

# Comparison of the Venom Peptides and Their Expression in Closely Related *Conus* Species: Insights into Adaptive Post-speciation Evolution of *Conus* Exogenomes

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**Data deposition:** Transcriptome Shotgun Assembly projects of *Conus tribblei* and *Conus lenavati* have been deposited at DDBJ/EMBL/GenBank under the accessions GCVM00000000 and GCVH00000000, respectively. The versions described in this article are the first versions: GCVM01000000 (*C. tribblei*) and GCVH01000000 (*C. lenavati*). The COI sequences of the specimens of *C. tribblei* and *C. lenavati* are available from GenBank under accessions KR107511–KR107522 and KR336542.

## Abstract

Genes that encode products with exogenous targets, which comprise an organism's "exogenome," typically exhibit high rates of evolution. The genes encoding the venom peptides (conotoxins or conopeptides) in *Conus sensu lato* exemplify this class of genes. Their rapid diversification has been established and is believed to be linked to the high speciation rate in this genus. However, the molecular mechanisms that underlie venom peptide diversification and ultimately emergence of new species remain poorly understood. In this study, the sequences and expression levels of conotoxins from several specimens of two closely related worm-hunting species, *Conus tribblei* and *Conus lenavati*, were compared through transcriptome analysis. Majority of the identified putative conopeptides were novel, and their diversity, even in each specimen, was remarkably high suggesting a wide range of prey targets for these species. Comparison of the interspecific expression patterns of conopeptides at the superfamily level resulted in the discovery of both conserved as well as species-specific expression patterns, indicating divergence in the regulatory network affecting conotoxin gene expression. Comparison of the transcriptomes of the individual snails revealed that each specimen produces a distinct set of highly expressed conopeptides, reflecting the capability of individual snails to fine-tune the composition of their venoms. These observations reflect the role of sequence divergence and divergence in the control of expression for specific conopeptides in the evolution of the exogenome and hence venom composition in *Conus*.

**Key words:** conotoxin, species-specific expression pattern, diversification, conopeptide.

## Introduction

The genes that encode the venom peptides in *Conus* (conopeptides or conotoxins) belong to a class of genes whose products act on exogenous targets (Olivera 2006). These genes, which may be referred to as "exogenes" and which collectively make up an organism's "exogenome" (Olivera 2006), share a common characteristic, namely their rapid evolution and diversification. The accelerated pace of evolution of conopeptide genes (Duda and Palumbi 2000; Conticello et al. 2001; Espiritu et al. 2001; Duda 2008; Puillandre et al. 2010; Chang and Duda 2012), together with the high diversity of species in *Conus sensu lato* (Puillandre et al. 2014, 2015), and

the evolution of biochemical mechanisms that introduce a variety of post-translational modifications to the peptides (Dutertre et al. 2013) collectively account for the remarkable chemical diversity of conopeptides in nature.

A number of mechanisms have been proposed to explain the rapid evolution of conopeptides. Gene duplication and positive selection have been shown to be important factors (Duda and Palumbi 2000; Duda 2008; Duda and Remigio 2008; Puillandre et al. 2010; Chang and Duda 2012, 2014). Several molecular mechanisms such as recombination (Espiritu et al. 2001), hypermutation (Olivera et al. 1999; Espiritu et al. 2001), presence of error-prone DNA polymerase, and even

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special molecules protecting cysteine codons (Conticello et al. 2001) have been suggested, although solid evidence in support of these mechanisms is still lacking.

Studies have also shown that the expression of conopeptides differs among *Conus* species with different diets: Worm-, mollusk-, and fish-hunting (Conticello et al. 2001; Duda and Palumbi 2004) indicating the importance of diet as a factor influencing the evolution of venom components. Furthermore, the diet of each vermivorous *Conus* species co-occurring in a habitat is dominated by a different type of polychaete where each *Conus* species have specialized on different prey taxa (Kohn 2001). Therefore, there seems to be a dietary specialization within worm-hunting species (Duda et al. 2001; Kohn 2001). In addition, *Conus* species that feed exclusively on specific types of prey express less diverse conopeptides than those with a broader range of prey (Remigio and Duda 2008; Elliger et al. 2011).

Little is known about how gene expression patterns differ between closely related species of *Conus*. In primates, variations in gene expression patterns are known to contribute to interspecies phenotypic variation and adaptation (Romero et al. 2012). Although the expression of majority of genes in closely related species is under purifying selection (Khaitovich et al. 2006), the expression of some genes has evolved more rapidly giving rise to species-specific gene expression patterns (Enard et al. 2002). A number of studies have shown that closely related species of *Conus* having similar diet also tend to produce conopeptides with similar structure and molecular targets (Espiritu et al. 2001; Olivera and Teichert 2007). For example, a few orthologous genes of the O1- and A-superfamilies were found to be expressed among closely related species of *Conus* (Duda and Remigio 2008; Chang and Duda 2012, 2014).

A crucial limitation of these previous studies was the insufficient sampling of expressed genes or conotoxin superfamilies partly owing to the technical limitations of the traditional cDNA library-based approaches (Duda and Remigio 2008; Chang and Duda 2012, 2014). Recent studies using next generation sequencing have shown the presence of 11–36 conopeptide gene superfamilies in each species (Hu et al. 2011, 2012; Lluisma et al. 2012; Terrat et al. 2012; Dutertre et al. 2013; Lavergne et al. 2013; Robinson et al. 2014; Barghi et al. 2015). In addition, the advent of high-throughput sequencing has made possible not only the identification but also the quantification of expression levels of conopeptide transcripts. Therefore, high-throughput sequencing could be leveraged to obtain deeper insights into the expression patterns of conopeptides and identifying the orthologous genes in all expressed conopeptide gene superfamilies. Similar to those in primate species, the differential expression of orthologous conotoxin exogenes may also contribute to the species-specific gene expression patterns and differentiation of venom composition in closely related species of *Conus*.

In addition to the interspecific variation in venom composition, characterization of intraspecific variability of the venom components may also provide clues to adaptive ability of the species as a whole. High intraspecific variability in the masses of venom peptides has been observed among the individuals of worm-, mollusk- (Davis et al. 2009), and fish-hunting (Jakubowski et al. 2005; Dutertre et al. 2010; Chun et al. 2012) cone snails. It was shown recently that variable post-translational processing could account for the observed large number (thousands) of peptide fragments (Dutertre et al. 2013). The expression of toxins in individuals of venomous animals such as snakes and scorpions varies with a number of factors such as sex, age, geographic location (Daltry et al. 1998; Menezes et al. 2006; Abdel-Rahman et al. 2009), and diet (Barlow et al. 2009; Casewell et al. 2009; Gibbs et al. 2013). The factors influencing the intraspecific variation in the expression patterns of conopeptides are still poorly known, and transcriptomic analysis of the venom duct of several individuals will be essential in characterizing such variability.

This study aims to investigate the expression and diversification of conopeptides in closely related species to provide insights into the processes that contribute to inter- and intraspecific differences in the venom composition and facilitate the evolution of *Conus* exogenome. This is the first comprehensive inter- and intraspecific transcriptome analysis of the venom duct in closely related *Conus* species using next generation sequencing. In this study, the expression patterns of conopeptide gene superfamilies were evaluated in several specimens of the closely related worm-hunting species, *Conus tribblei* and *Conus lenavati*. These two species belong to the same clade (VIII) (Espiritu et al. 2001). Recently, Puillandre et al. (2014) assigned these species to the subgenus *Splinoconus*. *Conus coffeae* and *Conus glans*, members of subgenus *Leporiconus*, the most closely related clade to the subgenus *Splinoconus* (Puillandre et al. 2014), are known to prey on errant polychaetes, mostly eunicidae (Duda et al. 2001). Furthermore, *Conus papuensis* another member of the subgenus *Splinoconus* (Puillandre et al. 2015) is also known to prey on polychaetes (Tucker and Tenorio 2009). As closely related *Conus* species generally have similar diet (Espiritu et al. 2001; Puillandre et al. 2014), *C. tribblei* and *C. lenavati* may also prey on polychaetes.

## Materials and Methods

### Sample Collection

A total of eight specimens of *C. tribblei* (~6 cm shell length) and five specimens of *C. lenavati* (~5–8 cm shell length) were collected using tangle nets from depth of 96–136 m in Sogod, Cebu province in the Philippines. The venom duct of each specimen was dissected, stored separately in RNAlater (Ambion, Austin, TX), and kept at  $-20^{\circ}\text{C}$ . The whole body

tissue and the shell of each specimen were stored in 95% ethanol and kept at room temperature.

### Phylogenetic Analysis

DNA was extracted from a small piece of the foot tissue of each specimen using DNeasy blood and tissue kit (Qiagen, USA) according to manufacturer's recommendations and stored at  $-20^{\circ}\text{C}$ . Using universal primers: LCO1490 and HCO2198 (Folmer et al. 1994), a fragment of cytochrome oxidase c subunit 1 (COI) gene segment of mitochondrial DNA was amplified. All PCR reactions were performed in 30  $\mu\text{l}$  containing 1  $\times$  Titanium Taq PCR Buffer, 0.4 mM dNTP, 0.2 mM of each primer, 0.15  $\mu\text{l}$  Titanium Taq DNA polymerase, and 0.5–4 ml of DNA template (total concentration of 100 ng in each PCR reaction). The PCR amplification consisted of one cycle of 1 min at  $94^{\circ}\text{C}$ ; five cycles of 1 min at  $94^{\circ}\text{C}$ , 1.5 min at  $45^{\circ}\text{C}$  and 1.5 min at  $72^{\circ}\text{C}$ ; 35 cycles of 1 min at  $94^{\circ}\text{C}$ , 1.5 min at  $50^{\circ}\text{C}$  and 1 min at  $72^{\circ}\text{C}$ , and a final cycle of 5 min at  $72^{\circ}\text{C}$  (Hebert et al. 2003). Each PCR product was gel purified using GF-1 AmbiClean Kit (vivantis) and sequenced in both directions using forward and reverse primers on an ABI 3730XL Capillary sequencer (Applied Biosystems). The manual inspection and cleaning of the sequences based on the chromatograms were performed using BioEdit version 7.0.0 (Hall 1999). Multiple sequence alignment was performed on the COI sequences of *C. tribblei* and *C. lenavati* specimens and other members of the subgenus *Splinoconus* downloaded from the National Center for Biotechnology Information (NCBI) (supplementary table S1, Supplementary Material online) using MEGA 5.2 (Tamura et al. 2011). Bayesian analysis was performed using six substitution categories, a gamma-distributed rate variation across sites approximated in four discrete categories, and a proportion of invariable sites in two parallel runs using MrBayes (Ronquist et al. 2012) each consisting of six Markov chains of 5 million generations with a sampling frequency of one tree in every 10,000 generations, and the chain temperature was set to 0.02. Convergence was evaluated using Tracer1.4.1 (Rambaut and Drummond 2007) to check that all effective sample size values exceeded 200. After stabilization of the log-likelihood scores, the first 25% of trees were omitted as burn-in, and a consensus tree was calculated. MrBayes was run on the Cipres Science Gateway (<https://www.phylo.org/portal2/>, last accessed June 15, 2015) (Miller et al. 2010) using MrBayes 3.2.2 on XSEDE.

