The Effects of Increasing the Number of Taxa on Inferences of Molecular Convergence

Gregg W. C. Thomas¹,*, Matthew W. Hahn¹, and Yoonsoo Hahn²

¹Department of Biology and School of Informatics and Computing, Indiana University, Bloomington, Indiana
²Department of Life Science, Research Center for Biomolecules and Biosystems, Chung-Ang University, Seoul, Republic of Korea

*Corresponding author: E-mail: grthomas@indiana.edu.
Accepted: January 1, 2017

Abstract

Convergent evolution provides insight into the link between phenotype and genotype. Recently, large-scale comparative studies of convergent evolution have become possible, but researchers are still trying to determine the best way to design these types of analyses. One aspect of molecular convergence studies that has not yet been investigated is how taxonomic sample size affects inferences of molecular convergence. Here we show that increased sample size decreases the amount of inferred molecular convergence associated with the three convergent transitions to a marine environment in mammals. The sampling of more taxa—both with and without the convergent phenotype—reveals that alleles associated only with marine mammals in small datasets are actually more widespread, or are not shared by all marine species. The sampling of more taxa also allows finer resolution of ancestral substitutions, revealing that they are not in fact on lineages leading to solely marine species. We revisit a previous study on marine mammals and find that only 7 of the reported 43 genes with convergent substitutions still show signs of convergence with a larger number of background species. However, four of those seven genes also showed signs of positive selection in the original analysis and may still be good candidates for adaptive convergence. Though our study is framed around the convergence of marine mammals, we expect our conclusions on taxonomic sampling are generalizable to any study of molecular convergence.

Key words: convergent evolution, molecular convergence, taxonomic sampling.

Introduction

From the beginning of the genome era, researchers have attempted to associate individual genomic features with the phenotypes present in sequenced species. Such features include unique amino acid substitutions (Kim et al. 2011), gene family expansions (Lespinet et al. 2002) or contractions (Aravind et al. 2000), and changes in gene expression (Khaitovich et al. 2005), to name a few. In order to increase the power to detect causative changes as more genomes were sequenced, researchers looked for convergent molecular changes associated with convergent phenotypes. Again, such changes were found among amino acid substitutions (Rokas and Carroll 2008; Foote et al. 2015), gene gains and losses (Pellegrini et al. 1999; McBride and Arguello 2009; Hiller et al. 2012; De Smet et al. 2013; Liebeskind et al. 2015), in sex chromosome morphology (Bellott et al. 2010), and in gene expression (Ogura et al. 2004; Pankey et al. 2014).

However, it has become evident that the number of species and genomes used in an analysis greatly influences the probability of finding unique genomic changes in only taxa with the phenotypes of interest. For example, the original description of the naked mole-rat genome found that a unique amino acid substitution may have been responsible for the rodent’s hairless phenotype (Kim et al. 2011). A subsequent study with a larger number of taxa found that this substitution was not, in fact, unique to naked mole-rats but is found throughout mammals, including in the closest living relative to the naked mole-rat, the guinea pig (Delsuc and Tilak 2015). Similarly, we expect that as more genomes are analysed in studies looking for uniquely convergent molecular changes—those shared only by the taxa with convergent phenotypes—it will be more and more likely that we find similar changes in nonphenotypically convergent species. While such analyses will certainly increase our power to identify truly causative convergent mutations, in some cases they may also reveal that convergent phenotypes are not underlain by convergent molecular mechanisms (Storz 2016).
Here we investigate how the number of taxa sampled affects inferences of convergent evolution among amino acid sequences. We find that as more lineages are added to an analysis, the number of unique convergent substitutions among a given set of species decreases rapidly. We also find that adding taxa can decrease the number of convergent substitutions along target lineages (unique to those lineages or not), as the added phylogenetic resolution can help to more accurately reconstruct ancestral sequences. Finally, we revisit results on convergent amino acid substitutions among marine mammals presented in Foote et al. (2015) to see how these substitutions hold up to the addition of more species to the analysis. Three separate mammalian lineages have transitioned to an aquatic lifestyle and in the process have experienced extensive phenotypic convergence (Kelley et al. 2016), making them a prime choice for the study of convergence at the molecular level.

