Siphonophore Phylogeny

Catriona Munro¹, Stefan Siebert^{1,2}, Felipe Zapata^{1,3}, Mark Howison⁴, Alejandro Damian Serrano^{1,9}, Samuel H Church¹, 5[^], Freya Goetz^{1,6}, Phil Pugh⁷, Steven H.D. Haddock⁸, Casey W. Dunn^{1,9}*

- ¹ Department of Ecology and Evolutionary Biology, Brown University, Providence, RI 02912, USA
- 2 Current address: Department of Molecular & Cellular Biology, University of California at Davis, Davis, CA 95616, USA
- ³ Current address: Department of Ecology and Evolutionary Biology, University of California Los Angeles, Los Angeles, CA 90095, USA
- ⁴ Brown Data Science Practice, Brown University, Brown University, Providence, RI 02912, USA
- 5 Department of Organismic and Evolutionary Biology, Harvard University, Cambridge, MA 02138, USA
- 6 Current address: Smithsonian Institution, National Museum of Natural History, Washington, DC 20560, USA
- ⁷ National Oceanography Centre, Southampton, SO14 3ZH, UK
- ⁸ Monterey Bay Aquarium Research Institute, Moss Landing, CA 95039, USA
- 9 Current address: Department of Ecology and Evolutionary Biology, Yale University, New Haven, CT 06520, USA
- * Corresponding author, casey dunn@brown.edu

Abstract

Introduction

Siphonophores are...

Methods

This manuscript is an executable document computed directly from the data, providing an explicit and reproducible description of all findings. All scripts for the analyses are available in a git repository at https://github.com/caseywdunn/siphonophore_phylogeny_2017. The most recent commit at the time of the analysis presented here was b4860437e8e3265a30aaa829f6997e70d484435d.

Collecting

Collection data on all examined specimens, a description of the tissue that was sampled from the colony, collection mode, sample processing details, mRNA extraction methods, sequencing library preparation methods and sequencing details are summarized in supplementary table 1. Monterey Bay and Gulf of California specimens were collected by remotely operated underwater vehicle (ROV) or during blue-water scuba dives. Chelophyes appendiculata and Hippopodius hippopus specimens were collected in the bay of Villefranche-sur-Mer, France, during a plankton trawl on 04/13/11. Available physical vouchers have been deposited at the Museum of Comparative Zoology (Harvard University), Cambridge, MA, and at the United States National Museum (Smithsonian Institution), Washington, DC. Accession numbers are given in supplementary table X. In cases where physical vouchers were unavailable we provide photographs to document species identity (table x).

Sequencing

When possible specimens were starved overnight in filtered seawater at temperatures close to ambient water temperatures at the time point of specimen collection (supplementary table 1). mRNA was extracted directly from tissue using a variety of methods (supplementary table x): Magnetic mRNA Isolation Kit (NEB, #S1550S), Invitrogen Dynabeads mRNA Direct Kit (Ambion, #61011), Qymo Quick RNA MicroPrep (Zymo #R1050), or from total RNA after Trizol (Ambion, #15596026) extraction and through purification using Dynabeads mRNA Purification Kit (Ambion, #61006)- in case of very small total RNA quantities, only a single round of bead purification was performed; or Trizol directly into the Illumina TruSeq Stranded Library Kit. Extractions were performed according to the manufacturer's instruction. Any resulting higher rRNA read counts were dealt with further downstream in the bioinformatics workflow. Libraries were prepared for sequencing using the Illumina TruSeq RNA Sample Prep Kit (Illumina, #FC-122-1001, #FC-122-1002), the Illumina TruSeq Stranded Library Prep Kit (Illumina, #RS-122-2101) or the NEBNext RNA Sample Prep Master Mix Set (NEB, #E6110S). We collected long read paired end Illumina data for de novo transcriptome assembly. In the case of large tissue inputs, libraries were sequenced separately for each tissue and subsequently were subsampled and pooled in silico. Libraries were sequenced on the HiSeq 2000, 2500, and 3000 sequencing platforms (supplementary table 1). Summary statistics for expression libaries are given in Table 1.

Analysis

New data were analysed in conjunction with 13 publically available datasets, with a total number of 43 species. Sequence assembly, annotation, Maximum Likelihood (ML) phylogenetic analysis were conducted with the tool Agalma (Dunn et al. 2013), v. 1.00, and Bayesian Inference (BI) analyses were conducted using Phylobayes(Lartillot et al. 2009) v. 1.7a-mpi. Source code for all analysis steps, sequence alignments, sampled and consensus trees, and voucher information are available in a git repository https://github.com/caseywdunn/siphonophore_phylogeny_2017.

In the final analyses, we sampled 1,071 genes to generate a supermatrix with 60% occupancy and a length of 378,468 amino acids. Two outgroup species, $Atolla\ vanhoeffeni$ and $Aegina\ citrea$, were removed from the final supermatrix and phylogeny due to low gene occupancy (gene sampling of 20.8% and 14.5% respectively in a 50% occupancy matrix with 2,203 genes). ML analyses were conducted on the unpartitioned supermatrix using the WAG+ Γ model of amino acid substitution, and bootstrap values were estimated using 1000 replicates. BI was conducted using two different CAT models, CAT-Poisson and CAT-GTR (Lartillot and Philippe 2004). Two independent MCMC chains were run under the CAT-GTR model, and four independent MCMC chains were run under the CAT-Poisson model. The CAT-GTR and CAT-poisson models did not converge after a long CPU time, and only the results from the CAT-poisson model are included here.

Morphological character data were obtained from the literature, or from direct observation of available voucher material.

Subsequent analyses were conducted in R and integrated into this manuscript with the knitr package. See Supplementary Information for R package version numbers.

Hypothesis testing

Results and Discussion

Sample collecting and sequencing

Species phylogeny

(XX This paragraph on sampling through Agalma analyses) The analyses presented here consider XXX siphonophore species and 8 outgroup species. This includes new data for XXX species. Summary stats on assemblies XX (Table XX). Matrix has XX genes, XX sites, and occupancy is XX.

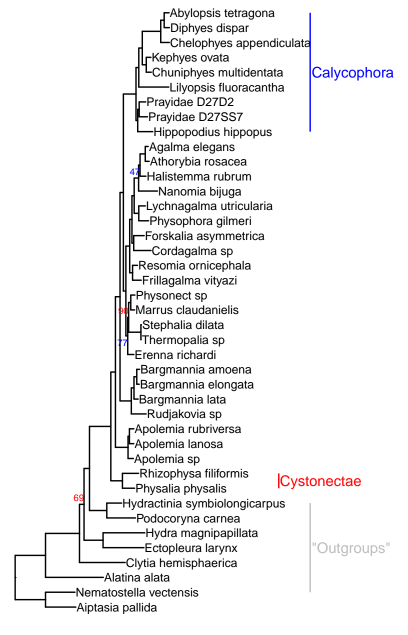


Figure 1: Phylogram of siphonophore relationships. Node labels indicate bootstrap support percent, unnumbered nodes have 100% support. The image was rendered with ggtree (Yu et al. 2016)

(XX This paragraph on summarizes phylogeny runs, apart from tree topology) Maximimum likelihood analyses had 1000 replicates. We ran 4 phylobayes chains, and visual inspection of the traces indicated that a burn in of 400 trees was sufficient for all runs. This left 15847 trees in the posterior. XXConvergence...

These findings are entirely consistent with a previous analysis based on two genes (16S and 18S ribosomal RNA) (Dunn et al. 2005). Cystonectae is the sister group to the remaining siphonophores, and Calycophorae is nested within the paraphyletic "Physonectae". In addition, multiple nodes that were not resolved in the previous two-gene analysis do recieve strong support in this 1,071-gene transcriptome analysis. These findings include XXX.

Character Evolution

Siphonohores have evolved a fascinating diversity of morphological features, zooid types, life history traits, and habitats. Here we explore the evolutionary history of some of these features.

In our previous siphonophore phylogenetic analysis (Dunn et al. 2005) there were several characters left with equivocal evolutionary histories, due to unresolved relationships between physonects. With our current cladistic resolution, we were able to answer some of the questions left open:

Evolution of Monoecy

[Citation needed] noticed for the first time that some siphonophores were monoeicous and others were dioiecious. Our analyses in 2005 reconstructed this character and found a great amount of phylogenetic conservatism, with an unambiguous resolution of the MRCA (most recent common ancestor) as dioecious, and the appearance of monoecy in several taxa and clades (including Calycophorae) within the polytomy. Figure X shows the evolution of sex distribution in siphonophores under the current better-resolved tree model, and it strongly indicates that monoecy in siphonophores from a dioecious ancestor occured twice, in the branch leading to Calycophorae and in the branch leading to Agalmatids (sensu lato). There is a small probability for an alternative scenario featuring a single gain of monoecy before the split of Calycophorae, with a subsequent derived shift back to dioecy in the *Marrus-Erenna* clade.

