Genetic load in marine animals: a review

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Abstract

Marine invertebrates and fish are well known for their remarkable genetic diversity, which is commonly explained by large population size and the characteristic dispersive nature of their early, planktonic life history. Other potential sources of diversity in marine animals, such as a higher mutation rate, have been much less considered, though evidence for a high genetic load in marine bivalves has been accumulating for nearly half a century. In this review, I examine evidence for a higher genetic load in marine animals from studies of molecular marker segregation and linkage over the last 40 years, and survey recent work examining mutational load with molecular evolution approaches. Overall, marine animals appear to have higher genetic load than terrestrial animals (higher dn/ds ratios, inbreeding load, and segregation distortion), though results are mixed for marine fish and data are lacking for many marine animal groups. Bivalves (oysters) have the highest loads observed among marine animals, comparable only to long-lived plants; however, more data is needed from other bivalves and more marine invertebrate taxa generally. For oysters, a higher load may be related to a chronically lower effective population size that, in concert with a higher mutational rate, elevate the number of deleterious mutations observed. I suggest that future studies use high-throughput sequencing approaches to examine (1) polymorphism in genome-scale datasets across a wider range of marine animals at the population level and (2) intergenerational mutational changes between parents and offspring in crosses of aquaculture species to quantify mutation rates.

Key words: dn/ds ratios, high fecundity, larval mortality, life history, mutation, oyster, segregation distortion.

Introduction

Marine fish and invertebrates have long been noted for their remarkable genetic diversity. Population genetic studies using allozyme or DNA markers have repeatedly demonstrated markedly higher diversity in comparisons of marine versus terrestrial species or marine versus freshwater fish (e.g., Smith and Fujio 1982; Ward et al. 1994; Bazin et al. 2006), which suggests something potentially unique about the biology or ecology of marine animals (Williams 1975; Bazin et al. 2006; Harrang et al. 2013; Plough et al. 2016). Explanations for such extreme diversity in marine animals have often emphasized the highly dispersive nature of marine environments (few physical barriers, strong ocean currents) and the long-lived dispersive larval stage of most fish and invertebrate species, which facilitate potentially high gene flow and the establishment of large populations capable of sustaining high levels of allelic or nucleotide variation (e.g., Smith and Fujio 1982; Mitton 1993; Ward et al. 1994; Small et al. 2007; Hellberg 2009).

Though the assumption of large population size may hold for some marine animal species, as a general explanation for high levels of observed diversity overall, it is insufficient for a few reasons (e.g., Harrang et al. 2013). First, features of marine animal life history and reproductive biology, particularly high variance in reproductive success among parents, can dramatically decrease the effective population size of marine species relative to census numbers (Nunney 1995; Hedrick 2005; Hedgcock and Pudovkin 2011). Indeed, a number of recent population genetic studies of fish and shellfish find that effective population sizes of marine species can be orders of magnitude below their census size and thus are not universally large
(e.g., Hedgecock 1994; Turner et al. 1999; Turner et al. 2002; Hedgecock and Pudovkin 2011). Second, population size is only one-half of the equation when considering the expected level of polymorphism in a population at equilibrium. Theta, $\theta = 4Neu$, the expected level of diversity at equilibrium under the neutral theory (Nei 1987), is a function not only of effective population size but also of mutation rate. Thus, a higher rate of mutation in marine animals could also contribute to the high polymorphism observed in marine populations in nature (e.g., Tsagkogeorga et al. 2012); however, this has not been sufficiently examined in the literature.

Previous work on the population genetics of marine animals has revealed important insights about connectivity and adaptive change within high gene flow species (e.g., Hedgecock et al. 2007; Hauser and Carvalho 2008; Hellberg 2009; Gagnaire et al. 2015), but a number of questions remain regarding the biological and demographic factors underlying high genetic diversity in these species. First, is higher diversity in marine animal species associated with a higher mutation rate and higher genetic load? Second, how do the life-history characteristics of marine animals impact diversity and shape how genetic load is expressed in populations? Recent advances in sequencing technology and the availability of genome-scale datasets for non-model marine species are providing novel insights into these questions, such as, the number and effect of mutations segregating in the Pacific oyster genome (e.g., Plough and Hedgecock 2011) or the distribution of effects for recent deleterious mutations in the flat oyster (e.g., Harrang et al. 2013). In this review, I examine the evidence for a higher genetic load in marine animals from the 40+ years of literature on segregation distortion in experimental crosses, which includes a comprehensive survey of recent linkage mapping data from marine (aquaculture) animals. I also examine evidence from studies of natural populations, recapping the heterozygote fitness correlation (HFC) literature in bivalves as well as addressing the small but significant molecular evolution literature for marine animals. In addition to reviewing the literature, I provide a background on the genetic load concept, discuss how newer sequencing technologies are advancing the field, and suggest what the next steps should be to continue advancing research in this field.

**Genetic Load: Definitions and A Brief History**

The origin of the genetic load concept can be traced to Haldane (1937), in which he examined the loss of population fitness associated with mutation-selection balance. Haldane argued that populations routinely harbor individuals with suboptimal fitnesses relative to an expected population maximum and that the number of these less-fit individuals would remain more or less constant despite the action of selection because the allelic variants underlying fitness loss are being continuously replenished via mutation (i.e., mutational load) or because specific combinations of those alleles may be more or less advantageous (i.e., segregation load). Haldane illustrated this mathematically, showing that the expected mean population fitness (at equilibrium) for a single locus with a normal and deleterious allele (where the frequency of the deleterious allele, $q$, is expected to be quite rare, i.e., $q \ll 1$) is: $
abla \approx 1 - 2q$bs, which simplifies to $\nabla \approx 1 - 2\mu$ at mutation-selection balance (see Wallace 1991; Agrawal and Whitlock 2012 for a review of calculations). The important and perhaps surprising result here is that the reduction in mean fitness at a locus depends only upon the rate of mutation and is thus independent of its effect size (i.e., Load or $L \approx \mu$). Under the assumption of no linkage disequilibrium or epistasis, this result can be extrapolated across the genome, and the total mean average fitness, $\nabla_T$, is estimated as the product of mean fitnesses at each locus,

$$\nabla_T = \prod \nabla_i = \prod (1 - 2\mu_i),$$

where $\nabla_i$ is the mean fitness with respect to locus $i$, and $\mu_i$ is the mutation rate for that locus. A similar result was arrived at by Crow (1958) as well as by Muller (1950), in his classic paper on the role of mutation in disease and defect among human populations, which is one of the publications to use the term "genetic load". Crow (1958) would formalize the definition of genetic load as "the proportional decrease in average fitness (or other measurable quantity) of a population relative to the genotype possessing the maximum or optimum value," result in the equation $L = \nabla_{\text{max}} - \nabla_{\text{avg}}/\nabla_{\text{max}}$ or $L = 1 - \nabla$, following the convention that maximum fitness = 1.00.

