with fishing opportunities gradually opening in a northward direction along the Cardigan Bay coast (*pers. obs.* and CBFA *pers comms*). Similarly, fishing opportunities decline earlier in the south relative to the north, with fishermen from Aberystwyth and Aberdovey continuing
to fish for months after fishing has ceased to be commercially viable in Fishguard and New Quay (pers. obs. and CBFA pers. comm.). Fishermen therefore hold the view that females migrate in a northerly direction, sustaining different rates of catch in different areas through the season. The scientific evidence presented here neither validates nor disproves this view on the migratory behaviour of prawn in the region. Further fisheries independent research (ideally using mark recapture methods) is required to determine if the observed patterns in female catch indicate sex-overfishing, decreasing catchability as a result of seasonal migration by females, or a cumulative response to both of these.

The potential for sex-overfishing identified by this study may have consequences on recruitment levels in the future, although the life-history of palaemonids (highly fecund and typically multiple broods per season) may safeguard it against depletion events. The data show female skewed catches in the early period of both fishing seasons (Emmerson et al., 2014), which indicates the population has a degree of resilience in sustaining size-selective fishing at present effort levels, the research presented here cannot draw a conclusion with regards to sex-overfishing in the absence of both long-term datasets and evidence pertaining to adult migration patterns.

Size at maturity

Crustacean fisheries are most commonly managed in the UK using a minimum landing size (MLS), appropriated by maturity characteristics. In order to determine a valid MLS in decapod crustaceans, maturity indicators such as morphological sexual maturity and functional maturity can be applied (Waddy and Aiken, 2005; Pardo et al., 2009). Size at maturity has been determined from allometric growth parameters (e.g. Hartnoll, 1974; Little & Watson III, 2005; Claverie & Smith, 2009) and specifically the CL:PLW relationship in Palaemonidae species (Cartaxana, 2003). In this study, the pleura has been shown to undergo allometric growth in female P. serratus, which expand the brood chamber in preparation for egg carriage at a size CW = 7.7 mm. At this point, females undergo an expansion in the PLW relative to males as they continue to grow. It is highly likely that this dynamic allometry amongst females represents a physical change of the abdomen in preparation for egg bearing and thus a sign of sexual maturity. Only 1.5% (n = 9) of ovigerous females were observed at a size below our estimate of morphological size at maturity, implying a high degree of confidence in the results of the iterative search procedure used. A total of 18.6% of females (n = 361) captured by scientific pots throughout this study had a CW < 7.7 mm and were assumed to be
sexually immature. With a CW < 10 mm, immature female prawn that have yet to develop their brood chamber and are released by Cardigan Bay fishers onto the fishing ground from where they were captured.

The size at morphological sexual maturity supports the results from this study’s estimate of functional maturity (L<sub>50</sub>), with the results implying that female prawn undergo a puberty moult at an estimated size CW = 7.7 mm, whilst 50% of females are able to contribute to the reproductive capacity of the population by the size CW = 9.9 mm (15.9 mm CL). In this way, the voluntary measure of releasing prawn below CW = 10 mm by CBFA fishers has been shown to be a potentially valuable conservation measure. The CL<sub>50</sub> reported here is greater than that reported in similar studies elsewhere for the species (Ireland; CL<sub>50</sub> = 12.5mm; Kelly, 2009), though similar to previous estimates for the Welsh population (CL<sub>50</sub> = 16.5; Huxley, 2011).

**Fecundity**

*P. serratus* were found to carry between 221 and 5,121 eggs at any one time (mean average = 1,916). This is similar to estimates published by Forster (1951), who found large prawn (TL = 105 mm) carry up to 4,282 eggs and within a similar range of other Palaemonidae species (Corey & Reid, 1991). The fecundity (number of eggs carried) of female prawns was positively correlated with body size (CL); however, there was a high degree of variability between individuals and CL only explained ~23% of the variation. Studies of similar species (*P. elegans*, *P. adspersus* and *P. xiphias*) report R<sup>2</sup> values > 0.95 (Guerao et al., 1994; Cartaxa, 2003; Bilgin & Samsun, 2006). Different methodologies for estimating fecundity may be the reason behind the variable R<sup>2</sup> values reported here and in the published literature. In particular, previous fecundity estimates were derived from the number of eggs at stage 1 (e.g Guerao et al., 1994) in order to account for egg loss during incubation, which can be the result of mechanical stress or parasites (Glamuzina et al., 2014) and has been reported to be as high as 38% in this species (Reeve, 1969 in Zimmermann et al., 2015). Egg counts by developmental stage were unavailable in this study, which is the likely explanation for the high variability in the fecundity estimate. Nonetheless, the results are within the range reported for the species, as shown in figure 6, and provide an important baseline from which to further understand the reproductive capacity of the Welsh *P. serratus* population by providing an estimate between the numbers of eggs laid on pleopods during spawning and the total that eventually hatched.

Fecundity can be influenced by temporal-spatial variations of environmental factors such as depth (e.g *P. naval*; Thessalou-Legakiand, 1992), mean bottom temperature (e.g *P. borealis*; Parsons & Tucker, 1986) and
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the extent of which is yet to be fully understood, affording a scientifically-validated level of protection to juvenile females via a MLS would be a valuable safeguard against recruitment failures in the future.

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FIGURE LEGENDS

**Fig 1** The homeports for the six active *Palaemon serratus* fishers in Cardigan Bay, Wales; who contributed monthly samples (when possible) during the prawn fishing seasons from 2013, 2014 and 2015. Ports are numbered north to south and are as follows: 1, Aberdovey (2 fishers); 2, Aberystwyth; 3, New Quay; 4, Cardigan; 5, Fishguard

**Fig 2** A length frequency histogram with a probability density function for male (above) and female (below) *Palaemon serratus* caught in science pots during the 2013-2015 *Palaemon serratus* research period in Cardigan Bay. The solid vertical red line represents the voluntary sorting size (10 mm CW) used by many fishers in Cardigan Bay
**Fig 3a** The sex-ratio of prawn (*Palaemon serratus*) caught in science pots from during the 2014 / 2015 in Cardigan Bay and **3b** the sex ratio of catches in localised datasets from New Quay and **3c** Aberystwyth.

**Fig 4** Inflection point indicating allometric growth based on morphometric variance between iterative tests on linear models of PLW and CL for the prawn *Palaemon serratus*. The dotted vertical line is the value with the lowest mean standard error (12.5 mm CL). Solid black line shows the linear male relationship. Hashed line shows the allometric female relationship after inflection event.
Fig 5 Functional maturity model fit for female prawn (*Palaemon serratus*) from Cardigan Bay (Wales) with 95% CIs as indicated by the presence or absence of eggs. The horizontal line represents L_{50} (15.9 mm CL) for the females sampled within period of peak spawning (April; n = 544).

Fig 6 Fecundity of gravid prawn (*Palaemon serratus*) from Cardigan Bay (Wales) with size (CL) (n = 273). The solid line shows the power relationship between the correlating variables CL and Fecundity (p < 0.001; R^2 = 0.222). The red triangle points and associated hashed trendline show the fecundity data available from Forster (1951).