The Evolution of Zooid Types

One of the most striking aspects of siphonophore biology is their diversity of unique zooid types. Other colonial cnidarians (such as Hydractinia) and some bryozoans (example) have been found to have up to X different zooid types [Citation 1 , Citation 2]. The siphonophore genus Forskalia has 6 basic zooid types (pneumatophore, nectophore, gastrozooid, palpon, bract, and gonophore), and a total of 10 counting subtypes (4 types of bract, male & female gonophores). Diphyomorphs have more than 1 type of nectophore, while Cystonects have none. Here we reconstruct the evolutionary origins of the different zooid types on the present transcriptome tree.

Nectophores are retained modified medusae that Codonophora use for coordinated colony-level swimming. The nectosome is the region of the colony that develops from the nectosomal growth zone. Unlike the siphosomal growth zone, the nectosome does not bud gastrozooids, but nectophores (and in the case of *Apolemia*, also palpons). If fact, with the exception of *Physalia physalis* (which grows small nectophores near the gonodendra), siphonophore nectophores are exclusively found on the nectosome. It is possible that the MRCA of siphonophores had a nectosome, which has lost on the branch leading to Cystonects. We cannot exclude with certainty the alternative hypothesis of a nectosome-less ancestor followed by a gain of the nectosome in the branch leading to the Codonophora. The nectosome probably arose as a duplication of the siphosome, followed by functional specialization in propelling the colony. The nectosome has been lost within Codonophora in the genus *Athorybia*.

Following the colony development orientation framework presented in (citation), the nectosome can be located in a dorsal or a ventral position. Our ancestral reconstructions for this character (Supp figure) show that

a ventrally-oriented nectosome was the ancestral form in siphonophores, and that a dorsal nectosome has evolved twice independently, in the braches leading to the Agalmatidae (sensu stricto) and the branch leading to the Bargmannia species. All Codonophora (with the exception of Athorybia species) have a nectosome, but the number and subtypes of nectophores present varies greatly between species. As shown in Figure X, most Codonophora presents the ancestral nectosome with multiple nectophores of the same subtype. However, Calycophorans evolved a different system with just 2 nectophores of one type. This shift may be associated with the loss of the pneumatophore. Not all Calycophorans remained with this arrangement. The Hippopodidae returned to bearing multiple identical nectophores, many of which are inactive and serve functions of defense (like a shell to retract in) and buoyancy. As in the rest of Calycophorans, the Hippopodids only use 2 nectophores to propel the colony. Another interesting shift occurs in the branch leading to Diphyomorpha, where the 2 nectophores specialize into 2 subtypes, associated with a shift into a vertically aligned position and pointed bell shapes. The 2 types function together in a coupled hydrodynamic system that allows very fast escape responses (Mackie 1964).

Bracts are highly reduced zooids unique to siphonophores, but they are only present in the Codonophora. As with the nectosome, we have ambiguity determining whether the MRCA of siphonophores had bracts or not. The MRCA of Codonophora and only one bract subtype, which was lost in Hippopodidae and in *Gymnopraia lapislazula*. Bracts are functional for protection of the delicate zooids and to help maintain neutral buoyancy. Some calycophorans are able to actively exclude suphate ions in their bracts to adjust their buoyancy along the colony Bidigare, R. R., & Biggs, D. C. (1980).

The ancestral siphonophore certainly had a pneumatophore (Supp Figure X). This unique zooid fills itself with gas, which helps the colony float and maintain its orientation in the water column. Recent evidence of neural arrangement in the pneumatophore of *Nanomia bijuga* (Church, 2013) suggests it could also gather information on relative pressure changes (and thus depth changes), helping regulate geotaxis. Despite its multiple biological functions, it was lost in the Calycophorae and never gained again in that clade. Calycophorans rely on the ionic balance of their gelatinous nectophores and bracts to retain posture and neutral buoyancy.

Palpons are modified mouthless gastrozooids used for digestion and circulation of the gastrovascular fluid. They were present in the MRCA of siphonophores (Supp Figure X), retained in most species, but lost twice independently in the branches leading to *Bargmannia* and Calycophorae. These taxa might have found other avenues to effectively circulate nutrients across the colony.

The Gain and Loss of Tentilla

The most complex nematocyst batteries of Cnidaria can arguably be found among the siphonophores, hanging in regularly spaced tentacle side branches called tentilla. Most hydrozoans, including the clade that contains siphonophores, bear simple tentacles (tentacles with no side branches). It is still an open question whether the MRCA of Siphonophora had simple or branched tentacles. The only siphonophores genera regarded as lacking tentilla are *Physalia physalis* and *Apolemia* spp., and *Bathyphysa conifera*. Since *B. sibogae* is the only member of the *Rhizophysidae* (and of the *Bathyphysa* genus) lacking tentilla, we can safely assume this is a case of secondary loss. When we reconstruct the evolution of this character on the current phylogeny, we find that 70% of simulations support an MRCA bearing tentilla, with two independent losses leading to *Physalia* and *Apolemia*. However, this leaves a 30% support for a simple-tentacled MRCA followed by 2 independent gains of tentilla in the branches leading to *Rhizophysidae* and *{Bargmannia, Diphyes}*.

A key issue here is how we code for absence of tentilla, especially for the case of *Physalia physalis*. The tentacles of this species, when uncoiled, show very prominent, evenly spaced, bulging buttons which contain on their ectoderm all active and functionally arranged nematocysts used by the organism for prey capture. Siphonophore tentilla are complete diverticular brachings of the tentacle ectoderm, mesoglea, and gastrovascular canal (lined by endoderm). Hessinger & Ford 1988 (in the Biology of Nematocysts) described *Physalia*'s buttons as enclosing individual fluid-filled chambers connected by narrow channels to the tentacular canal, lined by endoderm. This suggests they are not just ectodermal swellings, but probably are reduced tentilla. When we code *Physalia physalis* as tentilla bearing, the results for the character reconstruction lead to a

more robust support for a tentilla-bearing MRCA followed by a single loss of tentilla in the branch leading to *Apolemiidae* (Figure 5b).

Siphonophore tentilla present an astounding diversity of sizes, shapes, colors, and nematocyst complements, and some have been observed to rapidly uncoil in contact with prey. Future research should explore the evolutionary history of these unique structures.

The Evolution of Vertical Habitat Use

Siphonophores are abundant predators in the pelagic realm, ranging from the surface (*Physalia physalis*) to bathypelagic depths (ref , *Bargmannia sp* 3888m VARS unpublished). While there are some pleustonic (*Physalia*) and benthic (*Rhodaliidae*) siphonophores, the phylogeny suggests the siphonophore MRCA was planktonic, as most extant taxa are. Some interesting questions arise from these facts, including 1) what was the bathymetric niche of the siphonophore MRCA, and 2) how did siphonophore's vertical habitat use of the water columns evolve along the phylogeny. Our results indicate a mesopelagic MRCA, with several convergent transition events to epipelagic and bathypelagic waters. There was only a single transition to benthic lifestyle on the stem of *Rhodaliidae*.

Discussion

The strong phylogenetic signal in the characters traditinally used for taxonomic diagnostics is a positive indicator of the applicability and unambiguity of these characters.

Conclusions

Acknowledgements

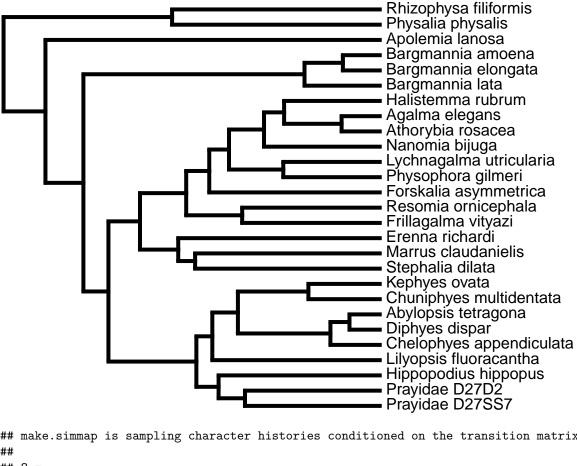
This work was supported by the National Science Foundation (DEB-1256695 and the Waterman Award). Sequencing at the Brown Genomics Core facility was supported in part by NIH P30RR031153 and NSF EPSCoR EPS-1004057. Data transfer was supported by NSF RII-C2 EPS-1005789. Analyses were conducted with computational resources and services at the Center for Computation and Visualization at Brown University, supported in part by the NSF EPSCoR EPS-1004057 and the State of Rhode Island. We also thank the MBARI crews and ROV pilots for collection of the specimens.

