The Finding of a Group IIE Phospholipase A₂ Gene in a Specified Segment of Protobothrops flavoviridis Genome and Its Possible Evolutionary Relationship to Group IIA Phospholipase A₂ Genes

Kazuaki Yamaguchi 1, Takahito Chijiwa 1,*, Naoki Ikeda 1, Hiroki Shibata 2, Yasuyuki Fukumaki 2, Naoko Oda-Ueda 3, Shosaku Hattori 4 and Motonori Ohno 1

1 Department of Applied Life Science, Faculty of Bioscience and Biotechnology, Sojo University, Kumamoto 860-0082, Japan; E-Mails: g1319d02@m.sojo-u.ac.jp (K.Y.); cavia_tschudii_xist@yahoo.co.jp (N.I.); mohno218@ybb.ne.jp (M.O.)
2 Medical Institute of Bioregulation, Research Center of Genetic Information, Kyushu University, Fukuoka 812-8582, Japan; E-Mails: hshibata@gen.kyushu-u.ac.jp (H.S.); yfukumak@gen.kyushu-u.ac.jp (Y.F.)
3 Department of Biochemistry, Faculty of Pharmaceutical Sciences, Sojo University, Kumamoto 860-0082, Japan; E-Mail: naoko@ph.sojo-u.ac.jp
4 Institute of Medical Science, University of Tokyo, Oshima-gun, Kagoshima 894-1531, Japan; E-Mail: shattori@ims.u-tokyo.ac.jp

* Author to whom correspondence should be addressed; E-Mail: chijiwa@life.sojo-u.ac.jp; Tel.: +81-96-326-3984.

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Abstract: The genes encoding group IIE phospholipase A₂, abbreviated as IIE PLA₂, and its 5' and 3' flanking regions of Crotalinae snakes such as Protobothrops flavoviridis, P. tokarenisis, P. elegans, and Ovophis okinavensis, were found and sequenced. The genes consisted of four exons and three introns and coded for 22 or 24 amino acid residues of the signal peptides and 134 amino acid residues of the mature proteins. These IIE PLA₂s show high similarity to those from mammals and Colubridae snakes. The high expression level of IIE PLA₂ in Crotalinae venom glands suggests that they should work as venomous proteins. The blast analysis indicated that the gene encoding OTUD3, which is ovarian
tumor domain-containing protein 3, is located in the 3' downstream of IIE PLA2 gene. Moreover, a group IIA PLA2 gene was found in the 5' upstream of IIE PLA2 gene linked to the OTUD3 gene (OTUD3) in the P. flavoviridis genome. It became evident that the specified arrangement of IIA PLA2 gene, IIE PLA2 gene, and OTUD3 in this order is common in the genomes of humans to snakes. The present finding that the genes encoding various secretory PLA2s form a cluster in the genomes of humans to birds is closely related to the previous finding that six venom PLA2 isozyme genes are densely clustered in the so-called NIS-1 fragment of the P. flavoviridis genome. It is also suggested that venom IIA PLA2 genes may be evolutionarily derived from the IIE PLA2 gene.

**Keywords:** group IIE phospholipase A2; venom; evolution; gene cluster; comparative genomics

1. Introduction

Protobothrops genus snakes (Crotalinae, Viperidae) are distributed in the southwestern islands of Japan, P. flavoviridis and Oophis okinavensis in Amami-Oshima, Tokunoshima, and the Okinawa islands, P. tokarensis in the Tokara islands, and P. elegans in the Sakishima islands. The venoms of Protobothrops snakes are produced and stored in the venom glands, which are assumed to share an original developmental organ with the mammalian submaxillary glands. The injection of the venom through tubular front fangs causes various severe lesions in humans, such as myonecrosis, hemorrhage, and edema [1–3].