### RNA-Sequencing and Transcriptome Assembly

The mRNA extraction was performed individually on the venom ducts of three specimens of *C. tribblei* (*C. tribblei*\_0317\_1 [trib1, female], *C. tribblei*\_0317\_3 [trib3, male], and *C. tribblei*\_0106\_21 [trib21, male]) and three specimens of *C. lenavati* (*C. lenavati*\_0317\_1 [lena1, male], *C. lenavati*\_0317\_2 [lena2, female], and *C. lenavati*\_0317\_3 [lena3, male]) (fig. 1) using Dynabeads mRNA DIRECT kit (Invitrogen

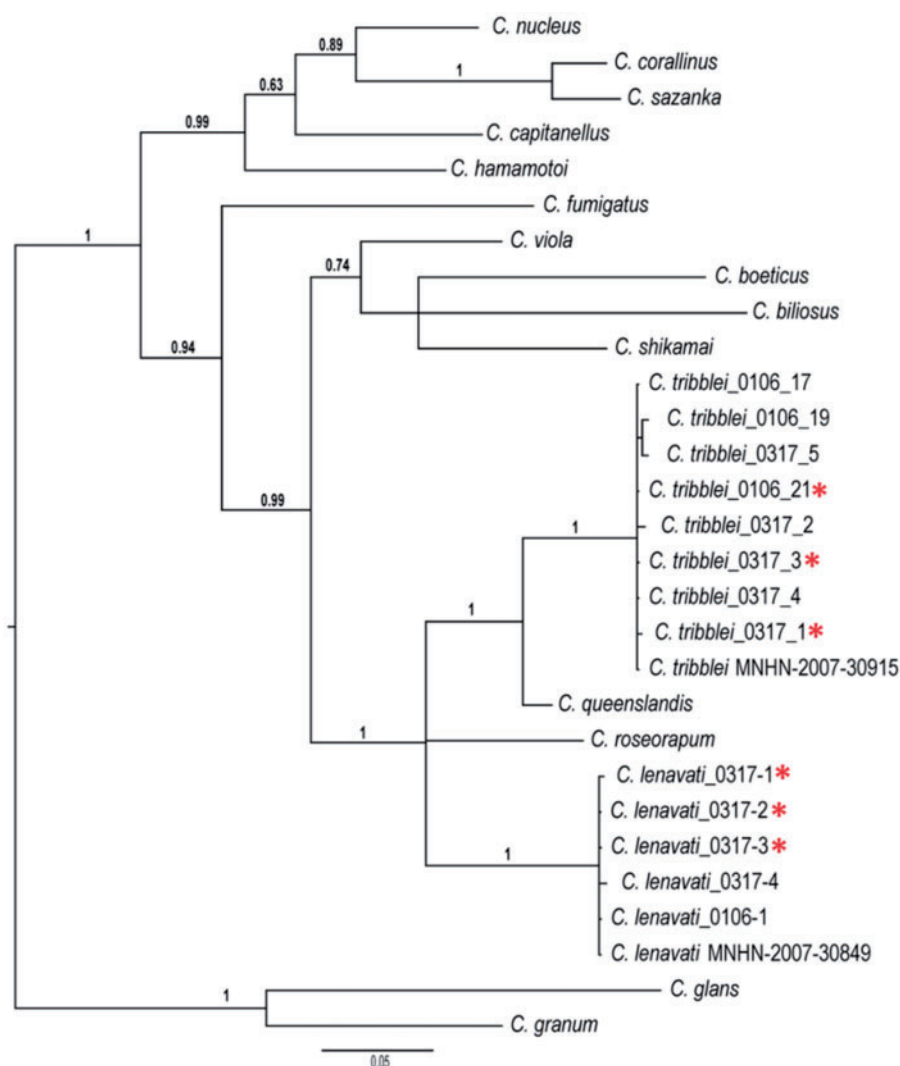
Dynal AS Oslo, Norway). For a more efficient extraction, each venom duct was homogenized using 0.5-mm Zirconia/Silica beads (Biospec Products, Inc.) in a bead beater (Precellys, Berlin Technologies), and the rest of the protocol was performed according to the manufacturer's recommendations. A cDNA library was constructed for each of the specimens, fragments with the average insert size of 200 bp were selected, and the libraries were multiplexed and paired-end (PE) sequenced using Illumina HiSeq 2000.

The quality of 90-bp long PE reads generated for each specimen was evaluated using FastQC software v0.10.1 ([www.bioinformatics.babraham.ac.uk/projects/fastqc/](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/), last accessed June 15, 2015). Trimming of the low quality bases ( $<20$  phred score) at 3'- and 5'-ends of the reads and removal of the low quality reads (reads in which less than 90% of bases had phred score  $\geq 20$ ) were performed using FASTX-Toolkit version 0.0.6 (Pearson et al. 1997), and the reads with no pairs were excluded using an in-house Python script. For each species, the sequencing reads from the three specimens were combined into a single data set and *de novo* assembled using Trinity (Grabherr et al. 2011) with default settings to generate the "reference assembly."

### Conopeptide Identification and Superfamily Classification

The conopeptide sequences in NCBI's nr protein database, UniProtKB/Swiss-Prot database (UniProt Consortium 2015), and ConoServer (Kaas et al. 2012) were downloaded and pooled in a data set. The putative conopeptides identified by Barghi et al. (2015) were also added, the redundant sequences of the data set were removed, and the remaining sequences were formatted into the "reference conopeptide database" using formatdb software of BLAST (Basic Local Alignment Search Tool) 2.2.29+ (Altschul et al. 1990). To identify putative conopeptides, the sequences in the reference assembly of *C. tribblei* and *C. lenavati* were searched against the reference conopeptide database using BLASTX. In each BLASTX result, the sequences with significant hits ( $e$  value  $< 10^{-5}$ ) were translated according to the identified reading frame and were manually inspected. The mature regions of the conopeptides were predicted using ConoPrec (Kaas et al. 2012), the redundant and truncated transcripts were excluded from the data set, and the good quality putative conopeptide precursors were collected into the "tribblei conopeptide data set" and "lenavati conopeptide data set."

The classification of the conopeptides in the tribblei conopeptide data set and lenavati conopeptide data set into gene superfamilies was performed as described in Barghi et al. (2015). Briefly, conopeptides were assigned into gene superfamilies based on the identification of the two highest-scoring full-length conopeptide precursor hits in the BLASTX search (in which the sequences in the reference assemblies of *C. tribblei* and *C. lenavati* were used to search for similar sequences in the reference conopeptide database). The signal regions of



**FIG. 1.**—Phylogenetic relationship of the species of subgenus *Spinoconus*. The phylogenetic tree is inferred from the partial COI sequences using Bayesian analysis. Posterior probabilities are indicated for each node. The specimens of *C. tribblei* and *C. lenavati* used for the transcriptome analysis are marked with red asterisks.

the putative conopeptide precursors were predicted using SignalP 4.1 (Petersen et al. 2011), and percentage sequence identity (PID) between the signal region of each putative conopeptide precursor and the highly conserved signal sequence of the known conopeptide gene superfamilies was computed using MatGAT 2.02 (Campanella et al. 2003). Originally, the PID value of 76 was chosen as the threshold for superfamily assignment (Kaas et al. 2010). However, some conopeptide gene superfamilies have less conserved signal regions. Therefore, based on the average PID of members of each superfamily, specific PID threshold values were set for different conopeptide gene superfamilies (Barghi et al. 2015). If the PID of a conopeptide's signal region was above the threshold of a superfamily, the conopeptide was assigned to that gene superfamily. Otherwise, the conopeptide was classified as a new conopeptide group. The assigned name of each new group

was "SF-" plus Arabic numbers such as "01." Multiple sequence alignment was performed on the precursors of the putative conopeptides and the reference conopeptide sequences using ClustalX version 2.1 (Larkin et al. 2007) followed by manual refinement using BioEdit.

### Comparison of Conopeptide Identity

For each conopeptide gene superfamily identified in both *C. tribblei* and *C. lenavati*, the conopeptides' mature regions were aligned using ClustalX, the pairwise distances were computed using MEGA, and the PID was computed for every conopeptide pair using the following formula:

$$\text{percentage sequence identity} = \frac{\text{identical residues} \times 2}{\text{sum of the length of a conopeptide pair}}$$

For every conopeptide in *C. tribblei*, the conopeptide in *C. lenavati* with the highest PID (and belonging to the same superfamily) was chosen as the most similar match, and the same criterion was applied for choosing the most similar match for the conopeptides in *C. lenavati*.

### Test of Positive Selection for Orthologous Conopeptide Genes

To identify the orthologous conopeptide genes in *C. tribblei* and *C. lenavati*, the nucleotide sequences of the conopeptide precursors of each superfamily identified in both species were aligned using ClustalX, and the best-fitted model was chosen using MEGA. The phylogenetic tree of the conopeptides of each superfamily was constructed using maximum-likelihood method based on the best-fitted model with bootstrap support value based on 500 replicates using MEGA. Those conopeptides that occurred in clades with strong bootstrap support containing only two sequences, one from *C. tribblei* and the other one from *C. lenavati*, and had identical cysteine framework were considered orthologous. For the superfamilies containing only one conopeptide in *C. tribblei* and one conopeptide in *C. lenavati*, the sequences were considered orthologs if, in addition to having identical length and cysteine framework, the PID of the mature region was above 78%. The pairwise estimates of nonsynonymous substitutions per nonsynonymous sites ( $d_N$ ) and synonymous substitutions per synonymous sites ( $d_S$ ) for the mature regions of the orthologous conopeptide pairs were computed using the maximum-likelihood approach implemented in PAML (runmode = -2, CodonFreq = 2) (Yang 1997).