Supplementary Information

% latex table generated in R 3.4.1 by xtable 1.8-2 package % Fri Nov 3 16:00:19 2017 Table 1 (at the end of file): Summary statistics for libraries.

Supplementary Analyses

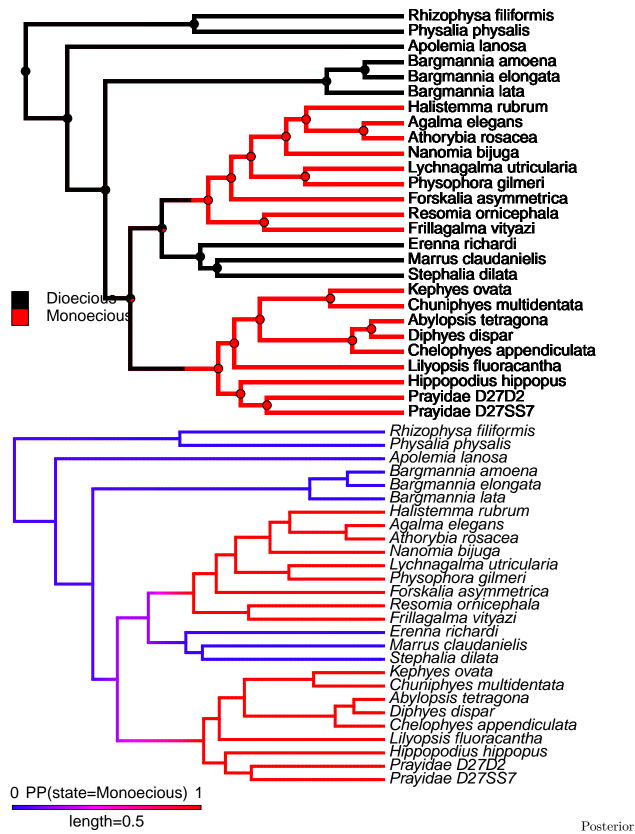
The Tree



```
## make.simmap is sampling character histories conditioned on the transition matrix
##
## Q =
## 0 1
## 0 -0.1892209 0.1892209
## 1 0.1892209 -0.1892209
## (estimated using likelihood);
## and (mean) root node prior probabilities
## pi =
## 0 1
## 0.5 0.5
```

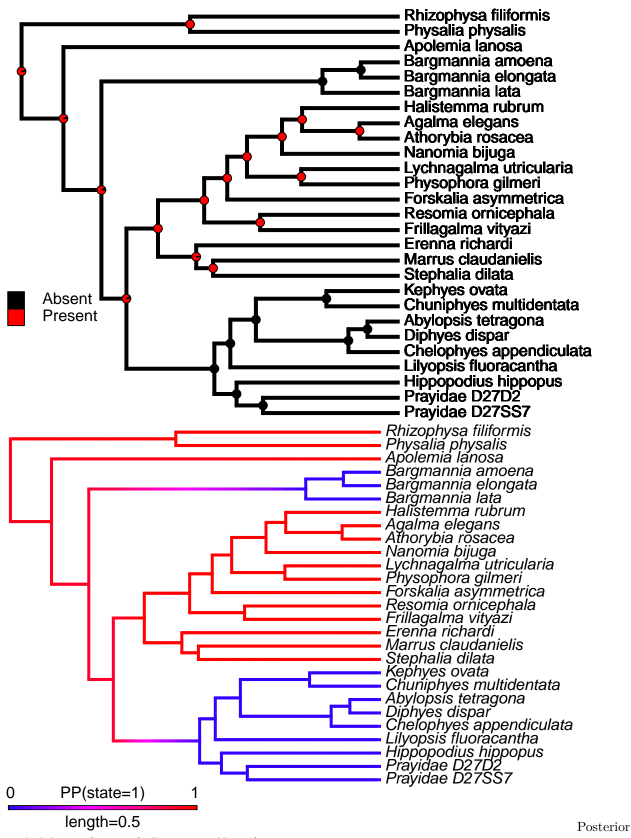
When we reconstruct the evolutionary history of different traits in siphonophore species, we obtain the following results:

SIMMAP Sex distribution



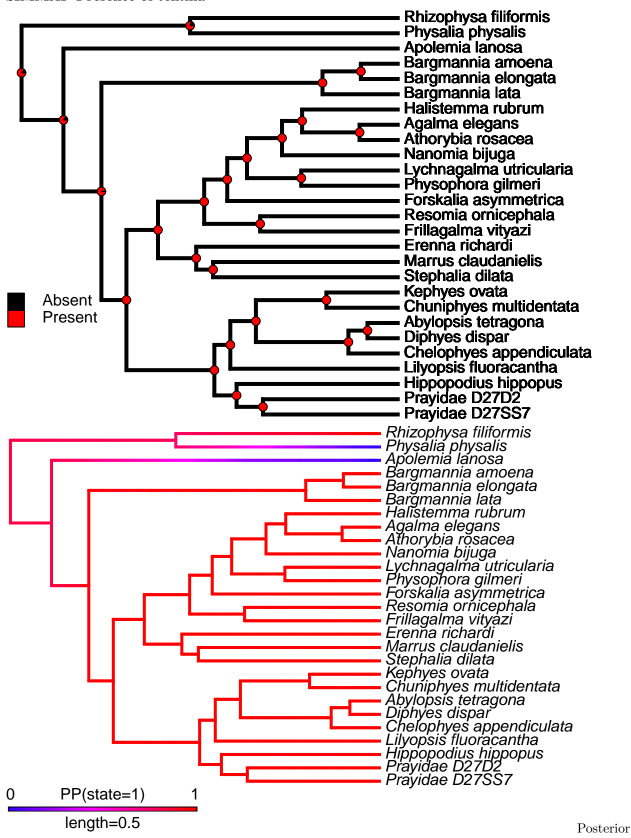
probabilities of states (0 Dioecious, 1 Monoecious).

SIMMAP Presence of palpons



probabilities of states (1 Present, 0 Absent).

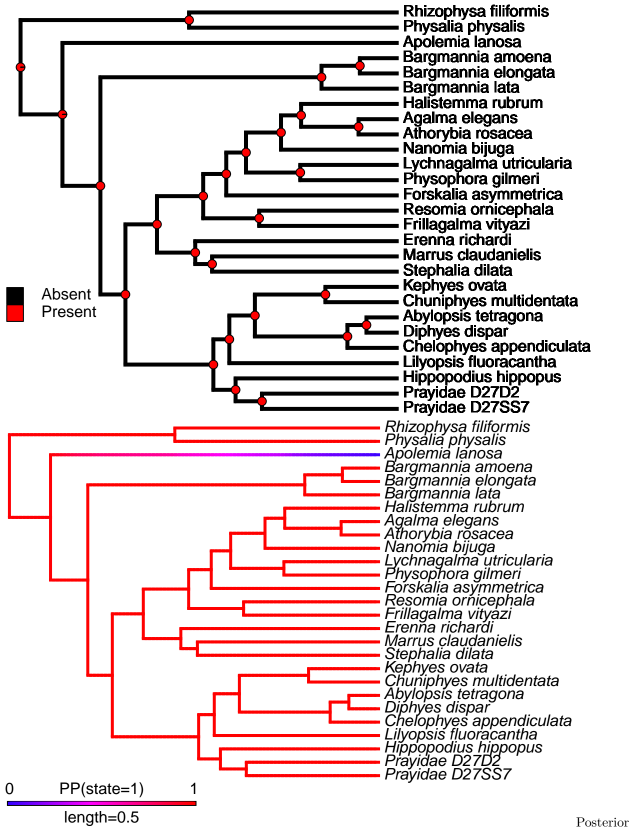
SIMMAP Presence of tentilla



probabilities of states (1 Present, 0 Absent).