Materials and Methods
To assess how the number of taxa influences inferences of molecular convergence, we began by downloading the 100 species hg19 human reference alignments from the UCSC Genome Browser (Kent et al. 2002; http://hgdownload.soe.ucsc.edu/goldenPath/hg19/multiz100way/; last accessed January 6, 2017). We pruned and filtered all but 59 mammal species (fig. 1). Of these 59 species, 5 are marine mammals. The West Indian manatee (Trichechus manatus latirostris) is from the Order Sirenia; two Cetacean species are present, the killer whale (Orcinus orca) and the bottlenose dolphin (Tursiops truncatus); and there are two Pinniped species, the walrus (Odobenus rosmarus) and the Weddell seal (Leptonychotes weddellii). These three orders represent three independent gains of phenotypes associated with aquatic lifestyles in mammals (fig. 1; Foote et al. 2015), and were the target lineages for our tests of convergence. In our analyses, we classify lineages in which we are searching for convergence as “target” lineages and all others within the phylogeny as “background” lineages. We filtered all but the longest isoform from the original 39,361 sequences giving us 19,215 orthologs from which to draw on for our analysis of unique convergent substitutions.

We define unique convergent substitutions as sites where all marine mammal species have an identical amino acid that is different from any amino acid found in the background species. We count unique convergent substitutions with varying numbers of background species in the following way: starting with the five marine mammal species, we randomly added n background species and counted unique convergent sites for each dataset. Values of n range from 11 to 54 (there were 11 background species in the original analysis of Foote et al. 2015), and we repeated the process 100 times for each n.

We also identified possible convergent substitutions by reconstructing ancestral amino acid states and looking for changes along the lineages leading to the three marine mammal clades (gray, dashed branches in fig. 1). For this analysis we restricted our set of 19,215 sequences to only those 10,521 in which all 59 mammalian species were present. We used codeml in PAML (Yang 2007) to reconstruct ancestral sequences and counted a site as a convergent substitution if a change from any ancestral state to the descendant state had occurred along all three of these branches and resulted in the same descendant state. We followed a similar replicate scheme as above, but with fewer values of n and fewer numbers of replicates due to computational constraints. Values of n for ancestral reconstructions are: 15, 20, 25, 30, 35, 40, 45, and 50 and the process was repeated 5 times for each value of n. For this experiment, background species are chosen randomly for every replicate at every value of n. We also tried various cutoffs for the posterior probability of inferred ancestral states, but found that they did not affect our main result (supplementary fig. S1, Supplementary Material online) so we report results with no cutoff in the main text.

Finally, we replicated the analysis in Foote et al. (2015) by performing ancestral reconstructions on the same species used in that study (highlighted and underlined species in fig. 1, excluding Weddell seal) and counting convergent substitutions along the marine mammal lineages. We then added Weddell seal to the analysis to see how another target species affects inferences of convergence. From there, we iteratively added background species (15, 20, 25, 30, 35, 40, 45, 50 species) and repeated the convergence inference process. In this iterative setup, the background species that are added are still chosen randomly, but all species in previous replicates are included in the current one. For example, in the replicate with 20 background species, 15 of them are preserved from the trial with 15 background species and 5 more are added randomly. At each replicate, the list of genes with convergent substitutions was compared with the list of 43 genes with convergent substitutions inferred by Foote et al. (2015).

Results
Our current study is motivated by the recent investigation of molecular convergence in marine mammals by Foote et al. (2015). In that study, the authors (including G.W.C.T. and M.W.H.) used the genomes of four marine mammals (highlighted in fig. 1, excluding the Weddell seal) as target species and 11 terrestrial mammals (underlined in fig. 1) as background species. Alignments of those 15 species were used to identify convergent substitutions by inferring ancestral sequences. In total, convergent substitutions in 43 genes were found along the lineages leading to all three marine mammal clades. We used the same number of species as the starting point for this study.

Sample Size Affects the Number of Uniquely Convergent Substitutions
We find that as the number of background taxa is increased, the number of uniquely convergent substitutions among a set
of target species decreases (these represent amino acid states found only in marine mammals). We observed this by counting unique convergent substitutions among the five marine mammal species highlighted in figure 1 and randomly adding other background mammalian taxa to the analysis in increasing numbers (fig. 2). Although there is a lot of variability in the exact number of initially unique convergent substitutions that are found in other lineages—and this is associated with the
random sampling of additional background lineages—there is a clear monotonic relationship that declines as more taxa are added. Our results imply that with enough additional taxa added there will be one, or possibly zero, unique convergent substitutions among the three marine mammal clades. Using all 54 background species, we find only 1 uniquely convergent substitution in one gene (ITGA8; NM_003638) (fig. 2).