### Shannon's Diversity Index of Venom Conopeptides

Diversity of the conopeptides in the tribblei and lenavati conopeptide data sets was computed as Shannon's diversity index,  $H'$ , where  $R$  is the number of conopeptide gene superfamilies and  $p_i$  is the proportion of conopeptides belonging to the  $i$ th superfamily in the data set. The evenness of conopeptide data sets was computed as Shannon's equitability,  $E_{H'}$ , whereas  $S$  is the data set richness based on the number of conopeptide gene superfamilies:

$$H' = - \sum_{i=1}^R p_i \ln(p_i) \quad E_{H'} = H' / \ln S$$

### Quantification of the Transcript Expression Level

In the recent transcriptome studies of the venom duct of several *Conus* species, the expression level of each conopeptide has been determined by the total number of sequencing reads generated from each conopeptide transcript (Hu et al. 2012; Terrat et al. 2012; Dutertre et al. 2013; Jin et al. 2013; Robinson et al. 2014). However, the transcript length and the sequencing depth affect the number of reads generated

from each transcript (Mortazavi et al. 2008). Therefore, for a more accurate comparison of the expression levels among conopeptides with different lengths, normalization of the number of reads by the transcript length and sequencing depth was performed (Mortazavi et al. 2008; Trapnell et al. 2010).

To determine how many of the conopeptides were expressed in each specimen of *C. tribblei* and *C. lenavati*, the sequencing reads of each specimen were aligned separately to the tribblei and lenavati reference assemblies, respectively, using Bowtie 1.0.1 (Langmead and Salzberg 2012). For each specimen, the alignment of the reads to the transcripts were visualized in Tablet (Milne et al. 2012) to ensure that reads align to the entire length of conopeptide precursor or at least to the full length of the mature region. Only those transcripts where the mature regions were entirely covered by reads were considered as "expressed." For each specimen, the number of reads representing a transcript was estimated as the maximum-likelihood abundances using expectation-maximization algorithm by RSEM (Li and Dewey 2011) using the default settings, and was represented as the "expected count." To enable comparison of the expression levels of transcripts among specimens, trimmed mean of M values (TMM) normalization factor was computed using edgeR (Robinson and Oshlack 2010). Then, the "effective library size" for each specimen was computed by normalizing the library size of each specimen (total number of aligned reads) using the TMM normalization factor. The "effective length" of each transcript which is the mean number of positions along the transcript sequence that a read may start (Li and Dewey 2011) was also computed using RSEM. Finally, the expression level of each transcript was computed as TMM-normalized Fragments Per Kilobase of transcript per Million fragments mapped (FPKM) (Trapnell et al. 2010; Haas et al. 2013) using the following formula:

$$\text{TMM-normalized FPKM} = \frac{\text{expected count} \times 10^9}{\text{effective length} \times \text{effective library size}}$$

For each specimen, the expression level of each gene superfamily is the sum of expression levels (TMM-normalized FPKM) of the conopeptides belonging to that superfamily represented as "sum TMM-normalized FPKM." As for each species, the expression level of each gene superfamily is the average of the expression levels of that superfamily among three specimens represented as "mean TMM-normalized FPKM."

### Functional Annotation

The functional annotation of the reference assemblies of *C. tribblei* and *C. lenavati* was performed using the pipeline version of BLAST2GO software (B2G4Pipe) (Götz et al. 2008). The transcripts of the reference assemblies were searched

against the UniProtKB/Swiss-Prot database using BLASTX, and the results containing the significant hits ( $e$  value  $< 10^{-5}$ ) were loaded to the pipeline. The GO terms for each blast hit were extracted from the local B2G MySQL database and assigned to each transcript. After refinement of the annotation results by GO-Slim function, the plot of transcripts GO classifications was constructed using Web Gene Ontology Annotation Plot (WEGO) software (Ye et al. 2006).

### Correlation Analysis of Transcripts' Expression Levels

The Pearson's correlation coefficient was computed for the expression levels of conopeptides and gene superfamilies among specimens of either *C. tribblei* or *C. lenavati* using "TMM-normalized FPKM" and "sum TMM-norm FPKM" values, respectively. In order to validate the correlation in the expression patterns of conopeptide superfamilies between *C. tribblei* and *C. lenavati*, the expression levels of conopeptide superfamilies, mean TMM-normalized FPKM, were used for the Pearson's correlation coefficient test. Additionally, to confirm the correlation of expression levels of the orthologous conopeptide genes between *C. tribblei* and *C. lenavati*, the mean expression level of each orthologous conopeptide was computed among specimens of each species, and these values were used for the correlation test. In all the correlation analyses, the expression level values were transformed to logarithmic scale ( $\log_2$ ), and the significance levels were calculated using Bonferroni correction.

## Results

### Taxonomic Identification of the Specimens and Their Phylogenetic Relationships

The phylogenetic reconstruction of the relationship of COI sequences of members of the subgenus *Splinoconus* (Puillandre et al. 2014) (supplementary table S1, Supplementary Material online) showed that *C. tribblei*, *Conus queenslandis*, *Conus roseorapum*, and *C. lenavati* clustered together in a clade (fig. 1). Specifically, *C. tribblei* and *C. lenavati* formed two distinct branches of this clade whereas all the specimens of each species were grouped together.

### Transcriptome Sequencing and Assembly

The mRNA sequencing of *C. tribblei* specimens generated 29,138,176 (trib1), 23,917,406 (trib3), and 29,916,262 (trib21) good quality PE reads, whereas the mRNA sequencing of *C. lenavati* specimens generated 32,335,757 (lena1), 28,637,629 (lena2), and 33,027,986 (lena3) good quality PE reads. The pooled reads of three *C. tribblei* specimens (82,971,844 PE reads) and three specimens of *C. lenavati* (94,001,372 PE reads) with the average size of 81 bp were *de novo* assembled using Trinity into "tribblei reference assembly" and "lenavati reference assembly," respectively.

The tribblei reference assembly contained 327,700 transcripts with an average length of 489 bp, whereas 298,481 transcripts with an average length of 476 bp were identified in the lenavati reference assembly (supplementary table S2a, Supplementary Material online). The size of the assembled transcriptome was 160.27 Mb in *C. tribblei* and 142.14 Mb in *C. lenavati*. The N50 of the transcripts was 567 bp in the tribblei reference assembly and 547 bp in the lenavati reference assembly. Around 64% of the transcripts in both assemblies were 201–400 bp long and the length of 23–24% of the transcripts was 401–800 bp (supplementary table S2b, Supplementary Material online). However, the maximum transcript length was 21,724 and 16,659 bp in the tribblei and lenavati reference assemblies, respectively.

### Conopeptide Diversity

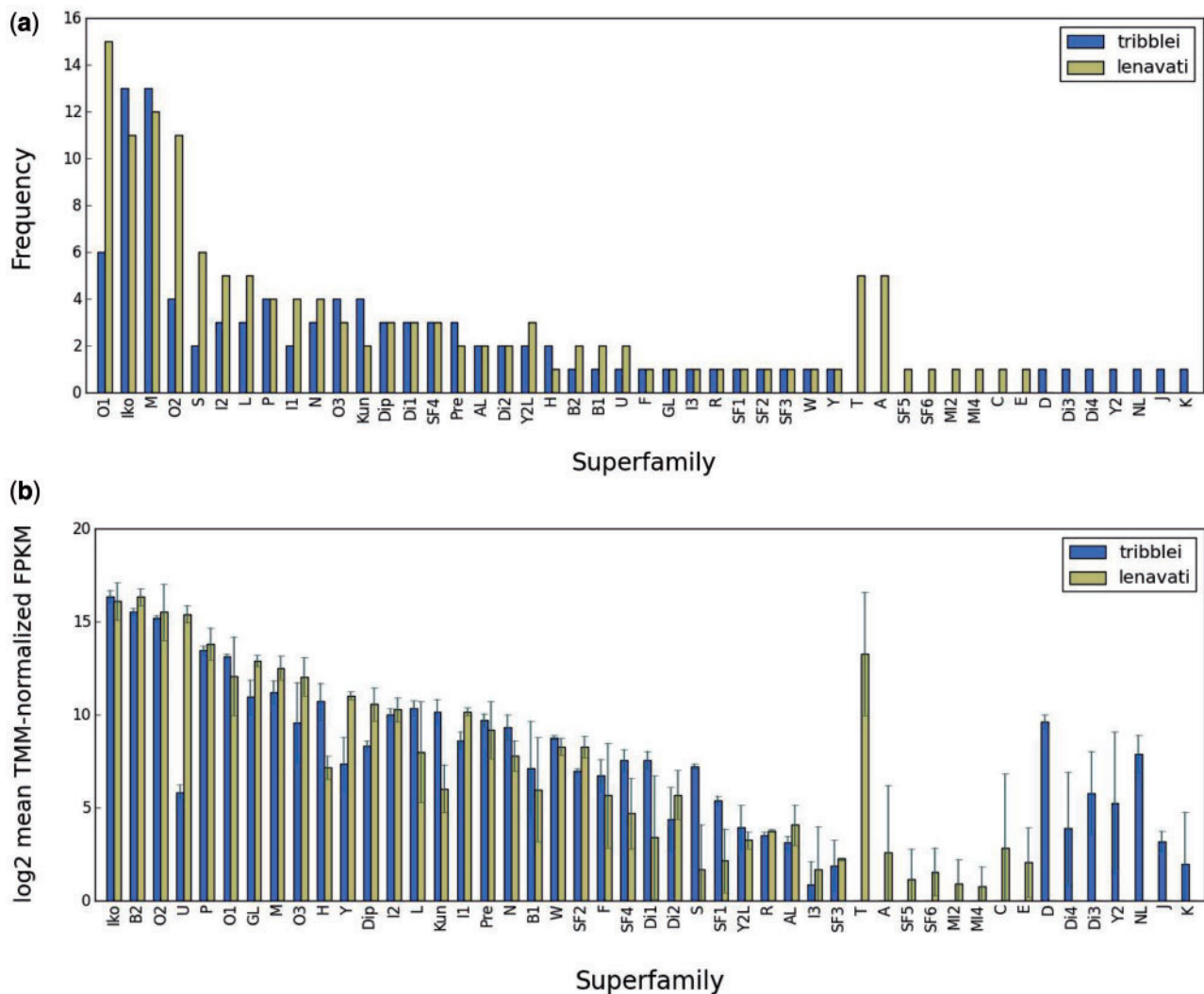
A total of 100 unique putative conopeptide precursors belonging to 39 gene superfamilies were identified in the tribblei conopeptide data set whereas the lenavati conopeptide data set contained 132 novel putative conopeptide precursors that were classified into 40 gene superfamilies (fig. 2a). A total of 55 conopeptides in the tribblei conopeptide data set were previously discovered in *C. tribblei* (accession GCJM000000000 in DDBJ/EMBL/GenBank; Barghi et al. 2015), whereas the rest of the identified conopeptides were new. The majority of the identified conopeptides was classified into the known gene superfamilies (fig. 2a). Moreover, some new conopeptide groups identified in recent transcriptome studies of the *Conus* venom duct were also discovered (fig. 2a); several sequences of the tribblei and lenavati conopeptide data sets showed high similarity to SF-mi2 and SF-mi4 in *Conus miles* (Jin et al. 2013), R, W, and Y2 in *Conus marmoratus* (Lavergne et al. 2013), U superfamily in *Conus victoriae* (Robinson et al. 2014), and A-like, G-like, N-like, Y2-like, and SF-01 groups in *C. tribblei* (Barghi et al. 2015) (supplementary fig. S1, Supplementary Material online). In addition, conopeptides with signal region PID values below the threshold for any conopeptide gene superfamily were classified into conopeptide groups SF-02 to SF-06 (fig. 3). Furthermore, seven putative conopeptides in the tribblei conopeptide data set and five conopeptides in the lenavati conopeptide data set showed similarity to the "divergent" superfamilies: Divergent MKFPLLIFSL, divergent M---L-LTVA, divergent MSTLGMSTLL-, and divergent MSKLVILAVL (fig. 2a). Divergent superfamilies refer to the conopeptides identified in early divergent species, *Conus californicus* and *Conus distans*. The conopeptide sequences identified in *C. tribblei* and *C. lenavati* are shown in supplementary figure S1, Supplementary Material online.