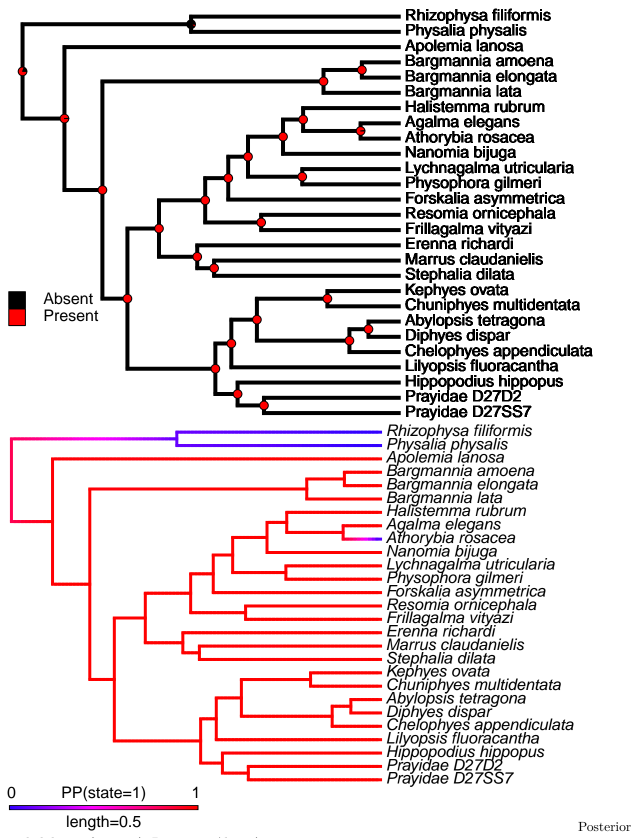
 $Physalia\ physalis\ may\ have\ reduced\ tentilla,$ which would indicate there was only one loss of tentilla at the branch leading to Apolemia.

SIMMAP Presence of tentilla - Physalia corrected



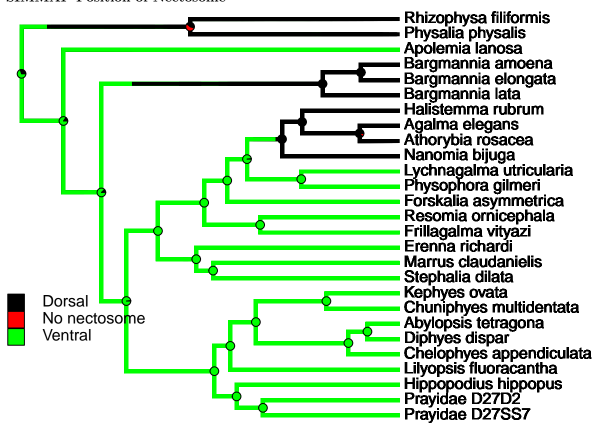
probabilities of states (1 Present, 0 Absent).

SIMMAP Presence of Nectophores

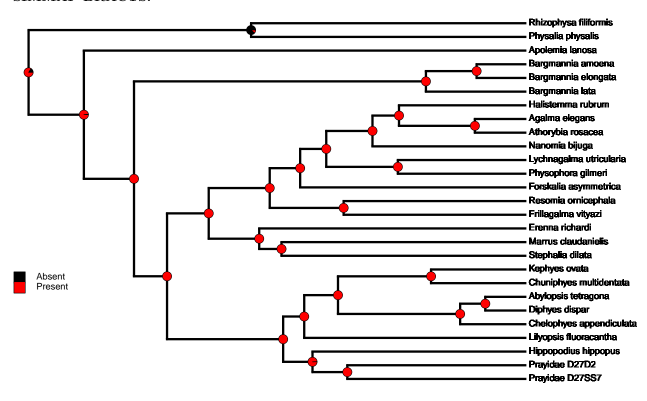


probabilities of states (1 Present, 0 Absent).

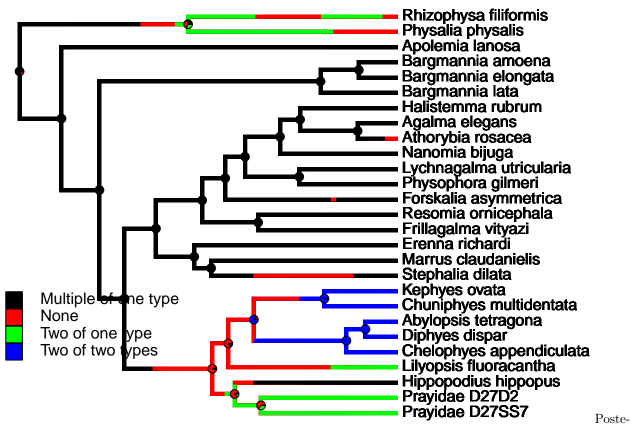
SIMMAP Position of Nectosome



SIMMAP BRACTS:

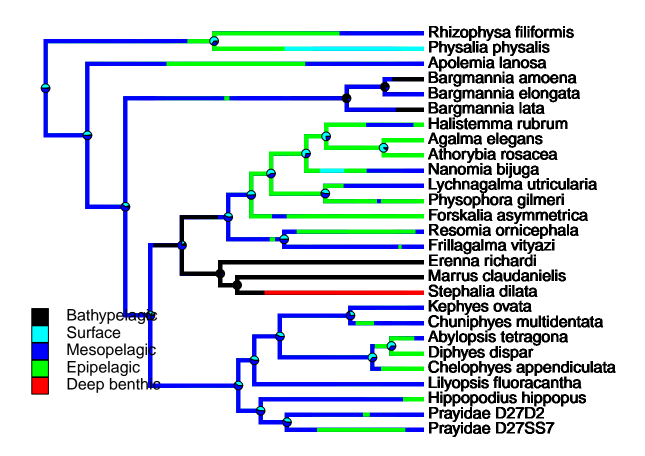


SIMMAP Nectophore number and types



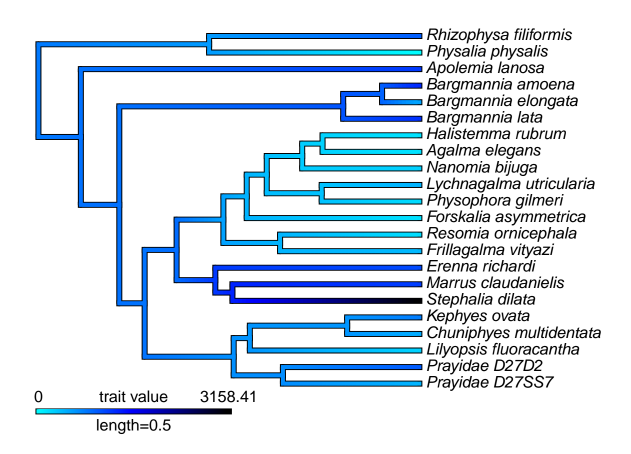
rior probabilities of states (1 Present, 0 Absent). It is not clear whether or not the stem group of siphonophores had a pneumatophore.

SIMMAP Habitat

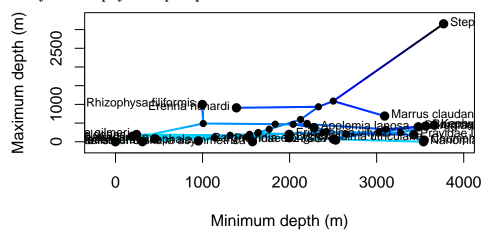


Continuous traits: Brownian motion reconstructions

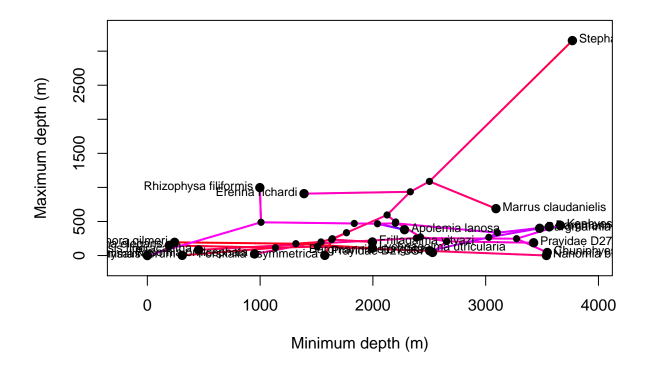
Median depth reconstruction:



