Optimizing hatchery practices for genetic improvement of marine bivalves

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Abstract
Aquaculture currently accounts for approximately half of all seafood produced and is the fastest growing farmed food sector globally. Marine bivalve aquaculture, the farming of oysters, mussels and clams, represents a highly sustainable component of this industry and has major potential for global expansion via increased efficiency, and numbers of, production systems. Artificial spat propagation (i.e. settled juveniles) in hatcheries and selective breeding have the potential to offer rapid and widespread gains for molluscan aquaculture industry. However, bivalves have unique life-histories, genetic and genomic characteristics, which present significant challenges to achieving such genetic improvement. Selection pressures experienced by bivalve larvae and spat in the wild contribute to drive population structure and animal fitness. Similarly, domestication selection is likely to act on hatchery-produced spat, the full implications of which have not been fully explored. In this review, we outline the key features of these taxa and production practices applied in bivalve aquaculture, which have the potential to affect the genetic and phenotypic variability of hatchery-propagated stock. Alongside, we compare artificial and natural processes experienced by bivalves to investigate the possible consequences of hatchery propagation on stock production. In addition, we identify key areas of investigation that need to be prioritized to continue to the advancement of bivalve genetic improvement via selective breeding.

Key words: gene-environment interactions, genomic selection, marine bivalve aquaculture, selective breeding, sustainable development.

The future of bivalve aquaculture relies on artificial propagation

With the global human population projected to exceed 9 billion by 2050, food production must increase by at least 59% to meet projected demand (Valin et al. 2014). Feeding this growing population, whilst maintaining biodiversity and good environmental stewardship, is one of the major global challenges of the 21st century. This issue is exacerbated further by the need to ensure that the future intensification of food production is sustainable, especially in the face of climate change (UN 2015; IPCC 2018).

Aquaculture is the fastest growing food production sector globally, expanding on average 6.4% per annum since 2001 (Subasinghe 2017). Nearly half of the current global finfish and shellfish production derives from aquaculture (FAO 2019a), with this sector expected to underpin most future growth in seafood production (Kobayashi et al. 2015). Currently, mollusc farming accounts for approximately 21% of world aquaculture production (Subasinghe 2017).
Important scientific advances in bivalve husbandry practices (i.e. optimization of diet, fertilization protocols and larval rearing) occurring within the last century (e.g. Galstoff 1938; Carriker 1956; Loosanoff & Davis 1963) led to the establishment of the first commercial bivalve mollusc hatcheries (Mann 1983) resulting in the global expansion of shellfish aquaculture. The ability to control environmental conditions in indoor facilities enables broodstock conditioning and spat production almost year-round. Most importantly, the development of a constant and reliable source of spat benefits the expansion of the bivalve aquaculture sector, facilitating the predictability of production and enabling the possibility of selective breeding.

Hatcheries are expected to play a key role in the continued expansion of bivalve aquaculture. The potential of hatchery production is highlighted in China, which accounts for 80% of global production of Pacific oysters (Crassostrea gigas) (Yang et al. 2014), and the sector now relies almost entirely on hatchery sourced spat (Li et al. 2011). Nonetheless, demand for hatchery-produced spat is often low in areas where natural (wild capture) spat is available and abundant. A similar situation occurs in France, which is responsible for 82% of Pacific oyster production in Europe (92 000 tonnes in 2018 (Eurostat 2020)) where over 60% of spat is captured from wild sources (Richez 2012). This contributes to a slow shift from a natural to hatchery production model (Adamson et al. 2017). This same production template is also true for mussels; currently, industries for two of the main farmed species, the blue mussel (genus Mytilus) in Europe (Kamermans et al. 2013) and the green-lipped mussel (Perna canaliculus) in New Zealand (Symonds et al. 2019) still rely primarily on natural spat. This process is an inexpensive but unreliable practice, which is vulnerable to habitat disturbances and restricts the development of cultivation technologies such as selective breeding.

The Food and Agriculture Organization (FAO) has recently proposed a number of key developments, which will assist the aquaculture industry in addressing several long-term sustainability challenges (FAO 2016). One priority area highlighted by the FAO is the use of stock management and selective breeding to produce lines with greater reliability and productivity in a wide range of environments (FAO 2019b). To date, encouraging responses to selection have been observed in aquatic species: the average gain in body weight per generation is 8.7% in shrimps, 10.3% in oysters and between 9% and 17.9% among finfish species (Gjedrem & Rye 2018). Although recent estimates show that in the 10 main farmed aquatic species 75% of production benefits from some form of selection (Houston et al. 2020), only a small percentage of global aquaculture production (<10% in 2012) utilizes genetically improved stock (Gjedrem et al. 2012).

Despite an increasing availability of genomic resources for bivalves, the mechanisms underlying domestication (i.e. adaptation to a farmed environment), and genotype-environment interactions (GxE) ongoing in cultured bivalve species remain poorly studied. The degree to which these processes influence the response to selection in these taxa, and consequently the potential to genetically improve organisms, represent two key knowledge gaps with respect to bivalve selective breeding (Figs 1 and 2). Accordingly, the potential for losses in genetic diversity during production is exacerbated and likely hinders the efficiency of existing hatchery management and selective breeding programmes, jeopardizing sustainable growth of this sector. There is a fundamental need to clarify the impacts of hatchery-management practices on the genetic and phenotypic constitution of cohorts, and the resulting long-term implications for bivalve production.

In this review, we explore the mechanisms by which production practices and life-history characteristics can influence the genetic variability and quality of spat during hatchery-propagation (Fig. 1). We describe the current status of selection in bivalve aquaculture globally and the different methods employed for production. Further, we discuss how management practices potentially benefit or hinder the optimization of selective breeding approaches, and how a greater control of hatchery-propagation processes can contribute to the sustainable intensification of bivalve aquaculture. There are considerably fewer studies investigating the consequences of domestication selection in bivalves in comparison with other aquatic species. Therefore, in order to infer the possible consequences of artificial propagation in these organisms, we compare the selection pressures acting in hatcheries with those acting in the wild, when applicable. By identifying the main gaps in knowledge and bringing awareness to this topic, we expect to contribute to increasing efficiency and accuracy of selection in these taxa and inspire future research which may contribute to increasing efficiency and accuracy of selection in marine bivalves.

**Selective breeding in bivalve aquaculture: current status and opportunities**

Successful breeding programmes have been established for bivalves worldwide and include those applying mass and family selection approaches (Table 1) (Hollenbeck & Johnston 2018). In mass selection, individuals are typically selected according to their performance in comparison to the population’s mean for a specific trait (e.g. growth) without fully accounting for family structure. This strategy can be effective but runs the risk of inbreeding depression and is only suitable for a focus on one or two traits. Alternatively, family selection is based on pedigree information,
and individuals from the top performing families are chosen to form the breeding populations, allowing for effective maintenance of genetic diversity. Family-based selection has been applied in a commercial *M. galloprovincialis* breeding programme. Here, the use of 77 full-sib families resulted in a heritability of 0.35 (SE = 0.09) for total weight and 0.23 (SE = 0.08) for meat yield as a ratio between meat weight and total weight, after 2 generations; both of which are commercially relevant traits (Nguyen et al. 2014).

Selection has also successfully improved traits such as growth rate (Hershberger et al. 1984; de Melo et al. 2016), disease resistance (Naciri-Graven et al. 1998; Dove et al. 1998).
2013a; Dégremont et al. 2015b) and resilience towards environmental perturbation (Parker et al. 2015). Despite these success stories, bivalve aquaculture production still relies greatly on wild type strains (Hollenbeck & Johnston 2018), that may not be adapted to the farming environment (Yáñez et al. 2015). Therefore, significant potential for genomic improvement exists, providing the opportunity to maximize productivity for bivalve aquaculture species worldwide.

