



Article Active Expression of Genes for Protein Modification Enzymes in Habu Venom Glands

Akiko Isomoto ^{1,2,*}, Eiichi Shoguchi ³, Kanako Hisata ³, Jun Inoue ³, Yinrui Sun ¹, Kenji Inaba ^{4,5}, Noriyuki Satoh ³, Tomohisa Ogawa ^{2,*} and Hiroki Shibata ^{1,*}

- ¹ Division of Genomics, Medical Institute of Bioregulation, Kyushu University, Fukuoka 812-8582, Japan; syr0221@gen.kyushu-u.ac.jp
- ² Department of Biological Chemistry, Graduate School of Agricultural Science, Tohoku University, Sendai 980-8572, Japan
- ³ Marine Genomics Unit, Okinawa Institute of Science, Technology Graduate University, Onna 904-0495, Japan; eiichi@oist.jp (E.S.); kanako@oist.jp (K.H.); jun.inoue@oist.jp (J.I.); norisky@oist.jp (N.S.)
- ⁴ Institute of Multidisciplinary Research for Advanced Material, Tohoku University, Sendai 980-8577, Japan; kenji.inaba.a1@tohoku.ac.jp
- ⁵ Department of Biomolecular Science, Graduate School of Life Sciences, Tohoku University, Sendai 980-8577, Japan
- * Correspondence: isomoto.akiko.p3@dc.tohoku.ac.jp (A.I.); tomohisa.ogawa.c3@tohoku.ac.jp (T.O.); hshibata@gen.kyushu-u.ac.jp (H.S.)

Abstract: Genes encoding snake venom toxins have been studied extensively. However, genes involved in the modification and functioning of venom proteins are little known. *Protobothrops* is a genus of pit vipers, which are venomous and inhabit the Nansei (Southwest) islands of Japan, Taiwan China, Vietnam, Thailand, Myanmar, Nepal, Bhutan, and India. Our previous study decoded the genome of *Protobothrops flavoviridis*, a species endemic to the Nansei Islands, Japan, and revealed unique evolutionary processes of some venom genes. In this study, we analyzed genes that are highly expressed in venom glands to survey genes for candidate enzymes or chaperone proteins involved in toxin folding and modification. We found that, in addition to genes that encode venom proteins and ribosomal proteins, genes that encode protein disulfide isomerase (PDI) family members (orthologs of human *P4HB* and *PDIA3*), Selenoprotein M (*SELENOM*), and Calreticulin (*CALR*) are highly expressed in venom glands. Since these enzymes or chaperones are involved in protein modification and potentially possess protein folding functions, we propose that *P4HB*, *SELENOM*, *CALR*, and *PDIA3* encode candidate enzymes or chaperones to confer toxic functions upon the venom transcriptome.

Keywords: venom protein modification enzyme; disulfide-bond; PDI; venom protein folding; *P4HB*; *PDIA3*

Key Contribution: *P4HB* and *PDIA3* from the *PDI* gene family, the selenoprotein, *SELENOM*, and the ER-resident protein, *CALR*, are highly expressed in venom glands of habus (*Protobothrops flavoviridis*), suggesting the most likely involvement of these proteins in protein folding and modification during snake venom production.

1. Introduction

Several crotalid snakes inhabit the Nansei (southwestern) Islands of Japan (Amami-Oshima, Tokunoshima, Okinawa, and Sakishima) and Taiwan [1]. They include *Protobothrops flavoviridis* (Amami-Oshima, Tokunoshima, and Okinawa islands), *Ovophis okinavensis* (Amami-Oshima, Tokunoshima, and Okinawa islands), *P. elegans* (Ishigaki and Iriomote islands of the Sakishima Islands), *P. mucrosquamatus* (Taiwan), and *Trimeresurus stejnegeri* (Taiwan) (The Reptile Database: http://www.reptile-database.org/ (accessed on 31 March 2019)). From the point of view of snake envenomation in Japan, the habu, *P. flavoviridis*, is



Citation: Isomoto, A.; Shoguchi, E.; Hisata, K.; Inoue, J.; Sun, Y.; Inaba, K.; Satoh, N.; Ogawa, T.; Shibata, H. Active Expression of Genes for Protein Modification Enzymes in Habu Venom Glands. *Toxins* **2022**, *14*, 300. https://doi.org/10.3390/ toxins14050300

Received: 14 January 2022 Accepted: 16 April 2022 Published: 22 April 2022

Publisher's Note: MDPI stays neutral with regard to jurisdictional claims in published maps and institutional affiliations.



Copyright: © 2022 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). the greatest threat to people, with more than 1000 people bitten per year, and a fatality rate of 1% [2].

To clarify snake venom repertoires, as well as the evolution and physiological function of these venoms, many proteomic and transcriptomic studies have been performed [3–5]. These have revealed that snake venom proteins are produced as isoforms with various physiological functions and considerable inter- and intraspecific variation. This diversification of venom proteins may be caused by various molecular evolutionary mechanisms, including the accelerated evolution of venom genes [6–9] and alternative splicing of transcripts [10].

Since genomic information is essential for further studies on snake venom, we decoded the *P. flavoviridis* genome in 2018 [9]. In that study, we identified 60 venom genes belonging to 18 protein families. Interestingly, those genes arose through multiple duplication processes, resulting in many paralogs for each gene [9]. The habu genome analysis also demonstrated that six protein families (metalloproteinase (MP), serine protease (SP), C-type lectin-like proteins (CTLP), phospholipase A₂ (PLA₂), three-finger toxins (3FTX), and cysteine-rich secretory proteins (CRISP)) show accelerated evolution [9].

However, there are many questions to be answered in order to understand snake venom evolution, physiology, and pharmacology. One important question concerns how venom proteins are processed post-translationally to acquire their toxic functions. For example, one of the salient characteristics of venom proteins and peptides from venomous snakes, scorpions, spiders, and sea snails is that they are rich in disulfide bonds that stabilize their 3D structures [11,12]. PLA₂s, which are major habu venom components, contain seven disulfide bonds. Type III svMPs, such as flavorase, HR1A, HR1B, and HV11, possess 17 disulfide bonds, and a metalloprotease domain with disintegrin-like and cysteine (Cys)-rich domains at the C-terminus. In addition, CTLP, 3FTXs, and CRISP family proteins also contain four or more disulfide bonds. Furthermore, a certain type of CTLP forms an interchain disulfide bond with type III MPs, resulting in type IV MPs [13]. Thus, most *P. flavoviridis* snake venom proteins are Cys-rich proteins. It is thought that venom proteins pass through several modification processes to assume fully functional forms.

As mentioned above, although studies of venom transcriptomes have been conducted for many venomous snakes [3–5], there is little information on non-toxin transcripts expressed in venom glands. Moreover, there are even fewer reports focused on genes for enzymes that modify protein structure. Rokyta et al. (2012) [14] carried out transcriptomic analyses of venom glands of the eastern diamondback rattlesnake (Crotalus adamanteus) and found that genes for protein disulfide isomerases (PDIs) are highly expressed in the venom glands. In addition, Luna et al. (2013) [15] carried out proteomic analysis of the venom glands of Bothrops jararaca and found that proteins from the endoplasmic reticulum (ER), such as PDI and GPR78, and cytoplasmic proteins, such as 40S ribosomal protein, are enriched in venom glands. We expect that more genes involved in protein-modification functions are also highly expressed in venom glands of habus. Thus, to identify genes for candidate enzymes or chaperones of venom protein modification in *P. flavoviridis*, we carried out comprehensive transcriptomic analyses of venom glands and other organs. Here, we propose P4HB, PDIA3, SELENOM, and CALR as candidate enzyme genes for regulating/modifying venom proteins, since these genes are suggested to be strongly associated with the oxidative folding of venom proteins with multiple disulfide bonds.

2. Results

2.1. Identification of Candidate Genes for Modification, Folding, and Quality Control of Venom Proteins

In the *Protobothrops flavoviridis* whole-genome decoding project, we carried out thorough transcriptomic analyses of 20 organs and tissues of an adult snake, which included the venom gland (sample name: venom gland-specimen#1), venom fang-forming tissue, lung, liver, kidney, small intestine, colon, stomach, pancreas, heart, masseter muscle, brain (brain-specimen#1), eye, nose, pit organ, spleen, and ovary (accession # DRA006600, see Supplementary Table S2 of Shibata et al. [9]). In addition, we also analyzed mRNAs of the venom gland (venom gland-specimen#2), brain (brain-specimen#2), and accessory gland of another adult, as well as the neonate venom gland, and fetal fibroblasts from a neonate. We carried out comprehensive comparative analyses of these data to identify genes for enzymes with protein modification functions. All mRNAs were identified and characterized from habu whole-genomic information [9]. We expected that genes for venom-protein modification enzymes would also be more highly expressed in venom glands than in other organs and tissues, such as in digestive organs.

We first identified the 50 genes that are most highly expressed in habu venom glands (Table 1 and Supplementary Table S1). Based on the number of transcripts per million reads (TPM), there were no differences between the expression profiles of the two brains. The two venom glands also showed similar profiles of highly expressed genes. However, there were different numbers of venom protein genes in these two specimens (Table 1 and Supplementary Table S1). The venom gland from the second adult specimen included more venom protein genes (19 genes) (Table 1) than that from the first specimen (12 genes) (Supplementary Table S1), suggesting that the second specimen (Table 1) may have been actively synthesizing venom (Supplementary Table S1). Therefore, we profiled the expression of protein modification enzymes in the venom gland from the second specimen as a main focus (Table 1).

