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Reappraisal of Vipera aspis Venom Neurotoxicity

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Background. The variation of venom composition with geography is an important aspect of intraspecific variability in the Vipera genus, although causes of this variability remain unclear. The diversity of snake venom is important both for our understanding of venomous snake evolution and for the preparation of relevant antivenoms to treat envenomations. A geographic intraspecific variation in snake venom composition was recently reported for Vipera aspis aspis venom in France. Since 1992, cases of human envenomation after Vipera aspis aspis bites in south-east France involving unexpected neurological signs were regularly reported. The presence of genes encoding PLA2 neurotoxins in the Vaa snake genome led us to investigate any neurological symptom associated with snake bites in other regions of France and in neighboring countries. In parallel, we used several approaches to characterize the venom PLA2 composition of the snakes captured in the same areas.

Methodology/Principal Findings. We conducted an epidemiological survey of snake bites in various regions of France. In parallel, we carried out the analysis of the genes and the transcripts encoding venom PLA2s. We used SELDI technology to study the diversity of PLA2 in various venom samples. Neurological signs (mainly cranial nerve disturbances) were reported after snake bites in three regions of France: Languedoc-Roussillon, Midi-Pyrénées and Provence-Alpes-Côte d’Azur. Genomes of Vipera aspis snakes from south-east France were shown to contain ammodytoxin isoforms never described in the genome of Vipera aspis from other French regions. Surprisingly, transcripts encoding venom neurotoxic PLA2s were found in snakes of Massif Central region. Accordingly, SELDI analysis of PLA2 venom composition confirmed the existence of population of neurotoxic Vipera aspis snakes in the west part of the Massif Central mountains. Conclusions/Significance. The association of epidemiological studies to genetic, biochemical and immunochemical analyses of snake venoms allowed a good evaluation of the potential neurotoxicity of snake bites. A correlation was found between the expression of neurological symptoms in humans and the intensity of the cross-reaction of venoms with anti-ammodytoxin antibodies, which is correlated with the level of neurotoxin (vaspin and/or ammodytoxin) expression in the venom. The origin of the two recently identified neurotoxic snake populations is discussed according to venom PLA2 genome and transcriptome data.

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INTRODUCTION

Snake venoms are complex mixtures of biologically active proteins. They contain several enzymes and toxins that act in synergy to fulfill the two main functions of the venom that are subduing and digesting prey. Diversity of snake venom within species has repeatedly been described over the last twenty years. The variation of venom composition with geography is an important aspect of this intraspecific variability. Several species of medical importance belonging to Viperidae and Elapidae families produce different clinical symptoms across the geographical range of their distribution [1,2,3,4,5]. The causes of this variability remain unclear [6,7]. In some cases the validity of the definition of the species is necessary and the apparent intraspecies variation is really interspecies variation. Interspecies hybridization may be a major mechanism of diversification of the composition of snake venoms. Whatever the origin, the diversity of snake venom is important both for our understanding of venomous snake evolution and for the preparation of relevant antivenoms to treat envenomations [5,6,8,9].