Phospholipase A2 (PLA2) [EC 3.1.1.4] catalyzes the hydrolysis of glycerophospholipid at the sn-2 position to produce free fatty acids and lysophospholipids [4]. As various forms of PLA2s work in almost whole organs in the body [5], they are divided into three categories: secretory, cytosolic, and Ca2+-independent PLA2s, based on the working modes [6]. Furthermore, novel transcriptome analysis in mammals showed that secretory PLA2s are classified into 11 groups: IB, IIA, IIC, IID, IIE, IIF, III, V, X, XIIA, and XIIB, according to the primary structures and the organs to be expressed [7]. Snake venoms also contain PLA2 isoforms as major toxic components. With regard to the primary structures and the modes of disulfide bond pairings [8], snake venom PLA2s are classified into group IA found in Elapidae (Elapinae and Hydrophiinae) venoms and group II found in Viperidae (Viperinae and Crotalinae) venoms [9]. Group II venom PLA2s are further divided into group IIA PLA2s ([Asp49]PLA2 forms) and group IIB PLA2s ([Lys49]PLA2 forms) [10,11]. P. flavoviridis (Crotalinae) group IIA venom PLA2 genes form a multi-gene family of 16–32 copies per haploid [12] and are located at two loci on a microchromosome [13]. The mathematical analysis of their nucleotide sequences delineated that they have evolved in an accelerated manner to acquire isozymes with diverse physiological activities [14–16]. Recently, the nucleotide sequence of the 31,348 bp genome fragment of P. flavoviridis was completely deciphered. It showed that six PLA2 isozyme genes are aligned in series and four of them are linked with the fragment of CR1 long interspersed nuclear element (LINE), named PcRTF (PLA2 gene-coupled reverse transcriptase fragment), at the 3’ terminus [13]. We call this fragment, composed of six PLA2 isozyme genes, NIS-1 (Figures 5 and 6). Fry et al. (2012) found
that group IIE PLA$_2$ was expressed in the venom glands of Colubridae snakes and proposed that it is a component of Colubridae snake venoms [17].

In the work reported here, we sequenced the segment-harboring novel IIE PLA$_2$ gene linked to OTUD3, which codes for the ovarian tumor domain-containing protein (OTUD) 3, in the P. flavoviridis genome. Moreover, the IIA PLA$_2$ gene was found in the 5' upstream of IIE PLA$_2$ gene. It became evident that the linear arrangement of the IIA PLA$_2$ gene, the IIE PLA$_2$ gene, and OTUD3, in this order, is common in the genomes of humans to snakes. It is also found that the clusters of the genes encoding various PLA$_2$s in the 5' upstream region of IIE PLA$_2$ gene in the genomes of humans to birds possibly correspond to those of six PLA$_2$ isozyme genes in NIS-1 fragment of P. flavoviridis genome. Possible conversion of the IIE PLA$_2$ gene to the IIA PLA$_2$ gene and its multiplication in Crotalinae snake genomes are discussed.

2. Materials and Methods

2.1. Materials

P. flavoviridis (Amami-Oshima Island, Japan), P. tokarensis, P. elegans, and O. okinavensis specimens were provided from the Institute of Medical Sciences of the University of Tokyo. High molecular weight genomic DNAs were prepared from the livers or the venom glands of the snakes according to the method of Blin and Stafford (1976) [18]. Total RNAs were prepared from various organs of the snakes according to the protocol of ISOGEN (Nippon Gene, Toyama, Japan). Restriction endonucleases and KOD plus DNA polymerase were purchased from Nippon Gene and TOYOBO (Osaka, Japan), respectively. The other reagents and antibiotics were from Nacalai Tesque (Kyoto, Japan) and TAKARA BIO (Shiga, Japan). Specific oligonucleotide primers were synthesized by GENNET (Fukuoka, Japan). All relevant ethical safeguards have been met in relation to animal experimentation.

2.2. Cloning and Sequencing of the Genome Segments Harboring IIE PLA$_2$ Gene and Its 5' and 3' Flanking Regions of Crotalinae Snakes and of Their IIE PLA$_2$ cDNAs