Overall, this relatively simple but powerful theoretical result (Load = mutation rate) would spur novel work by theorists and empiricists to quantify and compare loads across populations and species, to estimate and compare mutation rates across species (e.g., humans and *Drosophila*), and to consider the theoretical limits of load burden that could be absorbed by populations.

**Inbreeding load—a practical measure of the mutational load across species**

In the decades that followed Haldane’s original paper, the theory of genetic load would be widely tested and debated among empirical and theoretical geneticists (e.g., Wallace 1956, 1970, 1991; Mukai et al. 1972; Agrawal and Whitlock 2012) but it would ultimately recede from the front lines of population genetic research in the 1980s as workers, now armed with more advanced molecular techniques, could more directly estimate genetic variation at the protein and then DNA level. Perhaps the most challenging aspect of the load concept was that it was difficult, in practice, to measure load in natural populations. For example, how does one find and determine the fitness of a mutation-free, “optimal” reference genotype in a natural population, especially when it is very unlikely to exist in the first place? Though largely limited in its practical utility, load theory did provide a mathematical and theoretical framework to estimate the sources or components of fitness reduction compared to some base or “wild” (e.g., heterozygous) population. One of the most powerful experimental avenues for this work was the balanced-lethal system in *Drosophila* (e.g., Greenberg and Crow 1960, Simmons and Crow 1977), in which mutations or entire chromosomes from a wild population are made homozygous identical-by-descent through a series of controlled crosses with specialized stocks, and their viability is compared to heterozygotes or an index of relative heterozygote viability among crosses. The balancer technique (available only for *Drosophila* and a few other model systems; e.g., Steiner 1956; Simmons and Crow 1977; Herman 1978; Edgley et al. 1995) uses strains or stocks with special chromosomes that have recombination-suppressing inversions and morphological markers that allow researchers to assay the viability of the “captured” wild chromosome by the relative frequency of marked (heterozygous) versus non-marked (homzygous) flies in the offspring. Based on the relative viability of different chromosomes from nature made homozygous, the total impact of these mutants could then be partitioned into various components of load (e.g., lethal load vs. detrimental load) and the dominance of mutations (i.e., their effects on heterozygote viability) quantified. The balanced lethal system in *Drosophila* facilitated substantial progress in understanding the magnitude of mutation rates for various classes of mutants, their
dominance, and the relative effects of mild versus lethal mutations on overall fitness (Simmons and Crow 1977).

For researchers working on species that lack a balanced lethal system, mutational (inbreeding) load cannot be examined for entire chromosomes made homozygous identical-by-descent, but crosses among relatives can be carried out to examine the relative reduction in offspring fitness upon inbreeding, that is, Load = 1 – μ_{inbred}/μ_{outbred}. Because many deleterious mutations are largely hidden from selection in the heterozygous state, mating of close relatives exposes the fitness effects of a portion of these recessive mutations in the homozygous state, and the comparison of inbred versus outbred fitness, particularly the viability of early life stages, provides a measure of the recessive mutational load in a given population or species. This approach was formalized into a regression framework by Morton et al. (1956) in calculating the number of lethal equivalents, which were defined as: “... a group of mutant genes of such number that, if dispersed in different individuals, would cause on average one death, e.g., one lethal mutant or two mutants each with a 50% probability of death.” By performing linear regression of log survival across levels of inbreeding, the investigator can estimate the concealed load (the slope, B, or reduction in fitness with increasing inbreeding) and the expressed load (A, the intercept; Malogolowkin-Cohen et al. 1964; Anderson 1992; Lynch and Walsh 1998).

The experimental framework for measuring inbreeding load, primed by the original load theory, has guided much of the empirical work on genetic load over the last half-century and thus, much of the literature reviewed herein. In the following sections, a synthesis of trait- and molecular marker-based studies of inbreeding load provide a comprehensive view of differences among taxa, and then a survey of the molecular evolution literature provides a complementary view of genetic load from investigations of polymorphism in natural populations.

**Trait-based analysis of inbreeding load—lethal equivalents**

Lethal equivalents (LEs) have been measured for a wide variety of plants and animals since Morton et al. (1956) and thus provide an important and robust source of data to compare levels of genetic load across diverse taxa. Lynch and Walsh (1998; Tables 10.4 and 10.6) compiled LE estimates per gamete across a number of species and I have replotted these data, arranging them by broad taxonomic group (e.g., mammals, birds, angiosperms, Drosophila; Figure 1). Included in this figure are novel estimates of LE for a variety of marine fish, invertebrates, and salmon, calculated from published data on inbreeding depression (ID) for early survival or growth rate using a maximum likelihood method (Kalinowski and Hedrick 1998) or molecular marker-based estimates of the inbreeding load (see below; Supplementary Table S1). The literature search for additional ID studies was extensive, selective, and representative of a variety of marine animals, but was not exhaustive. Studies were only considered if they (1) provided raw or averaged data and (2) measured fitness of early life-history traits in lab-based experiments (see Supplementary Table S1, File S1 for more details). Estimates of LEs derived from molecular marker data (oysters) were made by counting the number of independent lethal loci per haploid parental genome within families, using partial (Launey and Hedgecock 2001) or whole-genome linkage data (quantitative trait locus [QTL] mapping of distortion; Plough and Hedgecock 2001, see next section). Specifically, LEs were tallied as the sum of the selection coefficients (often 0.9–1.0) calculated for each independent lethal or nearly lethal mutation detected. For example, two loci, each with a selection coefficient of 0.80 together would count as 1.6 LE. Note that molecular-based approaches may substantially underestimate the number of LEs because only the most lethal mutations are detected and the cumulative effect of many weakly deleterious alleles is likely to be missed. Furthermore, only ~25% of the recessive lethal alleles carried by a grandparent become homozygous identical by descent in the F2 generation (the typical cross examined) and thus many lethal alleles potentially remain hidden in the heterozygous state.

Figure 1 shows that marine invertebrates have the widest range of LEs, but appear to have the highest estimates among animal taxa. Within the invertebrate grouping, oysters have the highest estimates (top four data points; Supplementary Table S1), some of which are molecular-based (triangles); however, there is also an estimate of ~0 (no ID for larval survival in the Eastern oyster; Mallet and Haley 1983). The two trait-based data points for oysters come from the Eastern oyster *Crassostrea virginica* and it is not clear why such different levels of inbreeding load would be observed within the same species (0 vs. 6.02; Longwell and Stiles 1973; Mallet and Haley 1983). The molecular-based estimates, on the other hand, appear to group more closely (range 2.67–4.02; Figure 1, Supplementary Table S1). Overall, LE estimates for oysters appear to be most similar in magnitude to conifers, which have the highest estimates across the taxa examined. Long-lived conifers share a number of life-history and population biology characteristics with bivalves, such as high early mortality, early expression of inbreeding load, and high...
fecundity, which have been argued to result in a higher load (Williams 1975; Carr and Dudash 2003; Plough and Hedgecock 2011). Though the differences in load across taxa are intriguing, it is important to acknowledge how varied the data are in terms of quality, quantity, and approach, particularly among the studies of marine animals (Supplementary Table S1). For example, some LE estimates are based on fitness data measured across many families and at 4 or more inbreeding levels, whereas other studies use just 2 levels of inbreeding (outbred and full sib mating, $f = 0.25$) from a single crossing experiment. Moreover, fitness is measured for a variety of related but very different traits across studies, including larval viability, egg hatching rate, and relative genotype viability in the molecular-based studies. Thus, it is quite difficult in practice to directly compare load or LE estimates among taxa. Nevertheless, these data provide a rough view of the differences among taxonomic groups, and suggest that oysters and bivalves may have a higher inbreeding load than other animals.