Three species of venomous snakes are of medical importance in Europe: Vipera aspis, Vipera berus and Vipera ammodytes (Vam). The clinical features of Vipera aspis aspis (Vaa) and Vipera berus berus (Vbb) envenomations are mostly local, but can be associated with systemic signs (gastrointestinal and coagulation disorders, low blood pressure) in cases of severe envenomation [10,11]. Envenomation by Vipera ammodytes ammodytes (Vam), a species located in Eastern Europe, can cause neurological symptoms [12]. Vam venom contains neurotoxic presynaptic phospholipases A2 (PLA2), the three isoforms of single chain ammodytoxin (AtxA, AtxB, AtxC) [13], whereas the venom of Vipera ammodytes meridionalis (Vam), another Vam subspecies of northern Europe, contains vipoxin, an heterodimeric postsynaptic PLA2 neurotoxin [14,15,16]. A geographic intraspecific variation in snake venom composition was recently reported for Vaa venom in France. Since 1992, the Marseille Poison Center has regularly observed cases of human envenomation after Vaa bites in south-east France involving unexpected neurological signs [17,18]. In France, neurological symptoms had previously only been described in cases of envenomation by a subspecies of Vipera aspis, Vipera aspis zinnikeri (Vaz) located mainly in the south-west of France.
Vaz venom neurotoxicity is due to a postsynaptic neurotoxin, PLA2-I, homologous to vipoxin [20]. The viper responsible for one of these “neurotoxic” envenomations was classified as Vaa on the basis of its morphology [17]. The venom gland mRNA of this neurotoxic Vaa snake was sequenced and led to the identification of two neurotoxins, AtxB, an isoform of ammodytoxin previously described only in the venom of Vamam, and vaspin, a neurotoxin similar to PLA2-I and vipoxin [21], composed of two subunits, an acidic one called vaspin A (vasA) and a basic one, called vaspin B (vasB). Among the genes encoding the venom PLAr of the same neurotoxic snake were genes encoding ammodytin I1 (AmI1), ammodytin I2 (AmI2), vaspin and the three isoforms of Atx, as well as a retrotransposon reported to be a phylogenetic marker of the genome of Viper ammodytes species [22,23]; a true Vaa genome only contains genes encoding AmI1, AmI2 and vaspin from this list. These observations indicate that there is a previously undescribed population of Vaa snakes found only in areas of contact with other snake species is able to cause neurological symptoms. We also describe a geographical variation in the composition of Vaa venom in the southwestern part of “Puy-de-Dôme” department: it is indistinguishable from a pure Vaz venom.

MATERIALS AND METHODS

Material

Pooled Vaa and pooled Vaz venoms were purchased from Latoxan (Valence, France).

Methods

Epidemiological and clinical analysis of snake bites and ecological analysis of Vipera snakes. This study is not a clinical trial. The patients were treated as usual like every envenomed victims in France, explaining why no special approval was necessary. The sole difference was the clinical feature with a new observed neurotoxicity, but there were no treatment protocol modifications in the medical management. Analytical assays were performed in blood samples collected for the classical exploration of bitten patients in order to evaluate the clinical gradation. Snake bites were analyzed retrospectively in 7 regions of France (Aquitaine, Auvergne, Midi-Pyrénées, Languedoc-Roussillon, Pays de Loire, Provence-Alpes-Côte d’Azur, Rhône-Alpes) from 1990 to 2002 (Figure 1). Unusual cases of neurotoxic envenomation reported by the Marseille Poison Centre from 2003 to 2005 were also included. Patient data (age, sex, time of bite,
clinical manifestations and treatment) were collected. The severity of envenomation of each patient was determined according to the classification established by [24]. Grade 1 envenomations (mild envenomations) were identified by the presence of local signs and absence of systemic symptoms; grade 2 (moderate envenomations) by regional oedema and moderate systemic symptoms; grade 3 (severe envenomation) by extensive oedema and severe systemic symptoms. Epidemiological data from 110 patients that presented a minor, moderate or severe envenomation were collated using Excel and Epi-Info software. We then excluded from the study cases of envenomation for which no indication of department was mentioned.

Authorization for capture and transportation of Vipera aspis and Vipera berus snakes in France were obtained from the prefectures of the following French administrative departments: “Alpes de Haute-Provence” (code no. 04), “Alpes-Maritimes” (code no. 06), “Puy-de-Dôme” (code no. 63), “Ariège” (code no. 69), “Haute-Pyrénées” (code no. 65), “Gironde” (code no. 33), “Hérault” (code no. 34), “Loire-Atlantique” (code no. 44), “Haut-Savoie” (code no. 74), and “Seine-et-Marne” (code no. 77) (see Figure 1). For Vipera ursinii (Vf) and Vau capture, transportation authorizations were obtained from the Ministry of Ecology and Sustainable Development of France. Vipera ammodytes montandoni (Vannon) snakes were captured at Strazha (Bulgaria), Vipera aspis francescici (Vafi) in Piemont (Italy), Vipera aspis hungry (Vah) in Calabria (Italy) and Vipera latastei latasti (Vil) in the region of Burgos (Spain). Seventy-two snakes were collected. The scientists in charge of snake captured in the various regions filled in a questionnaire. The data collected, including the sex, age and biotope (precise geographical localization, altitude) for each animal captured in France, are listed in Supplementary Table S1. They were identified by use of classical keys (including the numbers of ventral and sub ocular plates, scales between the labial plates and the eyes). When the dorsal pattern was clearly distinguishable from that of other Vipera aspis snakes from the same department, the subspecies was not indicated and replaced by a question mark. Snakes were of other ventral and sub ocular plates, scales between the labial plates and the biotope (precise geographical localization, altitude) for each animal captured in France, and incubated with restriction enzyme HpaII for 4 hours at 37°C to test for the presence of AtxA. AtxA contains an extra HpaII site to AtxC, yielding four cDNA fragments (63, 51, 129 and 201 bp) instead of three (63, 100 and 201 bp).