The personal expressed sequence tags (ESTs) database was constructed to unite the snake ESTs collected from Genbank and the ESTs of P. flavoviridis venom glands supplied from the Medical Institute of Biodefense of Kyushu University (Fukumaki and Shibata, unpublished) by utilizing the NCBI C++ Toolkit (National Center for Biotechnology Information, Bethesda, MD, USA). The tblastx analysis of the database was carried out with the nucleotide sequences of Homo sapiens IIE PLA$_2$ gene (NM_014589) [19] and Mus musculus IIE PLA$_2$ gene (NM_012044) [20] as query. The 1085 bp candidate subject, named isotig03504, was acquired. Isotig is a subsequence of an isogroup. An isogroup is an assembled transcription sequence approximately equivalent to that of a gene. Based on the nucleotide sequences of its 5' and 3' ends, the sense primer named SPII-3, 5'-gTA gAC TgC gCg TAA TTT gTA g-3', and the antisense primer named SPII-2, 5'-ggC CgA gTC CgT CgT AgC T-3', were designed (Figure 1). Genomic PCRs with these primers were carried out against the P. flavoviridis, P. tokarensis, P. elegans, and O. okinavensis genomes as the templates. Thus, about 2.6 kbp DNA fragments, which contain four exons coding for IIE PLA$_2$s, were obtained. Moreover,
to acquire the nucleotide sequences of its 5' and 3' flanking regions, adaptor ligation PCR, designated as “Ligation-Mediated PCR (LM-PCR)” (Takara Bio, Shiga, Japan), was conducted. The genomic DNA obtained by digestion with Hind III or Pst I was ligated with the adaptor nucleotide fragments with Hind III- or Pst I-terminus, designated as a “cassette.” Then, PCR was done with a C1 primer, 5'-gTA CAT ATT gTC gTT AgA ACg CgT AAT ACg ACT CA-3', which can anneal to the “cassettes,” and SPII-2 primer to amplify the genome fragment containing the 5' flanking region or SPII-3 primer to amplify the genome fragment containing the 3' flanking region (Figure 1). Moreover, to ensure the validity of the PCR, another PCR was conducted with the C2 primer, 5'-gTg TAg AAC gCg TAA gTA gAC TCA CTA TAg ggA gA-3', which can anneal to the “cassette” at the internal portion of the C1 primer and SPII-8 primer, 5'-CAg TCC TTC CAT AAA gCT C-3', to amplify the genome fragment corresponding to the 5' flanking region, or SPII-10 primer, 5'-CTT gCA CgT CTC Cgg ATT gTg-3', to amplify the genome fragment corresponding to the 3' flanking region to be overlapped to the fragments prepared as described above (Figure 1). Amplified genome fragments were ligated to pCR™-Blunt II-TOPO® vector (Life Technologies, Carlsbad, CA, USA), and transformed with DH5α competent cells (Takara Bio). The nucleotide sequences were determined with an ABI 3130xl capillary sequencer. The nucleotide sequences of Crotalinae IIE PLA2 genes and their 5' and 3' flanking regions are available in the Genbank/EMBL/DDBJ databases under Accession Nos. KM488538-KM488542.

**Figure 1.** The schematic representation of the genome segment harboring the Crotalinae IIE PLA2 gene. The nucleotide positions are numbered. Closed boxes represent open reading frames (ORFs) and open boxes untranslated regions (UTRs). Vertical bars indicate the positions of restriction enzyme sites. Arrow heads show the positions of primers.

2.3. Acquisition of the Genome Segment Harboring IIA PLA2 Gene, IIE PLA2 Gene, and OTUD3 of *P. flavoviridis*

Long genomic PCR was carried out with CH05, 5'-gAT TCg ggA TgA ggA CTC TC-3' [21], which anneals to the 5' UTR of the IIA PLA2 gene, and OTUD3-1, 5'-CCT Tgg TAg CCT CTT TgC CAT CAg-3', which anneals to the middle portion of intron 7 of OTUD3, against the *P. flavoviridis* genome in order to confirm whether the IIA PLA2 gene is located in the 5' upstream of the IIE PLA2 gene linked to OTUD3.