We also tested how the number of target species affects counts of unique substitutions by specifying the same target and background species used in Foote et al. (2015) and either adding Weddell seal or removing killer whale or dolphin. With the original 15 species, we find 131 unique convergent substitutions. However, when Weddell seal is added as a target species this number drops to 87. This is because the seal has a different amino acid at a position where the other four marine mammals share one to the exclusion of all other species in the analysis. Likewise, when either killer whale or dolphin is removed from the analysis as target species, the number of uniquely convergent substitutions rises to 144 and 141, respectively. In this case the removal of target species has made it more likely to find amino acids shared among a smaller set of marine mammals.

Sample Size Affects Inferences of Convergent Evolution by Changing Ancestral Reconstructions

Molecular convergence can also be assessed by using ancestral sequence reconstruction to look for shared changes along specific branches of interest in a phylogeny. Following this procedure, we searched for convergent substitutions along the three branches leading to marine mammals in our dataset (gray, dashed branches in fig. 1) while varying the number of background species. We again find a sharp drop-off in convergent substitutions with increasing numbers of background species (fig. 3). Note that unlike in the previous analysis, here it is the reconstruction that has changed, not the uniqueness of these convergent substitutions. This analysis solely asks whether the substitutions are convergent along all three lineages based on ancestral sequence reconstruction, not whether additional substitutions to the same state have occurred elsewhere on the tree.

The apparent loss of convergent substitutions is caused when the inclusion of additional species changes the reconstructed ancestral amino acid states, leading the substitutions to be assigned to different branches. In the many cases we examined this occurs because the reconstructed state at the ancestral nodes of the target lineages is inferred to have the...
The Effects of Increasing the Number of Taxa on Inferences of Molecular Convergence

The Effects of Increasing the Number of Taxa on Inferences of Molecular Convergence

state shared among marine mammals. Therefore, no convergent substitution along these three branches is necessary. Figure 4 highlights an example in which a convergent substitution is found in marine mammals in an analysis with 11 background species (serine → lysine or alanine → lysine). However, with the addition of four more species, three of which have an observed lysine residue at that position, the ancestral reconstructions throughout the tree are changed, mostly to lysine to accommodate the new observed states. The original convergent substitution is therefore no longer counted, as it has been moved to a nonmarine mammal lineage.

Comparisons with the Original Analysis of Marine Mammal Convergence

Given these results, we revisited the original findings of amino acid convergence among marine mammals (Foote et al. 2015). In that article, the authors found 44 convergent substitutions in 43 genes along marine mammal lineages among a set of 5,893 orthologs. Several of these genes were filtered out in the current analysis because they were not present in all 59 species; only 33 of the original 43 are present in our final dataset. Replicating their analysis (using ancestral sequence reconstruction and the same set of 15 species) with our set of 10,521 genes, we find 246 convergent substitutions in 233 genes. We recovered 28 of the original 33 genes with convergent substitutions. The five unrecovered genes are accounted for either by the use of a different isoform in this study (compared with Foote et al. 2015) or because 1 of the 15 species was missing from the alignment in the original study, and the inclusion of this species here shows that these were not convergent. The pattern of decreased convergence shared by the marine mammals continues as we add additional background species to the analysis: the total number of convergent substitutions and the number of convergent genes recovered from the original study decreases (table 1).

When we add the fifth marine mammal, Weddell seal, to the analysis, the number of convergent substitutions drops to 161 and we recover only 20 of the original 43 convergent genes found by Foote et al. (2015). The decrease in convergent substitutions, even putatively adaptive ones, with more focal taxa is not unexpected. Foote et al. (2015) previously reported that some convergent substitutions in the four marine mammals studied were not present in further comparisons with the minke whale genome (Yim et al. 2013). However, because minke was not included in their phylogenetic analysis, it is possible that these substitutions represent reversions in minke whale.

We also replicated Foote et al.’s (2015) use of terrestrial mammals as an empirical null distribution to test if the amount of convergence in marine mammals exceeds expectations. In the original article, the authors counted convergent substitutions between dog, cow, and elephant as a control and found no excess of convergence in marine mammals compared with these land mammals. The authors in fact reported more convergence in land mammals, with 93 convergent substitutions in 90 genes. Similarly, we also find that convergence in marine mammals does not exceed that of convergence in the same set of land mammals, even when increasing the number of taxa in the dataset. With the original set of species we find convergent substitutions in 398 genes among land mammals, many more than the 233 in marine mammals. This pattern holds with increasing numbers of background species. With all 50 background species, we find 117 genes with convergent substitutions in land mammals compared with 78 in marine mammals (table 1). This confirms the original findings of Foote et al. (2015) that there is no excess amino acid convergence among marine mammals. This also implies that the pattern of decreasing convergence with increasing number of taxa remains regardless of the specific target species used in the phylogeny.