**Molecular marker-based measures of genetic load in marine animals**

Recent analyses of inbreeding load in marine animals have turned to molecular marker-based methods to uncover the number and effect of deleterious mutations carried by a set of parents. Among marine species, most available data come from studies of bivalves, perhaps due to the ease with which bivalves can be crossed and reared in the laboratory and the early adoption of molecular marker technologies (allozymes) among workers in the field. Similar to the LE approach (Morton et al. 1956), molecular-based estimates of inbreeding load require crosses to be made between related individuals (often full siblings from an F$_1$ cross), to expose the effects of homozygosis for identical-by-descent deleterious recessive alleles on offspring viability. However, the magnitude of load is measured not as phenotypic decline over successive levels of inbreeding, but as the deviation from expected Mendelian ratios of inheritance (segregation distortion) in the counts of genotypes at one or more molecular markers segregating in the offspring (e.g., Mallet et al. 1985; Bierne et al. 1998; Launey and Hedgecock 2001; Carr and Dudash 2003; Plough and Hedgecock 2011). The deficiencies of genotypes (relative to Mendelian expectation) at a given point in offspring development are either the result of differential survival of gametes (gametic selection) or zygotes (zygotic selection), though temporal studies of the Pacific oyster indicate that most if not all of the early selection is zygotic (see below; Plough and Hedgecock 2011). The observation of differential mortality among genotypes (genotype-dependent mortality) implies viability selection either at the focal marker or, more likely, at a closely linked, deleterious locus, the magnitude of which depends on the recombination distance ($c$) between marker and mutation and the strength of selection ($s$) against that mutation (two-locus linked selection model; Hedrick and Muona 1990; Launey and Hedgecock 2001).

**Marker segregation data in inbred crosses**

Distorted segregation ratios were first reported for marine bivalves in the 1970s from inbred crosses that produced substantial deficiencies of homozygous genotypes in offspring, indicating selection against recessive deleterious alleles (e.g., Wada 1975; Wilkins 1976; Beaumont et al. 1983). Various explanations for the observed distortions were proposed, including gametic selection (e.g., meiotic drive) and zygotic selection (e.g., Mallet et al. 1985; Foltz 1986), but null alleles could generally be ruled out because they could easily be detected as unexpected genotypes segregating in the offspring (e.g., Plough and Hedgecock 2011). Over the next 4 decades, similar observations of significant and sometimes extreme levels of segregation distortion were made primarily in inbred, but also outbred, crosses for a variety of marine bivalves including the blue mussel, eastern oyster, Pacific oyster, and European flat oyster (e.g., Beaumont et al. 1983, 1988, 1990; Gaffney and Scott 1984; Mallet et al. 1985; Foltz 1986; Hu et al. 1993; Hu and Foltz 1996; McGoldrick and Hedgecock 1997; Bierne et al. 1998; Launey and Hedgecock 2001; Bucklin 2003, Lallias et al. 2007a, 2007b; Lallias et al. 2009; Sauvage et al. 2010; Plough and Hedgecock 2011; Harrang et al. 2015). The proportion of markers exhibiting segregation distortion varies across studies, with some experiments exhibiting distortion in greater than 75% of markers assayed (Bierne et al. 1998), but commonly, ~15–50% of markers assayed within a family are distorted. Moreover, the frequency of distortion is exacerbated in parents from related aquaculture stocks or with higher levels of inbreeding, which connects the observed load and distortion from molecular data to inbreeding level (e.g., Wilkins 1976; McGoldrick and Hedgecock 1997).

Interestingly, segregation distortion is also observed in crosses of unrelated parents, though the patterns of selection are quite different and the results of outbred crossing experiments have been published less frequently. Mallet et al. (1985) was one of the first studies to report significant heterozygote deficiencies in the offspring of pair-crossing experiments of wild-caught blue mussels *Mytilus edulis*. After considering a number of explanations for heterozygote deficiencies, the authors concluded that they most likely were explained by zygotic selection. A more recent study of segregation in the offspring of four outbred pair-crosses of the Pacific oyster also revealed substantial deficiencies of heterozygous genotypes in ~40% of typed markers (Plough et al. 2016). That wild pair-crosses show characteristic deficiencies of heterozygous genotypes implies a load affecting early viability that is expressed in the absence of inbreeding, which would not be expected if most harmful mutations are primarily recessive (only expressed when made homozygous identical-by-descent). Partial dominance was demonstrated for a number of deleterious alleles identified in inbred Pacific oyster families (see below; e.g., Launey and Hedgecock 2001, Plough and Hedgecock 2011), which suggests that at least some portion of the load may reduce fitness in outbred crosses.

Researchers using molecular markers have also revealed details about inbreeding and genetic load in natural populations of marine bivalves through studies of heterozygosity–fitness correlation (HFC; e.g., David 1998, Szulkin et al. 2010). Correlations between multilocus heterozygosity (MLH; the proportion of heterozygous markers for a given individual) and fitness traits (e.g., growth, size) have been found in a variety of marine bivalves, which signals some level of inbreeding variance or drift (non-random mating) within a population (e.g., Singh and Zouros 1978; Koehn and Shumway 1982; Koehn and Gaffney 1984; David 1998). Though not as powerful an approach to quantify genetic load as genotyping markers in controlled crosses, HFCs in natural populations have the advantage of yielding estimates of both the potential genetic load and its actual impact in a population and can be performed in species for which crossing experiments may be impractical. The fairly consistent presence of HFCs across marine bivalves, though often of quite small magnitude, was initially surprising because marine bivalves are generally thought to have large randomly mating populations with little opportunity for inbreeding. Although confusion with HFC terminology and underlying theoretical mechanisms, as well as issues
related to power and significance have limited the scope of this literature in recent years (e.g., Szulkin et al. 2010), the population-level HFC data in bivalves does support the idea that these species may have an elevated genetic load.