2.4. Expression Analysis by Semi-Quantitative RT-PCR of Crotalinae IIE PLA2 mRNA

The first strand cDNA of snake body organs was synthesized by reverse transcription and primer extension of the SMART cDNA Library Construction Kit (Clontech Laboratories, Mountain View,
CA, USA). Based on the nucleotide sequences of the genes encoding the IIE PLA₂s of Crotalinae snakes, the sense primer SPIIRT-1, 5'-CAC ATC ATC RAa CAC TTg AC-3', which commonly anneals to the middle portion of exon 2, was designed. The antisense primers SPIIRT-2 (5'-TCC TTC gCA CAg gCg gTT A-3', which can anneal specifically to the middle portion of exon 4 of the P. flavoviridis IIE PLA₂ gene) and SPIIRT-3 (5'-TCC TTC gCA CAg gCg gTT A-3', which can anneal specifically to the middle portion of exon 4 of the O. okinavensis IIE PLA₂ gene) were designed. The cDNA of β-actin, designated as ACTB, was amplified as an internal standard with the sense primer SHU7, 5'-CAg AgC AAg AgA ggT AT C C N-3' (N = G, A, T, C), and the antisense primer SHU8, 5'-TAg ATg ggC ACA gTg Tgg gN-3', as described previously [22]. The intensities of the bands of the amplified DNA fragments were estimated with Image J (NIH, Bethesda, MD, USA) and corrected relative to those of ACTB. The vertical numerals of the histogram are the values relative to that of the lung of P. flavoviridis, taken as one.

2.5. Phylogenetic Analysis of Secretory PLA₂s

A phylogenetic tree was constructed based on the amino acid sequences of the mature proteins of the secretory PLA₂s from various organisms (Homo sapiens, Mus musculus, Gallus gallus, Ornithorhynchus anatinus, Macaca mulatta, Pan troglodytes, Oryctolagus cuniculus, Canis lupus familiaris, Bos taurus, Laticauda semifasciata, Leioheterodon madagascariensis, Dispholidus typus, P. flavoviridis, P. tokarensis, P. elegans, and O. okinavensis) with the maximum likelihood method of the RAxML program [23]. The degrees of confidence for internal lineage in the phylogenetic tree were determined by the bootstrap confidence [24] using Kimura’s (1969) method to compute a distance matrix with 1000 replicates [25].

2.6. Comparative Structural Analysis of the Cluster Domains of Secretory PLA₂ Genes in the Genomes

The BLAST analysis done with the nucleotide sequences of the cDNA encoding human secretory IIA, IIC, IID, IIE, IIF, and V PLA₂s [5], against the draft genome databases of H. sapiens (GRCh37p.p13), M. musculus (GRCm38.p2), and G. gallus (Gallus_gallus-4.0), deciphered that secretory PLA₂s are distributed within a 300 kb genome segment of H. sapiens chromosome 1 (NC_000001 GPC_000000025), 200 kb of M. musculus chromosome 4 (NC_000070 GPC_000000777), and 21 kb of G. gallus chromosome 21 (NC_006108 GPC_000000738) (Figure 5). The OTUD3 (NP_056022 for H. sapiens, NP_082729 for M. musculus, XP_424363 for G. gallus) was found in the 3' downstream of a series of PLA₂ genes in these regions. In the case of Ophiophagus hannah, all the draft genome data (AZIM00000000.1) [26] were downloaded and made it personal O. hannah genome database. Referring to these gene arrangements in H. sapiens, M. musculus, and G. gallus, the contig harboring secretory PLA₂ genes was constructed through tblastn analysis against the O. hannah personal genome database. Then, the chromosomal loci of the secretory PLA₂ genes and OTUD3 were mapped on their scaffolds and compared with the P. flavoviridis genome segments harboring the IIE PLA₂ gene and OTUD3 (this study) and the NIS-1 fragment composing of six consecutive IIA PLA₂ genes [13].
3. Results and Discussion