Interestingly, several of the genes originally found as convergent by Foote et al. (2015) continue to show signals of convergence despite the increased number of background species (table 2). However, even though the convergent substitutions remain, there is still a possibility that they are not adaptively convergent, but rather the result of neutral processes. An additional layer of testing is needed to confirm adaptive convergence. In Foote et al. (2015) some of these genes also show signs of positive selection (based on analyses of $d_{NS}/d_{S}$) and this extra test indicates that they may have possible functional implications for phenotypic convergence among marine mammals. $MGP9$ may play a role in bone formation, $MYH7B$ is involved in cardiac muscle development, and $SERPINC1$ regulates blood coagulation. $GCLC$ is another interesting gene that we still detect as convergent with a large number of species. This gene is involved in glutathione

![Fig. 3.](image-url)
metabolism, a molecule that has been shown to prevent oxidative damage in cetaceans during long underwater dives.

**Discussion**

The study of convergence at the molecular level is still relatively new. Although most recent papers have examined convergence in amino acid substitutions (Rokas and Carroll 2008; Foote et al. 2015), there can be many types of convergent molecular changes, including gene gains, losses, and shifts in gene expression (Ogura et al. 2004; De Smet et al. 2013; Pankey et al. 2014; Liebeskind et al. 2015). Researchers are also discovering many nuances that must be considered when measuring molecular convergence, such as choosing the correct null model (Thomas and Hahn 2015; Zou and Zhang 2015a) and accounting for physical constraints on convergence (Goldstein et al. 2015; Zou and Zhang 2015b).
number of taxa in a experiment can also affect any possible inferences of lineage-specific changes (Deluc and Tilak 2015) and this must also be considered in measures of molecular convergence.

Stayton (2008) showed that inferences of convergence in quantitative traits increased with increasing number of taxa, and this must also be true of molecular data. With the addition of each new species, there will also be more branches in the phylogeny on which substitutions of all types can occur. What we have shown here is that increased sample size decreases inferences of molecular convergence among a set of target species. These results illustrate the need for caution when making conclusions about molecular convergence. Molecular convergence that is responsible for adaptive phenotypic convergence appears to be much less common when additional species are included in any analysis. In many cases, the convergent amino acid state will be sampled in a nonphenotypically convergent species. In our study of marine mammals these species were not even semi-aquatic mammals, thus ruling out these changes as being adaptively convergent in tests that rely on unique convergent sites among a set of target species. However, sites such as these may still play a role in convergent phenotypes, just not a strong enough one to be detected by these stringent criteria. Similar patterns of ecologically diverse taxa sharing convergent states are found in studies of morphological characters (Zou and Zhang 2016).

We expect to see the same effect of taxonomic sampling when considering convergence in any type of molecular change. As more species are added, for instance, more gene duplications and losses will have occurred, making it likely that we observe such changes in nonphenotypically convergent lineages. There can also be different ways that changes in protein sequences can be convergent. Chikina et al. (2016) propose that molecular convergence may be the result of selection at the gene level rather than the amino acid level. These authors found an excess of genes with accelerated or decelerated rates of evolution across marine mammals when compared with terrestrial mammals. This model proposes that multiple different changes in common genes lead to a convergent phenotype rather than a single identical convergent amino acid change. We expect that this measure of convergence should still be sensitive to changes in the number of species represented, though we have not tested this here (note that Chikina et al. use at least 30 species in their analysis).

In this study, we used two methods for detecting convergent substitutions: unique substitutions among a set of target species and convergent changes inferred with ancestral reconstructions on a set of target lineages. Both methods showed a similar pattern of decreasing numbers of convergent sites with increasing taxa, though they may be affected by different features of the added taxa. The use of unique substitutions would seem to be a conservative method, as the presence of “convergent” amino acid states in nonphenotypically convergent species does not mean such changes are not involved in adaptation. For example, if the individual sequenced from a newly added species carries a deleterious allele at mutation-selection balance at a site that is truly convergent in a set of target species, we would eliminate this substitution as a candidate. Convergent substitutions inferred with ancestral reconstructions may be lost because additional taxa move the convergent substitution to a different branch of the phylogeny, or because they reverse the direction of the evolutionary change (e.g., fig. 4).