The recent development and increasing affordability of high-throughput sequencing technologies have facilitated the incorporation of genomic tools in breeding programmes of aquatic species (Zenger et al. 2019). This has enabled a step forward from family selection, particularly for traits which are difficult or impossible to measure directly on selection candidates, such as disease resistance. For such traits, family selection would only allow for family level breeding values, thereby missing out on within-family genetic variation. Genomic tools allow breeders to access and utilize the within-family component of genetic variation. This can be achieved in two main ways. Firstly, mapping of quantitative trait loci (QTL) allows the identification of genetic markers significantly associated with a specific trait within the species of interest. Selection for traits with large effect QTLs can be improved by applying marker-assisted selection (Zenger et al. 2019). Secondly,
genomic selection can be applied for selection of polygenic traits (Meuwissen et al. 2016). Such approach can cover a large number of loci across the genome and provides enough information to capture all causative variants for a given trait, as loci are expected to be in linkage disequilibrium with one or more common markers (Meuwissen et al. 2001). Besides, genomic selection captures the within-family variance as markers shared between individuals can be identified, increasing the accuracy of the estimated breeding values and response to selection (see Zenger et al. 2019 and references therein). Additionally, it enables retrospective parental assignment, which allows multiple families to be grown in mixed tanks and reduces the generation of confounding genetic and environmental effects. Genomic selection can also be designed to fit different levels of ploidy (Ashraf et al. 2016; Endelman et al. 2018) and can be a valuable asset to guarantee a high precision in breeding programmes utilizing broodstock with increased value through ploidy manipulation.

Having a set of tools which link high-resolution genetics with phenotypes is a main requirement for genomic selection. To date, genomes have been assembled for several of the main cultured species (Hollenbeck & Johnston 2018). In addition, the development of DNA markers including microsatellites (Li et al. 2003; Wang et al. 2016) and single nucleotide polymorphisms (SNPs) (Sauvage et al. 2007; Fleury et al. 2009; Nguyen et al. 2014; Wang et al. 2015; Vu et al. 2021) as well as the identification of genomic regions associated to traits of economic importance through QTL mapping (Sauvage et al. 2010; Guo et al. 2012; Jiao et al. 2014) and genome-wide association studies (GWAS) (Gutierrez et al. 2018; Meng et al. 2019), create a genomic toolbox which provides a backbone for future research. Ultimately, this information promotes the development of genomic based selection techniques and the fine-tuning of breeding programmes.

Marine bivalves share complex genomic and life-history features, including high levels of nuclear genetic diversity, high heterozygosity, and elevated numbers of deleterious mutations and null alleles (Bierne et al. 1998; Plough & Hedgecock 2011; Hollenbeck & Johnston 2018; Gerdol et al. 2019). In addition, reproductive attributes (broadcast spawning, high fecundity, high early mortality rates), and a high variance in reproductive success (Vk) among individuals (Hedgecock & Pudovkin 2011), are commonly described in these taxa. Variance in reproductive success can result in low effective population sizes (Ne) and low numbers of effective breeders (Nb) relative to census size, termed 'sweepstake reproduction', which has been observed in both wild and hatchery-propagated stock (Hedgecock & Sly 1990; Hedgecock 1994; Boudry et al. 2002; Plough & Hedgecock 2011). Heterozygous deficiencies relative to Hardy-Weinberg equilibrium, and segregation distortion of markers described in paired crosses, are also commonly reported in bivalves (Launey & Hedgecock 2001; Peñaloza et al. 2014).

These properties of the bivalve genome, together with specific life-history characteristics of these organisms may influence the efficiency and applicability of genomic resources in breeding programmes. Therefore, efforts to elucidate the role these features play in the selection process are vital to enhance production in this sector. Selection must focus on traits that enhance larval performance and productivity, whilst simultaneously selecting for traits which are relevant in later development. Thus, another key priority is to understand the genetic basis of these traits, as well as their genetic and developmental correlations.

### Table 1: Large-scale breeding programmes for cultured marine bivalve species (adapted from Hollenbeck & Johnston, 2018)

| Common name          | Species                  | Group       | Location  | Type of selection | Programme type | Founded | References                        |
|----------------------|--------------------------|-------------|-----------|-------------------|----------------|---------|-----------------------------------|
| Mediterranean mussel | *Mytilus galloprovincialis* | Mussel      | Australia | Family            | Industrial     | 2008    | Nguyen and Ingram (2012)          |
| Greenlip mussel      | *Perna canaliculus*      | Mussel      | New Zealand | Family           | Industrial     | 1999    | Camara and Symonds (2014)         |
| Pacific oyster       | *Crassostrea gigas*      | Oyster      | USA       | Family; mass      | Industrial     | 1996    | de Melo et al. (2016), Langdon et al. (2003) |
| Pacific oyster       | *Crassostrea gigas*      | Oyster      | Australia | Family; mass      | Industrial     | 1997    | Kube et al. (2011), Ward et al. (2005) |
| Pacific oyster       | *Crassostrea gigas*      | Oyster      | New Zealand | Mass             | Experimental   | 1999    | Camara and Symonds (2014)         |
| Pacific oyster       | *Crassostrea gigas*      | Oyster      | France    | Mass             | Experimental   | 2009    | Dégemont et al. (2015b)           |
| Sydney rock oyster   | *Saccostrea glomerata*   | Oyster      | China     | Mass             | Experimental   | 2007    | Li et al. (2011), Zhong et al. (2016) |
| Bay Scallop          | *Argopecten irradians*   | Scallop     | China     | Mass             | Unknown        | 2001    | Zheng et al. (2004), Zheng et al. (2006) |
Genomic resources have the potential to revolutionize aquaculture production, contributing to the rapid expansion and optimization of marine bivalve production. Nonetheless, socioeconomic factors also play a key role in the implementation of new technologies in existing production systems, and may slow down the pace of genomic breeding in aquaculture, especially in developing countries (Kumar et al. 2018). To date, industrial applications of genomic selection in aquatic species are limited, and largely restricted to finfish species (Zenger et al. 2019).

Broodstock conditioning and its implications on genetic variability

Contrary to natural ecosystems, hatcheries offer a largely uniform environment to cultivate broodstock, reducing sources of stress caused by sub-optimal or fluctuating conditions. In these artificial systems environmental conditions can be manipulated to trigger gametogenesis in broodstock throughout the year, extending the period through which mature breeders are available (Helm 2004). Overall, the process of induced gametogenesis, known as conditioning, aims to maximize the fecundity of progenitors whilst maintaining the high quality of gametes and larval viability (Lannan et al. 1980; Utting & Millican 1997). For aquaculture purposes, broodstock are either collected in their natural environment or taken from previous generations of hatchery stock and are held in flow-through systems (Helm 2004). During the conditioning process, quality and availability of food resources have a direct effect on adult fecundity levels and reproductive output (Utting & Millican 1997), with lipid and proteins obtained from food accumulated during oogenesis (Li et al. 2000). A significant correlation between biochemical content of oocytes and early developmental success (Massapina et al. 1999; Corporeau et al. 2012; Boulais et al. 2015), highlights the vital role that conditioning can play in production, and consequently, in the genetic makeup of cohorts (Fig. 1).

To date, standard conditioning protocols have been established for the main cultured bivalve species (Helm 2004). However, a large (up to twofold) variation in length of conditioning period is reported among strategies adopted by different hatcheries, and a quality check of broodstock gonad development is not consistently undertaken among hatcheries (de Reynaga-Franco et al. 2020). Without equal opportunity for success in breeding, Nb/N ratio is lowered. In addition, an unsynchronized response of broodstock to conditioning may reduce the potential number of breeding pairs, promote discrepancies of both Vk among individuals and performance among families (Boudry et al. 2002), with inbreeding levels within a breeding programme consequently increasing. Such issues rapidly nullify predictive ability of selective breeding methods and impose a challenge for the implementation of genomic selection in these taxa.

Genomic and phenotypic consequences of hatchery propagation

In hatcheries, spawning of broodstock can be triggered either by non-lethal techniques (thermal cycling, intermittent exposure to air and/or introduction of potassium chloride, hydrogen peroxide, steroids or neurotransmitters in the mantle cavity or adductor muscle) or by stripping (scarifying) the gonads of individuals (Helm 2004). The adoption of gonad-stripping or chemically induced spawning protocols can help to standardize the time of gamete release, reducing the deterioration of gametes. However, such approaches do not discriminate between mature and immature gametes present in the gonad. The lack of control of gamete quality during artificial spawning may lead to a high variability in developmental rate within a batch (Tan-yaros & Tarangkoon 2016). In fact, for some species such as M. edulis, gonad stripping is a non-viable approach which impairs production (Kamermans et al. 2013). Moreover, the required sacrifice of pedigreed broodstock individuals (where identified) may render this approach unfavourable for selective breeding.