**Advances in mapping of load in marine bivalves**

Subsequent studies of marker segregation in marine bivalves have employed a greater number of markers and linkage or QTL approaches to more precisely estimate the number, location, and effect of deleterious mutations in the genome. Biern et al. (1998) and Launey and Hedgecock (2001) made the first systematic attempts to estimate the total number of harmful recessive mutations in the genome, accounting for the proportion of the genome assayed by independent marker segregations and linkage information among markers. With data at 7 microsatellite markers, Biern et al. (1998) estimated between 15 and 38 deleterious mutations, whereas Launey and Hedgecock (2001), using 19 markers and a linked-selection model (Hedrick and Muona 1990), estimated between 8 and 14 mutations in the wild (grandparent) founders. With 1st-generation microsatellite linkage maps, Plough and Hedgecock (2011) and Plough (2012) employed a QTL mapping approach to scan the genome for deleterious mutations (termed viability loci or vQTL in the QTL mapping context), using the viability model of Luo and Xu (2003). These studies revealed a similar number of mutations (11–15 vQTL) scattered throughout the genome, some of which had significant dominance effects. By genotyping oysters throughout the life cycle (e.g., larval, juvenile, adult), Plough and Hedgecock (2011) also demonstrated that this load was first expressed (i.e., caused significant viability selection) during the larval stages and at metamorphosis to the juvenile stage, but was not gametic. In other words, inbreeding load was expressed early in the life cycle. Further, Plough (2012) showed that diet influenced the fitness effect of individual mutations. Rearing offspring in a more stressful, low-quality diet produced a significant increase both in selection against and dominance of deleterious alleles at vQTL. Finally, these studies attempted to estimate the total cost of deleterious mutations in the partially inbred offspring by calculating the total mortality associated with viability selection at all identified deleterious loci. By first testing for and ruling out interactions (epistasis) among vQTL, Plough and Hedgecock (2011) and Plough (2012) were then able to calculate the cumulative, egg-to-adult genetic mortality from the product of average relative survival across all independent viability loci. In these experiments, mortality attributed to viability selection ranged from 85% to 99% across the families examined. Thus, the real-world effect of high inbreeding load in marine bivalves is to significantly reduce the early survival of offspring from inbred pair-crosses.

**Load in Outbred Crosses of Marine Animals: Aquaculture Data**

The previous paragraphs have summarized numerous reports of segregation distortion in the progeny of inbred crosses of marine bivalves, and some data on HFCs in natural populations, with much work focused specifically on mussels or oysters. These studies establish that marine bivalves may possess a very high load of partially recessive deleterious mutations; however, they provide less information on the broader question of whether genetic load is higher across marine animals in general, and whether or not this load has serious fitness or adaptive consequences in natural populations. Fortunately, a great wealth of marker segregation data for marine animals, including fish and crustaceans, can be found in the recent aquaculture literature in the form of genetic linkage maps for outbred (F1) families using hundreds to thousands of markers. These maps are usually the first step in generating genomic resources for a species and are used for QTL mapping of economically important production traits. The number of genome-wide marker datasets for linkage analyses has increased substantially over the last decade due to the availability of novel marker technologies and high-throughput genotyping approaches that make large-scale genotyping of non-model species possible (e.g., Luikart et al. 2003; Davey et al. 2011). Examining the expression of genetic load (i.e., viability selection) in the progeny of outbred crosses of marine animals begins to address the question of how a high load in marine animals might directly impact natural population fitness (early offspring mortality), at least in an experimental context. If some proportion of the load in these species is only partially recessive, as has been shown in Pacific oysters (Launey and Hedgecock 2001; Plough and Hedgecock 2011; Plough 2012; Plough et al. 2016) it would be detected as a significant viability selection in the outbred offspring of experimental pair-crosses.

To compile and summarize available genome-wide segregation data from marine animals, I performed a Web of Science (WOS) search for articles with titles containing words about segregation/linkage mapping and names of marine animal groups (e.g., clam, crustacean or sea bass; see Supplementary Table S2 for the complete list of studies and more details of the search criteria). The initial search resulted in the identification of 89 published research articles, ~20 of which were removed because they examined segregation data from inbred (F2) families, or were off-topic (non-marine animal, or no report of linkage or segregation data). An additional 12 linkage studies of marine fish and shellfish not found in the WOS search were added; these studies were missed initially because they had specific common or scientific names in the title. Finally, a maximum of 3 studies per species were retained to reduce redundancy. Within bivalves, a few species had many published linkage studies (e.g., Pacific oyster, Zhiqong scallop, and Bay scallop), so culling to 3 per species reduced the apparent number of studies in this group. Overall, 67 studies from 43 species were examined, comprising a variety of marine animal groups including crustaceans, fish, echinoderms, gastropods, and bivalve mollusks. The primary reporting metric was the proportion of distorted marker loci within a cross or family, significant at the nominal, \( \alpha = 0.05 \), level. The tally of distorted markers was done at \( \alpha = 0.05 \) because not all studies explicitly stated what significance threshold they used to report or remove loci and in the absence of any information, it was assumed that significance threshold was at \( \alpha = 0.05 \) level. Some studies only reported the number of distorted markers after Bonferroni or FDR correction (noted in Supplementary Table S2); so, estimates may be conservative in some cases.

Two major trends emerged from the review of segregation distortion data in outbred crosses of marine animals. First, marine bivalves appear to have the highest proportion of distorted markers among the marine animal groups examined (Figure 2, Supplementary Table S2). The average proportion of distorted markers reported for bivalves was 24.74%, whereas averages for crustaceans and fish were lower (18.2% and 14.3%, respectively). The difference in the proportion of distorted markers between fish and bivalves was statistically significant (Wilcoxon rank sum test; \( P = 0.034 \)). Interestingly, there was a wide range in the level of distortion reported across bivalve species and, in some cases, substantial variation within species (Figure 2, Supplementary Table S2). The highest proportion of distorted markers among bivalves was...
reported for the Bay scallop (60%; Wang et al. 2012), the Pacific oyster (27–49%; Li and Guo 2004; Hedgecock et al. 2015; Plough et al. 2016) the flat oyster (32%; Lallias et al. 2007a) and the Zhikong scallop (35%, 6%, and 30%—Wang et al. 2003, 2005; Zhan et al. 2009, respectively). The high variability in distortion among studies of Zhikong scallop is not easy to explain. In the two studies with the most contrasting results (35% and 6% distortion), AFLP (amplified fragment length polymorphisms) markers were used and similar numbers of progeny were examined (51 and 60 in Wang et al. 2003 and 2005, respectively), so differences in power or marker class do not explain this. No effect of marker type on distortion was detected across the studies examined. Generally, the proportion of distorted markers was somewhat similar across studies of the same species (e.g., 10–15% in Asian sea bass, Wang et al. 2007, 2015 and 7–21% in Tiger shrimp, Staelens et al. 2008; Baranski et al. 2014), which would be expected if the level of distortion was associated with the biology of the species and not with marker type or other sources of error. However, some variation among estimates within species could be attributed to different husbandry techniques or rearing conditions, the latter of which has been shown to affect the magnitude of segregation distortion in Pacific oysters (Plough 2012). Among crustaceans, studies of Pacific white shrimp showed the greatest proportion of distorted loci (39%; Zhang et al. 2007) and for fish, the half smooth-tongued sole demonstrated the largest proportion of distorted markers (up to 33%, Liao et al. 2009), though one estimate was only 4% (Diopere et al. 2014). A few fish had very low or negligible numbers of markers with segregation distortion (white croaker, Dor et al. 2014; red drum, Hollenbeck et al. 2015; Japanese eel, Kai et al. 2014). Unfortunately, segregation results were reported for only a single echinoderm (of 2 studies retrieved; Yan et al. 2013), so it is difficult to infer any patterns for this group. Many data points were also missing for marine fish despite a large number of linkage studies examined (18 out of 38 studies; Figure 2, Supplementary Table S1). Clustering of distorted loci on specific linkage groups was reported for 4 fish species, (Asian sea bass, Wang et al. 2007; brill, Hermida et al. 2014; smooth sole, Liao et al. 2009; red drum, Portnoy et al. 2010) but was not addressed explicitly in most studies. Overall, segregation distortion in wild crosses appears to be lower in fish and crustaceans compared with bivalves (Figure 2).