Table 1 shows the 50 most highly expressed genes in the venom gland from the second specimen. Fourteen of the top sixteen genes were categorized as venom protein genes: including bradykinin-potentiating and C-type natriuretic peptides (*CNP01*), *svPLA2-02* and -03, *svMP06*, *svPLA2-01*, -07, and -08, *svMP07* and -08, *svCTLP06*, and -07, *svMP04*, *svCLTP08*, *svMP05*, *svSP04* and *LAAO01* (Table 1), which were all the same as in the venom gland from the other adult specimen except for *svSP04* and *LAAO01* (Supplementary Table S1). Moreover, other venom protein genes, such as *CRISP01* and *02*, *svSP11*, *svMP09* and -10, *svMP01*, and *sv5Nase*, were also included in the top 50 genes of only the second adult specimen (Table 1).

Interestingly, we found that a gene for protein disulfide isomerase (*PDI*), also known as the beta-subunit of prolyl 4-hydroxylase (*P4HB*), was the ninth most highly expressed gene in one adult venom gland (Table 1) and the sixth most highly expressed in the other specimen (Supplementary Table S1). In addition, genes for Selenoprotein M, Calreticulin, and protein disulfide isomerase A3 (*SELENOM*, *CALR* and *PDIA3*) were ranked as the 19th, 23rd, and 31st most highly expressed genes, respectively (Table 1). These four genes are likely involved in the folding and quality control of venom proteins. The high expression profiles of these genes were further confirmed with another venom gland from the first adult specimen and in neonate venom glands (Figure 1 and Table 2). The high expression of *PDI* genes in habu venom glands was consistent with the *PDI* expression profile of *Crotalus adamanteus* [14], suggesting that snake venom glands actively express genes not only for toxic proteins, but also for enzymes that fold and modify proteins in other vertebrate tissues.

Table 1 also shows that many genes associated with translation and protein biosynthesis, including 40S and 60S ribosomal proteins, elongation factor 1 (EF1)-alpha 1 and EF1-beta are highly expressed in habu venom glands, an observation reported previously [14,15].

Gene Model ID	Gene Name	Annotation by Blast2GO	TPM	
habu1_s258676_g24318.t1	CNP01	Bradykinin-potentiating and C-type	438,781.0	Venom protein
habu1_s47459_g23397.t1	svPLA2-02, svPLA2-03	phospholipase a?	88,799,1	Venom protein
habu1 s14911 g21429.t1	svMP06	p-ii partial	76,130.9	Venom protein
habu1 $s9571 \sigma 19434 t1$	s71PLA2-01	phospholipase a2	44 208 4	Venom protein
habu1 $s9570 g19432 t1$	s77PLA2-07	phospholipase a2	34 366 9	Venom protein
habu1 $s9571 \circ 19433 t1$	S71PLA2-08	phospholipase a2	31,067,9	Venom protein
habu1 s14911 g21430 t1	s71MP07 s71MP08	snake venom metalloproteinase	30 773 7	Venom protein
habu1_s10061_g19809.t1	svCTLP06	flavocetin-A beta	15,201.6	Venom protein Modification enzyme for
habu1_s3443_g11710.t1	P4HB	protein disulfide-isomerase	8572.4	proteins analyzed in this study
habu1_s10061_g19810.t1	svCTLP03, svCTLP04, svCTLP07	snaclec stejaggregin-a subunit alpha	6786.0	Venom protein
habu1_s3258_g11210.t1	svMP04	metalloprotease piia	4821.0	Venom protein
habu1_s3168_g10977.t1	svCTLP08	venom c-type lectin mannose binding isoform 1	3305.5	Venom protein
habu1 s3258 g11211.t1	svMP05	snake venom metalloproteinase	2659.2	Venom protein
habu1_s6789_g17480.t1	svSP04	snake venom serine protease 2a homolog	2416.4	Venom protein
habu1_s2955_g10552.t1		elongation factor 1-alpha 1	2283.4	1
habu1_s402940_g24950.t1	LAAO01	l-amino acid oxidase	2124.7	Venom protein
habu1_s158_g00895.t1		forkhead box protein j1	2102.3	1
habu1 s5309 g15576.t1		40S ribosomal protein Ś25	1678.5	Ribosomal protein
habu1_s2696_g09642.t1	SELENOM	Selenoprotein M	1514.7	Modification enzyme for
habu1_s68614_g23647.t1		60s ribosomal protein 17	1508.8	Ribosomal protein
habu1 $s1917 = 007050 t1$		60s ribosomal protein 17a	1458 7	Ribosomal protein
habu1_s4106_g13431_t1		serine partial	1457.8	Venom protein
habu1_s3471_g11839.t1	CALR	Calreticulin	1422.0	Modification enzyme for
habu1 s4158 g13542 t1		55 kda erythrocyte membrane protein	1406 7	proteins analyzed in this study
habu1_s4037_g13262.t1		low quality protein: glutathione	1337.9	
habu1 s3765 g12603 t1		40s ribosomal protein s3a	1259.4	Ribosomal protein
habu1 s22025 g22470 t1	CRISP01 CRISP02	cysteine-rich venom partial	1201 7	Venom protein
habu1_s22023_g22469 t1	s7/SP11	snake venom serine protease 2	1194 1	Venom protein
habul $s6274 g17028 t1$	500111	ubiquitin-40s ribosomal protein s27a	1159.9	Ribosomal protein
habu1 $s1466 g05403 t1$		peptidyl-prolyl cis-trans isomerase a-like	1149.4	rub obolinar proteini
habu1 s399953 g24864.t1	svMP09. svMP10	metalloproteinase precursor	1138.2	Venom protein
habu1_s1390_g04983.t1	PDIA3	protein disulfide-isomerase a3	1085.9	Modification enzyme for
habu1 e2087 g07635 t1		ribosomal protein	1075.8	Ribosomal protein
habu1 $s7284$ $g17906$ $t1$		40s ribosomal protein s13	1075.8	Ribosomal protein
habu1 $_{0}$		Flongation factor 1-bota	000 5	Ribbsoniai protein
habu1_s/833_g1/854_t1		tmsh4x partial	971.8	
habu1 s^{2449} σ^{08987} t^{1}		60s ribosomal protein 131	965.9	Ribosomal protein
habu1 s7991 g18513 t1		40s ribosomal protein partial	934.3	Ribosomal protein
habu1 s37466 g23178.t1		growth-related translationally	928.1	Ribbsoniai protent
		controlled tumor protein	006.0	P ¹ 1 1 1
habu1_s16120_g21649.t1		60s ribosomal protein 118a	906.8	Ribosomal protein
habu1_\$7978_g18472.t1	svMP01, svMP02, svMP03,	disintegrin and metalloproteinase	902.7	Venom protein
	svMP11, nvMP57	domain-containing protein 28	202.0	
habu1_s1408_g05043.t1		60s ribosomal protein 15	892.3	Ribosomal protein
habu1_s74_g00428.t1	- -	60s ribosomal protein l11	872.1	Ribosomal protein
habu1_s6028_g16570.t1	sv5Nase	ecto-5 -nucleotidase	852.8	Venom protein
habu1_s399842_g24584.t1		60s ribosomal protein l6	852.5	Ribosomal protein
habu1_s188_g01057.t1		60s ribosomal protein partial	852.1	Ribosomal protein
habu1_s1608_g06024.t1		40s ribosomal protein s4	837.9	Ribosomal protein
habu1 s1167 o04045 t1		polyadenylate-binding protein 1	834.0	
		isoform x1	010.2	
nabu1_s2592_g09366.t1		cytoplasmic 2	819.2	

Table 1. Highly expressed genes in venom glands from an adult habu (venom gland-specimen#2: amain adult specimen).

Transcripts are expressed as "Transcripts per million" (TPM). red: Venom protein genes, blue: Protein modification genes analyzed in this study.

	venom gland (specimen #1) Venom gland (specimen #2) Neonate venom gland Accessory gland Venom fang forming tissue Lung Liver	Kidney Small intestine Colon Stomach Pancreas Heart Masseter muscle Brain (specimen #1) Brain (specimen #2) Eye	Nose Pit, infrared sensing organ Spleen Ovary Fetal_fibroblast
P4HB ⇔			2 3 4 5 6 7 8 9 9 10 11 12 13 13 14 15
SELENOM \Rightarrow			16 18 18 19 20 21
$CALR \Rightarrow$			22 23 24 24 24 25 26 26 26 27 28 28
PDIA3 ⇒			230 31 31 32 33 33 34 35 36 37 38 39 39 44 41 41 41 42 43 44 45 45 46 46 46 47 47 48 50 50

Figure 1. Heat map representing genes predominantly expressed in habu venom glands and other organs. Numbers for each gene correspond to genes in Table 1. Genes for candidate modification enzymes or chaperones related to protein folding are indicated with yellow boxes.

2.2. Expression Profiles of Highly Active Genes, P4HB, SELENOM, CALR, and PDIA3, Are Specific to Venom Glands

Our analysis confirmed the highly active expression of P4HB, SELENOM, CALR, and PDIA3 genes in habu venom glands. However, since P4HB, SELENOM, CALR, or PDIA3 participate in protein folding and modification as part of the ER chaperone system, the active expression of these genes in the venom gland does not necessarily imply the lower expression of these genes in other organs and tissues. To examine the expression profiles of these genes in various organs, we compared the expression levels of the top 50 venom gland genes in other tissues (Figure 1). The expression profiles of those 50 genes were shared only by adult and neonate venom glands, but not by other tissues, clearly indicating that the venom gland gene expression profile is unique and specific and is quite distinct from those of other tissues (Figure 1 and Table 2). Interestingly, the expression profile of the neonate venom gland was similar to that of two adult venom glands. This similarity is especially clear for chaperone proteins, such as PDIs, Selenoprotein M, and Calreticulin (Figure 1). In the 50 most highly expressed venom gland genes, another common expression profile was seen among the lungs, kidneys, small intestine, colon, stomach, and pit organ (Figure 1). In the case of genes encoding ribosomal proteins, high expression profiles were observed not only in venom glands, but also in the lungs and small intestine.