3.1. The Structure of a 6436 bp P. flavoviridis Genome Segment Containing the IIE PLA₂ Gene

The tblastx analysis of the personal EST database gave three subjects, two of which were the wrong transcripts; one contained a stop codon with the redundant mutations and the other was fused with an irrelevant nucleotide fragment. As the deduced amino acid sequence encoded by the remaining subject, isotig03504, is similar to those of human (59%) and mouse IIE PLA₂ proteins (61%)—in particular, the positions of the half-cystine residues and the sequences of the Ca²⁺ binding site and the catalytic site are identical—this subject is thought to be the transcript derived from the gene encoding P. flavoviridis IIE PLA₂, designated as PfIIEPLA₂. Genomic PCR with SPII-2 and SPII-3 primers, which can anneal to the 5' and 3' terminal portions, respectively, of isotig03504, of the Amami-Oshima P. flavoviridis genome gave a 2616 bp fragment, which covers from the 5' portion of the first exon to the 3' terminal portion of the fourth exon of the PfIIEPLA₂ gene. In addition, LM-PCR of the Amami-Oshima P. flavoviridis genome gave a 6436 bp genome fragment harboring the PfIIEPLA₂ gene and its 5' and 3' flanking regions (Figure 1). Moreover, genomic PCR of P. tokarensis, P. elegans, and O. okinavenesis genome DNA also gave the genome fragments harboring the PtIIEPLA₂, PeIIEPLA₂, and OoIIEPLA₂ genes, together with their 5' and 3' flanking regions.

3.2. The Characteristic Primary Structures of Snake IIE PLA₂ Proteins

The deduced amino acid sequences of Crotalinae IIE PLA₂s are aligned with those of the IIE PLA₂s from two Colubridae genus snakes [17], from H. sapiens (NP_055404) [19] and M. musculus (NP_036174) [20], as well as those from the venom IIA PLA₂s from P. flavoviridis [27–29], venom IA PLA₂ from Laticauda semifasciata [30], and pancreatic IB PLA₂ from P. flavoviridis [31] (Figure 2). This alignment confirms that four PLA₂s, PfIIEPLA₂, PtIIEPLA₂, PeIIEPLA₂, and OoIIEPLA₂, from Crotalinae genus snakes are clearly classified into group IIE. Although the amino acid sequences of IIE and IIA PLA₂s are similar to one another, the C-terminal amino acid sequences from the 133th residue are distinct between them. The phylogenetic analysis including other group PLA₂s, such as IA, IB, IIC, IID, IIF, and V PLA₂, also shows that the IIE PLA₂s, including four novel Crotalinae PLA₂s, form an independent clade separated from other group PLA₂s (Figure 3). Moreover, IIE PLA₂s are further divided into those from snakes or mammals, in accordance with the differences in their C-terminal sequences.
**Figure 2.** The aligned amino acid sequences of IIE, IIA, IA and IB PLA from snakes and mammals. The positions are numbered from the first residue of the signal peptides. The half-cystines are shown in shaded letters. Abbreviations: **Dt**, *Dispholidus typus*; **Hs**, *Homo sapiens*; **Lm**, *Leioheterodon madagascariensis*; **Ls**, *Laticauda semifasciata*; **Mm**, *Mus musculus*; **Oo**, *Ovophis okinavensis*; **Pt**, *P. flavoviridis*; and **Pt**, *P. tokarenensis*. References: *Pf*IIEPLA₂ (this work); *Pt*IIEPLA₂ (this work); *Pf*IIEPLA₃ (this work); *Oo*IIEPLA₂ (this work); *Dt*Dis-1 (AFH69958) [17]; *Hs*IIEPLA₂ (NP_055404) [19]; *Lm*Lei-1 (AFH66960) [17]; *Ls*LsPLA₂cPm09 (BAB03302) [32]; *Mm*IIEPLA₂ (NP_036174) [20]; *Pf*PLA-B (BAG82670) [13]; *Pf*PLA-N (BAG82669) [13]; *Pf*PLA 6 (BAJ84552) [29]; and *Pf*PancPLA₂ (BAN08536) [31]. Numerals in parentheses show the identities of the amino acid sequences against those of the mature protein of *Pf*IIEPLA₂.
Figure 3. The phylogenetic tree constructed for the secretory PLA\(_2\)s of snakes and mammals, based on the amino acid sequences of their mature proteins. The numerals at the nodes represent bootstrap confidence values and the branch lengths represent the numbers of amino acid substitutions per site. Abbreviations: Bt, Bos taurus; Cf, Canis lupus familiaris; Gg, Gallus gallus; Mc, Macaca mulatta; Oa, Ornithorhynchus anatinus; Oc, Oryctolagus cuniculus; and Pn, Pan troglodytes. References: BtIIEPLA\(_2\) (NP_001179015) [33], C/IIEPLA\(_2\) (XP_544525) (automated computational prediction by GNOMON); DtDis-2 (AFH66959) [17]; GgIBPLA\(_2\) (NP_001138961) [34]; GgIIEPLA\(_2\) (NP_001171878) [35]; HsIBPLA\(_2\) (NP_000919) [36]; HsIAPLA\(_2\) (NP_001155199) [37]; HsIICPLA\(_2\) (NP_001099042) [38]; HsIIDPLA\(_2\) (NP_036532) [39]; HsIIFPLA\(_2\) (NP_073730) [40]; HsVPLA\(_2\) (NP_000920) [38]; LmLei-2 (AFH66961) [17]; LmLei-3 (AFH66962) [17]; McIIEPLA\(_2\) (XP_001094364) (automated computational prediction by GNOMON); MmIBPLA\(_2\) (NP_035237) [41]; MmIAPLA\(_2\) (NP_001076000) [42]; MmIICPLA\(_2\) (NP_032894) [41]; MmIIDPLA\(_2\) (NP_035239) [39]; MmIIFPLA\(_2\) (NP_036175) [20]; MmVPLA\(_2\) (NP_001116426) [41]; OaIIEPLA\(_2\) (XP_001505559) (automated computational prediction by GNOMON); OcIIEPLA\(_2\) (XP_002716050) (automated computational prediction by GNOMON); OoIIEPLA\(_2\) (this work); PfIIEPLA\(_2\) (this work); PfAPLA\(_2\) (this work); PfIIEPLA\(_2\) (this work); PfPLA-B (this work); PfPLA-N (this work); GgIBPLA\(_2\) (this work).
3.3. Venom Gland-Specific Expression of IIE PLA₂s in Crotalinae Snakes