The second major finding from the review of linkage data is that segregation distortion routinely goes unreported and/or distorted markers are removed before analysis in studies of fish and crustaceans, but not bivalves. For example, approximately half of the studies reviewed for fish and crustaceans failed to report segregation distortion or removed distorted markers before analysis (Figure 3). This was particularly pronounced in studies with single nucleotide polymorphism (SNP) data generated via next-generation sequencing methods (i.e., RADseq), in which distorted markers were treated as genotype error or marker artifact, and culled at the same stage of data filtering that low-coverage markers or markers with excessive missing data were removed (e.g., Palaiokostas et al. 2015, Wang et al. 2015). The number of distorted markers that were removed was reported in some cases (e.g., Hubert et al. 2010) but often, a quantitative estimate of segregation distortion was not possible (Figure 3). This is in sharp contrast to marker segregation studies of bivalves, for which the research community is highly attuned to issues surrounding non-Mendelian inheritance and segregation distortion was always addressed or reported explicitly. In some cases, removal of markers with segregation distortion may be necessary as distortion can produce erroneous marker order and map lengths (e.g., Lorieux et al. 1995, Lorieux 2012; Hedgecock et al. 2015). Moreover, cases of segregation distortion could be caused by systematic genotype errors, and the culling of markers with questionable inheritance patterns may be appropriate (e.g., Pompanon et al. 2005). Still, it appears that a number of studies do not treat instances of segregation distortion as relevant biological phenomena, and a reporting bias may depress observations in the literature for marine fish.

Overall, based on the 67 studies examined, segregation distortion appears to be a common feature in crosses of marine animals, particularly bivalves, which implies a potentially large load of mutations affecting early viability. However, while bivalves appear to

![Figure 2. Proportion of markers exhibiting segregation distortion (z = 0.05) in linkage mapping studies of marine animals in outbred crosses. Red bars represent the mean of the individual observations for each group.](https://example.com/figure2)

![Figure 3. Proportion of studies that reported segregation distortion results, plotted by broad marine animal group.](https://example.com/figure3)
have the highest levels of segregation distortion among marine animals, high variation within this group and the lack of linkage or segregation studies in most other marine taxa makes it difficult to draw any definitive conclusions at this point.

### Estimating Genetic Load from Molecular Data in Natural Populations

An alternative approach to estimate the influx of deleterious mutations (mutational load) at the population level is to quantify the rates of diversity in different classes of polymorphisms (e.g., synonymous vs. non-synonymous mutations) in natural population samples. Polymorphism of non-synonymous mutations is expected to be lower relative to synonymous or silent mutations, assuming amino acid changes generally have negative effects on fitness and are removed by purifying selection (depending on population size among other parameters; Kimura 1983). Thus, by examining the ratio of diversity (e.g., nucleotide diversity, $\pi$; Nei 1987) at amino acid changing versus silent SNPs ($dn/ds$ ratios), one can infer how selection is acting on mutations in a population (e.g., Fay et al. 2001; Eyre-Walker et al. 2006; Eyre-Walker and Keightley 2007). Further, by categorizing SNPs according to their minor allele frequency (e.g., low, moderate, or common), the ratio of non-synonymous to silent diversity within frequency classes can be used to infer the proportion of silent, weakly deleterious, and strongly deleterious mutations (Fay et al. 2001; Harrang et al. 2013). Averaged across many or all coding regions in the genome, the $dn/ds$ ratio can tell the investigator something about the mutation pressure in a given species: organisms with higher load (whatever the cause) would be expected to have a greater number of weak to moderately deleterious mutations segregating within populations (excess rare variation), which would manifest as a higher $dn/ds$ ratio (Fay et al. 2001). It is important to note that $dn/ds$ ratios were originally developed in a phylogenetic context to estimate the ratio of fixed non-synonymous to synonymous changes between highly diverged lineages (species), the magnitude of which (e.g., above or below unity) is used to infer the type of selection acting on that particular locus (e.g., Kimura 1977; Goldman and Yang 1994). In a population genetics context, where polymorphisms may not be fixed among populations, the interpretation of the $dn/ds$ ratio is not straightforward for inferring selection at a particular gene (e.g., Kryazhimskiy and Plotkin 2008; Mugal et al. 2014). However, in the context of inferring the fitness effects of new mutations and genetic loads, these ratios can be used in a comparative framework to examine the distributions of mutation fitness effects.

Studies of polymorphism with multi-marker $dn/ds$ datasets (at least 10s of loci) are relatively scarce for marine animals, but the few datasets that are available suggest that a high load of weakly selected mutations may be present in these species. In an analysis of polymorphism across 35 gene regions in the flat oyster *Ostrea edulis*, Harrang et al. (2013) found a relatively modest silent diversity ($\pi_{ds} = 0.0067$), but a high non-synonymous diversity ($dn = 0.0025$), which produces a remarkable $dn/ds$ ratio of 0.38, the highest such estimate among animals currently reported in the literature. The $dn/ds$ ratios appear to be somewhat similar across other oyster species, though perhaps are not as extreme as *O. edulis*. Using data from 41 coding genes spanning $\sim 10.5$ kb of the Pacific oyster *Crassostrea gigas* genome, Sauvage et al. (2007) estimated an average $dn/ds$ ratio of 0.16; silent diversity ($\pi$) was relatively high ($ds = 0.035$), but non-synonymous diversity was also rather substantial ($dn = 0.0056$). In another study of SNP variation in *C. gigas*, Kim et al. (2011) estimated a 10-fold higher silent versus non-synonymous ($\pi$ silent = 0.0254, $\pi$ non-synonymous = 0.0023), resulting in a $dn/ds$ ratio of 0.09. It is important to note that these 2 studies of *C. gigas* used hatchery stocks (Sauvage et al. 2007) or a mixture of wild and hatchery oysters (Kim et al. 2011), thus these estimates may not reflect the true level of polymorphism in a natural oyster population. The relatively high $dn/ds$ values observed in bivalves contrast with data from another highly polymorphic marine invertebrate, the sea squirt *Ciona*, which has similar levels of non-synonymous diversity, but very high silent diversity (e.g., Small et al. 2007; Tsagkogeorga et al. 2012) resulting in much lower $dn/ds$ ratios (*Ciona intestinalis* sp. B = 0.05; Tsagkogeorga et al. 2012). Overall, bivalves (or at least the oyster species examined so far) appear to have a high segregating load of deleterious mutations, reflected by relatively higher non-synonymous diversity.