Owing to its practicality, mass spawning (combining gametes from multiple females with an aliquot of pooled male gametes) is a common procedure for artificial fertilization (Helm 2004; Tetrault 2012). This approach does not control parental contribution and can result in reduced numbers of effective parents in the programme. Moreover, as best performing individuals may be excluded from crosses, mass selection can limit the accuracy of the breeding programme. However, fecundity levels observed in bivalves are high and fertilization is commonly successful, and a sufficient number of offspring is often achieved. Inbreeding load, as well as impaired development, can be concealed by management practices (e.g. culling) where the low performing individuals are eliminated from a batch by size selection (Taris et al. 2006). As genomic and marker assisted selection endeavour to capture favourable genetic variation, it is vital to identify and control for the possible impacts of spawning and fertilization protocols on the genetic variability and performance of cohorts (Fig. 2).

To overcome issues with parental contribution, pairs can be individually crossed. Paired crossing is less commonly adopted in hatcheries as it is a more laborious approach, requiring the control of fertilization rates of individual crosses and investment in personnel and equipment. This method demands additional physical space to separate mating pairs and subsequent offspring during larval development. Furthermore, the rearing of juveniles in family-specific tanks presents an issue with confounding of genetic
and common environmental effects, which would require multiple replicate tanks per family to resolve. Subsequently mixing families, and growing them together in a common environment, can mitigate against this issue. However, the gamete density used in artificial crosses is substantially higher than in nature. Empirical evidence demonstrates that mass spawning increases $V_k$ among males and pair crossing individuals increases the variance in reproductive success among females (Hornick & Plough 2019). Handling practices may additionally contribute to increase variation in family sizes, which often goes undetected. Long term, such practices can bottleneck genetic variability in artificially propagated stock and dramatically reduces the ability to predict success for selective breeding. Therefore, altered genetic diversity of hatchery-propagated stock is an inevitable consequence of the chosen fertilization approach (Fig. 1) (Hornick & Plough 2019).

Phenomena occurring at the gamete level may also play a role in determining parental contribution in crosses and act as an early selective pressure. For example, the distance which sperm must travel to reach oocytes, gamete phenotype (biochemical composition, sperm motility and behaviour, oocyte size and age) and gamete interactions influence the success of fertilization (Levitan 2006; Suquet et al. 2010; Boulais et al. 2015; Boulais et al. 2017). Genetic compatibility can influence fertilization success, favouring crosses between less related individuals (Lymbery et al. 2017). In sea urchins, low sperm densities favoured crosses between common genotypes which match at the gamete binding locus (oocyte-sperm compatibility locus) (Levitan & Ferrell 2006). Sperm-saturation, in turn, promoted reproductive success of individuals with less frequent genotypes. These findings highlight the putative role of gamete density in sperm choice behaviour. Factors such as affinity between crosses (Kekäläinen & Evans 2017) and sperm longevity (Crean et al. 2012) have been linked to increased postzygotic fitness. However, the extent to which interactions at the gamete-level, as well as gamete phenotype, influences fertilization success in external fertilization is not yet fully understood (Breed & Moore 2015).

The precise determination of oocyte-sperm ratios and controlled-crossing approaches may benefit fertilization success and enhance the contribution of individual broodstock (Song et al. 2009). Gamete density used in artificial crosses is substantially higher than in nature, increasing competition among individuals, and acts on Nb. Oocyte mechanisms acting against polyspermy are not 100% effective, thus, increased competition can lower the rates of fertilization success among crosses. Commonly, substantial variation in gamete phenotype and fertilization rates are commonly observed among and within individuals (Breed & Moore 2015). During hatchery propagation, gamete quality is assessed via crude visual observations of sperm motility and concentration, as well as shape (roundness), size and colouration of oocytes. Individuals classified with high quality gametes are selected for fertilization, whilst those not meeting the quality criteria are excluded from crosses. Correlations between gamete phenotypes (e.g. oocyte biochemical composition) and larval viability in artificially bred bivalves and other invertebrates have been previously described (Massapina et al. 1999; Crean et al. 2012; Boulais et al. 2015). Nonetheless, the extent to which gamete traits and gamete-level interactions influence $V_k$ and genetic variability of offspring is not fully understood. If such implications can be carried over throughout the individuals’ life, the expression of key genotypes may be modulated by pre-fertilization selection. However, further investigation is required to clarify how physiological and molecular mechanisms underlie gamete phenotype, and affinity of crosses during external fertilization (Fig. 2). Such knowledge can benefit the development of mating systems that maximize fertilization and homogenize $V_k$ among breeders.

Hatchery-propagated larvae are reared in a controlled environment, avoiding the risks imposed by oceanic drift and predation. In this environment, water quality parameters are maintained at, or close to, conditions considered optimal for the survival of the species being cultured. This optimized environment enables the levels of production to be improved, maximizing larval growth and settlement rates of the produced species. In the long term, domestication contributes to enhance performance under these artificial rearing conditions. However, domestication selection can lower environmental resilience when exposed to natural conditions. A lower fitness of individuals in the wild has been observed in fish species which are currently in transition to a domesticated status (Araki et al. 2008). Moreover, genomic footprints of domestication can vary greatly between populations from independent origins selected for the same trait (López et al. 2019), as a result of the specific characteristics of a rearing environment (Vandeputte et al. 2009). Recent findings indicate that selectively bred *C. virginica* larvae were less able to tolerate starvation compared with wild cohorts, experiencing significantly higher mortality rates (McFarland et al. 2020). However, the genomic mechanisms underlying domestication selection of marine bivalve species remains poorly investigated in comparison to finfish species. Optimization of selective breeding will require these factors to be better understood and controlled for.

Culling, or size selection, is commonly practised in hatcheries throughout larval development. Selection for similar growth rates under culture conditions generally improves overall spat production and reduces variation in development within a cohort (Taris et al. 2006), but may potentially mask the signs of inbreeding depression (Taris et al. 2007). Therefore, such a practice may benefit early
production stages. However, the effect of size selection on a stocks’ genetic variation are not clear (Fig. 2). Culling may result in accidental removal of individuals that may reach market size quickest in later development, individuals with alternative traits of interest (e.g. disease resistance) or traits that are relevant during later stages of production (e.g. robustness), directly impacting a breeding scheme. A practical example is seen in Mercenaria mercenaria larvae, where initially small individuals present in culture tanks are capable of surpassing the size of individuals that were initially larger, at later stages of development (Gionet et al. 2010). In addition, this process can reduce genetic variability of offspring (Taris et al. 2006) acting as a genetic bottleneck in hatcheries. Losses of entire cohorts can result from sudden shifts in conditions when GxE interactions mean that the animals selected as optimal under a hatchery production environment perform poorly in a subsequent grow-out environment.

Developmental plasticity (input during early development persisting in adult phenotype) can modify the performance of individuals in their later life. For example, exposure of quagga mussel larvae (Dreissena bugensis) to a range of temperatures has been correlated with the development of different shell morphotypes in adults (Peyer et al. 2010). If early exposure to stressors can imprint performance of organisms in later life, alternative culling strategies (e.g. application of a salinity or temperature shock during early development) would enable selection for robustness to future environmental conditions. Accordingly, the hatchery environment and management practices may themselves help or hinder spat development, potentially affecting the performance of individuals at grow-out sites (Reynaga-Franco et al. 2019).

Currently, research into the implications of hatchery practices on the genetic characteristics of bivalves is restricted to a few studies (Boudry et al. 2002; Taris et al. 2006; Taris et al. 2007; Lallias et al. 2010; Hornick & Plough 2019; McFarland et al. 2020). Unravelling the genomic basis of environmental resilience will allow the potential of selection towards robustness, or generalist phenotypes, and its association with other commercially relevant QTL to be determined (Vu et al. 2021). Additionally, the development of physiological indices of larval performance, and their association with the individual genotype, can contribute to improve selection in these taxa (Pan et al. 2016). Further studies clarifying the correlation between larval performance of hatchery selected stock and juvenile and adult performance during grow-out will contribute to the development of breeding strategies and optimization of production throughout the entire life cycle of these taxa.