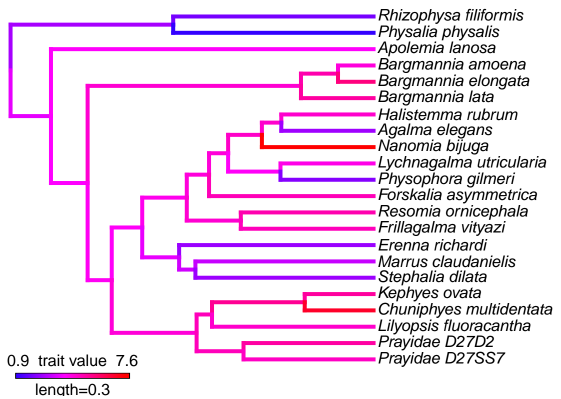
Bathymetrics phylomorphospace



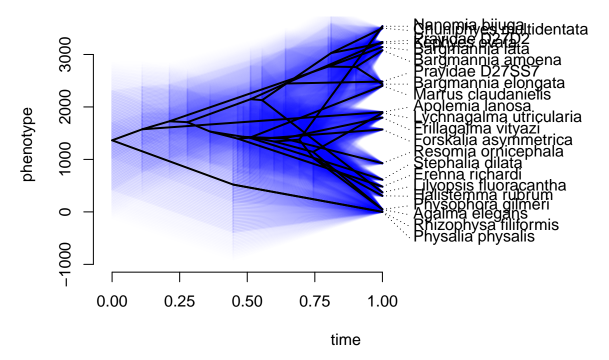
Bathymetrics phylomorphospace with time from root as color



Abundance of each taxon in the Monterey Bay sampling site

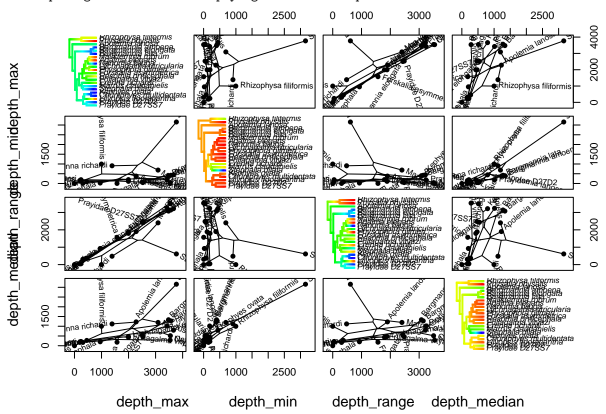


Uncertainty inclusive traitgram for depth range

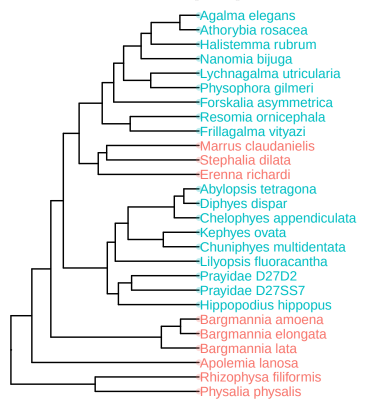


Bathymetrics scattergram array

Computing multidimensional phylogenetic scatterplot matrix...



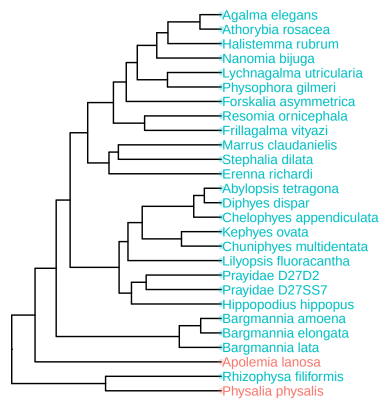
Distribution of sex across siphonophore taxa



Sex distribution

- Monoecious
- Dioecious

Tentilla presence across siphonophore taxa

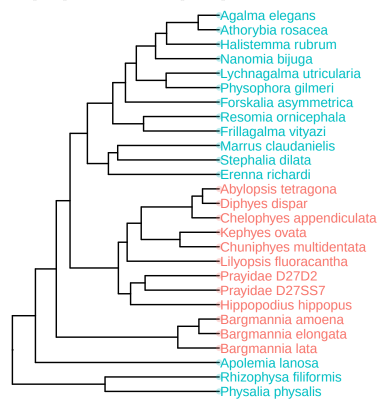


Tentilla

• 1

• 0

Palpon presence across siphonophore taxa



Palpons

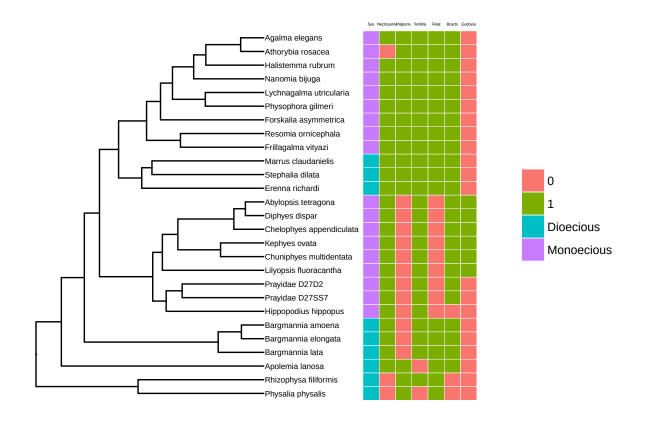
• 1

• 0

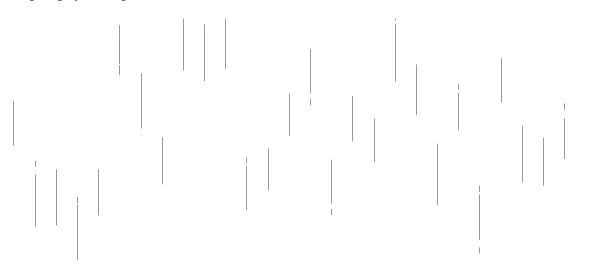
Overview grid of main binary traits

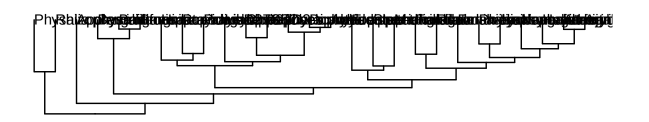
##		Sex	Nectosome	Palpons	Tentilla
##	Prayidae D27SS7	${\tt Monoecious}$	1	0	1
##	Prayidae D27D2	${\tt Monoecious}$	1	0	1
##	Hippopodius hippopus	${\tt Monoecious}$	1	0	1
##	Lilyopsis fluoracantha	${\tt Monoecious}$	1	0	1
##	${\tt Chelophyes\ appendiculata}$	${\tt Monoecious}$	1	0	1
##	Diphyes dispar	Monoecious	1	0	1
##	Abylopsis tetragona	Monoecious	1	0	1
##	Chuniphyes multidentata	${\tt Monoecious}$	1	0	1
##	Kephyes ovata	${\tt Monoecious}$	1	0	1
##	Stephalia dilata	Dioecious	1	1	1
##	Marrus claudanielis	Dioecious	1	1	1
##	Erenna richardi	Dioecious	1	1	1
##	Frillagalma vityazi	${\tt Monoecious}$	1	1	1
##	Resomia ornicephala	${\tt Monoecious}$	1	1	1
##	Forskalia asymmetrica	${\tt Monoecious}$	1	1	1
##	Physophora gilmeri	${\tt Monoecious}$	1	1	1
##	Lychnagalma utricularia	${\tt Monoecious}$	1	1	1
##	Nanomia bijuga	${\tt Monoecious}$	1	1	1
##	Athorybia rosacea	${\tt Monoecious}$	0	1	1
##	Agalma elegans	${\tt Monoecious}$	1	1	1
##	Halistemma rubrum	${\tt Monoecious}$	1	1	1
##	Bargmannia lata	Dioecious	1	0	1
##	Bargmannia elongata	Dioecious	1	0	1

##	Bargmannia amoena	Dioecious	1	0	1
	Apolemia lanosa	Dioecious	1	1	0
##	Physalia physalis	Dioecious	0	1	0
##	Rhizophysa filiformis	Dioecious	0	1	1
##		${\tt Pneumatophore}$	${\tt Bracts}$	${\tt Eudoxia}$	
##	Prayidae D27SS7	0	1	0	
##	Prayidae D27D2	0	1	0	
##	Hippopodius hippopus	0	0	0	
##	Lilyopsis fluoracantha	0	1	1	
##	Chelophyes appendiculata	0	1	1	
##	Diphyes dispar	0	1	1	
##	Abylopsis tetragona	0	1	1	
##	Chuniphyes multidentata	0	1	1	
##	Kephyes ovata	0	1	1	
##	Stephalia dilata	1	1	0	
##	Marrus claudanielis	1	1	0	
##	Erenna richardi	1	1	0	
##	Frillagalma vityazi	1	1	0	
	Resomia ornicephala	1	1	0	
	Forskalia asymmetrica	1	1	0	
##	Physophora gilmeri	1	1	0	
	Lychnagalma utricularia	1	1	0	
	Nanomia bijuga	1	1	0	
	Athorybia rosacea	1	1	0	
	Agalma elegans	1	1	0	
	Halistemma rubrum	1	1	0	
##	Bargmannia lata	1	1	0	
	Bargmannia elongata	1	1	0	
	Bargmannia amoena	1	1	0	
##	•	1	1	0	
	Physalia physalis	1	0	0	
	Rhizophysa filiformis	1	0	0	
,		_	· ·	•	