Venom Venom Venom Pit. In-Brain Brain Fang Fetal Gland Gland Small Massefrared Gene Stom-Pan-(Speci-(Speci-Gene Model ID Kidney Colon Nose Ovary (Speci-(Spec-Form-Lung Liver Intes-Heart ter Eye Sens-Spleen Fibrob-Name ach creas men men imen ing tine Muscle ing last men #1) #2) Organ #1) #2) Tissue 104.5 53.2 habu1_s2696_g09642 SELENOM 1928.2 1514.7 273.2 3.9 12.1 38.7 179.5 31.5 311.1 15.0 178.8 72.2 675.8 1184.5 407.1 213.1 4.1261.9 258.3 1564.9 71.7 167.4 139.7 319.0 159.2 habu1_s3471_g11839 CALR 361.1 1422.0 531.0 172.5 295.9 375.4 78.4141.8281.4 184.1 569.8 413.7 464.0 1859.1 8572.4 3765.2 1789.5 2265.3 349.7 habu1_s3443_g11710 P4HB13.303.9 955.2 1874.44083.9 2381.0 601.7 249.8 213.4 771.6 1268.3 1256.5 1916.5 645.9 917.0 habu1_s2789_g10033 PDIA2 0.1 0.2 0.40.7 0.1 0.0 0.40.3 59.1 0.1 0.2 0.6 0.5 0.3 0.5 0.5 0.2 0.6 1.1 0.1 habu1_s1390_g04983 427.0 134.7 PDIA3 2266.3 1085.9 601.2 856.4 420.2 614.1 1074.1 857.4 675.8 425.2 272.4 214.7 270.0 276.4 434.1 892.6 728.6 455.6 872.6 129.3 63.4 182.9 172.3 159.4 141.3 122.6 252.0 95.7 164.9 68.8 83.5 164.6 29.2 47.050.3 82.5 habu1_s10239_g19902 PDIA4 61.6 121.6 72.8 135.3 habu1_s2439_g08885 PDIA5 15.8 27.7 35.9 14.1 9.9 7.1 26.8 44.124.9 3.0 5.6 1.5 17.4 13.8 14.438.1 34.2 6.5 79.6 habu1_s604_g02858 PDIA6 181.2 388.5 146.1 71.4 18.6 33.8 35.9 146.4 40.3 29.6 34.4 30.5 68.4 66.3 62.9 195.6 149.5 83.5 541.2 habu1_s25599 g22719 ERp27 0.7 2.9 0.0 1.2 0.2 0.7 402.7 0.5 1.6 0.8 1.6 4.9 240.8 0.3 2.0 0.1 1.5 2.4 1.1 0.6 habu1 s66 g00353 ERp27 15.7 5.9 119.4 203.5 20.6 32.1 41.4 170.0 50.9 885.9 49.8 36.5 12.0 275.5 312.6 39.9 86.8 134.5 791.5 6.4 262.6 53.1 112.9 84.2 19.5 22.6 109.9 43.6 15.4 40.5 180.2 102.4 52.0 239.2 habu1_s399842_g24580 ERp29 38.4 100.1 10.1 44.9 48.3 32.8 habu1_s32648_g23040 ERp44 15.5 45.5 17.9 5.8 13.1 22.8 4.3 4.7 12.3 4.5 22.9 38.9 27.2 42.5 25.7 15.2 19.6 15.6 24.3 14.832.1 10.4 17.1 habu1_s1939_g07114 TMX1 23.1 25.4 22.9 23.8 21.4 11.9 46.5 26.3 6.9 5.0 20.3 11.2 30.5 12.9 21.1 21.6 17.8 25.9 5.4 habu1_s5624_g16107 TMX2 18.9 9.5 36.8 43.6 45.5 30.2 35.9 42.8 9.1 93.2 29.3 122.0 142.7 35.3 43.4 42.5 101.2 26.9 14.8 habu1_s4708_g14660 TMX3 8.0 3.8 43.3 23.1 19.7 13.0 19.3 13.6 2.2 20.5 4.5 42.7 43.6 15.3 54.6 33.7 12.5 25.6 56.0 51.4 25.7 0.5 10.8 habu1_s8507_g18763 TMX4 1.7 1.4 9.9 8.6 14.9 6.0 6.6 3.1 3.4 1.0 19.5 16.5 6.5 5.5 9.5 4.5 habu1_s3682_g12384 TXNDC5 53.1 112.9 84.2 19.5 22.6 100.1 109.9 43.6 10.1 36.5 15.4 48.3 40.5 180.2 102.4 32.8 52.0 239.2 38.4 44.9 habu1_s2744_g09787 49.0 TXNDC12 99.8 160.0 134.1 101.1 28.8 51.5 72.6 75.0 58.3 21.8 54.424.5 99.7 70.7 60.8 127.3 127.1 50.3 224.3 AGR2 22.4 3.0 406.3 183.2 0.3 77.9 199.0 777.2 649.1 1.8 0.5 12.3 1.4 33.8 884.0 356.9 0.7 0.7 0.2 habu1_s49803_g23445 6.8 habu1_s49803_g23446 AGR3 0.0 0.0 0.3 0.0 0.6 0.1 0.0 0.0 0.1 0.5 0.2 0.1 0.0 0.1 0.4 11.4 0.6 0.1 0.3 0.0 habu1_s2168_g07811 DNAJC10 2.2 5.7 10.8 3.3 0.6 4.84.5 6.8 9.6 0.4 11.0 5.2 18.1 16.3 3.8 10.0 10.7 2.9 102.0 58.3 habu1_s2370_g08604 CAŚO1 3732.7 97.1 485.9 6.5 4.7 12.9 118.6 5.1 16.0 1.2 19.3 42,708.5 61.8 59.8 142.0 114.3 1277.6 4.3 39.8 9.4 1.9 CASQ2 0.1 0.5 2.6 0.5 662.8 22.2 20.0 22.2 1.7 0.6 habu1_s8721_g18867 1.3 0.0 0.2 0.8 0.111.7 0.9 26.3 1.3 9.2 7.7 habu1_s1275_g04450 1.8 3.3 21.0 1.2 3.2 7.413.3 16.7 0.3 2.0 0.425.2 18.9 4.6 12.0 5.9 3.3 29.3 Ero1α habu1_s114618_g23966 125.2 255.9 117.9 Ero1β 75.1 247.6 91.8 44.0 50.424.2 20.7 37.1 45.1 182.8 21.5 8.5 49.6 59.8 60.1 498.9 200.8 habu1_s4277_g13858 PRX4 229.4 342.2 98.1 154.7 157.9 229.6 71.1 28.8 68.6 144.5 187.7 215.9 211.6 118.4 280.8 245.3 52.8 53.5 121.5 104.7 habu1_s27414_g22805

Table 2. Gene expression level of *P4HB*, *PDIA3*, *SELENOM*, *CALR*, *PDI* family genes and *PDI* family related genes by RNA-seq analysis of tissues and organs of *Protobothrops flavoviridis*.

2.3. Characterization of Genes for PDI Family Proteins and PDI-Related Proteins in Habu Venom Glands

As already mentioned, major venom proteins, such as PLA₂, MPs, CTLP, and 3FTXs contain many disulfide bonds. Genes for PDI are ultra-highly expressed in venom glands. Therefore, to identify and characterize genes for PDI family members (Figure 2) in the habu genome, we carried out molecular phylogenetic analysis, taking advantage of the habu whole genome. We constructed molecular phylogenetic trees using ORTHOSCOPE, a species tree-based ortholog group identification tool [16], with human PDIA3, PDIA4, P4HB, PDILT, PDIA2 (Figure 3), PDIA5 (Supplementary Figure S1), and PDIA6 (Supplementary Figure S2) as query sequences for each. In addition, genes for other PDI family-related proteins were also analyzed using ORTHOSCOPE to identify orthologs in P. flavoviridis. These PDI family-related proteins include oxidoreductases, such as ERP27 (Supplementary Figure S3), ERP29 (Supplementary Figure S4), and ERP44 (Supplementary Figure S5), anterior gradient protein 2 (AGR2) (Supplementary Figure S6), anterior gradient protein 3 (AGR3) (Supplementary Figure S7), and transmembrane oxidoreductases, such as TMX1, TMX2, TMX3, and TMX4 (Supplementary Figures S8–S11), CALR members (Figure 4), TXNDC5 (Figure S12), and TXNDC12 (Supplementary Figure S13), redox partner proteins of PDIs, and selenoproteins. Genes identified in the habu (P. flavoviridis) genome and in transcriptomes of a related snake (*P. mucrosquamatus*) are summarized in Figure 2 and Table 2.



Figure 2. Schematic representation of *PDI* gene family members in humans and their domain composition. Catalytically active domains (**a**) and (**a**') are shown in red; inactive domains in light blue (**b**) and dark blue (**b**'); Asp/Glu rich Ca²⁺-binding domains are designated in green; orange represents transmembrane domains; x-linker regions (brown); COOH terminal ER retention sequences (white). Habu *PDI* family Gene IDs for *P. flavoviridis* and *P. mucrosquamatus* that are orthologs of human *PDI* family genes, are shown on the right side. This figure on the left side was adapted and modified with permission from [17] (under the terms of the Creative Commons Attribution License).





Figure 3. Cont.

(B) Query sequence: PDIA4



Figure 3. Cont.





Figure 3. Cont.



Figure 3. Cont.



Figure 3. Molecular phylogenetic trees of *protein disulfide isomerases (PDIs)*. These trees were estimated using ORTHOSCOPE with human *PDIA3* (**A**), *PDIA4* (**B**), *P4HB* (**C**), *PDILT* (**D**), and *PDIA2* (**E**) as query sequences (highlighted in purple-colored), respectively. Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (*P. flavoviridis*) orthologs, and those in red boxes represent other snake species. Values beside branches represent the percentages of times that nodes were supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position.