**Determination of protein concentration** The protein concentration of venom samples was determined using the Coomassie plus method (Pierce, Rockford, IL, USA).

**Fractionation of pooled Vaa and pooled Vaz snakes** The fractionation of pooled Vaa and pooled Vaz venoms was performed on a column containing Sephadex G100 superfine and using ammonium formiate 1M, pH 3.5 as running buffer. Fractions of 450 µl were collected at a flow rate of 2.6 ml per hour. The protein concentration of each fraction was measured on a spectrophotometer at the wavelength of 280 nm. Fractious corresponding to the same peak were pooled and tested for their PLA2 activity by fluorimetry [28].

**Microsequencing** Venoms 406 and 508 were subjected to PAGE (20% SDS-polyacrylamide gel) in reducing conditions. For venom 406, the gel was rinsed three times with water and stained overnight with AmidoBlack. A piece of gel containing AtxA/AC was cut out and digested with 2 µg/ml of endoproteinase Lys-C (Boehringer Mannheim GmbH, Mannheim, Germany) for 18 h at 35°C. The digested material was injected onto a DEAE column linked to a C-18 reversed-phase HPLC column and eluted with a 2–70% acetonitrile gradient in 0.1% trifluoroacetic acid. The molecular weight of eluted peptides was determined by surface enhanced laser desorption ionization-time of flight-mass spectrometry (SELDI-TOF-MS) on a Au proteinChip and the peptide corresponding to the C-terminal part of Atx was sequenced on a protein sequencer (494; Applied Biosystems, Foster City, CA). For venom 508, after the end of the electrophoresis, the gel was transferred onto PVDF overnight at a constant voltage of 15 V. The PVDF membrane was then stained with AmidoBlack and the band containing low molecular weight PLA2s was excised and subjected to Edman degradation on an Applied Biosystems 494 sequencer.

**Analysis of snake venom PLA2 contents by SELDI-TOF-MS** Venoms were diluted to 1.25 mg/ml in distilled water and spotted onto a variety of chemical surfaces: reverse phase H4 or H50 (hydrophobic surface; C-16 or C-4 (long chain aliphatic), weak cation exchanger (WCX2: carboxylic) or strong anion exchanger (SAX2: quaternary ammonium). Best and similar profiles were obtained with WCX2 and H4. H4 chips were activated with 10% acetonitrile and air dried prior to sample incubation. Venom samples (2.5 µg, final volume: 2 µl) were spotted, allowed to dry and spots were washed four times with 5 µl of distilled water. The samples were then air dried. WCX2 chips were equilibrated twice for 10 min with 150 µl of 100 mM sodium acetate pH 3.8, using a bioprocessor. Venom samples (2.5 µg) were diluted in 100 µl of 100 mM sodium acetate pH 3.8 containing 0.1% Triton and incubated on the chips for 30 min at room temperature under shaking. Then, the chips were washed: first for 5 min with 100 µl of 100 mM sodium acetate pH 3.8 containing 0.1% Triton, second, twice with 100 µl of 150 µl of 100 mM sodium acetate pH 3.8 and finally twice with double-distilled H2O. The samples were then air dried. Then, 0.5 µl of sinapinic acid (Ciphergen) saturated in 0.5 µl of 50% acetonitrile–5% trifluoroacetic acid was applied twice on each spot (WCX2 or H4 ProteinChips) and the spots were air dried. Molecules retained on the surfaces were visualized by reading the spots of each array in a SELDI-TOF-MS reader (PBSII; Ciphergen). Spectra were generated with 175 shots averaged at 210 laser intensity with a detector sensitivity of 7 and an accelerating voltage of 20 kV in positive mode with automatic data collection software.
3.0 program. External mass calibration curve was performed on one spot of each array by using ubiquitin (8564.8 Da), cytochrome C (12230.9 Da) and beta-lactoglobulin A (18363.3 Da).