Depth phylo-barplot





Phylogenetic signal in the character data

Agnostic branch length tree generation:

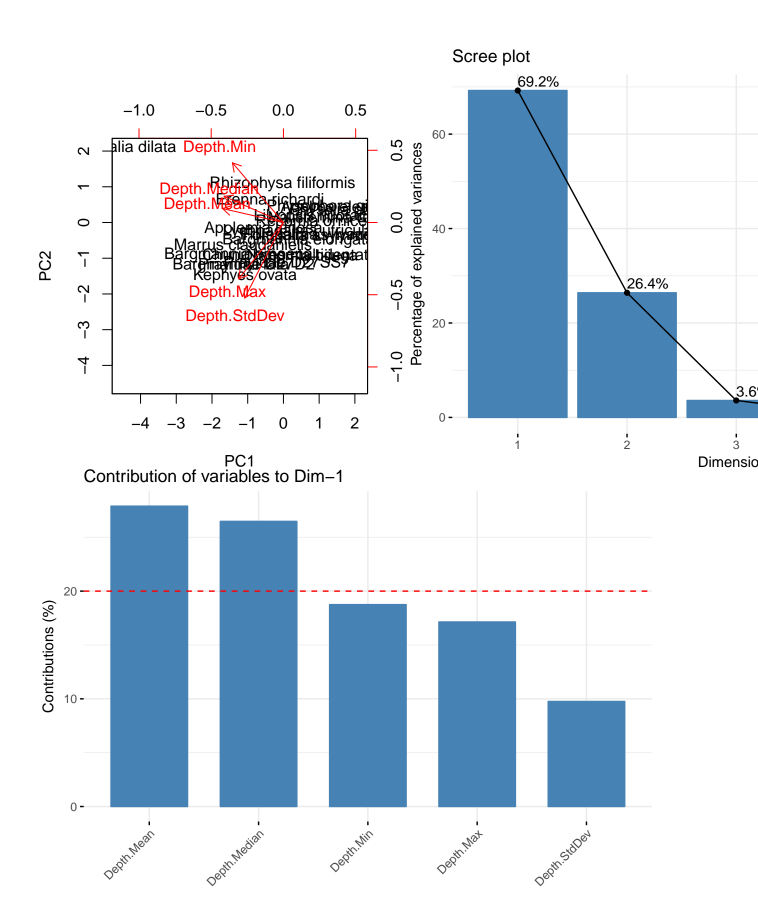
Phylogenetic signal in Binary Traits

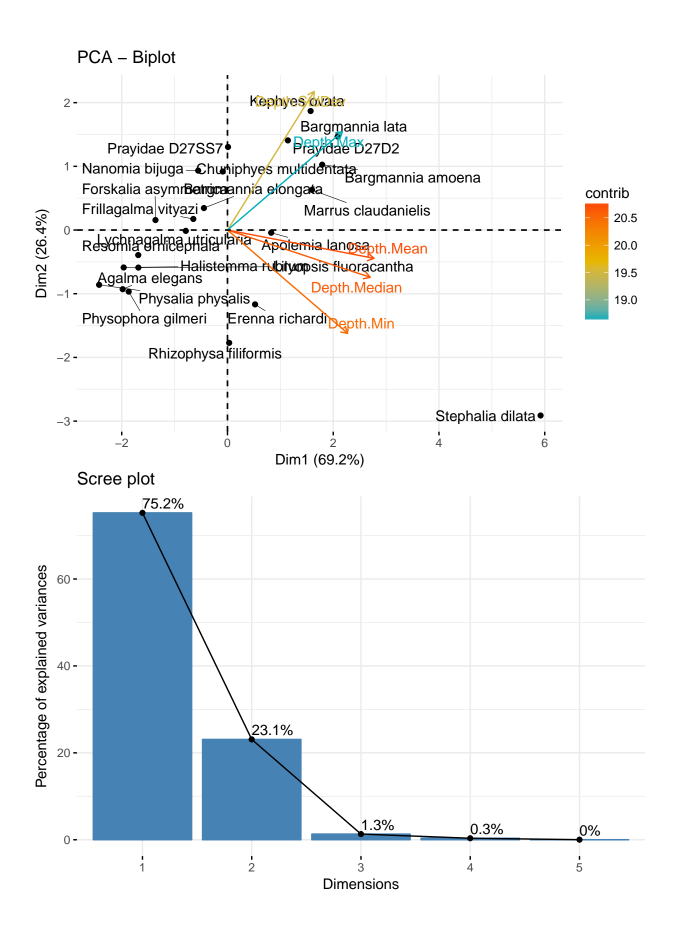
```
K PIC.variance.obs PIC.variance.rnd.mean
##
## Nectosome
                0.6773278
                                0.26773672
                                                       0.3507614
## Palpons
                3.3888711
                                0.10615819
                                                       0.8624120
## Tentilla
                2.1806411
                                0.07424251
                                                       0.2434057
## Pneumatophore 3.9000634
                                0.08521926
                                                       0.7768665
##
                PIC.variance.P PIC.variance.Z
## Nectosome
                         0.319
                                   -0.6680884
                         0.001
## Palpons
                                   -4.2989067
## Tentilla
                         0.003
                                   -1.5642236
## Pneumatophore
                         0.001
                                   -4.0370321
```

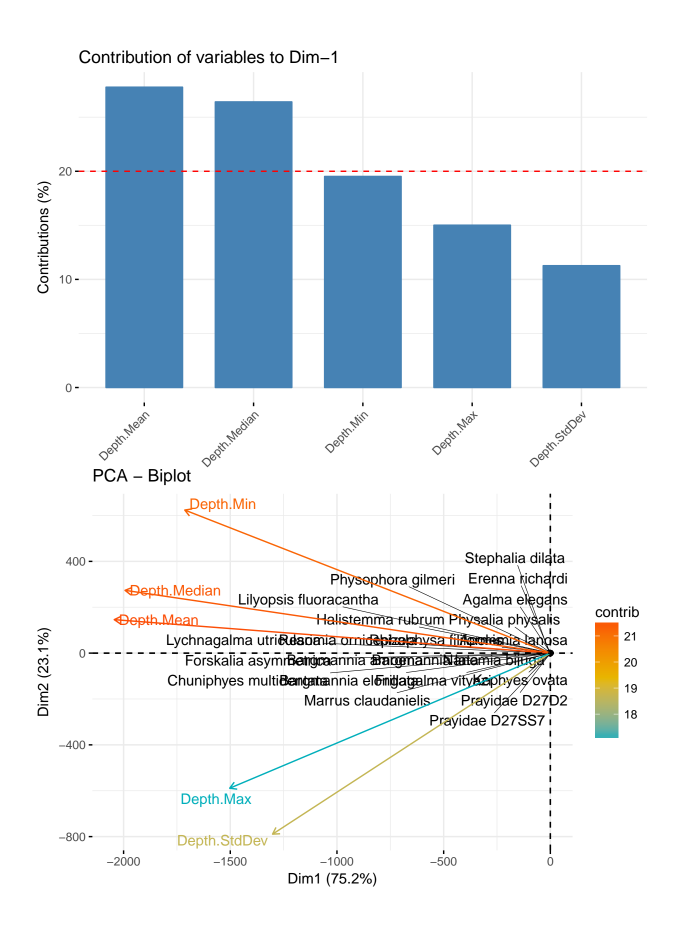
Phylogenetic signal in VARS bathymetrics