In both species, roughly half of the identified conopeptide precursors belonged to few superfamilies, and the most prominent superfamilies in both species were the con-ikot-ikot family and the M- and O1-, and O2-superfamilies (fig. 2a). Although the majority of gene superfamilies was represented

by more than one conopeptide, 19 and 16 superfamilies had only one sequence in the tribblei and lenavati conopeptide data sets, respectively. A total of 32 conopeptide gene superfamilies and groups were represented in both species. However, a few superfamilies such as the D-, J-, K- and Y2-superfamilies, the “divergent MSKLVILAVL” and “divergent MSTLGMTLL-” superfamilies, and the N-like group were only found in *C. tribblei*, whereas the T-, A-, E- and C-superfamilies, SF-05, SF-06, SF-mi2 and SF-mi4 groups were identified only in *C. lenavati*. On the other hand, the cysteine frameworks identified in the majority of conopeptide gene superfamilies were similar between the tribblei and lenavati conopeptide data sets (supplementary table S3, Supplementary Material

online). However, several conopeptide superfamilies such as the O1-, O2-, L-, M-, H- and S-superfamilies, the con-ikot-ikot family, and the Y2-like and SF-04 groups exhibited some cysteine patterns that were exclusively observed in either the tribblei or lenavati conopeptide data sets.

The diversity of putative conopeptides as indicated by the number of gene superfamilies was higher in *C. tribblei* and *C. lenavati* than in other previously studied species (table 1). Subsequently, both *C. tribblei* and *C. lenavati* had Shannon’s diversity index of 3.30, higher than in other species which ranges from 1.29 (*Conus bullatus*) to 2.35 (*C. victoriae*) (table 1). Additionally, specimens of *C. tribblei* and *C. lenavati* showed Shannon’s diversity indices ranging from 3.16 to 3.30



**Fig. 2.**—(a) The frequencies and (b) expression patterns of conopeptide gene superfamilies in *C. tribblei* and *C. lenavati*. The standard deviation ( $n = 3$ ) bars are shown, and the expression values are log<sub>2</sub>-transformed. The superfamily abbreviations are: Iko, con-ikot-ikot; Dip, conodipine; Kun, konkunitzin; Pre, conopressin/conophysin; Di1, divergent MKFPLLFISL; Di2, divergent M---L-LTVA; Di3, divergent MSTLGMTLL-; Di4, divergent MSKLVILAVL; SF1, SF-01; SF2, SF-02; SF3, SF-03; SF4, SF-04; SF5, SF-05; SF6, SF-06; MI2, SF-mi2; MI4, SF-mi4; NL, N-like; GL, G-like; AL, A-like; Y2L, Y2-like.

(table 2). However, the evenness of the conopeptides in *C. tribblei* and *C. lenavati* was slightly higher than other species (table 1).

### Expression Patterns of Conopeptide Gene Superfamilies

The expression patterns of conopeptides gene superfamilies in *C. tribblei* and *C. lenavati* were highly variable ranging from FPKM >85,000 to <10 (fig. 2b and supplementary table S4, Supplementary Material online). Superfamilies such as B2, O2, and P and con-ikot-ikot family were highly expressed (FPKM > 10,000) in *C. tribblei* and *C. lenavati*. Others, like M-, O1-, O3- and I2-superfamilies and G-like group also showed high expression levels (1,000 < FPKM < 10,000) in both species. Several conopeptide groups were moderately (100 < FPKM < 1,000) expressed: F-, N-, B1- and

W-superfamilies, conopressin/conophysin family, and SF-02 group. On the other hand, expression levels of “Divergent M---L-LTVA”- and R-superfamilies and A-like group in *C. tribblei* and *C. lenavati* were much lower (10 < FPKM < 100). Other conopeptide superfamilies had more variable expression levels between the two species (supplementary table S4, Supplementary Material online). Both *C. tribblei* and *C. lenavati* expressed seven to eight superfamilies that were not present in the other species (fig. 2b). Although, the majority of these conopeptide groups had relatively low expression levels, the T-superfamily in *C. lenavati* was highly expressed (FPKM > 10,000).

The expression patterns of all the conopeptide superfamilies were moderately correlated between *C. tribblei* and *C. lenavati* ( $r=0.608$ ,  $P$  value < 0.001) (fig. 4), but it was apparent that a great number of superfamilies had highly

### SF-02

1 76  
 Cln\_SF2\_1 MLFVFTVVWILTMVMIITDVTFQSTCNTDNKPS**CS**EDTRL**CG**KNN**SWG**N**C**VAL**CK**CP**NQQA**CTTDTDHKVQV**KRGP**  
 Ctr\_SF2\_1 MLFVFTVVWILTMVMIITDVTFQSTCNTDNKPS**CS**EDTRL**CG**KNN**SWG**N**C**VAL**CK**CP**NQQA**CTTDTDHKVQV**KRGP**

77 133  
 Cln\_SF2\_1 FQLTETYYT**CK**NVSTMSD**CQ**SNAKAMSGTSESTYKIM**CK**DDTYKPSAPT**NW**KFI**CG**  
 Ctr\_SF2\_1 FQLTETYYT**CK**NVSTMSD**CQ**SNAKAMSGTSESTYKIM**CK**DDTYKPSAPT**NW**KFI**CG**

### SF-03

Cln\_SF3\_1 MEAL**TIFRL**CLLVAL**TTS**VVV**S**APLNDK**VSD**Q**EGE**CPVGGGRNPFVLCMRACLTTST**PYQ**CEHEYCKHCP**R**GRYA**I**VGH**S**  
 Ctr\_SF3\_1 MEAL**TIFRL**CLLVAL**TTS**VVV**S**APLNDK**VSD**Q**EGE**CPVGGGRNPFVLCMRACLTTST**PYL**CEHEYCKN**C**-RGRYA**GL**GH**S**

### SF-04

Ctr\_SF4\_2 -----VLLLISTIAALYQ**EG**RATQ**RR**GK**TV**RTMS**NLL**NI**Q**KRE**CP**Y**G**CP**VV**CP**NG**DE**CC**DGIY**CS**---Q**G**DRRY**CV**G**CG**GG**RGE**  
 Ctr\_130\_T -----VLLLISTIAALYQ**EG**RATQ**RR**GK**TV**RTMS**NLL**NI**Q**KRE**CP**Y**G**CP**VV**CP**NG**DE**CC**DGIY**CS**---Q**G**DRRY**CV**G**CG**GG**RGE**  
 Cln\_SF4\_2 -----VLLLISTIAALYQ**EG**RATQ**RR**GK**TV**RTMS**NLL**NI**Q**KRE**CP**Y**G**CP**VV**CP**NG**DE**CC**DGIY**CS**---Q**G**DRRY**CV**G**CG**GG**RGE**  
 Ctr\_SF4\_1 **MN**CL**Q**LLL**V**LL**L**IST**T**T**A**L**Y**PD**V**R**K**TRRR**GG**SI**K**IL**R**LL**N**SE**K**RD**R**ST**G**CP**V**K**CP**K**H**TE**CC**SG**I**T**CP**Y**L**TP**GG**E**Y**F**C**IT**SG**GG**GG**E  
 Cln\_SF4\_1 -----TIAALYQ**D**GRATQ**RR**DG**NI**RTMS**DL**LNI**Q**KRE**CS**PD**CV**AT**CP**NG**NE****CC**DG**DL**CVYSSALE**KY**F**C**IG**CG**SG**GG**E  
 Ctr\_SF4\_3 -----TIAALYQ**D**GRATQ**RR**DG**NI**RTMS**DL**LNI**Q**KRE**CS**PD**CV**AT**CP**NG**NE****CC**DG**DL**CVYSSALE**KY**F**C**IG**CG**SG**GG**E  
 Ctr\_131\_T -----VLLLISTIAALYQ**D**GRATQ**RR**DG**NI**RTMS**DL**LNI**Q**KRE**CS**PD**CV**AT**CP**NG**NE****CC**DG**DL**CVYSSALE**KY**F**C**IG**CG**SG**GG**E  
 Cln\_SF4\_3 -----TITALYQ**V**DRAT**RR**EG**RT**IRAMS**NLL**K**IL**K**RR**CP**SP**D**CV**PY**CP**K**SE****CC**DG**DV**CVY**SE**TL**SK**H**F**CV**G**CG**-GG**GE

### SF-05

P01619 -----G**L**GR**CI**Y**NC**M**NS**GG**LS**F**I**Q**CK**T**MCY**  
 P02828 -----G**V**GR**CI**Y**NC**M**NS**GG**LN**F**I**Q**CK**T**MCY**  
 Cln\_SF5\_1 **MG**FK**V**AL**I**V**L**V**V**M**A**T**T**S**A**L**P**F**Q**F**S**E**F**A**E**T**S**V**G**Q**K**R**G**M**G**Q**C**I**H**D**C**M**NS**GG**LS**F**I**Q**CK**T**MCY**

### SF-06

Cln\_SF6\_1  
**ML**SM**F**AW**T**LM**T**AT**V**V**V**I**A**ER**Q**Y**C**PI**A**Q**T**CT**F**GS**D**L**CG**KE**S**GS**CS**PR**NC**K**N**ERM**CS**RS**D**HT**I**TV**V**RV**F**RR**RR**P**V**E**E**RY**T****C**VAL**S**GLE**E****C**SN**Q**K**A**L**T**D**L**  
**V**P**E**T**R**EL**N**S**V**EH**CK**CS**SP**K**V**Y**G**Y**H**MY**L**K**G**Y**F**CG**T**Y**E**R**S**