Figure 4. Molecular phylogenetic tree of *CALR*. This tree was estimated using ORTHOSCOPE with human *CALR* as the query sequence (highlighted in purple-colored). The tree includes *CALR*, *CALR*-*like*, *CALR3* and *CANX* clades. Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (*P. flavoviridis*) orthologs, and those in red boxes represent other snake species. Values beside branches represent the percentages of times that nodes were supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.07 substitutions per position.

(a) PDI family

Molecular phylogenetic trees of *PDI* family genes were estimated with ORTHO-SCOPE using human *PDIA3* (Figure 3A), *PDIA4* (Figure 3B), *P4HB* (Figure 3C), *PDILT* (Figure 3D), and *PDIA2* (Figure 3E) as query sequences, showing that these trees constitute discrete clades of *PDIA3*, *PDIA4*, and *P4HB*. Each clade contained one habu ortholog: habu1_s3443_g11710.t1 for the *P4HB* clade, habu1_s1390_g04983.t1 for the *PDIA3* clade, and habu1_s10239_g19902 for the *PDIA4* clade, indicating these habu gene model ID transcripts are the same genes as in humans. However, these molecular phylogenetic trees

failed to support *P. flavoviridis* orthologs for *PDIA2* (Figure 3E) and *PDILT* (Figure 3D). Interestingly, genes for *PDILT* were not found in snakes.

These *PDI* gene family phylogenetic trees also contain clades other than *PDIA3*, *PDIA4*, *P4HB*, *PDIA2*, and *PDILT*, so we next examined molecular phylogenetic trees of the other *PDI* gene family members, including the human and habu genomes. Molecular phylogenetic analysis using ORTHOSCOPE showed that the habu genome contains one copy of each of the same genes as humans in the *PDIA5* (Supplementary Figure S1) and *PDIA6* clades (Supplementary Figure S2), since in each clade there is only one habu ortholog of the human gene, without any habu paralogs. These results indicate that in contrast to the extensive duplication of habu venom protein genes, genes for the PDI family are present as only single copies in the habu genome, although they are actively expressed in venom glands.

(b) Other PDI family-related members

The molecular phylogenetic tree of *ERP44* analyzed by ORTHOSCOPE using other *PDI* family member human *ERP44* (Supplementary Figure S5) as query included the four main clades (*ERP44*, *PDIA4*, *PDIA3*, and *P4HB*) that contained one habu ortholog each, habu1_s32648_g23040.t2 for the *ERP44* clade, habu1_s10239_g19902.t1 for the *PDIA4* clade, and habu1_s1390_g04983 for the *PDIA3* clade.

Molecular phylogenies of *AGR2* and *AGR3* using human genes as queries are shown in Supplementary Figures S6 and S7, respectively. First, the *AGR2* and *AGR3* trees both consisted of three main clades (*AGR3*, *AGR2*, and *TXNDC12*), and each clade contained one habu ortholog, habu1_s49803_g23446.t1 for the *AGR3* clade, habu1_s49803_g23445.t1 for the *AGR2* clade, and habu1_s2744_g09787.t1 for the *TXNDC12* clade.

Molecular phylogenies of *TMX1*, *TMX2*, *TMX3*, and *TMX4* (Supplementary Figures S8–S11, respectively) using human genes as queries indicated that a single habu ortholog exists in each clade: habu1_s1939_g07114.t1 for the *TMX1* clade, habu1_s5624_g16107.t1 for the *TMX2* clade, habu1_s4708_g14660.t1 for the *TMX3* clade, and habu1_s8507_g18763.t3 for the *TMX4* clade.

Molecular phylogenetic trees for *TXNDC5* and *TXNDC12* using each human gene as a query (Supplementary Figures S12 and S13) also indicated that the tree for *TXNDC5* contains three main clades (*PDIA4*, *PDIA6*, and *TXNDC5*), each of which includes a single habu ortholog: habu1s_10239_19902.t1 for the *PDIA4* clade, habu1_s604_g02858.t1 for the *PDIA6* clade, and habu1_s3682_g12384.t1 for the *TXNDC5* clade. The tree for *TXNDC12* contains the three main clades (*AGR2*, *AGR3*, and *TXNDC12*) (Supplementary Figure S13), and each clade contains a single ortholog: habu1_s49803_g23446.t1 for the *AGR3* clade, habu1_s49803_g23445.t1 for the *AGR2* clade.

Molecular phylogenetic trees for *CASQ1* and *CASQ2* each using the human gene as a query showed that only a single habu ortholog exists in each: habu1_s2370_g08604.t2 for the *CASQ1* clade and habu1_s8721_g18867.t1 for the *CASQ2* clade (Supplementary Figures S14 and S15).

(c) Calreticulin members

In a similar fashion, to examine how many genes are present for *CALR* members in the habu genome, a molecular phylogenetic tree of *CALR* was constructed using ORTHOSCOPE, with human *CALR* as a query sequence (Figure 4). We found that a gene annotated as habu *CALR*, habu1_s3471_g11839.t1, was included in the human *CALR* clade (Figure 4), indicating that the gene annotation for habu *CALR* is correct, since the habu gene annotated as *CALR* is an ortholog of human *CALR*. In addition, we found *CALR3*, *CANX* and *CLGN* homologs in the habu genome, habu1_ s2279_g08248.t1 for *CALR3* gene, habu1_s3471_g11839.t1 for *CALR* gene, habu1_ s4907_g14941.t1 for *CANX* gene (Supplementary Figure S17), habu1_ s2744_g09792.t1 for *CALRL* gene, and habu1_ s67827_g17445.t1 for *CLGN* gene (Figure 4). These observations were confirmed by molecular phylogenies of *CALR3* (Supplementary Figure S16), *CANX* (Supplementary Figure S17), and *CALRL* (Supplementary Figure S18). As with *PDI* and related families, members of the *CALR* family exist as single copies in the habu genome. (d) Redox partner proteins of PDI family proteins, $Ero1\alpha$ and $Ero1\beta$

Ero1, which has two kinds of paralogs, Ero1 α and Ero1 β , is the re-oxidizing enzyme specific for P4HB in the *PDI* gene family (Figure 5) [18–20]. The *Ero1* gene is conserved evolutionarily from yeast to humans. Although the functional difference between Ero1 α and Ero1 β has not yet been clarified, it is well known that they show different expression profiles. *Ero1\alpha* is expressed ubiquitously in various organs, while *Ero1\beta* is expressed only in secreting tissues [21]. We examined the number of *Ero1* copies in the habu genome and their expression profiles in different organs.



Figure 5. The relationship between PDI and Ero1 during protein folding by PDI. Redox regulation of PDI by Ero1 produces H₂O₂. PDI^{OX}: oxidized PDI; PDI^{RED}: reduced PDI; PDI^{MIXED}: PDI with the bound substrate; Ero1^{OX}: oxidized Ero1; Ero1^{*RED*}: reduced Ero1. PDI domains are colored as shown in Figure 2. This figure was adapted and modified with permission from Ref. [22]. (2012, Claudio Hetz et al., John Wiley and Sons).

The molecular phylogeny of *Ero1* (Figure 6) indicated two distinct clades of *Ero1* α and *Ero1* β , each of which contains habu orthologs, habu1_s1275_g04450.t1 (*Ero1* α) and habu1_s4277_g13858.t1 (*Ero1* β). Although the habu ortholog of *Ero1* β was a single copy, the gene was divided into two scaffolds tentatively annotated as habu1_s114618_g23966 (N-terminal side) and habu1_s4277_g13858 (C-terminal side).



Figure 6. Molecular phylogenetic tree for $Ero1\alpha$ and $Ero1\beta$. This tree was estimated using ORTHOSCOPE with human $Ero1\alpha$ and $Ero1\beta$ as query sequences (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (*P. flavoviridis*) orthologs, and those in red boxes represent other snake species. Values beside branches represent the percentages of times that nodes were supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.09 substitutions per position.

(e) TXN and PRX members

The molecular phylogeny of thioredoxin (*TXN*) members using ORTHOSCOPE, with human *TXN* as query sequences is shown in Figure 7. The tree consists of four main clades, namely *TXN2*, *TXN*, *TXNL*, and *TXNL1*, and clades *TXN2*, *TXN*, and *TXNL1* each contained one habu ortholog: habu1_s6210_g16942.t1 for *TXN2* clade, habu1_s3891_g12904.t1 for *TXN* clade, and habu1_s4406_g14149 for *TXNL1* clade. However, we failed to find an ortholog for *TXNL*.



Figure 7. Molecular phylogenetic tree for TXN. This tree was estimated using ORTHOSCOPE with human *TXN* as the query sequence (highlighted in purple). The tree includes the clades *TXN2*, *TXN*, *TXNL*, and *TXNL1*. Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (*P. flavoviridis*) orthologs, and those in red boxes represent other snake species. Values beside branches represent the percentages of times that nodes were supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.1 substitutions per position.

Molecular phylogenic trees of peroxiredoxin (*PRX*) isoforms genes were analyzed, with human *PRX1* (Figure 8A), *PRX2* (Figure 8B), *PRX3* (Figure 8C) and *PRX4* (Figure 8D) as query sequences. The tree comprises four discrete clades of *PRX1*, *PRX2*, *PRX3*, and *PRX4*. Each *PRX* clade contained a habu ortholog, habu1_s1518_g05675.t1 for the *PRX1* clade, habu1_s2388_g08681.t1 for the *PRX2* clade, habu1_s1132_g03944.t1 for the *PRX3* clade, and habu1_s27414_g22805.t1 for the *PRX4* clade.



Figure 8. Cont.

PRX3



Figure 8. Cont.

(B) PRX2

(C) PRX3



Figure 8. Cont.

(**D**) PRX4



Figure 8. Molecular phylogenetic tree for *PRX1*, *PRX2*, *PRX3*, and *PRX4*. These trees were estimated using ORTHOSCOPE with human *PRX1* (**A**), *PRX2* (**B**), *PRX3* (**C**), and *PRX4* (**D**) as query sequences (highlighted in gray-colored), respectively. Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (*P. flavoviridis*) orthologs, and those in red boxes represent other snake species. Values beside branches represent the percentages of times that nodes were supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.1 substitutions per position.