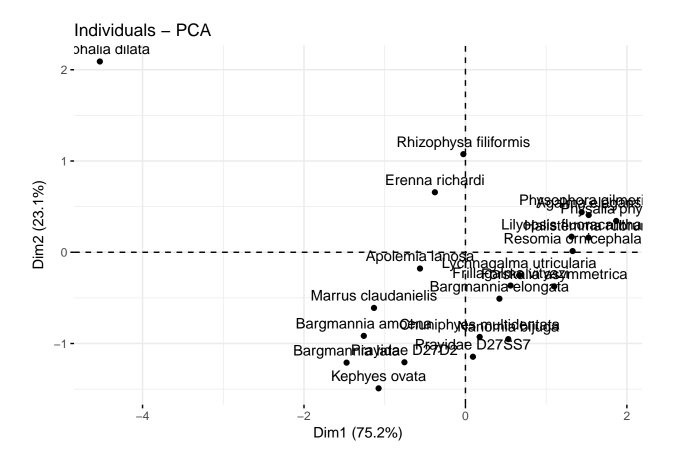
```
##
## Call:
## physignal(A = dpruned_data[, c(14, 15, 16, 18, 19)], phy = dpruned_tree)
##
##
##
##
## Observed Phylogenetic Signal (K): 0.7424
##
## P-value: 0.039
##
## Based on 1000 random permutations
##
                        K PIC.variance.obs PIC.variance.rnd.mean
## Depth.Median 0.7830816
                                  797529.1
                                                       1276559.4
## Depth.Mean
               0.7388410
                                  941754.2
                                                       1453919.6
## Depth.StdDev 0.5929636
                                 136382.4
                                                       164134.2
## Depth.Max
                                3265732.2
                                                       4754543.8
                0.7181495
## Depth.Min
                0.8396274
                                 723182.9
                                                       1258465.5
##
                PIC.variance.P PIC.variance.Z
## Depth.Median
                    0.065
                                  -1.0427676
## Depth.Mean
                        0.057
                                   -1.0214493
## Depth.StdDev
                        0.237
                                   -0.7314187
## Depth.Max
                        0.059
                                   -1.4636342
## Depth.Min
                        0.091
                                   -0.9867170
```

Phylogenetic PCA for bathymetrics









Phylogenetic ANOVA for depth

Each phylogenetic ANOVA compares variance of depth (continuous variable) within and between groups clustered by their categorical biological traits.

```
## [1] "Depth vs Sex Distribution"
## $F
   [1] 10.48139
##
## $Pf
   [1] 0.081
##
##
## $T
##
              Dioecious Monoecious
## Dioecious
               0.000000
                           3.237497
## Monoecious -3.237497
                           0.000000
##
   $method
##
##
   [1] "holm"
##
## $Pt
              Dioecious Monoecious
##
                   1.000
                              0.081
## Dioecious
## Monoecious
                   0.081
                              1.000
## [1] "Depth vs Nectosome"
```

```
## $F
## [1] 0.2358588
##
## $Pf
## [1] 0.784
##
## $T
##
            0
## 0 0.000000 -0.485653
## 1 0.485653 0.000000
##
## $method
## [1] "holm"
##
## $Pt
##
         0
## 0 1.000 0.784
## 1 0.784 1.000
## [1] "Depth vs Nectophore Types and Number"
## $F
## [1] 0.156464
##
## $Pf
## [1] 0.982
##
## $T
                         Multiple of one type None One of one type
## Multiple of one type
                                            NA
                                                 NA
## None
                                            NA
                                                 NA
                                                                  NA
## One of one type
                                            NA
                                                 NA
                                                                  NA
## Two of one type
                                            NA
                                                 NA
                                                                  NA
## Two of two types
                                            NA
                                                 NA
                                                                  NA
                         Two of one type Two of two types
## Multiple of one type
                                       NA
## None
                                       NA
                                                         NA
## One of one type
                                       NA
                                                         NA
## Two of one type
                                       NA
                                                         NA
## Two of two types
                                       NA
##
## $method
## [1] "holm"
##
## $Pt
                         Multiple of one type None One of one type
## Multiple of one type
                                            ΝA
                                                 NA
                                                                  NA
## None
                                            NA
                                                 NA
                                                                  NA
                                                 NA
                                                                  NA
## One of one type
                                            NA
## Two of one type
                                            NA
                                                 NA
                                                                  NA
## Two of two types
                                            NA
                                                                  NA
##
                         Two of one type Two of two types
## Multiple of one type
                                       NA
## None
                                       NA
                                                         NA
## One of one type
                                                         NA
```

```
## Two of one type
                                   NA
                                                    NA
## Two of two types
                                   NA
                                                    NA
## [1] "Depth vs Nectosome Position"
## [1] 0.237949
##
## $Pf
## [1] 0.909
##
## $T
##
              Dorsal
                       None
                                 Ventral
## Dorsal 0.0000000 0.2240508 -0.4988957
## None -0.2240508 0.0000000 -0.5640384
## Ventral 0.4988957 0.5640384 0.0000000
##
## $method
## [1] "holm"
##
## $Pt
##
          Dorsal None Ventral
## Dorsal
           1 1 1
              1
## None
                    1
                            1
## Ventral
             1
## [1] "Depth vs Palpons"
## $F
## [1] 0.05567054
##
## $Pf
## [1] 0.888
##
## $T
       0
## 0 0.0000000 0.2359461
## 1 -0.2359461 0.0000000
##
## $method
## [1] "holm"
##
## $Pt
##
       0 1
## 0 1.000 0.888
## 1 0.888 1.000
## [1] "Depth vs Tentilla"
## [1] 0.06009463
##
## $Pf
## [1] 0.856
##
## $T
##
            0
```

```
## 0 0.0000000 -0.2451421
## 1 0.2451421 0.0000000
##
## $method
## [1] "holm"
##
## $Pt
       0 1
##
## 0 1.000 0.856
## 1 0.856 1.000
## [1] "Depth vs Pneumatophore"
## [1] 0.1796993
##
## $Pf
## [1] 0.817
##
## $T
##
          0
## 0 0.0000000 -0.4239096
## 1 0.4239096 0.0000000
##
## $method
## [1] "holm"
## $Pt
       0
## 0 1.000 0.817
## 1 0.817 1.000
PGLS - Sex distribution
## -----
## | **Warning:
## | User reports suggest that this method may frequently
## | fail to find the ML solution. Please use with caution.
## phylolm(formula = depth_median ~ dp_sex, data = as.data.frame(cbind(depth_median,
##
      dp_sex)), phy = dpruned_tree, model = "BM", measurement_error = TRUE,
##
      boot = 100)
##
##
     AIC logLik
## 347.7 -169.9
##
## Parameter estimate(s) using ML:
## sigma2: 217361.9
## sigma2_error: 142822
##
## Coefficients:
## (Intercept)
                 dp_sex
## 192.7735 958.4204
```

```
## Generalized least squares fit by REML
##
     Model: dp_sex ~ depth_median
##
     Data: NULL
##
     Log-restricted-likelihood: -20.52555
##
## Coefficients:
## (Intercept) depth median
## 0.0978229509 0.0004267556
##
## Degrees of freedom: 22 total; 20 residual
## Residual standard error: 0.4176998
Software versions
This manuscript was computed on Fri Nov 03 16:03:02 2017 with the following R package versions.
R version 3.4.1 (2017-06-30)
Platform: x86_64-apple-darwin15.6.0 (64-bit)
Running under: macOS Sierra 10.12.2
Matrix products: default
BLAS: /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRblas.0.dylib
LAPACK: /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRlapack.dylib
locale:
[1] en_GB.UTF-8/en_GB.UTF-8/en_GB.UTF-8/C/en_GB.UTF-8/en_GB.UTF-8
attached base packages:
[1] grid
              parallel stats
                                   graphics grDevices utils
                                                                 datasets
[8] methods
              base
other attached packages:
 [1] bindrcpp_0.2
                      phylolm<sub>2.5</sub>
                                        geomorph_3.0.5
                                                         rgl_0.98.1
 [5] adephylo_1.1-10 ade4_1.7-8
                                        phylobase_0.8.4
                                                         geiger_2.0.6
 [9] phangorn_2.2.0
                      phytools_0.6-20
                                        picante_1.6-2
                                                         nlme_3.1-131
[13] vegan_2.4-4
                      lattice_0.20-35
                                       permute_0.9-4
                                                         ape_4.1
[17] hutan_0.5.0
                      FactoMineR_1.38
                                        factoextra_1.0.5 gridExtra_2.3
[21] seriation_1.2-2 fields_9.0
                                        maps_3.2.0
                                                         spam_2.1-1
[25] dotCall64_0.9-04 ggtree_1.8.2
                                        treeio 1.0.2
                                                         cowplot 0.8.0
                                                         digest_0.6.12
[29] xtable_1.8-2
                      jsonlite_1.5
                                        knitr_1.17
[33] magrittr 1.5
                      forcats_0.2.0
                                        stringr_1.2.0
                                                         dplyr 0.7.4
[37] purrr_0.2.3
                      readr_1.1.1
                                        tidyr_0.7.1
                                                         tibble_1.3.4
[41] ggplot2_2.2.1
                      tidyverse_1.1.1
loaded via a namespace (and not attached):
  [1] readxl 1.0.0
                              uuid 0.1-2
  [3] backports_1.1.1
                              fastmatch_1.1-0
  [5] plyr_1.8.4
                              igraph_1.1.2
  [7] lazyeval_0.2.0
                              sp_1.2-5
  [9] splines_3.4.1
                              rncl_0.8.2
                              htmltools_0.3.6
 [11] foreach_1.4.3
 [13] viridis_0.4.0
                              gdata_2.18.0
 [15] cluster_2.0.6
                              gclus_1.3.1
```