**Fig. 3.**—The putative conopeptide precursors of new conopeptide groups. The conopeptides identified in *C. tribblei* and *C. lenavati* are shown in black. The name of each conopeptide is presented as Ctr/Cln\_#:#: Ctr, *C. tribblei*; Cln, *C. lenavati*; #, superfamily (the abbreviations are indicated in fig. 2); #, arbitrary assigned number. The reference sequences are shown in green, and cysteine residues are shown in bold italic red. The signal regions are highlighted, and the mature regions are underlined. The reference sequences are Ctr\_130\_T, Ctr\_131\_T (Barghi et al. 2015), P01619, and P02828 (ConoServer database).

correlated expression (32 shared superfamilies,  $r=0.820$ ,  $P$  value  $< 0.001$ ). Although the expression patterns of majority of these superfamilies were apparently conserved, another subset consisting of the S-, H-, and U-superfamilies, and the conkunitzin family had very different expression levels in *C. tribblei* and *C. lenavati* (fig. 4 and supplementary table S4, Supplementary Material online). There is moderate correlation between the abundance and expression of different conopeptide gene superfamilies in *C. tribblei* ( $r=0.602$ ,  $P$  value  $< 0.001$ ) and *C. lenavati* ( $r=0.597$ ,  $P$  value  $< 0.001$ ) (supplementary fig. S2, Supplementary Material online).

### Interspecific Genetic Divergence of Conopeptide Superfamilies

Although, the total number of identified conopeptides in the lenavati conopeptide data set (132) was higher than in the tribblei conopeptide data set (100), the number of superfamilies in both species was similar (39–40 superfamilies). More than half of the conopeptides of the superfamilies shared in *C. tribblei* and *C. lenavati* had PID  $> 81\%$ , and these sequences were distributed across all gene superfamilies (except for H-, S-, and F-superfamilies) (fig. 5). Additionally, 12.02% (12 superfamilies) and 8.65% (11 superfamilies) of conopeptides in *C. tribblei* and *C. lenavati* showed 61–80% and 41–60% PID, respectively. Interestingly, a large portion

(19.71%) of conopeptides belonging to 12 gene superfamilies had only 21–40% PID. 2.88% of the conopeptides exhibited PID  $< 20\%$ , whereas the minimum sequence identity was 10%.

### Divergence of Orthologous Conopeptide Genes

A total of 67 pairs of orthologous conopeptide genes belonging to 31 conopeptide gene superfamilies and groups were identified in *C. tribblei* and *C. lenavati* (supplementary table S5, Supplementary Material online). More than half of the orthologous gene pairs (37 pairs) had  $d_N$  to  $d_S$  ratios  $> 1$ , whereas nine gene pairs showed  $d_N$  to  $d_S$  ratios  $< 1$ . Interestingly, 21 orthologous gene pairs exhibited identical mature regions. Although 15 of these identical orthologs also contained identical signal and pre regions ( $d_N = d_S = 0$ ), six gene pairs had 1–10 amino acid differences in the prepro regions.

The conopeptides with identical mature regions belonged to 14 superfamilies (B1-, N-, O1-, O2-, P-, W- and Y-superfamilies, “divergent MKFPLLFISL” superfamily, con-ikot-ikot and conopressin/conophysin families, and A-like, Y2-like, SF-02 and SF-04 groups) (fig. 6a). The expression levels of all orthologous conopeptide gene pairs in *C. tribblei* and *C. lenavati* were moderately correlated ( $r=0.640$ ,  $P$  value  $< 0.001$ , data not shown), and the correlation of the expression of identical orthologous conopeptide genes was slightly lower

**Table 1**  
Diversity of Conopeptides in the Studied *Conus* Species through Next Generation Sequencing

Species	Richness (S)	No. of Conopeptides	Shannon's Diversity Index	Evenness (E)	Sequencing Platform	References
<i>Conus bullatus</i>	6	30	1.29	0.72	Roche 454	Hu et al. 2011
<i>Conus geographus</i>	16	63	2.30	0.83	Roche 454	Hu et al. 2012
<i>Conus pulicarius</i>	14	82	2.02	0.77	Roche 454	Lluisma et al. 2012
<i>Conus consors</i>	11	61	2.04	0.85	Roche 454	Terrat et al. 2012
<i>Conus marmoreus</i>	26	263	2.17	0.67	Roche 454	Dutertre et al. 2013; Lavergne et al. 2013
<i>Conus victoriae</i>	20	117	2.35	0.78	Roche 454	Robinson et al. 2014
<i>Conus tribblei</i>	36	136	3.13	0.87	Roche 454, Illumina	Barghi et al. 2015
<i>Conus tribblei</i> <sup>a</sup>	39	100	3.30	0.90	Illumina	This study
<i>Conus lenavati</i> <sup>a</sup>	40	132	3.30	0.89	Illumina	This study

NOTE.—Richness (S) is the number of identified conopeptide gene superfamilies.

<sup>a</sup>The number of conopeptides of *C. tribblei* and *C. lenavati* in this study is the total nonredundant conopeptides identified in three specimens for each species.

**Table 2**  
Diversity of Conopeptides in the Specimens of *Conus tribblei* and *Conus lenavati*

Species	Specimen	Richness (S)	No. of Conopeptides	Shannon's Diversity Index	Evenness (E)
<i>Conus tribblei</i>	trib1	36	87	3.23	0.90
	trib3	36	85	3.23	0.90
	trib21	38	89	3.32	0.91
<i>Conus lenavati</i>	lena1	31	71	3.19	0.93
	lena2	31	71	3.16	0.92
	lena3	40	125	3.30	0.90

( $r=0.614$ ,  $P=0.003$ ) (fig. 6a). Notably, the expressions of three outliers (conopeptides of O1- and O2- and P-superfamilies) (fig. 6a) in the correlation test of identical orthologs were highly divergent that upon their exclusion, the correlation increased to  $r=0.924$  ( $P$  value  $< 0.001$ ). The expression of orthologs having higher rate of  $d_S$  to  $d_N$  was moderately though not significantly correlated ( $r=0.749$ ,  $P=0.02$ ) (fig. 6b), whereas the expression of orthologous pairs under positive selection showed weak correlation ( $r=0.585$ ,  $P$  value  $< 0.001$ ) (fig. 6c).

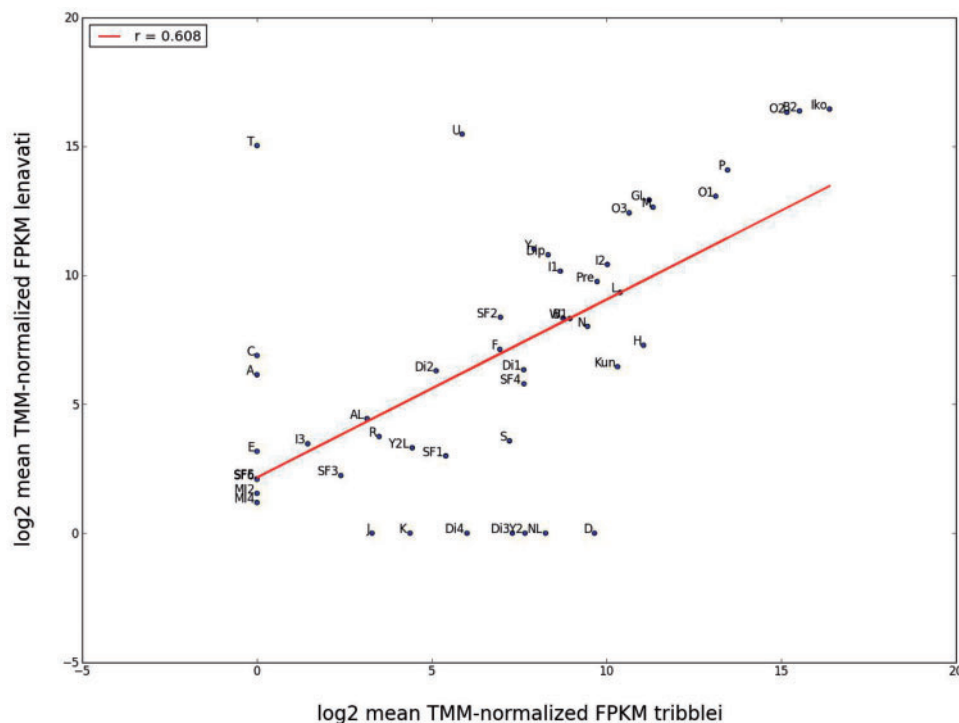
### Intraspecific Patterns of Conopeptide Expression

Most (75) of the identified conopeptide precursors in *C. tribblei* were found in all specimens, but some sequences were observed in either one or two individuals only (fig. 7a and supplementary table S6, Supplementary Material online). Similarly, although 34 superfamilies were identified in all specimens of *C. tribblei*, a few superfamilies were expressed in only one or two individuals (fig. 7a and supplementary table S6, Supplementary Material online). Interestingly, only less than half of the identified conopeptides in lenavati conopeptide data set were expressed in all three specimens (fig. 7b and supplementary table S6, Supplementary Material online). Specimen lena3 expressed 54 more conopeptides than the other two individuals (fig. 7b and supplementary table S6, Supplementary Material online). In addition, although 29