Ancestral reconstructions are also dependent on the tree topology. The addition of new taxa may change the species tree topology and drastically alter inferred ancestral states. Though we have not explored the effect of additional taxa on improving the underlying topology, this will clearly affect inferences of convergence. Incorrect topologies—either due to error in species tree inference or biological differences in the histories among different genes—can lead to truly convergent substitutions being missed, but can also lead to many incorrect inferences of convergence (Mendes and Hahn 2016; Mendes et al. 2016).

Both tests used to identify convergent genes in this study require an identical change on multiple lineages within a phylogeny. However, the requirement of an identical change in all target lineages may actually exclude highly nonrandom instances of amino acid convergence. Especially as more target lineages are added, these tests may be too stringent. For instance, Zhen et al. (2012) show an excess of parallel and convergent amino acid changes in the alpha subunit of the sodium pump Na+,K+-ATPase (ATP$_2$α) among subsets of 14 insects that can metabolize toxic cardenolides. With so many target lineages, most substitutions are not found in all species, and would therefore fail our tests for convergence.

In order to relax the strict criteria for identifying convergent genes, we propose to break the target lineages into smaller groups, counting convergent substitutions within each group. Interesting genes could then be identified as those with a convergent substitution (not necessarily the same one) in two or more groups of target species. Especially when there are large numbers of target lineages (Zhen et al. 2012; Natarajan et al. 2016), this method may recover instances
missed by requiring all taxa to share the same amino acid. We applied this method to the data presented above by splitting the three marine mammal lineages into three pairs of target lineages: Sirenia-Pinnipedia, Sirenia-Cetacea, and Pinnipedia-Cetacea. We then inferred convergent substitutions in each pair separately using ancestral reconstruction, with the remaining marine mammal lineage as a background lineage. A gene is classified as convergent if there are convergent substitutions at any position in two or more of the pairs. When using this paired analysis we observe far more putatively convergent genes than with the original tests that require an identical change in all three lineages, indicating that this is indeed a less stringent test (supplementary fig. S2A, Supplementary Material online). The pattern of diminishing convergence with increased taxa is still observed, however, as is the presence of more molecular convergence in land mammals than in marine mammals (supplementary fig. S2B, Supplementary Material online). In addition, even with relaxed criteria, it is still challenging to determine which convergent substitutions are important, as not all will prove to be functional (Natarajan et al. 2016).

Aside from the quantification of molecular convergence, an important consideration in any study is quantifying whether there is an excess amount of convergence. Because molecular convergence can be the result of neutral processes (Zhang and Kumar 1997), appropriate null comparisons must be made. The best way to account for this background convergence is with the use of an empirical null model, such as comparisons among nonconvergent species that are closely related to the target lineages. For example, Foote et al. (2015) found no evidence of excess convergence in marine mammals given levels of background convergence in the terrestrial species elephant, dog, and cow. We find similar results here with our larger taxonomic sample. And while the signal for convergence remained in several of the interesting genes originally found by Foote et al. (table 2), we must point out that even with 50 species the trend does not seem to have levelled out. This could indicate that perhaps with even more species some of these genes would no longer be convergent.

We recommend that future studies of molecular convergence should be performed so that they maximize the number of taxa represented for more confident inferences. Our results show that adding either target or background species can change the outcome of convergence analyses, so it is currently difficult to determine which is more valuable. However, it is likely that adding background species will be easier in most cases since target species sharing a convergent trait will be less common.

Supplementary Material

Supplementary data are available at Genome Biology and Evolution online.

Acknowledgments

We would like to thank Andy Foote, Jay Storz, and two anonymous reviewers for helpful feedback with the article. We acknowledge computational resources provided by the National Center for Genome Analysis Support (National Science Foundation grant DBI-1458641).