Raw spectra were processed and analysed with the Ciphergen Express data manager software version 3.0 (CE; Ciphergen Biosystems). Spectra were externally calibrated with ubiquitin (bovine) (8564.8 + 1H), cytochrome C (bovine) (12230.9 + 1H), and beta-lactoglobulin A (bovine) (18363.3 + 1H). The baseline was established using default parameters and spectral intensities were normalized by total ion current (TIC). Consistent peak sets of similar mass across the spectra were generated with Ciphergen Express Cluster Wizard. This Wizard operates in three passes across the spectra. The first pass performs peak detection at high signal-to-noise ratio (s/n) to pick out well-defined peaks as starting points for forming clusters. A second pass selects lower s/n ratio peaks, within a mass window defined around the first pass peaks. The algorithm completes the clusters in a third pass by creating artificial peaks where none were detected in the first two passes, at the exact centre of clusters. In this analysis, the first pass was performed with an s/n threshold of 5, and the second pass with an s/n threshold of 2, in a 0.2% width mass window. Estimated peaks were added to complete clusters. Clusters were assembled between 13000 and 14000 Da, the expected size of venom toxins. The cluster lists contained normalized peak intensity values for each sample within a group and P-values were calculated between the medians of the peak intensities to detect significant differences in abundance for particular proteins.

Preparation of anti-ammodytoxin antibodies

Rabbit polyclonal antibodies against AtxA (Latoxan, Rosans, France) were prepared as described by [29].

Determination of venom immunological crossreactions by ELISA

We determined by western blot the cross-reactivity of anti-AtxA antibodies (data not shown). They react with vasB and Atx but don’t bind AmI1 or AmI2. Microtitration plates (Nunc, Roskilde, Denmark) were coated by incubation overnight at 4°C or 1 hr at 37°C with 100 μl of venom solutions (5 μg/ml) diluted in carbonate buffer (0.01M, pH 9.5). The plates were washed and incubated for 1 hr at 37°C with 100 μl of various dilutions of rabbit polyclonal anti-AtxA antibodies in PBS containing 3% BSA (PBS-BSA), then washed again. Peroxidase-labeled anti-rabbit IgG (100 μl of 1/1000 dilution in PBS-BSA) (Cappel/ICN Biomedicals, Aurora, Ohio, USA) were added to the wells and the plates were incubated for 1 hr at 37°C. After washing, substrate medium (100 μl of 10 mM sodium phosphate, pH 7.3, containing 2 mg/ml of OPD and 0.06% of perhydrol) was added to each well and the plates were incubated for 7 min in the dark at room temperature. The reaction was stopped by adding 50 μl of 0.5% sodium sulfite in 2N sulfuric acid to each well. Absorbance was measured at 490 nm using a microtitration plate reader (MR5000, Dynatech, France). Optical densities measured at dilutions of anti-AtxA IgG of 1/16,000 were used for cross-reactivity gradation: level 0: OD # 0.1, level 1: 0.1 < OD # 0.2; level 2: 0.2 < OD # 0.3; level 3: 0.3 < OD # 0.4; level 4: OD > 0.4.