Hatchery bred spat, which have reached the settlement stage, are often induced to settle. This practice not only facilitates efficient husbandry but avoids any adverse consequences (e.g. depleted energy reserves) of spending too long in the pediveliger stage. Uniformity in settlement time can be achieved by manipulating environmental stimuli such as temperature shocks, or via the addition of fine shell particles or other material to induce settlement in tanks (Helm 2004). Alternatively, settlement of larvae can be chemically induced by exposure to neurotransmitters (Sánchez-Lazo & Martínez-Pita 2012; Grant et al. 2013; Joyce & Vogeler 2018). Further investigation is needed to elucidate the role such approaches play as a selective pressure in the hatchery environment and whether these can be used to select or induce favourable characteristics (Fig. 2).

Settlement and metamorphosis are critical moments in the life cycle of bivalves. Substantial mortalities occur during these stages in both natural populations and artificially propagated stock (Hunt & Scheibling 1997; Plough & Hedgecock 2011; Plough 2016), with survival at the post-settlement stage reaching only 2.8% of the original population in some cases (Plough 2016). Genotype-dependent mortality linked to deleterious recessive mutations can occur immediately before or during metamorphosis (Plough & Hedgecock 2011; Plough 2016). Insights on genotype-dependent mortality during settlement have opened the opportunity to investigating the applicability of QTLs to select for uniformity of settlement timing (Plough 2016). In contrast, mortality in the period immediately post-settlement is lower, with no indication of being genotype-dependent (Dégremont et al. 2007; Plough 2016).

**Genotype by environment responses to the grow-out environment**

All spat, both wild and hatchery propagated, are exposed to environmental variability experienced within the coastal and estuarine zones in which grow-out occurs, and are thus susceptible to this daily and seasonal variability (Fig. 1). To thrive in such demanding environments, individuals must either be genetically adapted to extreme conditions, or possess highly plastic physiological responses which allow them to regulate internal mechanisms.

Accordingly, Pacific oysters have demonstrated the ability to regulate genes involved in stress response pathways when facing abiotic stress conditions, including elevated temperature and air exposure (Zhang et al. 2012). These findings suggest that a high level of plasticity is a strategy which has allowed these sessile organisms to successfully colonize stressful environments. The expansion of gene families that function as part of the organism’s response against biotic and abiotic stress, as well as immune response, suggest that this group has adapted to a sessile life in fluctuating environments (i.e. intertidal coastal and estuarine waters). A better understanding of plasticity mechanisms in bivalves can contribute to the development of...
culturing conditions which improve performance in desirable traits. The selection processes experienced during early development in hatcheries could contribute to direct effects on the performance of cohorts, as well as increasing the likelihood of stochastic GxE interactions.

GxE is an important factor dictating performance of aquaculture species (see review by Sae-Lim et al. 2016 and references therein). Where animals from a breeding programme are reared in different environments, it can result in a re-ranking of families or genotypes. This can negatively impact genetic gain and the effectiveness of a breeding programme. These effects have been observed in previous studies investigating the variation in C. gigas performance among families across grow-out sites (Langdon et al. 2003; Evans & Langdon 2006). In other cases, selected genotypes outperform in certain conditions, but become poor performers when exposed to a different set of conditions – that is re-ranking of genotypes (Langdon et al. 2003; Dégremont et al. 2005; Evans & Langdon 2006; Wang et al. 2013). In marine bivalves, between-family variance described for traits such as growth, survival and environmental resilience (Dégremont et al. 2005; Dégremont et al. 2013a; Scanes et al. 2020) indicates the genetic basis of traits associated with performance (Vu et al. 2021). The re-ranking of genotypes, in turn, highlights the intrinsic effect of GxE on overall performance of a family or cohort. There is also genetic variation in how well animals perform across diverse environmental conditions, and this robustness of genotypes to diverse conditions can be analysed using reaction norms, and potentially incorporated into breeding goals to help tackle the impact of GxE (Hill & Mulder 2010).

Epigenetic mechanisms (e.g. DNA methylation, histone modifications, non-coding RNAs) are a relevant component of GxE interactions, through exposure mediated GxE. These mechanisms can modify a phenotype without changing the DNA sequence and can have long-lasting effects (Jablonska & Lamb 2002). In the last decades, new technologies have facilitated the study of epigenetics, providing insights into the contribution of the epigenome to the expressed phenotypes in response to the environment. Among the wide scale of techniques available to study epigenetic regulation, DNA methylation has received the most attention in marine bivalves. In C. gigas, DNA methylation patterns have been associated with gene function (Gavery & Roberts 2010) and have been linked to gene regulation (Riviere et al. 2013; Olson & Roberts 2014). Environmental heterogeneity has been associated with divergent DNA methylation patterns among C. virginica populations (Johnson & Kelly 2020). In Mytilus galloprovincialis and the New Zealand pygmy mussel Xenostrobus secures, methylation patterns of invasive populations differ from populations in their native range (Ardura et al. 2018). Whilst epigenetics can contribute to rapid and transient plasticity in response to stress and environment in marine bivalves, future studies combining genomic and epigenomic information are needed to elucidate the processes underlying GxE interactions and phenotype expression in these taxa.

Recent evidence also underlines the adaptive nature of phenotypic plasticity in traits involved in environmental resilience (Li et al. 2018). Domestication, or the reduction in environmental variation in early life stages, is unlikely to select for plasticity and may lead to epigenetic profiles that are less suited to the farm environment. However, the impact of artificial-breeding and early life hatchery conditions on the epigenome of marine bivalve species remains unresolved. Selective pressures acting during hatchery-propagation most likely favour domestication rather than adaptation towards variable natural environments. Therefore, the potential of hatchery-propagated stock to cope with environmental stress may be reduced during breeding and hatchery processes. Indeed, the epigenome of artificially bred Atlantic salmon differs greatly from wild populations, and the reduced fitness of hatchery-propagated stock in comparison to wild populations is likely a consequence of such variation (Le Luyer et al. 2017). Accordingly, signs of lower tolerance to environmental stress in C. virigina have been linked to domestication selection (Mcfarland et al. 2020). Here, we emphasize that domestication selection in early life stages could result in high discrepancies in performance and lower the mean performance of cohorts through GxE interactions.

For a single species, the commercial market may expand across multiple production environments, which may have to be reflected in the data collection for a breeding programme. In a simple breeding programme design, the breeding candidates and the test animals are the same and held in one environment. In a more advanced design, the selection decisions are also based on the performance records of pedigreed full- and half-sibs of the candidates held at test stations (sib-testing design). The breeding candidates are normally reared at a single breeding nucleus farm, where strict biosecurity and sanitary restrictions are imposed to prevent serious pathogens from entering the breeding nucleus. However, the breeding candidates may as well be reared at a few locations from which the families are produced and at a later and safer stage are transported to the central breeding nucleus. In both cases, this structure may induce GxE effects which may have to be accounted for.

To understand the role of GxE interactions in the expression of phenotypes, the performance of different lines needs to be tested in a range of environments. Strong GxE interactions could be countered by the creation of specific breeding programmes targeting specific grow-out environments (Dégremont et al. 2007). However, it would first be wise to understand if any of the potential hatchery stressors or selection events described herein are contributors to GxE.
events, and if they can be mitigated through alteration in early life selection. As a crude hypothesis, growth in *C. gigas*, a trait which has been under selection pressure in the hatchery, seems to be highly dependent on the environment, whereas other traits such as survival in the presence of disease seem dependent on the family (Dégremont *et al.* 2005; Evans & Langdon 2006). Expression of phenotypes can be maintained within families across a range of environments by epigenetic mechanisms (Gavery & Roberts 2017; Uren Webster *et al.* 2018).

Other omic techniques, such as proteomics and metabolomics, provide a direct measurement of expressed phenotypes, and are therefore valuable tools to explore the genotype-phenotype link and evaluate performance (Laudicella *et al.* 2020). Further studies investigating the relation between specific environmental conditions utilizing a holistic omic approach may allow to understand and control for GxE in bivalve breeding programmes and are critical to improve aquaculture (Fig. 2).

**Implications on selective breeding under a changing climate**

Shifts in sea surface salinity, temperature and ocean chemistry (e.g. ocean acidification), alterations in precipitation patterns as well as stronger and more frequent heat waves, are some of the main consequences of climate change to the marine environment predicted for the coming decades (IPCC 2018).