(A)

2.4. Characterization of the Habu SELENOM Gene

Although the *PDI* gene family has been studied in detail, *SELENOM* has been poorly characterized, including its exact function. Selenocysteine (U) is a unique, rare amino acid encoded by the codon, UGA. It is highly conserved in selenoproteins. We therefore performed an alignment with habu Selenoprotein M and confirmed that the position of selenocystein (U) is highly conserved in all gene sequences, including habu Selenoprotein M (Figure 9A).

In addition, to confirm whether the habu gene annotated as *SELENOM*, habu1_s2696_g09642.t1, is a human *SELENOM* ortholog, a molecular phylogenetic tree of *SELENOM* was constructed using ORTHOSCOPE, with human *SELENOM* as the query sequence (Figure 9B). The resulting phylogenetic tree includes a single habu gene annotated as *SELENOM* in the same clade as human *SELENOM*, and we found no paralogs of habu *SELENOM*.

	Signal Sequences	β1	a 1
		} _	
H. sapiens_SELENOM	MSLLLPPLALLLLAALVAPATA	ATAYRPDWNRLSGLTRARVETCGGU	QLNRLKEVKAFVTQD
M. musculus_SELENOM	MSILLSPPSLLLLLAALVAPATS	TTNYRPDWNRLRGLARGRVETCGGU	QLNRLKEVKAFVTED
Ga. gallus_SELENOM	MRRAALAALLLLAAAAG	-IERRPPRGLARGKVETCGGU	RLSRLPEVKAFVSQD
A. carolinensis_SELENOM	MAARVALRLLPVLLLAGPGPLLG	AFQLHPARLEGLARGKVESCRGU	RLNRLKEVKAFITED
P. flavoviridis_SELENOM	METPLLWLPLLLGLLSAVAPLRA	VQLDRSRLQWLARGKVESCGGU	RLNRLPEVKAFLNED
P. mucrosquamatus_SELENOM	METPLLWLPLLLGLLSAVAPLRA	VQLDRSRLQWLARGKVESCGGU	RLNRLPEVKAFLNED
T. sirtalis_SELENOM	MGTPLLLPLLLGLLSAVAPLRA	VQIDRSRLQWLARGKVESCGGU	QLNRLPEVKAFINED

Selenocystein (U)

a 2 —	β2	β3	β4	a 3	C-terminal Extension
IPFY	HNLVMKHLP	GADPELVLLGRR	YEELERIPLS	SEMTREEINALVQE	ELGFYRKAAPDAQVPPEYVWAPAKPPEETSDHADL
IQLY	HNLVMKHLP	GADPELVLLSRN	YQELERIPLS	SQMTRDEINALVQE	ELGFYRKSAPEAQVPPEYLWAPAKPPEEASEHDDL
IPLY	HNLEMKHLP	GADPELVLLSFR	YEELERIPLS	SDMTREEINQLVQE	ELGFYRKETPEAPVPEEFQFAPAKPLPTLTPRRAPAADGKTLSEQDKKDHPDL
LPLY	HNLVMKHLPO	GSDPELVLLSFQ	YKELERIPLS	SDMTREEINHLVQE	ELGFYRKEAPDAPVPKEYQLSPARPDTQATKAKAEGKGAAGNEL
LPLY	HNMDLKYLA	GADPELILLNIK	FEELQRIPLS	SDMSREEINQLMQE	ELGFYRKDTPDSPVPDAFQMAPANSLPSDVEAMKNRHAKEKKGAG
LPLY	HNMDLKYLAG	GADPELILLNIK	FEELQRIPLS	SDMSREEINQLMQE	ELGFYRKDTPDSPVPDAFQMAPANSLPSDVEAMKNRRAKEKKGAGSPDL
LPLY	HNMDLKYLA	GADPELILLNIK	FEELQRIPLS	SDMSRDEINQLMQQ	QLGFYRKETPDSPVPDAFQMAPANSLPPDVEAMKGKPGKQKKGAGGSDL

Figure 9. Cont.



Figure 9. Aligned amino acid sequences (**A**) and a molecular phylogenetic tree (**B**) for *SELENOM*. The conserved selenocysteine (U) residue, which is characteristic of selenoproteins, is shown in red. Secondary structures, α -helix and β -strand, are shown with blue boxes and orange arrows, respectively. The signal sequence is also shown in gray. This tree was estimated using ORTHOSCOPE with human *SELENOM* as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (*P. flavoviridis*) orthologs, and those in red boxes represent other snake species. Values beside branches represent the percentages of times that nodes were supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.06 substitutions per position.

2.5. Expression Profiles of Genes Encoding PDI and PDI-Related Proteins in the Habu

To determine the functions of candidate enzymes or chaperones of venom protein modification more specifically, we next investigated expression profiles of not only PDI family genes, but also their redox partner proteins in various organs and tissues of *P. flavoviridis*, including venom glands (Table 2). First, we focused on organ-specific genes in venom glands. Four genes for venom protein folding candidate enzymes or chaperones, *P4HB*, *SELENOM*, *CALR*, and *PDIA3*, showed higher expression levels in venom glands (TPM > 1000) than in other organs. However, except for *SELENOM*, they all showed high and ubiquitous expression in all organs (Table 2).

For example, *P4HB* showed a considerably higher expression level in venom glands (TPM (venom gland1/venom gland2) = 8572.4/13,303.9) than in other organs, in the range of (TPM =) 213–4084 (40 times higher than brain). *CALR* and *PDIA3* also showed high expression levels in venom glands (361.1/1422 and 2266.3/1085.9, respectively) and in other organs. *SELENOM* showed high expression levels in venom glands (1514.7/1928.2), and in other organs, except for the liver, ovary, kidney, stomach, and small intestine. In addition to these genes of the four protein-modification enzyme candidates, *PDIA4*, *PDIA6* (*P5*),

TXNDC12, and *CASQ1*, also showed high expression levels in venom glands (95.7/172.3, 181.2/388.5, 99.8/160, and 3732.7/97.1, respectively), while the expression levels of other protein-modification enzymes genes, such as *PDIA2*, *PDIA5*, *ERp46*, *ERp27*, *ERp29*, *TMX1*, *TMX2*, *TMX3*, *TMX4*, *AGR2*, *AGR3*, *DNAJC10*, and *CASQ2*, in the venom gland were lower than those in other organs.

Among the redox partner proteins of PDIs, PRX4 is an ER-localized PRX isoform that reduces hydrogen peroxide produced during Ero1-mediated oxidation of P4HB [23–25]. PRX4 oxidizes PDIA6 and TXNDC5 most efficiently [19,24,26]. In habus, *PDIA6* tends to have ubiquitously high expression, and especially in venom glands and fetal fibroblasts, specifically (TPM = 388.5 and 541.2, respectively). Additionally, in habus, all organs, including venom glands, showed high expression of *PRX4* (TPM = 229.4).

On the other hand, the expression level of *Ero1* in various habu organs showed that two kinds of paralogs of *Ero1*, *Ero1* α , and *Ero1* β , exhibit different expression profiles. *Ero1* α was ubiquitously expressed in all organs, but at low levels (TPM = 0.3–29.3). The expression level of *Ero1* α was especially low (TPM = 3.3) in venom glands, whereas *Ero1* β was comparatively highly expressed in venom glands (TPM = 75.1 to 247.6) as well as in the ovary, spleen, pancreas, and nose (TPM = 182.8–498.9). Thus, the expression levels of *Ero1* α and *Ero1* β differ, depending on organs and tissues, as noted in other vertebrates [27].

2.6. No Accelerated Evolution of Genes for Candidate Venom Protein Modification Enzymes or Chaperones

Our previous study of the habu genome revealed that most venom protein genes have been extensively duplicated and that major venom genes including *svMPs* exhibit accelerated evolution when examined by K_A/K_S analysis. In contrast, as shown here, genes involved in the modification of venom proteins are present as single copies in the habu genome. However, since these genes are also highly expressed in the venom gland, we examined whether accelerated evolution occurred in these genes by calculating K_A/K_S values using the Nei–Gojobori method.

We first examined the possible accelerated evolution of *P*4HB and *PDIA*3. However, K_A/K_S analysis against *P*4HB and *PDIA*3 genes showed no accelerated evolution of the *P*4HB and *PDIA*3 genes (Supplementary Figure S19). Further, K_A/K_S analysis indicated that habu *CALR* and *SELENOM* also showed no accelerated evolution (Supplementary Figure S20). K_A/K_S values for *PDI* family genes, *P*4HB and *PDIA*3, were almost 0, indicating that these two genes are highly conserved.

3. Discussion

In this study, in addition to genes that encode venom and ribosomal proteins, genes that encode PDI family proteins (human *P4HB* and *PDIA3* orthologs), Selenoprotein M (*SELENOM*), and Calreticulin (*CALR*), are highly expressed in habu venom glands. Since these enzymes or chaperones are involved in protein modification and folding in the cellular secretory pathway, the present results suggest the most likely involvement of PDI, PDIA3, Selenoprotein M, and Calreticulin in protein folding and modification during snake venom production as candidate modification enzymes or chaperones. Therefore, it is likely that PDI, PDIA3, Selenoprotein M, and Calreticulin are critical to confer toxic functions upon the disulfide bond-rich venom transcriptome.