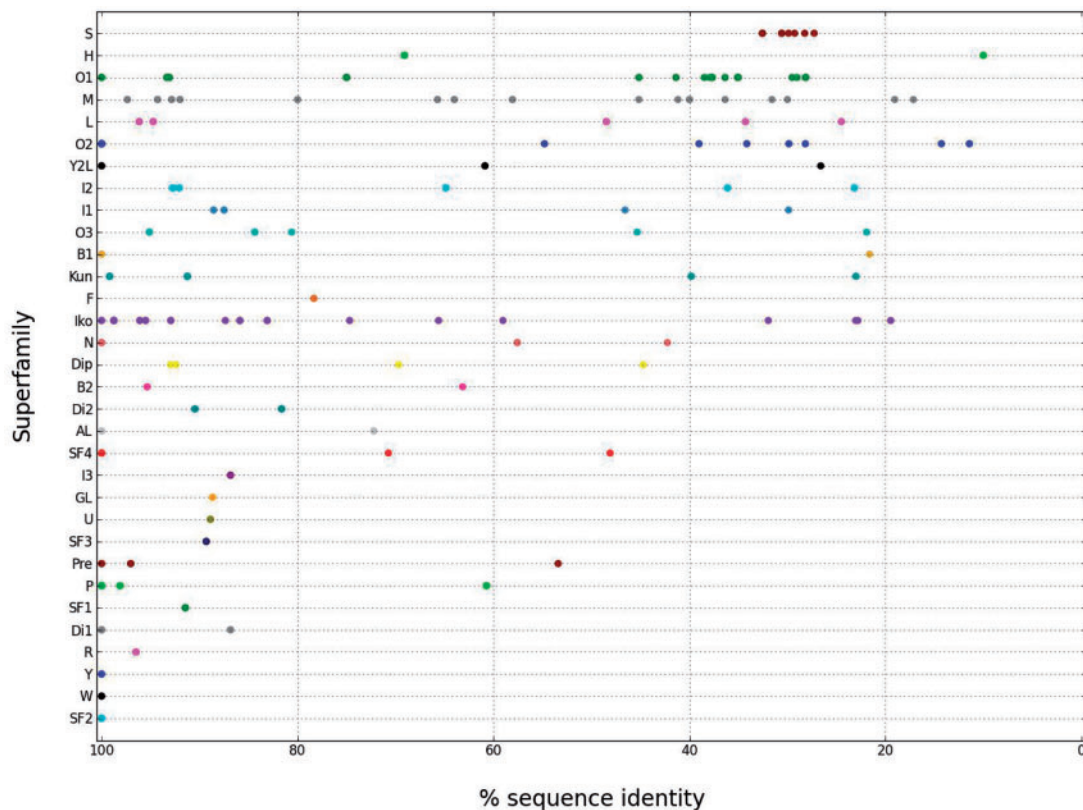
superfamilies were identified in all three individuals of *C. lenavati*, 7 superfamilies including A, C, I3, S and SF-05, SF-mi2, and SF-mi4 groups were expressed only in specimen lena3. Moreover, the number of orthologous conopeptide genes with identical mature regions among the individuals of *C. tribblei* and *C. lenavati* varied from 12 to 16 (fig. 7c).

The expression patterns of conopeptides among the specimens of *C. tribblei* were moderately correlated, whereas a higher correlation in the conopeptide expression was observed among the *C. lenavati* specimens (table 3). In addition, the expression levels of conopeptide gene superfamilies were highly correlated among individuals of *C. tribblei* as well as among specimens of *C. lenavati* (table 3). Despite the high correlation, the levels of expression of each conopeptide (and also each superfamily) were not the same among the specimens of each species (supplementary table S4, Supplementary Material online).

The similarity of the expression patterns of conopeptides across individuals of each species was evaluated using the ten most highly expressed conopeptides as reference (supplementary tables S7 and S8, Supplementary Material online). It was found that of the ten most highly expressed conopeptides in each specimen of *C. tribblei*, seven conopeptides were common among all individuals whereas other highly expressed conopeptides were observed in one or two specimens only (fig. 8a). Likewise, six of the ten most highly expressed conopeptides in each *C. lenavati* individual were similar in all



**Fig. 4.**—Correlation of the expression levels of conopeptide gene superfamilies in *C. tribblei* and *C. lenavati*. The Pearson's correlation coefficient ( $r$ ) is 0.608 ( $P$  value  $< 0.001$ ). The abbreviations of superfamily names are noted in figure 2.



**FIG. 5.**—Distribution of the conopeptides in each gene superfamily based on the highest PID (the most similar match) of the conopeptides' mature regions in *C. tribblei* and *C. lenavati*. The abbreviations of superfamily names are noted in figure 2.

three individuals whereas the other most highly expressed conopeptides were different among specimens of *C. lenavati* (fig. 8b). The presence of a combination of highly expressed conopeptides that are common in all individuals and other conopeptides which are highly expressed in one or two specimens resulted in a distinct set of highly expressed conopeptides in each individual (fig. 8 and [supplementary tables S7 and S8, Supplementary Material](#) online). The majority of the highly expressed conopeptides in both species belonged to superfamilies and groups (O1, O2, B2, P, con-ikot-ikot, and G-like) with high expression levels (fig. 2b). Notably, conopeptides of several superfamilies such as O3, M, T, and U were among the highly expressed conopeptide in individuals of *C. lenavati* but not in *C. tribblei* (fig. 8 and [supplementary tables S7 and S8, Supplementary Material](#) online). Similarly, one H-superfamily conopeptide was highly expressed only in the individuals of *C. tribblei*. Interestingly, six of the ten highly expressed conopeptides in the individuals of *C. tribblei* and *C. lenavati* were orthologous conopeptide pairs identified between these species (fig. 8).

### Functional Analysis of Transcriptomes

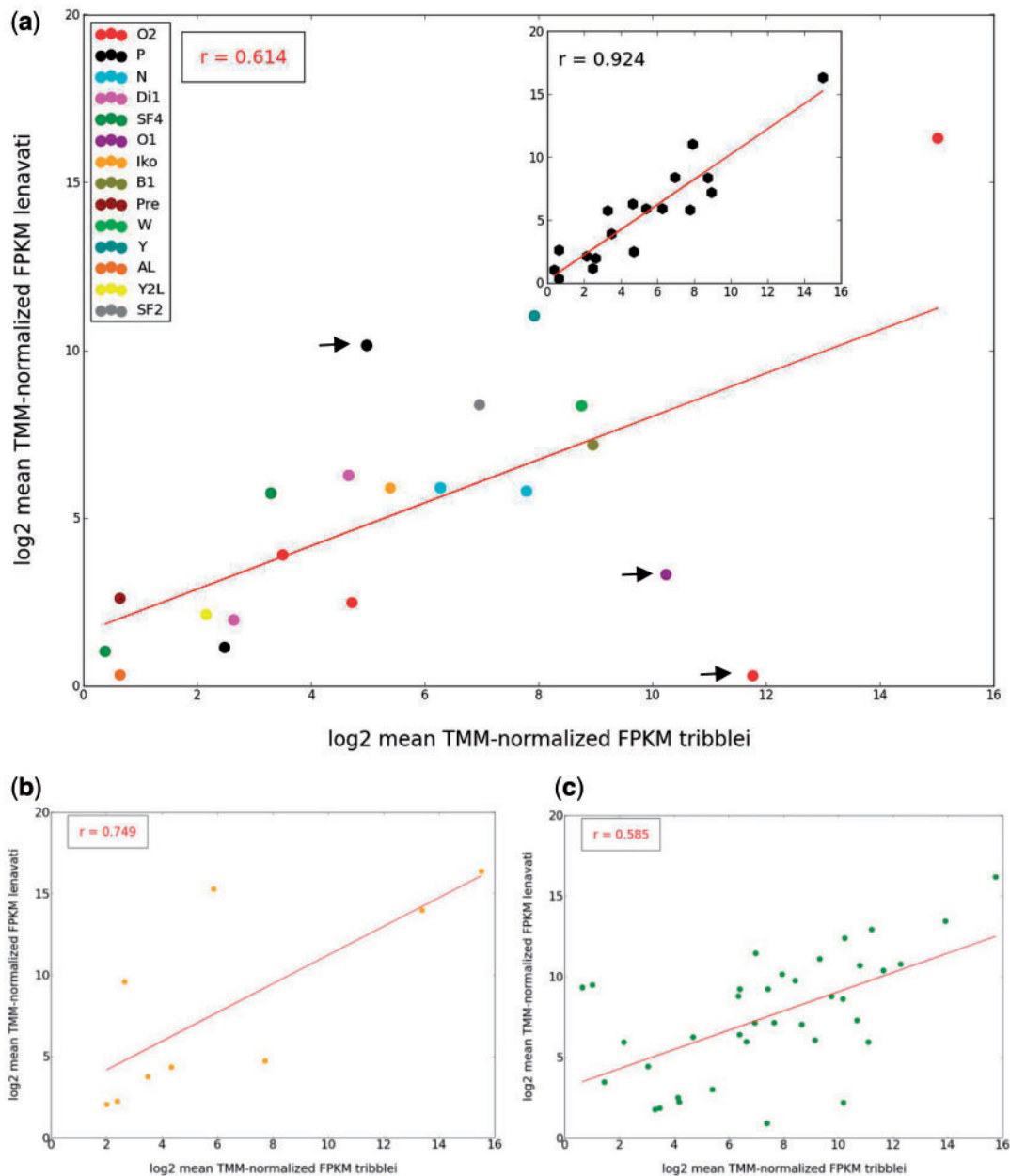
In addition to the conopeptides, a total of 41,875 and 39,516 transcripts in the tribblei and lenavati reference assemblies,

respectively, exhibited high similarity to the proteins in the UniProtKB/Swiss-Prot database. Gene ontology terms were assigned to 34,876 transcripts in *C. tribblei*, and also to 32,778 transcripts in *C. lenavati* data sets. In both species, transcripts having binding and catalytic activities comprised a high percentage of the GO terms in the molecular function category, whereas cellular and metabolic processes and biological regulation were the most prominent categories in the biological process category in *C. tribblei* and *C. lenavati* ([supplementary fig. S3, Supplementary Material](#) online). Also, cell and organelle were the most abundant terms in the cellular component category in the tribblei and lenavati data sets.