As ectothermic calcifying organisms, marine bivalves are particularly vulnerable to climate change. Shell dissolution and decreased shell growth caused by ocean acidification have been described in marine bivalves (Melzner *et al.* 2011). Higher sea surface temperatures, especially in summer months, may challenge species with lower thermal tolerance (Steeves *et al.* 2018). Fluctuating sea surface salinities may have deleterious implications for shell growth (Riisgard *et al.* 2012) whilst the interactions of this factor with increased temperature or hypercapnia (elevated CO₂) can increase mortality (Rybovich *et al.* 2016) and reduce hardness and resistance of shells (Dickinson *et al.* 2012). Phytoplankton communities are likely to be impacted by climate change (Käse & Geuer 2018), and temporal shifts in species abundance and composition may impact the nutrient uptake in marine bivalves, limiting physiological and biological processes. Climate change may also contribute to lowering the immune response of bivalves (Mackenzie *et al.* 2014), and modify host–pathogen interactions, increasing sensitivity towards diseases (Asplund *et al.* 2014).

The grow-out phase of bivalve aquaculture takes place in the natural environment. Therefore, the implications of climate change are not restricted to wild populations. Strong changes in local environmental conditions may limit production and force the relocation of grow-out sites to suitable areas. Environmental changes and increased disease outbreaks might lead to severe mortality and considerable economic losses in this industry and restrictions in spat commercialization may be needed to avoid the further spread of diseases. Thinner and weaker shells will facilitate their rupture during transportation. Hatchery propagation may also be impaired by climate change to a certain degree, as those rely on natural sea water supply. Therefore, there is an imminent need for research to develop bivalve strains robust to climate change and resilient towards diseases.

Epigenetic processes can contribute to rapid adaptation towards environmental stressors generated by climate change. In *S. glomerata*, short-term exposure to elevated CO₂ concentration not only increases resilience of exposed individuals, but can also be passed through generations (Parker *et al.* 2015). Such resilience has been associated with a change in regulation of genes associated with stress related functions (Goncalves *et al.* 2016). Accordingly, empirical evidence demonstrates that low pH stress (pH 7.4) can modify the methylation patterns of *Crassostrea hongkongensis* pediveliger larvae (Lim *et al.* 2021). Genomic processes, in turn, are involved in long term adaptation to environmental changes.

Identifying the mechanisms acting behind GxE interactions which increase performance of a species under climate change-associated stressors is an important step to characterize the genomic and epigenomic profile of robust genotypes. Accordingly, GxE can be exploited in breeding programmes to increase environmental resilience and the application of genomic selection can fast-track the development of such lines (Mulder 2016). The growing body of high quality assembled genomes facilitate the precise identification of genomic regions linked to traits responsible for environmental resilience. The application of genomic selection, or gene editing approaches, can then facilitate the development of robust lines, or lines able to withstand suboptimal environmental conditions relevant to a certain grow-out region (e.g. elevated temperature, low pH). As hatcheries allow for the control of genetic stocks, indoor propagation is undoubtedly an essential asset to guarantee the development of lines able to thrive under future predicted environmental scenarios.

**Summary and future perspectives**

A gap in knowledge remains on how domestication selection and husbandry practices can constrain genetic variability of hatchery-propagated stock during early life stages in marine bivalves (Fig. 2). Despite the negative implications of inbreeding load on performance, the control of reproductive output, differential performance of genotypes and genetic variability of stock remains relatively low in bivalve
production. Such lack of control may hinder spat performance and consequently aquaculture production. Future advances in bivalve production and selective breeding require an understanding and optimization of hatchery production processes in order to maximize genetic gain.

Selection pressures acting in hatcheries differ from those acting on the populations in their natural environment. In contrast to wild populations that face fluctuating environmental conditions, farmed stocks are produced under relatively stable, benign, conditions but exposed to other stresses such as elevated densities and handling practices. Organisms reared in hatcheries, naïve to the wild, might lack resilience to environmental variation due to domestication selection and/or epigenetic mechanisms. In addition, negative GxE interactions can be detrimental for production and can be compounded by artificial bottlenecks or epigenetic alterations caused by the hatchery environment. However, it is still not fully understood whether selection for phenotypes that enhance hatchery production contribute to adult performance during the grow-out phase. Therefore, whilst performance of larval stages must remain as an important component of breeding programmes and hatchery production, it is key to consider traits related to the challenges these larvae will face during the grow-out and production phases.

Genomic resources will contribute with the understanding of evolutionary and adaptive processes, as well as those which are linked to domestication (Yáñez et al. 2015). Elucidating the (epi)genomic mechanisms which underpin the expressed phenotypes will allow the divergence of selection from classic commercial traits towards broad environmental resilience (either outperforming or generalist genotypes). Integrating robustness as a founding criterion for selection can potentially contribute to increase grow-out productivity, especially in light of climate change.

Genomic selection can favour the development of genetically improved lines for multiple traits, facilitate the management of genetic variability (D’Ambrosio et al. 2019) and potentially reduce environmental sensitivity accounting for GxE (Mulder 2016). Most importantly, the implementation of such selection approaches is key to the sustainable optimization of bivalve aquaculture production, particularly in the light of climate change. It is crucial to focus resources on developing environmentally robust lines. However, progress of marker assisted and genomic selection in bivalve aquaculture will require a greater control of hatchery practices to allow sources of unaccounted genetic variation to be minimized and genetic gain to be maximized.

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References

Adamson E, Syvret M, Woolmer A (2017) Shellfish seed supply for aquaculture in the UK report on views collected from the industry in 2017. London.
Araki H, Berejikian BA, Ford MJ, Blouin MS (2008) SYNTHESIS: fitness of hatchery-reared salmonids in the wild. Evolutionary Applications 1: 342–355.
Ardura A, Clusa L, Zaiko A, García-Vazquez E, Miralles I. (2018) Stress related epigenetic changes may explain opportunistic success in biological invasions in Antipode mussels. Scientific Reports 8: 1–10.
Asphalt ME, Baden SP, Russ S, Ellis RP, Gong N, Herrnoth BE (2014) Ocean acidification and host–pathogen interactions: blue mussels, Mytilus edulis, encountering Vibrio tubiashii. Environmental Microbiology 16: 1029–1039.
Bierne N, Launey S, Naciri-Graven Y, Bonhomme F (1998) Early effect of inbreeding as revealed by microsatellite analyses on Ostrea edulis larvae. Genetics 148: 1893–1906.
Boudry P, Collet B, Cornette F, Hervouet V, Bonhomme F (2002) High variance in reproductive success of the Pacific oyster (Crassostrea gigas, Thunberg) revealed by microsatellite–based parentage analysis of multifactorial crosses. Aquaculture 204: 283–296.
Boulais M, Charlotte C, Arnaud H, Ismaïl B, Claudie Q, Virgile Q et al. (2015) Assessment of oocyte and trochophore quality in Pacific oyster, Crassostrea gigas. Aquaculture 437: 201–207.
Boulais M, Chenevert KJ, Demey AT, Darrow ES, Robison MR, Roberts JP et al. (2017) Oyster reproduction is compromised by acidification experienced seasonally in coastal regions. Scientific Reports 7: 1–9.
Breed MD, Moore J (2015) Animal Behaviour, 2nd edn. Elsevier, San Diego.
Camara MD, Symonds JE (2014) Genetic improvement of New Zealand aquaculture species: programmes, progress and
prospects. *New Zealand Journal of Marine and Freshwater Research* **48**: 466–491.

Carriker MR (1956) Biology and propagation of young hard clams, *Mercenaria mercenaria*. *Journal of the Elisha Mitchell Scientific Society* **72**: 57–60.

Corporeau C, Vanderplancke G, Boulais M, Suquet M, Quéré C, Boudry P et al. (2012) Proteomic identification of quality factors for oocytes in the Pacific oyster *Crassostrea gigas*. *Journal of Proteomics* **75**: 5554–5563.

Creen AJ, Dwyer JM, Marshall DJ (2012) Fertilization is not a new beginning: the relationship between sperm longevity and offspring performance. *PLoS One* **7**: e49167.