To put the high non-synonymous diversity of marine bivalves into biological context, it is instructive to compare these data to similar indices from a variety of other marine and non-marine taxa. In their analysis of non-synonymous diversity in the flat oyster *O. edulis*, Harrang et al. (2013) compiled and compared diversity data ($dn/ds$ values) from the plant, terrestrial animal, and model animal literature. These data are replotted here in Figure 4 by broad taxonomic group, with some additional data points for marine animals included. Estimates of $dn$ and $ds$ ($\pi$) for the blue crab *Callinectes sapidus* are taken from Yednock and Neigel (2014) and data from the aforementioned *C. gigas* studies are also plotted (Sauvage et al. 2007; Kim et al. 2011). Overall, it appears that marine animals (red squares) group more closely with terrestrial plants (green circles) than they do with other terrestrial animals (blue triangles) in terms of $dn/ds$ diversity data (Figure 4). Plotting simple least square regression lines of $dn$ versus $ds$ for each group (dashed lines colored according to group), slopes appear to be steeper for the plants and marine animals compared with terrestrial animals, which reflects greater non-synonymous diversity relative to silent diversity. Bivalves appear to have particularly elevated average non-synonymous diversity. One of the polymorphism datasets for *C. gigas* shows the highest value of $dn$ across all taxa (0.0056; Sauvage et al. 2007) and the $dn/ds$ ratio for the flat oyster *O. edulis* is the largest among all other animal species examined (0.38). However, data from the blue crab show a much more modest level of non-synonymous diversity, and data for *C. intestinalis* show a very large silent diversity and much lower relative non-synonymous diversity. Based on these data, bivalves appear to stand out somewhat among marine animals, but marine animals do group fairly well together overall, with a slightly higher slope ($dn/ds$) compared to terrestrial animals. As noted above, however, data from the 2 studies of Pacific oyster SNP diversity may be biased by the inclusion of hatchery stock, and the data for blue crab are based on just 5 loci (Yednock and Neigel 2014). Thus, more studies of marine animals with large or genome-scale datasets will be needed before any definitive conclusions about diversity can be drawn from these data.

One possible explanation for the extreme $dn/ds$ ratios observed in oysters, which imply a larger mutational load for this species, is a chronically low effective population size. Given that purifying selection is expected to be more efficient in larger populations (Nei 1987), populations with lower effective size may retain weakly deleterious variants at higher frequency for a longer duration, thus increasing the $dn/ds$ ratio. Indeed, a number of population genetic studies of oysters have demonstrated sweeps of reproductive success and a much lower effective population size than expected given census estimates (e.g., Hedgecock 1994; Hedgecock et al. 2007;
Taris et al. 2007; Lallias et al. 2010). A lower effective size and substantial mutation pressure could drive a larger number of weakly deleterious mutations to segregate in populations, increasing the dn/ds ratio. For *O. edulis*, this idea is supported by examining diversity across frequency classes of SNPs, which reveals that approximately 33% of segregating mutations are weakly deleterious (cf. estimates for humans and *Drosophila*, ~20–23%; Fay et al. 2001; Harrang et al. 2013).

An alternative explanation for the high dn/ds ratios in oysters is that the populations examined are actually very far from demographic equilibrium and that the excess of low frequency deleterious variants is due to demographic forces such as a bottleneck or recent population expansion. While these are valid hypotheses, Harrang et al. (2013) did not detect any systematic deviation from mutation–drift equilibrium (Wright–Fisher model) in diversity-based tests such as Tajima’s D, thus, a demographic explanation may not be sufficient. Moreover, oysters (and other marine animals) may not adhere to assumptions of the Wright–Fisher model due to their high fecundity life-history strategy, which can result in a high variance in reproductive success among parents (Eldon and Wakeley 2006, 2009; Sargsyan and Wakeley 2008; Hedgcock and Pudovkin 2011; Harrang et al. 2013). Alternatives to the Wright–Fisher or Kingman coalescent model framework that account for multi-fucrating genealogies and multiple merger coalescent events may better reflect the reproductive biology of highly fecund marine animals (Eldon and Wakeley 2006; Steinrücken et al. 2013; Tellier and Lemaire 2014). Under the multiple merger coalescent, an excess of high-frequency and low-frequency variants (singletons) is actually expected at equilibrium and may not be a true signal of population bottleneck (e.g., Hoban et al. 2013) or recent expansion as would be indicated by a negative Tajima’s D under a standard coalescent model. Recent population genetic analyses of the putatively selected *Ckma* locus in Atlantic cod using a multiple merger coalescent framework demonstrated a much better-fit than the Kingman coalescent, modeling the excess of singletons and capturing the relatively fast coalescent timescales expected (Árnason and Halldórsdóttir 2015).

Overall, examining molecular diversity data in a limited number of marine animals suggests that bivalves (oysters) may have elevated levels of non-synonymous diversity compared with other marine and terrestrial taxa. However, diversity data are quite variable within groups and even within the same species (i.e., the Pacific oyster), which highlights the need for more datasets. Potential differences in effective population size among marine invertebrate species may in part explain differences in silent (neutral) diversity (e.g., *Ciona* spp. vs. *O. edulis*), and the lower effective size of the oyster species examined may contribute to the increased number of low frequency deleterious variants segregating in their populations. However, until more large-scale (e.g., tens to hundreds of genes) datasets of molecular diversity are available for marine invertebrates and fish species, the molecular evolution literature will be too limited to draw any major conclusions about differences in load among animal or marine taxa.