RESULTS

Epidemiological study of envenomations in France: detailed symptomatology of neurotoxic envenomations according to geographical region

A hundred and ten cases of envenomation were reported between 1990 and 2005 in various regions of France (Aquitaine, Auvergne,
Midi-Pyrénées, Languedoc-Roussillon, Pays-de-Loire, Provence-Alpes-Côte d’Azur, Rhône-Alpes). Thirteen cases of envenomation for which no indication of department was mentioned were then excluded from the study and ninety-seven cases are reported in Supplementary Table S2. Envenomation cases were recorded from March to October (Figure 2A). They were distributed evenly from May to August in Languedoc-Roussillon (Figure 2B) and from March to October in Provence-Alpes-Côte d’Azur (PACA) (Figure 2C) whereas in Auvergne, peaks of envenomation were observed in May, July and August (Figure 2D). The age distribution was from 3 to 91 years. The severity of envenomation of each patient was determined according to the classification established by [24]. Fifteen cases of grade 1, 58 of grade 2 and 24 of grade 3 were reported. Neurological symptoms were only observed in grades 2 and 3 and constituted 17% of all moderate and severe envenomation cases. They were reported from April to October in four departments, “Alpes de Haute-Provence” (code no. 04), “Alpes-Maritimes” (code no. 06), “Aveyron” (code no. 12) and “Hérault” (code no. 34) (Supplementary Table S2). Figure 3 shows the prevalence of neurological symptoms in grade 2 and 3 envenomations for each department. The neurological signs are detailed in Table 1. They were mainly cranial nerve disturbances (ptosis, ophthalmoplegia, paresthesia, diploplia and dysphagia). Ptosis was observed in 100% of the cases. Ophthalmoplegia, drowsiness and dysphagia were recorded in all four departments. The most varied clinical symptoms were observed in “Alpes-Maritimes” and “Aveyron” with 9 different neurological signs (Table 1). We also analyzed the kinetics of appearance of each symptom. Drowsiness and diplopia were the most precocious symptoms, with mean appearance times of 5.83±2.63 hrs, 4.75±2.96 hrs after the envenomation, respectively. Ptosis and ophthalmoplegia were delayed, occurring after 9.03±5.6 hrs and 10±6.12 hrs.

Venom PLA2 screening at the genome and transcriptome level

To correlate the neurological signs observed with the venom composition, snakes were captured in the geographical regions in which the epidemiological analyses were conducted. Neurotoxins previously identified in the venoms of “neurotoxic” French vipers were phospholipases A2 or their homologs of 13 to 14 kDa. At least one snake of each region was sacrificed. cDNAs and genes encoding venom PLA2s were amplified from the venomous glands of V. berus snakes and probes specific of European viper venom PLA2s (AmI1, AmI2, vasA, vasB, AtxA/C and AtxB, AmL) were used for PLA2 profiling.

With regard to the French snake population, four classes can be identified based on their PLA2 genome composition (Table 2). The first class is characterized by the presence of genes encoding AmI1 and AmI2 and is only composed of V. laetata snake 651 and V. aspis snake 564. Interestingly, this latter snake belongs to an isolated population of V. laetata whose biotope is composed of garrigues and unlike other “classical” V. aspis snakes has lost vaspin encoding genes. The second class, characterized by genes encoding AmI1, AmI2, vasA and vasB, is composed of V. aspis snakes 502 (Puy-de-Dôme) and V. ursinii (Vu) snake 620 (Vaucluse). Although they differ in their genome composition,
class 1 and 2 have in common the unique expression of AmI1 and AmI2 in their venom, whereas class 3 includes only AmI1 and AmI2 as does that of Vipera ammodytes from Bulgaria.

The genome of V. latastei contains only AmI1 and AmI2 as does that of V. ammodytes from Bulgaria. All the snakes of this class present the same transcriptome profile (Puy-de-Dôme) and possesses genes encoding AmI1 and vaspin.

The third class is composed of V. aspis snakes 613 (Gironde) and 508 (Hautes-Pyrénées) and V. atra 456 (Puy-de-Dôme) and possesses genes encoding AmI1 and vaspin. All the snakes of this class present the same transcriptome profile for which all the venom PLA2 genes of the genome are expressed. The fourth class is composed of V. aspis snakes 406, 513, 514 (Alpes-de-Haute-Provence) and 603 (Alpes-Maritimes) and possess genes encoding AmI1, AmI2, vasA, vasB, AtxA, AtxB and AtxC as well as the Bov-B line retroposon. Interestingly, their transcriptome profiles differ at the level of vaspin and ammodytoxin isoforms expression, viper 603 being the only one that expressed all neurotoxins genes. This viper was captured after having bitten a human, who thereafter suffered from ptosis, ophthalmoplegia, agueusia, paresthesia and muscle weakness [17,18]. Snake 459 is the only V. atra specimen, which expressed AmI2 in its venom at the difference of V. aspis and V. atra, whereas vipers 456, 513 and 514 have the same transcriptome profile.