D’Ambrosio J, Phocas F, Haffray P, Bestin A, Brard-Fudulea S, Boudry P et al. (2019) Genome-wide estimates of genetic diversity, inbreeding and effective size of experimental and commercial rainbow trout lines undergoing selective breeding. *Genetics Selection Evolution* **51**: 26.

de Reynaga-Franco FJ, Grijalva-Chon J-M, Castro-Longoria R, Barraza-Guardado R-H, Arreola-Lizarraga J-A, Chávez-Villalba J (2020) Designing a protocol to evaluate *Crassostrea gigas* spat production in hatcheries: identification of critical aspects. *Aquacultural Engineering* **89**: 102055.

Dégrement L, Bédier E, Soletchnik P, Ropert M, Huvet A, Moal J et al. (2005) Relative importance of family, site, and field placement timing on survival, growth, and yield of hatchery-produced Pacific oyster spat (*Crassostrea gigas*). *Aquaculture* **249**: 213–229.

Dégrement L, Ermande B, Bédier E, Boudry P (2007) Summer mortality of hatchery-produced Pacific oyster spat (*Crassostrea gigas*). I. Estimation of genetic parameters for survival and growth. *Aquaculture* **299**: 21–29.

Dégrement L, Garcia C, Allen SK (2015a) Genetic improvement for disease resistance in oysters: a review. *Journal of Invertebrate Pathology* **131**: 226–241.

Dégrement L, Nourry M, Maurouard E (2015b) Mass selection for survival and resistance to OsHV-1 infection in *Crassostrea gigas* spat in field conditions: response to selection after four generations. *Aquaculture* **446**: 111–121.

Dickinson GH, Ivanina AV, Matoo OB, Pörtner HO, Lannig G, Bock C et al. (2012) Interactive effects of salinity and elevated CO2 levels on juvenile eastern oysters, *Crassostrea virginica*. *Journal of Experimental Biology* **215**: 29–43.

Dove MC, Neill JA, Mcorrie S, Wayne AO (2013a) Assessment of Qx and winter mortality disease resistance of mass selected Sydney rock oysters, *Saccostrea glomerata* (Gould, 1850), in the Hawkesbury River and Merimbula Lake, NSW Australia. *Journal of Shellfish Research* **32**: 681–687.

Dove MC, Neill JA, O’Connor WA (2013b) Evaluation of the progeny of the fourth-generation Sydney rock oyster *Saccostrea glomerata* (Gould, 1850) breeding lines for resistance to QX disease (*Martelia sydneyi*) and winter mortality (*Bomania roughleyi*). *Aquaculture Research* **44**: 1791–1800.

Endelman JB, Carley CAS, Bethke PC, Coombs JJ, Clough ME, da Silva WL et al. (2018) Genetic variance partitioning and genome-wide prediction with allele dosage information in autotetraploid potato. *Genetics* **209**: 77–87.

Eurostat (2020). Fisheries statistics https://ec.europa.eu/eurostat/web/fisheries/data/database

Evans S, Langdon C (2006) Effects of genotype × environment interactions on the selection of broadly adapted Pacific oysters (*Crassostrea gigas*). *Aquaculture* **261**: 522–534.

FAO (2016) The state of world fisheries and aquaculture 2016. Contributing to food security and nutrition for all. Rome, 2016.

FAO (2019a) FAO yearbook. Fishery and Aquaculture Statistics 2017/FAO annuaire. Statistiques des pêches et de l’aquaculture 2017/ FAO anuario. Estadísticas de pesca y acuicultura 2017. Rome, 2019.

FAO (2019b). The state of the World’s aquatic genetic resources for food and agriculture. Rome, 2019.

Fleury E, Huvet A, Lelong C, de Lorgeril J, Boulo V, Gueguen Y et al. (2009) Generation and analysis of a 29,745 unique expressed sequence tags from the Pacific oyster (*Crassostrea gigas*) assembled into a publicly accessible database: the GigaSBase. *BMC Genomics* **10**: 1–15.

Galstoff PS (1938) Physiology of reproduction of *Ostrea virginica*. II. Stimulation of spawning in the female oyster. *The Biological Bulletin* **75**: 286–307.

Gavery MR, Roberts SB (2010) DNA methylation patterns provide insight into epigenetic regulation in the Pacific oyster (*Crassostrea gigas*). *BMC Genomics* **11**: 1–9.

Gavery MR, Roberts SB (2017) Epigenetic considerations in aquaculture. *PeerJ* **5**: e4147.

Gerold M, Moreira R, Cruz F, Gómez-Garrido J, Vlasova A, Rosani U et al. (2019). Massive gene presence/absence variation in the mussel genome as an adaptive strategy: first evidence of a pan-genome in *Metazoa*. *bioRxiv* 781377.

Gionet C, Mayrand E, Landry T (2010) Elimination of animals with best growth potential as a possible effect of the culling of *Mercenaria mercenaria* notata (L.) larvae in hatchery procedure. *Aquaculture International* **18**: 801–812.

Gjedrem T, Robinson N, Rye M (2012) The importance of selective breeding in aquaculture to meet future demands for animal protein: a review. *Aquaculture* **350–353**: 117–129.

Gjedrem T, Rye M (2018) Selection response in fish and shellfish: a review.

Goncalves P, Anderson K, Thompson EL, Melwani A, Parker LM, Ross PM et al. (2016) Rapid transcriptional acclimation following transgenerational exposure of oysters to ocean acidification. *Molecular ecology* **25**: 4836–4849.

Grant MN, Meritt DW, Kimmel DG (2013) Chemical induction of settlement behavior in larvae of the eastern oyster *Crassostrea virginica* (Gmelin). *Aquaculture* **402**: 84–91.

Guo X, Li Q, Wang QZ, Kong LF (2012) Genetic mapping and QTL analysis of growth-related traits in the pacific oyster. *Marine Biotechnology* **14**: 218–226.

Gutierrez AP, Hooper C, Bean T, Stenton CA, Sanders MB, Paley RK et al. (2018) A genome-wide association study for...
host resistance to ostreid herpesvirus in Pacific oysters (Crassostrea gigas). G3: Genes Genomes Genetics 8: 1273–1280.

Hedgecock D (1994) Does variance in reproductive success limit effective population sizes of marine organisms? Genetics and Evolution of Aquatic Organisms 122: 122–134.

Hedgecock D, Pudovkin AI (2011) Sweepstakes reproductive success in highly fecund marine fish and shellfish: a review and commentary. Bulletin of Marine Science 87: 971–1002.

Hedgecock D, Sly F (1990) Genetic drift and effective population sizes of hatchery-propagated stocks of the Pacific oyster, Crassostrea gigas. Aquaculture 88: 21–38.

Helm MM (2004). Hatchery Culture of Bivalves: A Practical Manual, Vol. 639.2. Food and Agriculture Organization of the United Nations, Rome.

Hershberger WK, Perdue JA, Beattie JH (1984) Genetic selection and systematic breeding in Pacific oyster culture. Aquaculture 39: 237–245.

Hill WG, Mulder HA (2010) Genetic analysis of environmental variation. Genetics Research 92: 381–395.

Hollenbeck CM, Johnston IA (2018) Genomic tools and selective breeding in molluscs. Frontiers in Genetics 9: 1–15.

Hornick KM, Plough LV (2019) Tracking genetic diversity in a large-scale oyster restoration program: effects of hatchery propagation and initial characterization of diversity on restored vs. wild reefs. Heredity 123: 92–105.

Houston RD, Bean TP, Macqueen DJ, Gundappa MK, Jin YH, Jenkins TL et al. (2020) Harnessing genomics to fast-track genetic improvement in aquaculture. Nature Reviews Genetics 21: 389–409.

Hunt HL, Scheibling RE (1997) The relationship between egg size, genotype frequency, and plasticity shape adaptive potential of the Pacific oyster. Evolution of Aquatic Organisms 45: 105–114.

Jabalokka E, Lamb MJ (2002) The changing concept of epigenetics. Annals of the New York Academy of Sciences 981: 82–96.

Jiao W, Fu X, Dou J, Li H, Su H, Mao J et al. (2014) High-resolution linkage and quantitative trait locus mapping aided by genome survey sequencing: building up an integrative genomic framework for a bivalve mollusc. DNA Research 21: 85–101.