In general, mammalian IIE PLA₂s are non-venomous somatic molecules. On the other hand, as mRNA-encoding IIE PLA₂s, abbreviated as Lei-1, 2, and 3 and Dis-1 and 2, were found in the venom glands of Colubridae snakes Leioheterodon madagascariensis and Dispholidus typus, respectively, it was proposed that the IIE PLA₂ of Colubridae snakes may work as a venom protein [17]. In the case of Crotalinae genus snakes, the expression analysis by semi-quantitative RT-PCR performed on several organs of P. flavoviridis and O. okinavensis showed that the IIE PLA₂s of P. flavoviridis and O. okinavensis are expressed at remarkably high levels in the venom glands (Figure 4). These results suggest that Crotalinae IIE PLA₂s are also venom proteins. Its expression in the lungs, though at a low level, may show that they act as an immune factor to neutralize bacteria infected through the air [43,44].

3.4. The Genome Structures Harboring Secretory PLA₂ Genes Are Conserved from Human to Snake

The BLAST analysis showed that OTUD3 is located in the 3’ downstream of the PfIIEPLA₂ gene in the P. flavoviridis genome (Figure 5). Based on the linear arrangement of the IIE PLA₂ gene and OTUD3 and the nucleotide sequences of the cDNA encoding human secretory IIA, IIC, IID, IIE, IIF, and V PLA₂s [5], BLAST analysis was made against the H. sapiens draft-genome database [45]. Then, it was found that the secretory PLA₂ genes are aligned in the 5’ upstream of OTUD3 within the 300-kb genome segment of H. sapiens chromosome 1. Interestingly, similar genome structures were found in the M. musculus, G. gallus, and O. hannah genomes (Figure 5). Particularly, it should be noted that the linear arrangement of the IIA PLA₂ gene, the IIE PLA₂ gene, and OTUD3, that is, the triplet genes, in this order is common in the genomes of human, mouse, and snake. In the case of O. hannah, it was found that three PLA₂ genes, that is, the PfPLA 6-like gene, the IID PLA₂ gene, and the IIF PLA₂ gene, are aligned in the 5’ upstream of the IIE PLA₂ gene and OTUD3. The PfPLA 6 gene is contained in the P. flavoviridis NIS-1 fragment [13,29]. In the case of the G. gallus genome, the IIA PLA₂ gene is found in the 5’ upstream of IIE PLA₂ gene, but the IIA and IIE PLA₂ genes are interrupted by the V PLA₂ gene. This unexpected location of the V PLA₂ gene in G. gallus is thought to be specific to birds.
The alignment of secretory PLA₂ genes in the 5' upstream of OTUD3 should be highly conserved among the vertebrates. In this work, we also acquired an 11 kb genome fragment of *P. flavoviridis*, which encompasses from the gene encoding venom IIA PLA₂, called PLA-B', at the 5' terminus to intron 7 of OTUD3 at the 3' terminus, by genomic PCR with CHO5, which anneals to the 5' UTR of the venom IIA PLA₂ gene, and OTUD3-1, which specifically anneals to the middle portion of intron 7 of OTUD3 (data not shown). The alignment of six IIA PLA₂ isozyme genes in the *P. flavoviridis* NIS-1 fragment is shown in Figure 5 [13]. *PfPLA 6* codes for a novel basic [Asp⁴⁹]PLA₂ [29], *PfPLA 1* (Ψ) is 91% similar in sequence to *PfPLA 6* with 10 nucleotide deletions, *PfPLA 2* [Lys⁴⁹]PLA₂ called BPII, *PfPLA 3* (Ψ) is a fragment from the second intron to the fourth exon of the *G6D49PLA₂* gene found in the *Trimeresurus stejnegeri* snake [46], *PfPLA 4* is a neurotoxic [Asp⁴⁹]PLA₂ called PLA-N, and *PfPLA 5* is a basic [Asp⁴⁹]PLA₂ called PLA-B. PLA-B and PLA-B' are the same isozymes with only one amino acid substitution at position 53, Glu or Gly, respectively [47,48]. It could be assumed that a cluster of IIA PLA₂ isozyme genes like NIS-1 is located in the 5' upstream of the triplet genes, that is, the PLA-B' gene, the IIE PLA₂ gene, and OTUD3, in the *P. flavoviridis* genome.

**Figure 5.** Diagrammatic representation of secretory PLA₂ genes in human, mouse, chicken, and snake genomes. The names of the organisms and the numbers of chromosomes are shown at left. Bold arrows indicate the areas of the genes in the chromosomes and the direction of arrows indicates the transcribing direction of the genes. Dashed lines indicate the regions where the nucleotide sequences are not determined. Organisms and genome information: *H. sapiens* chr. 1 (NC000001.10); *M. musculus* chr. 4 (NC000070.6); *G. gallus* chr. 21 (NC006108.3); *O. hannah* scaffold 1015.1 (AZIM01001014); *P. flavoviridis* NIS-1 (AB440236), *PfPLA 6* (AB588615), and *PfIIEPLA₂* (this work, KM488539).

3.5. The Structural Relationship between the IIE PLA₂ Gene and IIA PLA₂ Genes in the *P. flavoviridis* Genome

Two-BLAST analysis showed that the three highly homologous nucleotide segments, named Alpha, Beta, and Chai, are commonly contained in both the *PfIIEPLA₂* gene (Figure 6A) and venom IIA
PLA₂ isozyme genes clustered in the NIS-1 fragment (Figure 6B). Alpha, Beta, and Chai segments are about 0.4, 0.3, and 1.4 kbps in length with 69%–94% aligned scores. Their locations in the genes are distinctive. In the PfIIIEPLA₂ gene (Figure 6A), the Alpha segment is found in the 5' flanking region, the Beta segment in the 3' downstream of the Alpha segment in the 5' flanking region, and the Chai segment encompasses from the middle portion of intron 1 to the posterior portion of intron 3. The NIS-1 fragment consists of a series of IIA PLA₂ isozyme genes with or without PcRTF segment in the 3' terminus, each of which is bracketed as a unit in Figure 6B. Here, the Alpha and Chai segments are located in the 3' flanking region in this order and the Beta segment is in the anterior portion of intron 2 (Figure 6B). Therefore, it could be thought that after the prototype of venom IIA PLA₂ gene containing the three segments had been formed, its multiplication occurred as seen in NIS-1 fragment. Since the Alpha, Beta, and Chai segments are found at the particular locations, it is hard to imagine that the three segments had been introduced after multiplication of the venom IIA PLA₂ genes. On the other hand, it could be thought that the IIA PLA₂ gene had been converted from a IIE PLA₂ gene as a precursor with unknown mechanism.