Although studies of venom transcriptomes or proteomes have been conducted for many venomous snakes [3–5], there have been only a few reports focused on non-toxin transcripts or proteins, especially genes for enzymes that are involved in venom protein modification. In transcriptomic analysis of venom glands of the eastern diamondback rattlesnake (*Crotalus adamanteus*), Rokyta et al. (2012) [14] demonstrated that genes for PDIs are highly expressed in venom glands. Furthermore, proteomic analysis of *Bothrops jararaca* venom glands [15] found that proteins from the endoplasmic reticulum (ER), such as PDI and GPR78, and cytoplasmic proteins, such as 40S ribosomal protein, are enriched in venom glands. Our present transcriptome analysis indicated that *P4HB*, *PDIA3*, *SELENOM*, and

CALR, which are non-toxin genes, are among the 50 most highly expressed venom gland genes. Consistently, the 20 most highly expressed non-toxin transcripts listed by Rokyta et al. (2012) [14] included PDI, calreticulin, and PDIA3. Thus, our findings for habu venom glands support results for other pit vipers, which manifest compositional similarities.

In this context, with transcriptomic data for the Indian cobra (*Naja naja*), venom of which is rich in post-synaptic neurotoxins, Suryamohan et al. (2020) [28] showed the high expression of *PDI* (log (CPM) = 13.27~15.65 (*n* = 6)), as well as the moderately high expression of *PDIA3* [28]. While their transcriptomic data for 109 differentially upregulated genes (DUGs) in cobra venom glands did not include *P4HB* and *PDIA3*, these genes were more highly expressed in venom glands than in other organs. This seems reasonable since *PDIs* are expressed ubiquitously due to their essential functions as protein modification enzymes or chaperones. DUG analysis is undoubtedly useful to find genes of venom proteins since venom proteins are exclusively expressed in venom glands. However, this type of analysis may not always be useful to find genes that are ubiquitously and highly expressed, but more so in specific organs. When focusing on essential genes encoding nontoxic proteins, we need to consider the most appropriate analytical method. Nevertheless, all data shown herein for various snake species allow us to speculate that PDIs are essential for the modification and folding of venom proteins with abundant cysteines and multiple disulfide bonds.

Given that PDI works together with Ero1 as the primary catalyst for disulfide bond formation (Figure 5), we surmised that high expression of habu *PDI* accompanies high expression of habu *Ero1*. Indeed, our transcriptomic data from habu venom glands indicate high co-expression of *Ero1* β and *PDI*, whereas *Ero1* α is expressed to a much lesser extent. This was an unexpected result. However, previous studies demonstrated that *Ero1* β is constitutively and abundantly expressed in professional secretory tissues and can be strikingly induced in the course of the unfolded protein response [27,29]. Moreover, recombinant human Ero1 β is twice as active as Ero1 α in enzymatic assays [30]. Thus, it is possible that PDI is oxidized by Ero1 β more efficiently than by Ero1 α , leading to the efficient catalysis of oxidative protein folding by the PDI-Ero1 β combination [30]. The present finding of the high expression of *PDI* together with *Ero1\beta* supports our notion that habu PDI could be the primary venom protein modification enzyme.

In this study, ORTHOSCOPE, the leading tool for phylogenetic analysis, enabled us to estimate phylogeny for all PDI family orthologs and paralogs and their redox partner genes, SELENOM and CALR. This program is a species tree-based ortholog or paralog relationship identification tool, and all orthologs and paralogs for a specific gene are estimated automatically using this software [16]. Our previous data showed that each habu venom protein gene has many paralogs and undergoes accelerated evolution. In contrast, habu genes encoding candidate venom protein modification enzymes are orthologs for those of humans, and consistently there are no paralogs. Furthermore, our K_A/K_S analysis indicates that these genes never undergo accelerated evolution. In contrast to habus, proteomic analysis of cone snails, which have conotoxin, by Safavi-Hemami et al. (2011) demonstrated multiple PDI isoforms in cone snail venom glands, suggesting that cone snail PDI does undergo accelerated evolution [31]. Therefore, this contrast suggests that in habus, venom modification enzymes modify a wide variety of venom proteins, without requiring multiple isoforms derived from gene duplication or accelerated evolution. This difference between habus and cone snails may also help clarify molecular mechanisms for venom protein modification.

Variables relative to venom gland sampling conditions can also affect results. Luna et al. (2013) [15] indicated that a venom production cycle has two stages: a quiescent stage and an active stage. In *Bothrops jararaca*, this cycle continues for about 30–50 days [15]. Moreover, they demonstrated that the kinds and quantities of proteins detected in these two steps were totally different, suggesting that although there are differences in the length of this cycle among snake species, timing is quite important for venom gland sampling for transcriptomic or proteomic analysis. Results of venom gland proteomic analysis change

dramatically depending on the timing of sampling [15]. Thus, in this study, the number of samples was insufficient for statistical analysis. However, our results are consistent with those of other studies. Time-course single-cell RNA-seq analysis might allow for a better understanding of the accurate transition of the venom glands expression profile according to the entire timing of the cycle, as well as mechanisms of venom protein modification.

4. Conclusions

In this study, we identified four non-toxin genes quite highly expressed in venom glands of *P. flavoviridis*, namely *P4HB*, *PDIA3*, *SELENOM*, and *CALR*, and in general all of which encode protein folding enzymes or molecular chaperones in the ER. These are single-copy genes and are highly conserved among *Protobothrops* species, in contrast to the multiple copies and accelerated evolution of venom protein genes. Gene for a redox partner protein of PDI, *Ero1* β is also highly expressed in venom glands. The highly active expression of *P4HB*, *PDIA3*, *SELENOM*, and *CALR* in habu venom glands, which is also supported by previous reports, together with the disulfide bond-rich nature of venom proteins and the high co-expression of PDIs' redox partner genes, suggests that PDI, PDIA3, Selenoprotein M, and Calreticulin are strongly associated with venom protein modification. It is most likely that folding enzymes or molecular chaperones such as habu PDIs are critical to endow them with toxic functions.

5. Materials and Methods

5.1. Transcriptomic Data and Comparative Analyses

For transcriptomic analyses, a total of 22 samples were used as follows: 20 samples for 18 tissues and organs from two adult animals, and 2 samples for 2 tissues and organs from one neonate, including two venom glands specimens from two adult (sample name: venom gland-specimen#1, venom gland-specimen#2); two brains specimens from two adult (brainspecimen#1, brain-specimen#2); one specimen of eye, nose, infrared sensing pit organ, and venom fang-forming tissue, from one female adult; one specimen of lungs, liver, kidneys, small intestine, colon, stomach, pancreas, heart, masseter muscle, spleen, ovary, and accessory glands from the other female adult; one neonate venom gland tissue, as well as fibroblast cells, from one neonate. We extracted total RNA using a standard TRIzol protocol (Thermo Fisher Scientific, Waltham, MA, USA), and prepared cDNA libraries using an NEBNext[®] UltraTM Directional RNA Library Prep Kit for Illumina (New England Biolabs, Ipswich, MA, USA). RNA quality was checked with an Agilent Technologies 2100 Bioanalyzer using an Agilent RNA 6000 Nano Kit. Sequencing was performed using an Illumina Hiseq2500. De novo assembly of whole RNA sequence reads was performed using a de Bruijn graph-based program, Trinity [32,33]. Assembled transcripts were annotated with BLASTX against UniProt. All Illumina reads are available from DRA under accession No. DRA006600.

Raw Illumina reads were quality filtered and trimmed with Trimmomatic (v0.36) [34]. After adapter trimming and quality filtering of RNA-seq reads, index files for Kallisto [35] psudoalignment were generated with transcriptome information of our habu gene model. Gene expression levels (TPM) were calculated with Kallisto.

The 50 most highly expressed genes in venom glands were identified, and high-expression profiles of genes in the venom gland are listed in order (Table 1 and Supplementary Table S1).

5.2. Molecular Phylogenetic Analysis

We performed molecular phylogenetic analysis using ORTHOSCOPE, which is a novel pipeline for molecular phylogenetic analysis [16]. This unique pipeline, which is a species tree-based ortholog identification tool, automatically estimates the phylogenetic trees, which can detect all orthologs and paralogs automatically from whole genomic databases of selected species among all registered species. The software judges the range of the ortholog clade, which is the monophyletic group (clade), including the operational taxonomic unit (OTU) of interest, based on the topology of the species trees, which are given as quide trees. All phylogenetic trees were rearranged with Fig-TREE using the Newick format results of ORTHOSCOPE analysis.

To identify orthologs or paralogs of not only P4HB, PDIA3, SELENOM, and CALR, but also the other PDI family genes and PDI related redox partner genes, phylogenetic analysis was conducted with protein-coding DNA sequences of human PDIA3, PDIA4, P4HB, PDILT, PDIA2, CALR, both of Ero1α and Ero1β, TXN, PRX1, PRX2, PRX3, PRX4, SELENOM, PDIA5, PDIA6, ERP27, ERP29, ERP44, AGR2, AGR3, TMX1, TMX2, TMX3, TMX4, TXNDC5, TXNDC12, CASQ1, CASQ2, CALR3, CANX, and Danio rerio CALRL/CALR3b as queries, respectively, using ORTHOSCOPE, a species tree-based ortholog identification tool [16] with the following settings: analysis group, all Lepidosauria species and the default setting species for vertebrata analysis; E-value threshold for reported sequences, 1×10^{-3} ; number of hits to report per genome, 3; aligned site rate threshold within unambiguously aligned sites, 0; data set, DNA (Exclude 3rd); rearrangement BS (bootstrap) value threshold, 60%. In ORTHOSCOPE analysis, to construct neighbor joining (NJ) trees of all the above genes, all of the following analysis were carried out: amino acid sequences of each query gene were aligned with MAFFT (v7.221) [36] using the –auto strategy. Unaligned regions were trimmed with TrimAl (v1.2rev59) [37] using the -gappyout option. To generate nucleotide alignments, the corresponding DNA sequences were converted into the amino acid alignment using PAL2NAL [38]. For phylogenetic analyses, the NJ method [39] implemented in APE in R [40] for DNA alignments and FastME [41] for amino acid alignments were applied. Analyses of DNA alignments employ the most parameter-rich model in the program, the TN 93 model [42], with a gamma-distributed rate for site heterogeneity [43]. Analyses of amino acid alignments employ a widely used substitution model for nuclear gene analysis, the WAG model [44], with the gamma model. To evaluate the robustness of internal branches, 100 bootstrap replications are calculated for each data set.