Johnson KM, Kelly MW (2020) Population epigenetic divergence exceeds genetic divergence in the Eastern oyster Crassostrea virginica in the Northern Gulf of Mexico. Evolutionary Applications 13: 945–959.

Joyce A, Vogeler S (2018) Molluscan bivalve settlement and metamorphosis: neuroendocrine inducers and morphogenetic responses. Aquaculture 487: 64–82.

Kamermans P, Galley T, Boudry P, Fuentes J, McCombie H, Batista FM et al. (2013). Blue Mussel Hatchery Technology in Europe, Vol. 2009, pp. 339–373. Woodhead Publishing Limited.

Käse L, Geuer JK (2018) Phytoplankton Responses to Marine Climate Change – An Introduction BT - YOUNARES 8 – Oceans Across Boundaries: Learning from each other. Springer Nature pp. 55–71.

Kellalainen J, Evans JP (2017) Female-induced remote regulation of sperm physiology may provide opportunities for gamete-level mate choice. Evolution 71: 238–248.

Kobayashi M, Miang S, Batka M, Vannuccini S, Dey MM, Anderson JL (2015) Fish to 2030: the role and opportunity for aquaculture. Aquaculture Economics and Management 3: 282–300.

Kube P, Cunningham M, Dominik S, Parkinson S, Henshall J, Finn B et al. (2011). Enhancement of the Pacific Oyster Selective Breeding Program. FRDC and Seafood CRC, Hobart. pp. 177 (Project no. 2006/227).

Kumar G, Enge C, Tucker C (2018) Factors driving aquaculture technology adoption. Journal of The World Aquaculture Society 49: 447–476.

Lallias D, Boudry P, Lapègue S, King JW, Beaumont AR (2010) Strategies for the retention of high genetic variability in European flat oyster (Ostrea edulis) restoration programmes. Conservation Genetics 11: 1899–1910.

Langdon C, Evans F, Jacobson D, Blouin M (2003) Yields of cultured Pacific oysters Crassostrea gigas Thunberg improved after one generation of selection. Aquaculture 220: 227–244.

Lannan JE, Robinson A, Breese WP (1980) Broodstock management of Crassostrea gigas. II. Broodstock conditioning to maximize larval survival. Aquaculture 21: 337–345.

Laudicella VA, Whitfield PD, Carboni S, Doherty MK, Hughes AD (2020) Application of lipidomics in bivalve aquaculture, a review. Reviews in Aquaculture 12: 678–702.

Launey S, Hedgecock D (2001) High genetic load in the Pacific oyster Crassostrea gigas. Genetics 158: 253–265.

Le Luyer J, Laporte M, Beacham TD, Kaukinen KH, Withler RE, Leong JS et al. (2017) Parallel epigenetic modifications induced by hatchery rearing in a Pacific salmon. Proceedings of the National Academy of Sciences 114: 12964–12969.

Levitan DR (2006) The relationship between egg siz. Integrative and Comparative Biology 46: 298–311.

Levitan DR, Ferrell DL (2006) Selection on gamete recognition proteins depends on sex, density, and genotype frequency. Science 312: 267–269.

Li G, Hubert S, Bucklin K, Ribes V, Hedgecock D (2003) Characterization of 79 microsatellite DNA markers in the Pacific oyster Crassostrea gigas. Molecular Ecology Notes 3: 228–232.

Li J, Li A, Song K, Meng J, Guo X, Li S et al. (2018) Divergence and plasticity shape adaptive potential of the Pacific oyster. Nature Ecology and Evolution 2: 1751–1760.

Li Q, Osada M, Mori K (2000) Seasonal biochemical variations in Pacific oyster gonadal tissue during sexual maturation. Fisheries Science 66: 502–508.

Li Q, Wang Q, Liu S, Kong L (2011) Selection response and realized heritability for growth in three stocks of the Pacific oyster Crassostrea gigas. Fisheries Science 77: 643–648.
Lim Y-K, Cheung K, Dang X, Roberts SB, Wang X, Thiyagarajan V (2021) DNA methylation changes in response to ocean acidification at the time of larval metamorphosis in the edible oyster, *Crassostrea hongkongensis*. *Marine Environmental Research* **163**: 105217.

Loosanoff VL, Davis HC (1963) *Rearing of Bivalve Mollusks*. In Vol. 1, ed. FSBT–A in MB Russell, pp. 1–136. Academic Press.

López ME, Benestan L, Moore J-S, Perrier C, Gilbey J, Di Genova A et al. (2019) Comparing genomic signatures of domestication in two Atlantic salmon (*Salmo salar* L.) populations with different geographical origins. *Evolutionary Applications* **12**: 137–156.

Lymbery RA, Kennington WJ, Evans JP (2017) Egg chemoattraction and disease status in a commercial shellfish species, *Mytilus edulis* L. *PLoS One* **9**: e99712.

Mann R (1983) Bivalve mollusc hatcheries: a critical appraisal of their development and a review of their potential value in enhancing the fisheries of developing nations. *Mem. Asoc. Latinoam. Acuicultura* **5**: 97–105.

Massapina C, Joaquim S, Matias D, Devauchelle N (1999) Oocyte and embryo quality in *Crassostrea gigas* (Portuguese strain) during a spawning period in Algarve, South Portugal. *Aquatic Living Resources* **12**: 327–333.

McFarland K, Plough LV, Nguyen M, Hare MP (2020) Are bivalves susceptible to domestication selection? Using starvation tolerance to test for potential trait changes in eastern oyster larvae. *PLoS One* **15**: 230222.

de Melo C, Durland E, Langdon C (2016) Improvements in desirable traits of the Pacific oyster, *Crassostrea gigas*, as a result of five generations of selection on the West Coast, USA. *Aquaculture* **460**: 105–115.

Melzer F, Stange P, Trübenbach K, Thomsen J, Casties I, Panknin U et al. (2011) Food supply and seawater pCO2 Impact calcification and internal shell dissolution in the blue mussel *Mytilus edulis*. *PLoS One* **6**: e24223.

Meng J, Song K, Li C, Liu S, Shi R, Li B et al. (2019) Genome-wide association analysis of nutrient traits in the oyster *Crassostrea gigas*: genetic effect and interaction network. *BMC Genomics* **20**: 625.

Meuwissen THE, Hayes BJ, Goddard ME (2001) Prediction of total genetic value using genome-wide dense marker maps. *Genetics* **157**: 1819–1829.

Meuwissen T, Hayes B, Goddard M (2016) Genomic selection: a paradigm shift in animal breeding. *Animal Frontiers* **6**: 6–14.

Mulder HA (2016) Genomic selection improves response to selection in resilience by exploiting genotype by environment interactions. *Frontiers in genetics* **7**: 178.

Naciri-Graven Y, Martin A-G, Baud J-P, Renault T, Gérard A (1998) Selecting the flat oyster *Ostrea edulis* (L.) for survival when infected with the parasite *Bonamia ostreae*. *Journal of Experimental Marine Biology and Ecology* **224**: 91–107.

Nell JA, Sheridan AK, Smith IR (1996) Progress in a Sydney rock oyster, *Saccostrea commercialis* (Iredale and Roughley), breeding program. *Aquaculture* **144**: 295–302.

Nell JA, Smith IR, Sheridan AK (1999) Third generation evaluation of Sydney rock oyster *Saccostrea commercialis* (Iredale and Roughley) breeding lines. *Aquaculture* **170**: 195–203.

Nguyen TTT, Hayes BJ, Ingram BA (2014) Genetic parameters and response to selection in blue mussel (*Mytilus galloprovincialis*) using a SNP–based pedigree Potential broodstock candidates. *Aquaculture* **420**: 295–301.

Nguyen TTT, Ingram BA (2012) Progress on Selective Breeding Program for Blue Mussel in Victoria. Department of Primary Industries, Australia.

Olson CE, Roberts SB (2014) Genome-wide profiling of DNA methylation and gene expression in *Crassostrea gigas* male gametes. *Frontiers in Physiology* **5**: 224.

Pan TCF, Applebaum SL, Lentz BA, Manahan DT (2016) Predicting phenotypic variation in growth and metabolism of marine invertebrate larvae. *Journal of Experimental Marine Biology and Ecology* **483**: 64–73.