**Figure 6.** The schematic representation of the locations of three typical nucleotide segments, named Alpha, Beta, and Chai, in the PfIIIEPLA₂ gene (A); and in six IIA PLA₂ genes in the NIS-1 fragment [13,29] of *P. flavoviridis* (B). Alpha, Beta, and Chai segments are shown by closed circle, closed star, and closed box, respectively. Gray boxes indicate exons of the PLA₂ gene and their numbers are shown as Roman numerals below the boxes. Boxes filled with oblique lines indicate the retroelements named PcRTFs [13]. The nucleotide position numbers are the same as those in Figure 1 and those reported previously [13]. The open star and open box mean the antisense nucleotide segments of Beta and Chai segments, respectively. The genome fragment, which encompasses from the venom IIA PLA₂ isozyme genes with or without PcRTF segment in the 3' terminus to the Alpha and Chai segments, is bracketed as a unit.
**Figure 7.** The constructed stem-loop structures of Chai-1 (A) and Chai-2 segments (B). The secondary structures are deduced based on their nucleotide sequences via DNA folding form of the mfold Web Server. The numerals at both termini of the segments are the position numbers of the corresponding nucleotides in Figure 6A.

3.6. Different Multiplication Processes between Non-Venomous Secretory PLA2 Genes and *P. flavoviridis* Venom IIA PLA2 Genes

As *OTUD3* is a single-copy gene and codes for an ordinary non-venomous protein, structural and functional boundaries must exist between the IIE PLA2 gene and *OTUD3*. The genome domain harboring the cluster of various PLA2 genes seems to be easily multiplied, unlike that harboring *OTUD3*. Thus, it could be assumed in the human to snake genomes that a series of secretory PLA2 genes have multiplied toward the 5' upstream direction from the IIE PLA2 gene as the ancestor and diversified to various PLA2 gene species (Figure 5). However, the multiplication pattern of *P. flavoviridis* venom IIA PLA2 isozyme genes in the NIS-1 fragment is considerably different from those of non-venomous secretory PLA2 genes. The IIA PLA2 isozyme genes of the NIS-1 fragment are periodically and densely repeated, whereas the non-venomous secretory PLA2 genes are considerably scattered and the proteins encoded are structurally diversified so as to be classified into IIA, IIC, IID, IIF,
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and V PLA2s (Figure 5). The two mechanisms may be considered for multiplication of PLA2 genes. As the two nucleotide sequences in Chai segments, named Chai-1 and Chai-2, can be predicted to form stem-loop structures (Figure 7A,B), which could be the scaffolding of the gene recombination [49,50], it appears that such gene recombination might have been involved in the multiplication of non-venomous secretory PLA2 genes. On the other hand, Castoe et al. (2011) pointed out that the quantities of retroelements like SINEs and LINEs in venomous snake genomes are much higher than those in nonvenomous snake genomes [51]. In fact, the associated forms between PLA2 genes and CR1 LINEs were found in the P. flavoviridis NIS-1 fragment as mentioned above [13]. This suggests that retrotransposition, such as 3'-transduction [52,53], with CR1 LINE has participated in the multiplication of venom IIA PLA2 isozyme genes.

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Author Contributions

Kazuaki Yamaguchi and Takahito Chijiwa conceived and designed the experiments; Kazuaki Yamaguchi performed the experiments and analyzed the data; Naoki Ikeda, Hiroki Shibata, Yasuyuki Fukumaki, Naoko Oda-Ueda, and Shosaku Hattori contributed reagents/materials/analysis tools; Kazuaki Yamaguchi, Takahito Chijiwa, and Motonori Ohno wrote the paper.

Abbreviations

CR 1 chicken repeat 1
EST expressed sequence tag
LINE long interspersed nuclear element
OTUD3 ovarian tumor domain-containing protein 3
Oo Ovophis okinavensis
ORF open reading frame
P Protobothrops
PLA2 phospholipase A2
SINE short interspersed nuclear element
UTR untranslated region

Conflicts of Interest

The authors declare no conflict of interest.

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