Literature Cited

Aravind L, Watanabe H, Lipman DJ, Koonin EV. 2000. Lineage-specific loss and divergence of functionally linked genes in eukaryotes. Proc Natl Acad Sci U S A. 97(21):11319–11324.
Bellott DW, et al. 2010. Convergent evolution of chicken Z and human X chromosomes by expansion and gene acquisition. Nature 466(7306):612–616.
Chikina M, Robinson JD, Clark NL. 2016. Hundreds of genes experienced convergent shifts in selective pressure in marine mammals. Mol Biol Evol. 33(9):2182–2192.
De Smet R, et al. 2013. Convergent gene loss following gene and genome duplications creates single-copy families in flowering plants. Proc Natl Acad Sci U S A. 110(8):2988–2990.
Delsuc F, Tilak MK. 2015. Naked by not hairless: the pitfalls of analyses of molecular adaptation based on few genome sequence comparisons. Genome Biol Evol. 7(3):768–774.
Foote, et al. 2015. Convergent evolution of the genomes of marine mammals. Nat Genet. 47(3):272–275.
Goldstein RA, Pollard ST, Shah SD, Pollock DD. 2015. Nonadaptive amino acid convergence rates decrease over time. Mol Biol Evol. 32(6):1373–1381.
Hiller M, et al. 2012. A “forward genomics” approach links genotype and phenotype using independent phenotypic losses among related species. Cell Rep. 2(4):817–823.
Kelley JL, Bornw AP, Therkildsen NO, Foote AD. 2016. The life aquatic: advances in marine vertebrate genomics. Nat Rev Genet. 17(9):523–534.
Kent WJ, et al. 2002. The human genome browser at UCSC. Genome Res. 12(6):996–1006.
Khaitovich P, et al. 2005. Parallel patterns of evolution in the genomes and transcriptomes of humans and chimpanzees. Science 309(5742):1850–1854.
Kim EB, et al. 2011. Genome sequencing reveals insights into physiology and longevity of the naked mole rat. Nature 479(7372):223–227.
Lespinet O, Wolf YI, Koonin EV, Aravind L. 2000. The role of lineage-specific gene family expansion in the evolution of eukaryotes. Genome Res. 12:1048–1059.
Liebeschkind BJ, Hills DM, Zakon HH. 2015. Convergence of ion channel genome content in early animal evolution. Proc Natl Acad Sci U S A. 112(8):E846–E851.
McBride CS, Arguello JR. 2009. Five Drosophila genomes reveal nonneutral evolution and the signature of host specialization in the chemoreceptor superfamily. Genetics 177(3):1395–1416.
Mendes FK, Hahn MW. 2016. Gene tree discordance causes apparent substitutions rate variation. Syst Biol. 65(4):711–721.
Mendes FK, Hahn Y, Hahn MW. 2016. Gene tree discordance can generate patterns of diminishing convergence over time. Mol Biol Evol. 33(12):3299–3307.
Natarajan C, et al. 2016. Predictable convergence in hemoglobin function has unpredictable molecular underpinnings. Science 354(6310):336–339.
Ogura A, Ikeo K, Gajobori T. 2004. Comparative analysis of gene expression for convergent evolution of camera eye between octopus and human. Genome Res. 14(8):1555–1561.
Pankey MS, Minin VN, Imholte GC, Suchard MA, Oakley TH. 2014. Predictable transcriptome evolution in the convergent and complex
bioluminescent organs of squid. Proc Natl Acad U S A. 111(44):E4736–E4742.
Pellegrini M, Marcotte EM, Thompson MJ, Eisenberg D, Yeates TO. 1999. Assigning protein functions by comparative genome analysis: protein phylogenetic profiles. Proc Natl Acad Sci U S A. 96(8):4285–4288.
Rokas A, Carroll SB. 2008. Frequent and widespread parallel evolution of protein sequences. Mol Biol Evol. 25(9):1943–1953.
Stayton CT. 2008. Is convergence surprising? An examination of the frequency of convergence in simulated datasets. J Theor Biol. 252(1):1–14.
Storz JF. 2016. Causes of molecular convergence and parallelism in protein evolution. Nat Rev Genet. 17(4):239–250.
Thomas GWC, Hahn MW. 2015. Determining the null model for detecting adaptive convergence from genomic data: a case study using echolocating mammals. Mol Biol Evol. 32(5):1232–1236.
Yim, et al. 2013. Minke whale genome and aquatic adaptation in cetaceans. Nat Genet. 45(1):88–92.
Zhang J, Kumar S. 1997. Detection of convergent and parallel evolution at the amino acid sequence level. Mol Biol Evol. 14(5):527–536.
Zhen Y, Aardema ML, Medina EM, Schumer M, Adolffatto P. 2012. Parallel molecular evolution in an herbivore community. Science 337(6102):1634–1637.
Zou Z, Zhang J. 2015a. No genome-wide protein sequence convergence for echolocation. Mol Biol Evol. 32(5):1237–1241.
Zou Z, Zhang J. 2015b. Are convergent and parallel amino acid substitutions in protein evolution more prevalent than neutral expectations? Mol Biol Evol. 32(8):2085–2096.
Zou Z, Zhang J. 2016. Morphological and molecular convergences in mammalian phylogenetics. Nat Commun. 7:12758.

Associate editor: Jay Storz