**Summing It Up: A Higher Load in Marine Animals?**

Reviewing 40+ years of literature on segregation distortion in experimental inbred crosses of marine bivalves, genome-wide linkage datasets from a range of marine aquaculture species, and empirical molecular evolution literature for marine animals, paints an interesting but largely incomplete picture of genetic load in marine animals. Overall, genetic load appears to be higher in marine versus terrestrial animals, but a general lack of data for many marine taxa (e.g., echinoderms), substantial variation within taxa, and a possible reporting bias of segregation distortion in studies of marine fish (Figure 3), complicates the emergence a clear trend from the available literature. What does emerge, however, and quite clearly, is that marine bivalves, and perhaps oysters in particular, stand apart from most marine species in their apparent level of mutational load. Marine bivalves consistently show higher inbreeding load, higher segregation distortion, and, from the few datasets available, elevated non-synonymous variation relative to neutral diversity (Sauvage et al. 2007; Kim et al. 2011; Harrang et al. 2013). Marine bivalves have received far greater and focused attention on topics like genetic load, segregation distortion, and molecular diversity compared with other marine animals, so we may yet see similar patterns emerging in other marine species groups as more studies are conducted. The finding of a high load in marine bivalves is consistent with predictions made by Williams (1975) in his "elm-oyster" model, in which he drew parallels between the population biology of oysters and long-lived, highly fecund plants. Williams hypothesized that the high and variable early mortality observed in these species resulted in part from a high segregating load. Indeed, across-species comparisons of LEs (Figure 1) and dn/ds ratios (Figure 4) illustrate that bivalves group more closely with plants and are rather distinct from terrestrial animals and even other marine animals.

**Potential origins of a high load in marine bivalves**

Based on our understanding of marine bivalve biology and life history, two major factors could explain the higher load observed in these organisms: (1) chronically low-effective population size driven
by a sweepstakes life history and (2) substantially elevated mutation pressure. These potential causes are not mutually exclusive and, in nature, may interact to increase the load observed in bivalves. Considering effective size and life history first, the idea is that a chronically lower effective size \( (N_e) \) could allow deleterious mutations to be retained for longer time periods or at higher frequencies than would be expected if the \( N_e \) was larger (closer to census size, \( N_c \)), where selection would be more efficient at removing deleterious mutations (Nei 1987). Reduced effective size in marine bivalves could be driven by characteristic features of their life history, such as prolific fecundity and substantial mortality of the larval stages, which can result in high variance in reproductive success among parents and a reduction in the effective population size relative to census size (e.g., Hedrick 2005; Hedgecock and Pudovkin 2011). Indeed, a lower relative effective size and evidence of sweepstakes reproductive success have been documented in multiple oyster species (e.g., Hedgecock 1994; Hedgecock et al. 2007; Taris et al. 2007; Lallias et al. 2010; Hedgecock and Pudovkin 2011). Thus, perhaps high fecundity, which is associated with low and variable recruitment in marine fish as well (e.g., Rickman et al. 2000), could be a driver of low effective size that may contribute to higher genetic load in these species. Fecundity may also affect the mutation rate more directly. Plough et al. (2016) hypothesized that the very large number of meiosis required to produce millions of eggs in oysters could increase the likelihood of clustered germ line mutations that would elevate the initial frequency and perhaps residence time of new deleterious mutations (similar to the male-driven evolution concept; e.g., Ellegren and Fridolfsson 1997; Makova and Li 2002). Support for this idea will require laboratory experiments to assess more directly the nature and rate of mutational change in high fecundity species.

If a chronically reduced effective size driven by a high fecundity life history is linked to higher load in marine bivalves, then we might expect a similar phenomenon in other highly fecund marine animals such as marine fish. This idea does not appear to be strongly supported based on the data in this review, however. Whereas many of the marine fish with linkage or segregation data are highly fecund (e.g., Atlantic cod, turbot, red drum), they generally demonstrate much lower levels of segregation distortion than bivalves (Figure 2, Supplementary Table S2). For example, Atlantic cod has fecundity in the millions (1–7 million eggs/female; Pinhorn 1984; Wrobleswksi et al. 1999) and thus high potential for sweepstake reproductive success, but exhibits minimal segregation distortion (Hubert et al. 2010). Similarly, red drum Sciaenops ocellatus has very high fecundity (tens of millions of eggs/female; Overstreet 1983), shows some evidence of sweepstakes reproductive success (Turner et al. 1999), and has a reduced effective population size (Turner et al. 1999, 2002), but exhibited almost no segregation distortion in crosses (1–5%; Portnoy et al. 2010; Hollenbeck et al. 2015). To the extent that segregation distortion in wild pair-crosses is indicative of the magnitude of genetic load for a given species (see next section and Plough et al. 2016), the lack of distortion in a number of highly fecund fish suggests that fecundity may not be a primary driver of genetic load, or at least not the only factor. However, the marker segregation and genetic mapping literature for marine fish (from which this inference is primarily drawn) appear to be somewhat biased toward non-reporting or culling of distorted markers, so more data is needed before we can conclusively rule out fecundity as a major factor underlying elevated genetic load in marine animals.

The other potential explanation for high genetic load observed in marine bivalves is a much higher mutation rate relative to other terrestrial and marine animals. This explanation takes us back to the beginning of the review, where it is suggested that the relatively higher genetic diversity observed in marine animals might be driven not just by large population size but also by an increased rate of mutation. Though there is currently very little empirical data on mutation rates for marine bivalves, recent work in the Pacific oyster suggests that the mutation rate in this species may be elevated. After finding high levels of segregation distortion in a study of outbred crosses of the Pacific oyster, Plough et al. (2016) attempted to estimate the mutation rate that would account for the frequency of numerous, partly dominant deleterious mutations in the families examined (~7.25 lethal alleles per parent). Using classical, mutation-selection equilibrium theory for partially dominant lethal alleles, they estimated a per gene mutation rate of ~3.6 × 10^{-4} or a genomic mutation rate, \( U \), of 1.81, which is approximately 90 times that of Drosophila, but quite similar to estimates of the mutation rate in conifers (10^{-4} to 10^{-5}; Lande 1994). This estimate establishes, at the very least, the biological plausibility of a mutation rate that would account for the presence of the high load observed in wild oysters. Nevertheless, an empirical estimate of mutation rate will be needed for oysters and other marine bivalves to verify this.

As to the potential source(s) of elevated mutation pressure in oysters (and possibly other marine bivalves), one can only speculate at this point, but transposable element (TE) activity appears to be an intriguing possibility. Novel classes of active TEs unique to bivalves have been reported previously (e.g., Gaffney et al. 2003) and the draft genome of the Pacific oyster revealed abundant repetitive sequences and active transposable elements, which could shape genome variation and contribute to high rates of structural variation and polymorphism observed in oysters (Zhang et al. 2012). The potential effect of TEs on indel variation in the Pacific oyster genome may be profound. Re-sequencing of a wild oyster aligned to the reference genome identified thousands of deletions 100 bp or greater, more than 80% of which overlapped with known transposable elements (Zhang et al. 2012). Thus, TE activity could contribute to an increased rate of large-effect deleterious mutations in critical genes or regulatory regions. Future work examining the relationship between TE activity and proximal viability mutations and segregation distortion could be a very fruitful area of future research on this topic.