5.3. Alignment and Molecular Evolutionary Analysis

We conducted the alignment of amino acid sequences of Selenoprotein M in a representative vertebrate using MUSCLE (v.3.8.31) alignment algorithms [45]. For K_A/K_S analysis, DNA sequences were aligned by using the ClustalW multiple alignment program (http: //clustalw.ddbj.nig.ac.jp (accessed on 31 August 2021)) [46]. Due to the complicated structure of the genes for this K_A/K_S analysis, all alignments were manually curated. For pairwise comparisons of nucleotide sequences of each the organism genes, the number of nucleotide substitutions per synonymous site (K_S) and per non-synonymous site (K_A) for proteincoding regions was computed according to the Nei–Gojobori method [47] using the Sqdif Plot online (http://www.gen-info.osaka-u.ac.jp/~uhmin/study/sqdifPlot/index.html (accessed on 31 August 2021)).

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10 .3390/toxins14050300/s1, Figure S1: Molecular phylogenetic tree estimated using ORTHOSCOPE with human PDIA5 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S2: Molecular phylogenetic tree estimated using ORTHOSCOPE with human PDIA6 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S3: Molecular phylogenetic tree estimated using ORTHO-SCOPE with human *ERP27* as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S4: Molecular phylogenetic tree estimated using ORTHOSCOPE with human ERP29 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.1 substitutions per position; Figure S5: Molecular phylogenetic tree estimated using ORTHO-SCOPE with human ERP44 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.1 substitutions per position; Figure S6: Molecular phylogenetic tree estimated using ORTHOSCOPE with human AGR2 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S7: Molecular phylogenetic tree estimated using ORTHO-SCOPE with human AGR3 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.09 substitutions per position; Figure S8: Molecular phylogenetic tree estimated using ORTHOSCOPE with human TMX1 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S9: Molecular phylogenetic tree estimated using ORTHO-SCOPE with human TMX2 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHO-SCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S10: Molecular phylogenetic tree estimated using ORTHOSCOPE with human TMX3 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.1 substitutions per position; Figure S11: Molecular phylogenetic tree estimated using ORTHOSCOPE with human TMX4 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S12: Molecular phylogenetic tree estimated using ORTHOSCOPE with human TXNDC5 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S13: Molecular phylogenetic tree estimated using ORTHOSCOPE with human TXNDC12 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S14: Molecular phylogenetic tree estimated using ORTHO-

SCOPE with human CASQ1 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S15: Molecular phylogenetic tree estimated using ORTHOSCOPE with human CASQ2 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S16: Molecular phylogenetic tree estimated using ORTHO-SCOPE with human CALR3 as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S17: Molecular phylogenetic tree estimated using ORTHOSCOPE with human CANX as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S18: Molecular phylogenetic tree estimated using ORTHO-SCOPE with Danio rerio CALRL as the query sequence (highlighted in purple). Sequences highlighted in blue indicate human orthologs. Those highlighted in pink represent habu (P. flavoviridis) orthologs, and those in red boxes represent other snake species. Values beside branches represent percentages of times that a node was supported in 100 bootstrap pseudoreplications implemented in ORTHOSCOPE. The scale bar indicates an evolutionary distance of 0.2 substitutions per position; Figure S19: K_A/K_S plot for major venom protein modification candidate enzyme or chaperone genes P4HB (left) and *PDIA3* (right). K_A and K_S values were calculated by the Nei-Gojobori method for each pairwise genes: venomous snake vs venomous snake (•), venomous snake vs other animal (including human) (Δ), and other animal vs other animal (\bigcirc). Venomous snakes include *Protobothrops flavoviridis* and *P. mucrosquamatus*, and other animals include human, mouse, chicken, anole and non-venomous snakes (*Python bivittatus* and *Thamnophis sirtalis*); Figure S20: K_A/K_S plot for major venom protein modification candidate enzyme or chaperone genes, CALR (left) and SELENOM (right). K_A and K_S values were calculated by the Nei-Gojobori method for each pairwise genes: venomous snake vs venomous snake (•), venomous snake vs other animals (including human) (**A**), and other animals vs other animals (\bigcirc). Venomous snakes include *Protobothrops flavoviridis* and *P. mucrosquamatus*, and other animals include human, mouse, chicken, anole and non-venomous snakes (Python bivittatus and *Thamnophis sirtalis*); Table S1: Genes highly expressed in venom glands from the first adult habu (venom gland-specimen#1: a sub-adult specimen).

Author Contributions: Conceptualization, E.S., N.S. and K.I.; methodology, E.S., K.H., A.I. and N.S.; software, K.H. and J.I.; formal analysis, A.I., K.H. and Y.S.; data curation, K.H.; investigation, A.I., E.S. and K.H.; resources, T.O.; writing—original draft preparation, A.I. and E.S.; writing—review and editing, A.I., E.S., H.S., T.O., K.I. and N.S.; supervision, E.S., K.I. and N.S.; project administration, E.S. and N.S.; funding acquisition, T.O., H.S. and N.S. All authors have read and agreed to the published version of the manuscript.

Funding: This research was supported by OIST Internal Fund to Marine Genomics Unit (N.S.) and by Grants-in-Aid of MEXT, Japan (#25440214 and #18H02498 to H.S., #24651130 and #23107505 to T.O.).

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: Not applicable.

Acknowledgments: We thank members of the Marine Genomics Unit and DNA sequencing sections of OIST. We also thank Steven D. Aird for technical editing of the manuscript. Supercomputing resources were provided by the Human Genome Center, the University of Tokyo (http://sc.hgc.jp/

shirokane.html (accessed on 31 March 2019)), and the National Institute of Genetics (https://sc2.ddbj.nig.ac.jp/index.php/en/ (accessed on 31 March 2019)). This work was partly performed in the Cooperative Research Project Program of the Medical Institute of Bioregulation, Kyushu University.

Conflicts of Interest: The authors declare no conflict of interest.

References

- Shibata, H.; Chijiwa, T.; Hattori, S.; Terada, K.; Ohno, M.; Fukumaki, Y. The taxonomic position and the unexpected divergence of the Habu viper, Protobothrops among Japanese subtropical islands. *Mol. Phylogenet. Evol.* 2016, *101*, 91–100. [CrossRef] [PubMed]
 Miyagi, Y. Dokujyakousyoh. *Clin. All-Round* 2004, *53*, 1245–1249.
- Aird, S.D.; Watanabe, Y.; Villar-Briones, A.; Roy, M.C.; Terada, K.; Mikheyev, A.S. Quantitative high-throughput profiling of
- snake venom gland transcriptomes and proteomes (*Ovophis okinavensis* and *Protobothrops flavoviridis*). BMC Genom. 2013, 14, 790. [CrossRef] [PubMed]
- 4. Aird, S.D.; Aggarwal, S.; Villar-Briones, A.; Tin, M.M.; Terada, K.; Mikheyev, A.S. Snake venoms are integrated systems, but abundant venom proteins evolve more rapidly. *BMC Genom.* **2015**, *16*, 647. [CrossRef] [PubMed]
- Castoe, T.A.; Hall, K.T.; Guibotsy Mboulas, M.L.; Gu, W.; de Koning, A.P.; Fox, S.E.; Poole, A.W.; Vemulapalli, V.; Daza, J.M.; Mockler, T.; et al. Discovery of highly divergent repeat landscapes in snake genomes using high-throughput sequencing. *Genome Biol. Evol.* 2011, *3*, 641–653. [CrossRef] [PubMed]
- Ohno, M.; Ménez, R.; Ogawa, T.; Danse, J.M.; Shimohigashi, Y.; Fromen, C.; Ducancel, F.; Zinn-Justin, S.; Le Du, M.H.; Boulain, J.C.; et al. Molecular evolution of snake toxins: Is the functional diversity of snake toxins associated with a mechanism of accelerated evolution? *Prog. Nucleic Acid Res. Mol. Biol.* 1998, 59, 307–364.
- Ohno, M.; Ogawa, T.; Oda-Ueda, N.; Chijiwa, T.; Hattori, S. Accelerated and Regional Evolution of Snake Venom Gland Isozymes; Ménez, A., Ed.; Perspectives in Molecular Toxinology; John Wiley & Sons: New York, NY, USA, 2002; pp. 387–400.
- Ohno, M.; Chijiwa, T.; Oda-Ueda, N.; Ogawa, T.; Hattori, S. Molecular evolution of myotoxic phospholipases A2 from snake venom. *Toxicon* 2003, 42, 841–854. [CrossRef] [PubMed]
- 9. Shibata, H.; Chijiwa, T.; Oda-Ueda, N.; Nakamura, H.; Yamaguchi, K.; Hattori, S.; Matsubara, K.; Matsuda, Y.; Yamashita, A.; Isomoto, A.; et al. The habu genome reveals accelerated evolution of venom protein genes. *Sci. Rep.* **2018**, *8*, 11300. [CrossRef]
- Ogawa, T.; Oda-Ueda, N.; Hisata, K.; Nakamura, H.; Chijiwa, T.; Hattori, S.; Isomoto, A.; Yugeta, H.; Yamasaki, S.; Fukumaki, Y.; et al. Alternative mRNA Splicing in Three Venom Families Underlying a Possible Production of Divergent Venom Proteins of the Habu Snake. *Toxins* 2019, *11*, 581. [CrossRef] [PubMed]
- Reeks, T.A.; Fry, B.G.; Alewood, P.F. Privileged frameworks from snake venom. *Cell Mol. Life Sci.* 2015, 72, 1939–1958. [CrossRef] [PubMed]
- 12. Mobli, M.; King, G.F. NMR methods for determining disulfide-bond connectivities. Toxicon 2010, 56, 849–854. [CrossRef] [PubMed]
- 13. Takeda, S.; Takeya, H.; Iwanaga, S. Snake venom metalloproteinases: Structure, function and relevance to the mammalian ADAM/ADAMTS family proteins. *Biochim. Biophys. Acta* **2012**, *1824*, 164–176. [CrossRef]
- 14. Rokyta, D.R.; Lemmon, A.R.; Margres, M.J.; Aronow, K. The venom-gland transcriptome of the eastern diamondback rattlesnake (*Crotalus adamanteus*). *BMC Genom.* **2012**, *13*, 312. [CrossRef] [PubMed]
- 15. Luna, M.S.; Valente RHe Perales, J.; Vieira, M.L.; Yamanouyea, N. Activation of Bothrops jararaca snake venom gland and venom production: A proteomic approach. *J. Proteom.* **2013**, *94*, 460–472. [CrossRef]
- Inoue, J.; Satoh, N. ORTHOSCOPE: An Automatic Web Tool for Phylogenetically Inferring Bilaterian Orthogroups with User-Selected Taxa. *Mol. Biol. Evol.* 2019, 36, 621–631. [CrossRef] [PubMed]
- 17. Galligan, J.J.; Petersen, D.R. The human protein disulfide isomerase gene family. Hum. Genom. 2012, 6, 6. [CrossRef] [PubMed]
- 18. Inaba, K.; Masui, S.; Iida, H.; Vavassori, S.; Sitia, R.; Suzuki, M. Crystal structures of human Ero1α reveal the mechanisms of regulated and targeted oxidation of PDI. *EMBO J.* **2010**, *29*, 3330–3343. [CrossRef]
- Sato, Y.; Kojima, R.; Okumura, M.; Hagiwara, M.; Masui, S.; Maegawa, K.; Saiki, M.; Horibe, T.; Suzuki, M.; Inaba, K. Synergistic cooperation of PDI family members in peroxiredoxin 4-driven oxidative protein folding. *Sci. Rep.* 2013, *3*, 2456. [CrossRef] [PubMed]
- 20. Okumura, M.; Kadokura, H.; Inaba, K. Structures and functions of protein disulfide isomerase family members involved in proteostasis in the endoplasmic reticulum. *Free Radic. Biol. Med.* **2015**, *83*, 314–322. [CrossRef]
- 21. Araki, K.; Inaba, K. Structure, mechanism, and evolution of Ero1 family enzymes. *Antioxid. Redox Signal.* **2012**, *16*, 790–799. [CrossRef]
- 22. Andreu, C.I.; Woehlbier, U.; Torres, M.; Hetz, C. Protein disulfide isomerases in neurodegeneration: From disease mechanisms to biomedical applications. *FEBS Lett.* **2012**, *586*, 2826–2834. [CrossRef]
- Zito, E.; Melo, E.P.; Yang, Y.; Wahlander, Å.; Neubert, T.A.; Ron, D. Oxidative protein folding by an endoplasmic reticulumlocalized peroxiredoxin. *Mol. Cell* 2010, 40, 787–797. [CrossRef] [PubMed]
- 24. Tavender, T.J.; Springate, J.J.; Bulleid, N.J. Recycling of peroxiredoxin IV provides a novel pathway for disulphide formation in the endoplasmic reticulum. *EMBO J.* **2010**, *29*, 4185–4197. [CrossRef] [PubMed]
- 25. Sato, Y.; Inaba, K. Disulfide bond formation network in the three biological kingdoms, bacteria, fungi and mammals. *FEBS J.* **2012**, 279, 2262–2271. [CrossRef] [PubMed]