## Discussion

### High Diversity of Expressed Conopeptide Genes

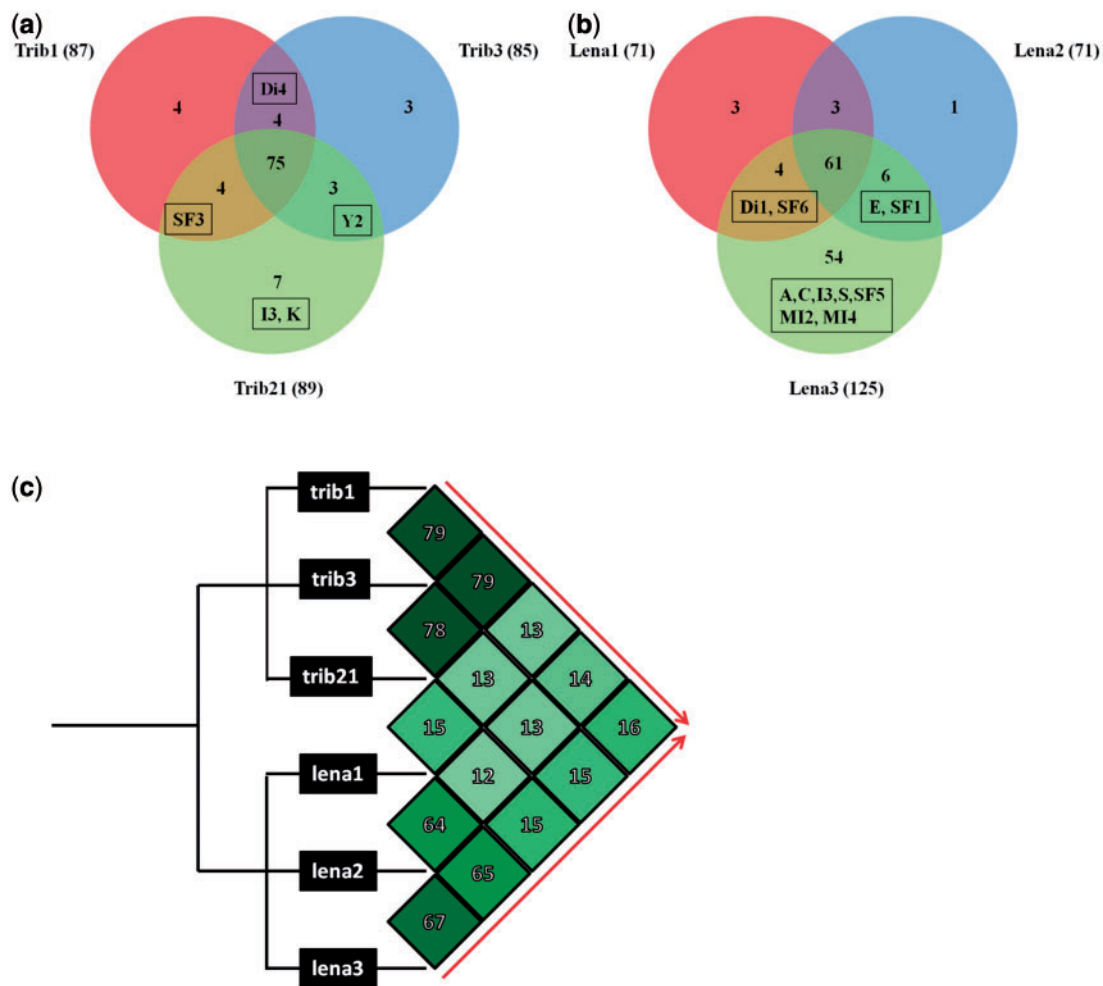
The venom of *C. tribblei* and *C. lenavati* had the highest number of conopeptide gene superfamilies and the most diverse venom complement observed in any *Conus* species thus far. Moreover, the conopeptide diversity even in the individuals of *C. tribblei* and *C. lenavati* was higher than in other *Conus* species indicating that the high conopeptide diversity and the venom complexity were consistent across the specimens. These observations also suggest that although using



**FIG. 6.**—The expression of orthologous conopeptide pairs in *C. tribblei* and *C. lenavati*. (a) The correlation of the expression levels of identical orthologous conopeptide pairs ( $r = 0.614$ ,  $P = 0.003$ ). The correlation of the expression of orthologs pairs except for the three conopeptides with divergent expression patterns (marked with black arrows) ( $r = 0.924$ ,  $P$  value  $< 0.001$ ) is shown in insert. The correlation of orthologous conopeptide pairs (b) showing  $d_N$  to  $d_S$  ratios  $< 1$  ( $r = 0.749$ ,  $P = 0.02$ ) and (c)  $d_N$  to  $d_S$  ratios  $> 1$  ( $r = 0.585$ ,  $P$  value  $< 0.001$ ). The abbreviations of superfamily names are noted in figure 2.

pooled sample of several individuals in the transcriptome studies may increase the total number of identified conopeptides, it may not significantly affect the inferred diversity of the conopeptide superfamilies. The diversity of the conopeptide genes in cone snails reflects dietary specialization and prey types (Duda and Palumbi 2004; Duda et al. 2009); species with more exclusive prey types produce less diverse conopeptides (Remigio and Duda 2008). The remarkably high diversity of conopeptide gene superfamilies in *C. tribblei* and *C. lenavati*

may thus indicate a relatively wide range of prey targets for these closely related species. In addition, identification of divergent superfamilies that have only been reported for the generalist feeder *C. californicus* (Elliger et al. 2011) is consistent with the hypothesis that the high diversity of conopeptides in *C. tribblei* and *C. lenavati* may be an adaptation to a wide taxonomic prey field. The conopeptide diversity in *C. tribblei*, *C. lenavati*, and other *Conus* species is compared in table 1; Although these estimates may to some degree be



**Fig. 7.**—Shared and distinct conopeptides and gene superfamilies for specimens of (a) *C. tribblei* and (b) *C. lenavati*, (c) the number of identical conopeptide (considering only the mature regions) among the specimens of *C. tribblei* and *C. lenavati*. The total number of conopeptides in each specimen is shown in parenthesis. The superfamilies that are found in only one or two specimens are shown in box. The abbreviations of superfamily names are noted in figure 2. The number of identical conopeptides between a pair of specimens is shown in each diamond. To find the diamond corresponding to one pair of specimens, the lines extended from the dark boxes containing the specimen’s name should be followed. The diamond, in which the columns leading from a pair of individuals intersect, shows the number of identical mature regions. For example, the diamond at the intersection of the two red arrows corresponds to trib1 and lena3. The phylogenetic relationship of the specimens shown here is not based on the actual Bayesian tree. The intensity of the colors of the diamonds corresponds to the number of identical conopeptides.

subject to inaccuracies (owing to such factors as sequencing platforms and bioinformatics tools used), the data clearly indicate the relatively high diversity of conopeptides in *C. tribblei* and *C. lenavati*.

The conopeptide gene superfamilies expressed in *C. tribblei* and also in *C. lenavati* had variable frequencies (fig. 2a). Similar observations were previously reported (Hu et al. 2011, 2012; Lluisma et al. 2012; Terrat et al. 2012; Dutertre et al. 2013; Lavergne et al. 2013; Robinson et al. 2014; Barghi et al. 2015). Surprisingly, the computed Shannon’s evenness indices did not reflect such variability (tables 1 and 2). This incongruence may have arisen because except for a few predominant superfamilies, 19 and 16

conopeptide groups in *C. tribblei* and *C. lenavati*, respectively, were each represented by only one sequence. Hence, despite the high diversity of gene superfamilies, the frequencies of most of the conopeptide superfamilies were similar.

### Patterns of Sequence Divergence in Conopeptide Superfamilies

The members of different conopeptide gene superfamilies have diverged differently after separation of closely related species *C. tribblei* and *C. lenavati* so that several divergence patterns have emerged (fig. 5). At one end, the divergence of

**Table 3**

Correlation of the Expression Levels of Conopeptides and Gene Superfamilies among Specimens of *Conus tribblei* and *Conus lenavati*

<i>Conus tribblei</i>			
	trib1	trib3	trib21
trib1	1	0.751	0.644
trib3	0.884	1	0.682
trib21	0.835	0.833	1
<i>Conus lenavati</i>			
	lena1	lena2	lena3
lena1	1	0.913	0.798
lena2	0.936	1	0.824
lena3	0.858	0.883	1

NOTE.—The Pearson's correlation coefficient for the conopeptides is shown in the cells above the diagonal and for the gene superfamilies below the diagonal. For all the correlation coefficients ( $r$ ), the  $P$  value is less than 0.001.

conopeptides of some gene superfamilies with identical or highly similar conopeptides has occurred slowly. At the other end, few superfamilies (S and H) had diverged more. Between these two groups, several superfamilies (O1, O2, M, con-ikot-ikot, and conodipine) with high frequencies (fig. 2a) showed a wide range of divergence, from identical conopeptides to the conopeptides showing as low as 11% PID between *C. tribblei* and *C. lenavati*. Similarly, the sequence identity of O1-superfamily conotoxins in *Conus abbreviatus* and *Conus miliaris* ranged from 51.66% to 96.3% (Duda 2008).

### Conserved and Species-Specific Expression Patterns of Conopeptide Genes

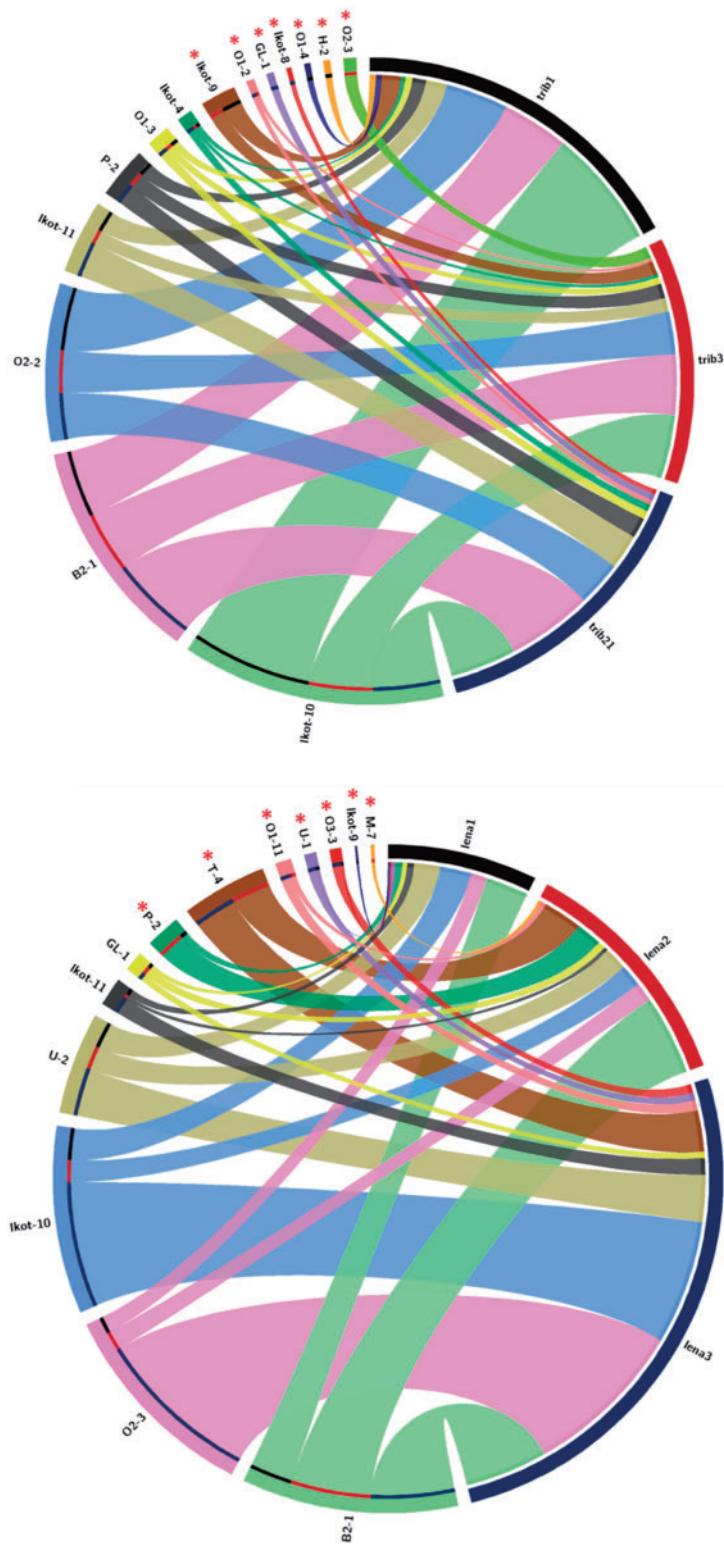
In both *C. tribblei* and *C. lenavati*, the expressions of different gene superfamilies had order-of-magnitude differences. Although half of the venom transcriptome consisted of transcripts from few highly expressed superfamilies, the majority of the conopeptide groups had moderate to low expression levels. Similarly, high variations in the expression patterns of different superfamilies have been observed in other species of *Conus* (Hu et al. 2011, 2012; Terrat et al. 2012; Dutertre et al. 2013; Robinson et al. 2014). High correlation ( $r=0.938$ ) between the frequency and the expression level of conopeptide superfamilies in *C. marmoreus* was observed (Dutertre et al. 2013). On the contrary, the frequency and the expression of different superfamilies in *C. tribblei* and *C. lenavati* were only moderately correlated (supplementary fig. S2, Supplementary Material online). This is because although O2-superfamily and con-ikot-ikot family were predominant components of the venom in terms of both the number of conopeptides and also the expression levels, other superfamilies such as P and B2 had extremely high expression levels despite containing only 1–4 conopeptides.