**Alternative explanations**

Finally, it is important to consider some alternative interpretations of the widespread segregation distortion observed in crosses of marine bivalves, the analysis of which forms the basis of much of the inference of high load in this review. As summarized in previous sections, a high mutational load in bivalves can be inferred from observations of substantial deviations from Mendelian inheritance (segregation distortion) at markers across the genome, which implies fitness or viability differences among genotypes due to selection against linked deleterious mutations (e.g., Plough and Hedgecock 2011). Yet, while this explanation appears to be supported by a number of studies (e.g., Bierne et al. 1998; Launey and Hedgecock 2011; Plough et al. 2016), other types of selection that are related to extrinsic or environmental factors may be important as well. One intriguing possibility is that the selection observed in the offspring actually reflects a dynamic of local adaptation or the lack of local adaptation for a given set of genotypes. Bivalves have a widely dispersing larval stage that promotes high gene flow among populations, but once settled as juveniles, individuals are fixed in a certain
location for the remainder of their life and must contend with local or highly variable environmental conditions. Thus, perhaps the selection that we are observing in the early life stages is related to fitness differences or fitness mismatch between the parental environment and the offspring environment. A modeling study by Yeaman (2015) found that local adaptation could develop in populations with high gene flow, driven primarily by alleles of small effect that could withstand "swamping" (high gene flow) and cause divergence in different environments, given sufficient genetic variance and genetic redundancy in the underlying trait. Though the selection observed in cross-experiments of bivalves is quite strong and involves alleles of rather large effect, the idea that incompatibilities between genotypes and the environment could drive the observed patterns of selection is interesting. If distinct genotypes are more or less fit in different environments, one could hypothesize that the production of such a diverse array of genotypes in the offspring of high fecundity bivalves is a response to the variable and unpredictable nature of recruitment and the eventual "fixed" environment of the settled juvenile, that is, a "bet-hedging" strategy. Though this is difficult to demonstrate empirically, some evidence of genotype–environment interaction was observed among viability loci (i.e., among different genotypes) in Pacific oysters reared in different diet environments (Plough 2012). A potentially complex dynamic of selection such as this could be playing out during recruitment of bivalve species and could contribute to observed patterns of chaotic genetic patchiness that are often described in genetic studies of marine invertebrate populations (e.g., Eldon et al. 2016). Future studies of the dynamics of selection during the larval period with field or mesocosm-based sampling could begin to address this.

Future Directions

It is an exciting time to be studying the evolutionary genetics of marine animals as genomic data becomes less expensive and easier to generate (e.g., Andrews and Luikart 2014) and fundamental population genetic theory is being revised to accommodate the extreme life history and demographic features of high fecundity marine species (e.g., Sargisyan and Wakeley 2008; Eldon and Wakeley 2009; Tellier and Lemaire 2014). Driven by the availability of genomic datasets for non-model marine species and advances in our understanding of the underlying causes of genetic load in marine species, I suggest two major research foci going forward. First, there is a definite need for more studies of molecular diversity focusing on coding regions at the genome scale, to increase the available data on dn/ds ratios, site frequency spectra (SFS), and the distribution of mutation fitness effects (Eyre-Walker and Keightley 2007) in marine species. Though RAD sequencing is a powerful technique to survey diversity across the genome in virtually any species (e.g., Davey et al. 2011; Andrews et al. 2016), it largely profiles non-coding regions, and therefore provides little information on the frequency and polymorphism of amino acid changing mutations. Other approaches, including re-sequencing of whole genomes, (Andrews and Luikart 2014), non-model exon-capture techniques (e.g., Cosart et al. 2011; Bi et al. 2012), or the analysis of RNAseq data across multiple (>15) individuals, will provide much needed data to address this gap. There is also the potential to mine the vast, online repositories of RNAseq data for SNP variation, and tools exist to estimate synonymous and non-synonymous diversity from pooled samples (e.g., Nelson et al. 2015). With more genome-scale data on the molecular diversity of marine species, especially from taxa that are poorly represented currently (e.g., fish, echinoderms, annelids, crustaceans, gastropods), more robust taxonomic comparisons of genetic load can be made across marine animals groups and to other non-marine taxa.

The second major research focus should be to generate high-resolution, genome-wide segregation data for a select number of taxonomically diverse "target" marine animal species, employing QTL mapping methodologies to characterize viability loci and sequence comparison between parents and offspring to estimate mutation rates more directly. This work could be focused on a select group of marine animal species for which husbandry protocols are well established and the generation of many offspring (hundreds to thousands) would be routine. However, this effort should not simply be limited to oysters, which may turn out to be a rather extreme case, but must also examine other bivalves (e.g., blue mussels, quahogs, dwarf surf clam), marine fish (e.g., cod, turbot, pollock) and crustaceans (e.g., Pacific white shrimp, blue crab, American lobster). High-resolution linkage maps (tens of thousands of markers) are now relatively routine to generate using RADseq data of parents and offspring; however, care should be taken with linkage map construction because it can be complicated significantly by many loci and segregation distortion (e.g., Lorieux et al. 1995; Lorieux 2012; Hedgecock et al. 2015). Statistical software to detect and characterize the number and effect viability loci through QTL mapping is available and open source (e.g., Luo and Xu 2003; Hu and Xu 2009; Zhan and Xu 2011, see Plough et al. 2016).

To examine de novo mutations and their frequency distribution in the offspring (e.g., clustered; Gao et al. 2011), sequences at RADseq loci could be compared between parents and offspring to detect any single base changes, deletions, or insertions, following methods and techniques developed for analysis of parent–offspring trios or quartets in human disease research (e.g., Roach et al. 2010; Sanders et al. 2012). Human parent–offspring sequencing studies often employ exon or whole-genome sequencing, approaches that are possible for some species with a sequenced genome like the Pacific oyster (though impractical for large numbers of offspring) but not for most "model" marine species. Instead, generation of genotype data with a RADseq approach would strike a balance between sequencing effort and cost, producing relatively unbiased genome-wide coverage at a cost that would allow the genotyping of hundreds of offspring, which increases the likelihood of detecting rare, novel variants. Paired-end RADseq would be particularly appropriate here, as the assembly of paired-end reads recovers larger overlapping stretches of the genome that may traverse coding regions. RADseq data sets may already be available from some of the linkage map data reviewed here (e.g., Kai et al. 2014; Wang et al. 2015; Palaiokostas et al. 2015) and these data could be reanalyzed for this purpose. Overall, novel work focused on either an experimental or molecular evolution approach in multiple, tractable, marine animals, will significantly advance our understanding of genetic load in these species and help continue the very exciting and theory-challenging work up to now.

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Supplementary Material
Supplementary material can be found at http://www.cz.oxfordjournals.org/

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