Parker LM, O’Connor WA, Raftos DA, Pörtner HO, Ross PM (2015) Persistence of positive carryover effects in the oyster, *Saccostrea glomerata*, following transgenerational exposure to ocean acidification. *PLoS One* **10**: 132276.

Peñaloza C, Bishop S, Toro J, Houston RD (2014). RAD Sequencing reveals genome-wide heterozygote deficiency in pair crosses of the Chilean mussel *Mytilus* spp. *10th World Congr. Genet. Appl. to Livest. Prod.*

Peyer SM, Hermanson JC, Lee CE (2010) Developmental plasticity of shell morphology of quagga mussels from shallow and deep-water habitats of the Great Lakes. *Journal of Experimental Biology* **213**: 2602–2609.

Plough LV (2016). Fine-scale temporal analysis of genotype-dependent mortality at settlement in the Pacific oyster *Crassostrea gigas*. bioRxiv 84616.

Plough LV, Hedgecock D (2011) Quantitative trait locus analysis of stage-specific inbreeding depression in the Pacific oyster *Crassostrea gigas*. *Genetics* **189**: 1473–1486.

Reymaga-Franco F-J, Grijalva-Chon J-M, Castro-Longoria R, Baraza-Guardado R-H, Arreola-Lizarraga J-A, Chávez-Villalba J (2019) Biological performance of *Crassostrea gigas* stocks produced at different hatcheries and cultivated under same environmental conditions. *Aquaculture Research* **50**: 621–633.

Richez F (2012) Report on the impact of recent *Crassostrea gigas* mortality in France and its consequences to oyster farming in Northern Ireland. Aquaculture Initiative, Department of Agriculture and Rural Development (NI) and European Fisheries Fund. Department of Agriculture and Rural Development (Northern Ireland), Belfast, UK.

Rissgård HU, Bettiger L, Pleissner D (2012) Effect of salinity on growth of mussels, *mytilus edulis*, with special reference to great belt (Denmark). *Open Journal of Marine Science* **2**: 167–176.
Riviere G, Wu G-C, Fellous A, Goux D, Sourdaire P, Favrel P (2013) DNA methylation is crucial for the early development in the oyster Crassostrea gigas. Marine Biotechnology 15: 739–753.

Rybovich M, La Peyre MK, Hall SG, La Peyre JF (2016) Increased temperatures combined with lowered salinities differentially impact oyster size class growth and mortality. Journal of Shellfish Research 35: 101–113.

Sae-Lim P, Gjerde B, Nielsen HM, Mulder H, Kause A (2016) A review of genotype-by-environment interaction and micro-environmental sensitivity in aquatic species. Reviews in Aquaculture 8: 369–393.

Sánchez-Lazo C, Martínez-Pita I (2012) Induction of settlement in larvae of the mussel Mytilus galloprovincialis using neuroactive compounds. Aquaculture 344: 210–215.

Sauve C, Bierne N, Lapègue S, Boudry P (2007) Single Nucleotide polymorphisms and their relationship to codon usage bias in the Pacific oyster Crassostrea gigas. Gene 406: 13–22.

Sauve C, Boudry P, De Koning D-J, Haley CS, Heurtebise S, Lapègue S (2010) QTL for resistance to summer mortality and OsHV-1 load in the Pacific oyster (Crassostrea gigas). Animal Genetics 41: 390–399.

Scanes E, Parker LM, O’Connor WA, Dove MC, Ross PM (2020) Heatwaves alter survival of the Sydney rock oyster. Saccostrea glomerata. Marine Pollution Bulletin 158: 111389.

Song YP, Suquet M, Quéau I, Lebrun L (2009) Setting of a procedure for experimental fertilisation of Pacific oyster (Crassostrea gigas) oocytes. Aquaculture 287: 311–314.

Steeves LE, Filgueira R, Guyondet T, Chassé J, Comeau L (2018) Past, present, and future: performance of two bivalve species under changing environmental conditions. Frontiers in Marine Science 5: 184.

Subasinghe R (2017) World aquaculture 2015: a brief overview. FAO Fisheries and Aquaculture Report Vol. 1140.

Suquet M, Labbe C, Brizard R, Donval A, Le Coz JR, Quere C et al. (2010) Changes in motility, ATP content, morphology and fertilisation capacity during the movement phase of tetraploid Pacific oyster (Crassostrea gigas) sperm. Theriogenology 74: 111–117.

Symonds JE, Clarke SM, King N, Walker SP, Blanchard B, Sutherland D et al. (2019) Developing successful breeding programs for new zealand aquaculture: a perspective on progress and future genomic opportunities. Frontiers in Genetics 10: 1–7.

Tanyaros S, Tarangkoon W (2016) Variability in larval period, post-settling growth and survival of the oyster Crassostrea belcheri produced by gamete stripping method. Agriculture and Natural Resources 50: 295–298.

Tarit N, Batista FM, Boudry P (2007) Evidence of response to unintentional selection for faster development and inbreeding depression in Crassostrea gigas larvae. Aquaculture 272: 69–79.

Tarit N, Sauve C, Batista FM, Baron S, Ernande B, Haffray P et al. (2006) Phenotypic and genetic consequences of size selection at the larval stage in the Pacific oyster (Crassostrea gigas). Journal of Experimental Marine Biology and Ecology 333: 147–158.

Tetrault K (2012) Reference manual for SPAT oyster gardeners. SPAT—Southold Project in Aquaculture Training. Cornell Cooperative Extension of Suffolk Marine Program pp 75.

United Nations (2015) Transforming our world: The 2030 agenda for sustainable development. General Assembly 70 session.

Uren Webster TM, Rodriguez-Barreto D, Martin SAM, Van Oosterhout C, Orozco-terWengel P, Cable J et al. (2018) Contrasting effects of acute and chronic stress on the transcriptome, epigenome, and immune response of Atlantic salmon. Epigenetics 13 (12): 1191–1207.

Utting SD, Millican PF (1997) Techniques for the hatchery conditioning of bivalve broodstocks and the subsequent effect on egg quality and larval viability. Aquaculture 155: 45–54.

Valin H, Sands RD, van der Mensbrugghe D, Nelson GC, Ahammad H, Blanc E et al. (2014) The future of food demand: understanding differences in global economic models. Agricultural Economics 45: 51–67.

Vandeputte M, Dupont-Nivet M, Haffray P, Chavanne H, Cenadelli S, Parati K et al. (2009) Response to domestication and selection for growth in the European sea bass (Dicentrarchus labrax) in separate and mixed tanks. Aquaculture 286: 20–27.

Vu SV, Gondro C, Nguyen NTH, Gilmour AR, Tearle R, Knibb W et al. (2021) Prediction accuracies of genomic selection for nine commercially important traits in the portuguese oyster (Crassostrea angulata) using DArT-seq technology. Genes 12: 210.

Wang C, Chai X, Wang H, Tang B, Liu B (2013) Growth performance of the clam, Meretrix meretrix, breeding-selection populations cultured in different conditions. Acta Oceanologica Sinica 32: 82–87.

Wang J, Qi H, Li L, Que H, Wang D, Zhang G (2015) Discovery and validation of single nucleotide polymorphisms in the Pacific oyster Crassostrea gigas. Molecular Ecology Resources 15: 123–135.

Wang X, Li Q, Yu H, Kong L (2016) Genetic variation assessed in larvae of the Sydney rock oyster. Theriogenology 84: 295–298.

Wang X, Li Q, Yu H, Kong L (2016) Genetic variation assessed in larvae of the Sydney rock oyster. Theriogenology 84: 295–298.
Zhang G, Fang X, Guo X, Li L, Luo R, Xu F et al. (2012) The oyster genome reveals stress adaptation and complexity of shell formation. *Nature* **490**:49–54.

Zheng H, Zhang G, Liu X, Zhang F, Guo X (2004) Different responses to selection in two stocks of the bay scallop, *Argopecten irradians* irradians Lamarck (1819). *Journal of Experimental Marine Biology and Ecology* **313**:213–223.

Zhong X, Feng D, Yu H, Kong L, Li Q (2016) Genetic variation and breeding signature in mass selection lines of the Pacific oyster (*Crassostrea gigas*) assessed by SNP markers. *PLoS One* **11**: e0150868.