Comparison of the expression levels of conopeptide gene superfamilies between the two species revealed the presence of both conserved and species-specific expression patterns. The expression levels of superfamilies common in both species were highly correlated whereas several conopeptide superfamilies appeared to be species-specific and are only expressed in either *C. lenavati* or *C. tribblei*. The complement of gene superfamilies identified in each *C. tribblei* specimen was a relatively robust representative of the superfamilies characteristically expressed in this species. Of the 36 gene superfamilies identified in the transcriptome analysis of the pooled mRNA from the venom ducts of 20 *C. tribblei* specimens (Barghi et al. 2015), 32 were identified in all the three individuals of *C. tribblei* in this study, and 3 gene superfamilies were expressed in either 1 or 2 specimens. Moreover, none of the species-specific superfamilies of *C. lenavati* (T, A, C, E, SF-05, SF-06, SF-mi2, and SF-mi4) was identified in the pooled sample transcriptome of *C. tribblei* (Barghi et al. 2015). Whether similar observation is relevant for *C. lenavati* requires an analysis of the transcriptome of a pooled sample of *C. lenavati*. The conopeptides identified in this transcriptome analysis represent the genes expressed at a specific time and in a specific developmental stage of the organism. The sequencing depth in this study has allowed detection of a wide range of expression level (10–85,000 FPKM). Hence, if some conopeptides were not detected in this study, they must have been expressed at extremely low level. Furthermore, some conopeptide genes might be present in the genome but not expressed in the specific individuals sequenced in this study.

In general, the diet of each worm-hunting cone snail is dominated by a different polychaete, and even *Conus* species coexisting in a habitat specialize on different prey types (Kohn 2001). The distribution of these species in the Philippines is not well documented, but both species used in this study were collected in deep waters at the same location in Eastern Cebu, Sogod. *Conus tribblei* was collected at lower depth (100 m) than *C. lenavati* which is usually found at the depth of 130 m. Hence the evolution of species-specific expression patterns in *C. tribblei* and *C. lenavati* may indicate their divergence in range of conopeptide targets (prey, predators, or competitors) in their habitat.

### Divergent Expression of Orthologous Conopeptide Genes

The expression pattern of identical (hence, orthologous) conopeptide genes in *C. tribblei* and *C. lenavati* reveals strong conservation of regulatory modes in the two species, although instances of divergence were noted (fig. 6a). On the other hand, the conopeptides under positive selection showed greater dissimilarity in expression in *C. tribblei* and *C. lenavati* and showed only slight correlation (fig. 6c). It appears that the conservation of sequence in the orthologous conopeptides of *C. tribblei* and *C. lenavati* is correlated with conservation of expression pattern. Our results elucidated that even among



**Fig. 8.**—The ten most highly expressed conopeptides among the individuals of (a) *C. tribblei* and (b) *C. lenavati*. The thickness of ribbons corresponds to the expression level of conopeptides. The nomenclature of the conopeptide names is noted in figure 3, and the abbreviations of superfamily names are noted in figure 2. The conopeptides that are present in the highly expressed conopeptides of only one or two individuals are marked with red asterisks. The figures were constructed using Circos (Krzywinski et al. 2009). The orthologous conopeptide pairs are: B2-1, GL-1, P-2, Ikot-10, Ikot-11, Ikot-8 (*C. tribblei*), and Ikot-9 (*C. lenavati*).

the orthologous conopeptide pairs, the divergence in pattern of expression can contribute to differences and uniqueness of venom composition in each species.

This study discovered a large number of orthologous conopeptide pairs (a total of 67) belonging to a wide range (31) of gene superfamilies in *C. tribblei* and *C. lenavati*. Duda and Remigio (2008) identified 12 sets of orthologous O1-superfamily genes among closely related worm-hunting *Conus* species. No orthologous pair was identified between recently diverged sister species *C. abbreviatus* and *Conus aristophanes*, eight pairs were identified in *C. abbreviatus* and *C. miliaris* which is separated from the lineage of *C. abbreviatus* and *C. aristophanes* (Duda and Remigio 2008). Additionally, few orthologs of A-superfamily genes were expressed in four closely related worm-hunting species (Chang and Duda 2012, 2014). Identification of very few orthologous conopeptide genes in the previous studies (Duda and Remigio 2008; Chang and Duda 2012, 2014) could be partly explained by the fact that only 1–2 conopeptide gene superfamilies were investigated. It must also be mentioned that because the previous studies (Duda and Remigio 2008; Chang and Duda 2014) used the traditional cDNA library method, the accuracy of the measurement of gene expression could have been limited, and some conopeptides with low expression levels could not have been detected.

### Intraspecific Variation of Conopeptide Expression

The intraspecific variation of the number of conopeptides among the specimens of *C. tribblei* was low. This observation may be due to the similar size of *C. tribblei* individuals (~6 cm), which implies their similar developmental stage. A more pronounced intraspecific variation was observed among the specimens of *C. lenavati* mainly because of the expression of significantly higher number of conopeptides and the exclusive expression of several superfamilies in one specimen, lena3. This specimen (shell length 5 cm) was smaller than the other two specimens (8 cm) which may indicate an earlier developmental stage. Ontogenic changes in the expression of conopeptides have been previously documented in *Conus* (Safavi-Hemami et al. 2011). However, further analysis of the conopeptide expression in individuals of different developmental stages is needed to corroborate our results. The observed correlation of the conopeptide expression patterns among individuals of *C. tribblei* and of *C. lenavati* (table 3) indicates species-characteristic mechanism for regulating conopeptide expression, but the observed quantitative differences in levels of gene expression (supplementary table S4, Supplementary Material online) and specimen-specific set of highly expressed conopeptides (fig. 8) suggest that the expression levels of a subset of conopeptides can also be independently regulated by individual snails. All the previous studies describing intraspecific variation in *Conus* only studied the peptide masses in the venom (Jakubowski et al. 2005; Davis et al.

2009; Dutertre et al. 2010; Chun et al. 2012). Despite the variability in the peptide masses among individuals, the venom profile of the individuals for each species remains consistent (Jakubowski et al. 2005; Dutertre et al. 2010), and even the predominant peptides remain the same over time (Chun et al. 2012). These observations suggest that each species apparently evolves species-specific regulatory mechanisms for its venom peptides.

### Evolution of *Conus* Exogenomes

Identification of positive selection in the conopeptide orthologs and the presence of several divergence patterns among the conopeptide superfamilies specifically those exhibiting very high divergence between *C. tribblei* and *C. lenavati* (fig. 5) provide further evidence that multiple molecular mechanisms synergistically act to cause differentiation of venom composition between species and drive the evolution of *Conus* exogenome. These extend previous observations on the molecular evolution and extremely high diversification rate of conopeptides through gene duplication and positive selection in piscivorous and vermivorous species (Duda and Palumbi 2000; Duda 2008; Duda and Remigio 2008; Puillandre et al. 2010; Chang and Duda 2012, 2014). The role of altered gene expression in evolution has been demonstrated, for example in morphological and behavioral changes in mammals (Enard et al. 2002). Divergence of gene expression and the protein sequence divergence both contribute to the adaptive evolution (Nuzhdin et al. 2004; Lemos et al. 2005; Khaitovich et al. 2006). Modifications of the expression level of toxin transcripts together with the diversity of isoforms in each toxin family were shown to account for the differentiation of the venom affecting prey specificity in species of vipers (Casewell et al. 2009). In this study, we showed that divergence in the control of conopeptide expression is another factor that may contribute to the differentiation of *Conus* venom composition. The divergent expression patterns of positively selected conopeptide orthologs in *C. tribblei* and *C. lenavati* (fig. 6) and the species-specific expression patterns of some superfamilies (figs. 2b and 4) contribute to the presence of unique venom complement in these species. Our results identified the divergence in the conopeptide expression as an important adaptation strategy in these two species; further studies will be necessary to uncover the molecular mechanisms of conotoxin gene regulation and exogenome evolution in *Conus*.

### Conclusion

The results of this study highlight the role of sequence divergence, positive selection, and selective changes in the patterns of gene expression in the generation of conopeptide diversity and modification of venom composition over evolutionary time, and how these processes reflect the evolution of *Conus* exogenomes. These genome-level processes are among the factors that underlie the capacity of *Conus* species

to exploit their taxonomic prey field or adapt to changing biotic conditions (e.g., changes in prey availability, or type of predators or competitors in their environment). However, other related processes (e.g., gene duplication, hypermutation, recombination, or chromosome-level mutations) are likely to play a significant role in this evolutionary process. To obtain deeper insights into the evolution of *Conus* exogenomes, it will be important to investigate how the processes described in this study and those related processes interact.

## Supplementary Material

Supplementary figures S1–S3 and tables S1–S8 are available at *Genome Biology and Evolution* online (<http://www.gbe.oxfordjournals.org/>).

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