gmodels_2.16.2

[17] modelr_0.1.1

5		
	prettyunits_1.0.2	jpeg_0.1-8
	colorspace_1.3-2	rvest_0.3.2
	ggrepel_0.7.0	haven_1.1.0
	bindr_0.1	survival_2.41-3
	iterators_1.0.8	glue_1.1.1
	registry_0.3	gtable_0.2.0
	seqinr_3.4-5	kernlab_0.9-25
	prabclus_2.2-6	DEoptimR_1.0-8
	scales_0.5.0	mvtnorm_1.0-6
[37]	DBI_0.7	Rcpp_0.12.13
	plotrix_3.6-6	viridisLite_0.2.0
[41]	progress_1.1.2	spdep_0.6-15
[43]	flashClust_1.01-2	foreign_0.8-69
[45]	subplex_1.4-1	bold_0.5.0
[47]	mclust_5.3	deSolve_1.20
[49]	stats4_3.4.1	animation_2.5
[51]	htmlwidgets_0.9	httr_1.3.1
	gplots_3.0.1	fpc_2.1-10
	modeltools_0.2-21	pkgconfig_2.0.1
	reshape_0.8.7	XML_3.98-1.9
[59]	flexmix_2.3-14	deldir_0.1-14
	nnet_7.3-12	crul_0.4.0
	tidyselect_0.2.0	labeling_0.3
	rlang_0.1.2	reshape2_1.4.2
	munsell_0.4.3	cellranger_1.1.0
	tools_3.4.1	broom_0.4.2
	evaluate_0.10.1	yaml_2.1.14
	robustbase_0.92-7	caTools_1.17.1
	dendextend_1.5.2	mime_0.5
	whisker_0.3-2	taxize_0.9.0
	adegenet_2.1.0	leaps_3.0
	xml2_1.1.1	compiler_3.4.1
	curl_2.8.1	clusterGeneration_1.3.4
	RNeXML_2.0.7	stringi_1.1.5
	highr_0.6	trimcluster_0.1-2
	Matrix_1.2-11	psych_1.7.8
	msm_1.6.4	LearnBayes_2.15
[93]	combinat_0.0-8	data.table_1.10.4
	bitops_1.0-6	httpuv_1.3.5
	R6_2.2.2	TSP_1.1-5
	KernSmooth_2.23-15	codetools_0.2-15
	boot_1.3-20	MASS_7.3-47
	gtools_3.5.0	assertthat_0.2.0
	rprojroot_1.2	mnormt_1.5-5
	diptest_0.75-7	mgcv_1.8-22
	-	-
	expm_0.999-2	hms_0.3
	quadprog_1.5-5	coda_0.19-1
	class_7.3-14	rmarkdown_1.6
	rvcheck_0.0.9	ggpubr_0.1.5
	shiny_1.0.5	numDeriv_2016.8-1
[119]	scatterplot3d_0.3-40	<pre>lubridate_1.6.0</pre>

References

Dunn C., Pugh P., Haddock S. 2005. Molecular Phylogenetics of the Siphonophora (Cnidaria), with Implications for the Evolution of Functional Specialization. Systematic biology. 54:916–935.

Dunn C.W., Howison M., Zapata F. 2013. Agalma: an automated phylogenomics workflow. BMC Bioinformatics. 14:330.

Lartillot N., Lepage T., Blanquart S. 2009. PhyloBayes 3: A bayesian software package for phylogenetic reconstruction and molecular dating. Bioinformatics. 25:2286–2288.

Lartillot N., Philippe H. 2004. A bayesian mixture model for across-site heterogeneities in the amino-acid replacement process. Molecular Biology and Evolution. 21:1095–1109.

Yu G., Smith D.K., Zhu H., Guan Y., Lam T.T.-Y. 2016. ggtree: an r package for visualization and annotation of phylogenetic trees with their covariates and other associated data. Methods in Ecology and Evolution.

Species	Mean insert size (bp)	Insert size stan- dard devia- tion	Number of read pairs	Percent kept after rRNA removal	Percent kept after as- sembly	Adapter fails	Quality fails	Base Compo- sition fails	Total assembled transcripts	Coding tran- scripts
Marrus	253.03	53.30	35866092	96.8	62.1	113298	15594637	4981676	105439	22511
clau- danielis										
Apolemia rubriversa	290.81	80.64	19228660	91.7	77.8	40594	5179800	516100	87701	17540
Chuniphye multi-	es284.46	35.74	17999147	77.0	74.3	24263	4941872	335297	84341	22084
dentata Apolemia	167.41	53.46	16761717	81.3	79.4	1332063	3062035	41943	51470	14752
sp Apolemia	167.74	55.30	18444476	96.4	80.9	923922	3882968	439569	70184	15579
lanosa Bargmann elon-	i 2 71.43	40.17	40008913	79.8	78.5	342825	8084832	2017709	152019	23661
gata Diphyes	223.14	58.73	80000000	78.7	63.4	14412120	21322624	690159	210406	50868
dispar Aiptasia	183.18	53.88	74558341	98.4	79.3	210730	23826421	307520	83950	33523
pallida Stephalia	267.82	43.33	30953585	98.0	63.9	86068	15731787	1435119	107925	23984
dilata Physalia	233.10	54.37	36481773	96.6	88.5	18553	5491979	259497	74994	23705
physalis Bargmann	i a 61.57	53.77	20195498	84.6	80.1	1049110	3996085	274520	66726	17975
amoena Frillagalma	a283.11	44.91	79902051	54.0	70.4	668400	13591301	5558767	181508	29293
vityazi Alatina	173.96	60.33	96259870	52.1	77.9	1236251	15771112	468662	166584	28743
alata Clytia hemis- phaer-									11476	6642
pnaer- ica Ectopleura	220.05	37.75	109024653	95.9	74.6	432309	40629127	897590	84034	28015
larynx Athorybia		93.65	28696930	99.9	86.9	691916	4026467	633700	100839	24543
rosacea Forskalia		35.04	25275184		65.1	41922	7463840	841930	82483	18419
asym- metrica										
Prayidae D27D2	267.32	45.59	38233199	85.2	66.4	59773	15338562	1878732	144909	28065
Lilyopsis fluora-	259.39	39.08	51855968	76.5	65.3	72416	19809124	1388875	114662	29854
cantha Hydractini symbi- olongi-	ial 75.22	57.84	60462724	49.8	85.2	162430	6678373	124392	71450	24639
carpus Rhizophys fili-	a269.74	46.52	26937827	92.5	65.3	66010	13238701	361376	110109	24833
formis Agalma	238.94	57.54	40007833	99.6	86.0	31957	7177806	748977	122053	26601
elegans Chelophye appen-	s275.93	39.43	21103284	89.0	74.2	63548	6777022	344708	110612	32309
diculata Prayidae	303.39	50.83	25233164	66.7	65.6	39061	8616289	347189	94917	24211
D27SS7 Rudjakovi:	a213.88	89.71	20582477	99.1	79.7	489520	3936086	1546686	91695	24184
sp Abylopsis tetrag-	292.16	38.32	21575176	73.3	73.8	38764	5759144	370604	102946	25268
ona Nematoste vecten-	ella								26511	18080
sis Hippopodi hippo-	u 2 66.99	45.11	35638254	84.5	66.3	56152	15490291	535026	137341	35031
pus Physophor gilmeri	a214.97	93.78	19847377	99.3	81.3	936617	3464079	1007839	52639	17381
gumeri Bargmann lata	i 4 66.71	62.80	16984483	98.7	71.1	3118496	4227140	868584	59051	16451
Nanomia	296.69	44.20	39983229	85.7	75 .3 5	155093	10854240	795226	165792	28927
bijuga Erenna richardi	291.53	36.76	29776547	64.2	66.6	39775	8958200	1245523	89423	18236
richardi Physonect sp	261.09	35.96	38243543	24.3	66.7	12494	4517982	274838	57746	12596