- Kojima, R.; Okumura, M.; Masui, S.; Kanemura, S.; Inoue, M.; Saiki, M.; Yamaguchi, H.; Hikima, T.; Suzuki, M.; Akiyama, S.; et al. Radically different thioredoxin domain arrangement of ERp46, an efficient disulfide bond introducer of the mammalian PDI family. *Structure* 2014, 22, 431–443. [CrossRef] [PubMed]
- Dias-Gunasekara, S.; Gubbens, J.; van Lith, M.; Dunne, C.; Williams, J.A.G.; Kataky, R.; Scoones, D.; Lapthorn, A.; Bulleid, N.J.; Benham, A.M. Tissue-specific expression and dimerization of the endoplasmic reticulum oxidoreductase Ero1 beta. *J. Biol. Chem.* 2005, 280, 33066–33075. [CrossRef]
- Suryamohan, K.; Krishnankutty, S.P.; Guillory, J.; Jevit, M.; Schröder, M.S.; Wu, M.; Kuriakose, B.; Mathew, O.K.; Perumal, R.C.; Koludarov, I.; et al. The Indian cobra reference genome and transcriptome enables comprehensive identification of venom toxins. *Nat. Genet.* 2020, 52, 106–117. [CrossRef]
- Pagani, M.; Fabbri, M.; Benedetti, C.; Fassio, A.; Pilati, S.; Bulleid, N.J.; Cabibbo, A.; Sitia, R. Endoplasmic reticulum oxidoreductin 1-lbeta (ERO1-Lbeta), a human gene induced in the course of the unfolded protein response. *J. Biol. Chem.* 2000, 275, 23685–23692. [CrossRef]
- 30. Wang, L.; Zhu, L.; Wang, C.C. The endoplasmic reticulum sulfhydryl oxidase Ero1β drives efficient oxidative protein folding with loose regulation. *Biochem. J.* **2011**, 434, 113–121. [CrossRef]
- Safavi-Hemami, H.; Siero, W.A.; Gorasia, D.G.; Young, N.D.; Macmillan, D.; Williamson, N.A.; Purcell, A.W. Specialisation of the venom gland proteome in predatory cone snails reveals functional diversification of the conotoxin biosynthetic pathway. *J. Proteome Res.* 2011, 10, 3904–3919. [CrossRef]
- Grabherr, M.G.; Haas, B.J.; Yassour, M.; Levin, J.Z.; Thompson, D.A.; Amit, I.; Adiconis, X.; Fan, L.; Raychowdhury, R.; Zeng, Q.; et al. Full-length transcriptome assembly from RNA-Seq data without a reference genome. *Nat. Biotechnol.* 2011, 29, 644–652. [CrossRef] [PubMed]
- Haas, B.J.; Papanicolaou, A.; Yassour, M.; Grabherr, M.; Blood, P.D.; Bowden, J.; Couger, M.B.; Eccles, D.; Li, B.; Lieber, M.; et al. De novo transcript sequence reconstruction from RNA-seq using the Trinity platform for reference generation and analysis. *Nat. Protoc.* 2013, *8*, 1494–1512. [CrossRef] [PubMed]
- Bolger, A.M.; Lohse, M.; Usadel, B. Trimmomatic: A flexible trimmer for Illumina sequence data. *Bioinformatics* 2014, 30, 2114–2120. [CrossRef] [PubMed]
- 35. Bray, N.L.; Pimentel, H.; Melsted, P.; Pachter, L. Near-optimal probabilistic RNA-seq quantification. *Nat. Biotechnol.* **2016**, *34*, 525–527. [CrossRef] [PubMed]
- Katoh, K.; Misawa, K.; Kuma, K.; Miyata, T. MAFFT: A novel method for rapid multiple sequence alignment based on fast Fourier transform. *Nucleic Acids Res.* 2002, 30, 3059–3066. [CrossRef]
- Capella-Gutiérrez, S.; Silla-Martínez, J.M.; Gabaldón, T. trimAl: A tool for automated alignment trimming in large-scale phylogenetic analyses. *Bioinformatics* 2009, 25, 1972–1973. [CrossRef]
- Suyama, M.; Torrents, D.; Bork, P. PAL2NAL: Robust conversion of protein sequence alignments into the corresponding codon alignments. *Nucleic Acids Res.* 2006, 34, W609–W612. [CrossRef]
- 39. Saitou, N.; Nei, M. The neighbor-joining method: A new method for reconstructing phylogenetic trees. *Mol. Biol. Evol.* **1987**, *4*, 406–425. [CrossRef]
- 40. Popescu, A.A.; Huber, K.T.; Paradis, E. ape 3.0: New tools for distance-based phylogenetics and evolutionary analysis in R. *Bioinformatics* **2012**, *28*, 1536–1537. [CrossRef]
- Lefort, V.; Desper, R.; Gascuel, O. FastME 2.0: A Comprehensive, Accurate and Fast Distance-Based Phylogeny Inference Program. *Mol. Biol. Evol.* 2015, 32, 2798–2800. [CrossRef]
- 42. Tamura, K.; Nei, M. Estimation of the number of nucleotide substitutions in the control region of mitochondrial DNA in humans and chimpanzees. *Mol. Biol. Evol.* **1993**, *10*, 512–526. [CrossRef]
- 43. Yang, Z. Maximum likelihood phylogenetic estimation from DNA sequences with variable rates over sites: Approximate methods. *J. Mol. Evol.* **1994**, *39*, 306–314. [CrossRef] [PubMed]
- 44. Whelan, S.; Goldman, N. A general empirical model of protein evolution derived from multiple protein families using a maximum-likelihood approach. *Mol. Biol. Evol.* **2001**, *18*, 691–699. [CrossRef] [PubMed]
- 45. Edgar, R.C. MUSCLE: Multiple sequence alignment with high accuracy and high throughput. *Nucleic Acids Res.* 2004, 32, 1792–1797. [CrossRef]
- 46. Larkin, M.A.; Blackshields, G.; Brown, N.P.; Chenna, R.; McGettigan, P.A.; McWilliam, H.; Valentin, F.; Wallace, I.M.; Wilm, A.; Lopez, R.; et al. Clustal W and Clustal X version 2.0. *Bioinformatics* **2007**, *23*, 2947–2948. [CrossRef]
- Nei, M.; Gojobori, T. Simple methods for estimating the numbers of synonymous and nonsynonymous nucleotide substitutions. *Mol. Biol. Evol.* 1986, *3*, 418–426. [PubMed]