We also analyzed the PLA2 genome composition of European snakes. The genome of V. aspis snake captured in Piemont is similar to that of V. ammodytes from Italy contains Atx and AmL but not vaspin, as does that of V. ammodytes from Bulgaria. V. latastei latastei contains only AmI1 and AmI2 as V. aspis 651 and V. atra 564.

In addition to the complexity of the transcriptome profiles, the sequencing of transcripts encoding PLA2 genes in V. aspis and V. atra snake venom glands has shown that three isoforms of AmI1 and two of AmI2 are expressed in the venom of these snakes: the molecular weight of AmI1 isoforms ranges between 13676 and 13694 Da and that of AmI2 isoforms between 13526 and 13553 Da [30]. Two isoforms of vasA were identified with molecular weights of 13655 and 13665 Da whereas only one isoform of vasB was characterized (13841 Da) [30]. All these molecular weights are indicated assuming that the 14 cysteine residues are engaged in disulphide bonds.

**PLA2 proteome of snake venoms from various French regions using SELDI-TOF-MS proteinchip**

SELDI-TOF-MS ProteinChip technology was used to capture and analyze proteins of 13 to 14 kDa from crude venom samples. Several ProteinChips were tested to identify the chemical interaction able to bind proteins of the expected molecular weights. Hydrophobic (H4, H50) and hydrophilic (SAX2 and WCX2) arrays were used to differentiate venom samples from V. aspis (French department “Loire-Atlantique”) and V. atra (French department “Hautes-Pyrénées”) snakes (4 individual snakes for each subspecies). Best and similar profiles were obtained using cation-exchange (WCX2) and H4 (hydrophobic) chromatography surfaces. V. aspis venoms gave 3 peaks at molecular weights between 13500 and 13900 Da, and V. atra venom gave 3 major peaks between 13650 and 14060 Da (Figure 4). According to the the PLA2 transcriptome data of venom of this two subspecies, we would expect to detect by SELDI, in this range of molecular weights, AmI1 and AmI2 in V. aspis venoms and AmI1, vasA and vasB in V. atra venoms. To verify whether PLA2s were indeed immobilized on the SELDI chips, we first resolved pooled V. aspis and pooled V. atra venom samples by gel filtration on a Sephadex G75 column (Figure 5A and 5B). The main fractions presenting PLA2 activity (13 kDa for V. aspis and 30 kDa for V. atra venom) were isolated. Proteins contained in these two fractions were separated by SDS-PAGE (not shown) and transferred onto PVDF, and stained with Amidoblack. N-terminal sequences were determined by Edman degradation. Three sequences were identified in the 13 kDa V. aspis PLA2 fraction: HLSQF (AmI1, 13673 Da, gi:33187128), NLYQF (AmI2, 13500 Da, gi:33187128), NLYQF (AmI2, 13500 Da, gi:33187128).
Da; gi:33187122) and RDRPM (no hit found by Blast searches).
Two sequences were identified in the 30 kDa Vaz PLA2 fraction:
NLFQFGDMILQK (vasB, 13655 Da, gi:33187114) and
NLFQFGDMILQK (vasA, 13841 Da, gi:33187118). The profile
of these fractions was also analyzed by SELDI (Figure 6A). The
PLA2s are effectively bound onto the ProteinChip surface. The
profile matched those of the complete venom samples; this confirms that
vasA (NLFQFG) (Figure 6B).
Two sequences were identified in the 30 kDa PLA2 fraction, was thin (Figure 6A) whereas the peaks
characterized by the presence of two main peaks consistent with
the molecular weights of AmI2 and AmI1 (Figure 7). The relative
abundance of these two proteins differed between the venoms of
this group. Venoms collected in the Puy-de-Dôme were classified
into two groups (Figure 8): the PLA2 profiles of 451, 456, 521, 552,
553, 554 and 560 were similar to those of Vaz venom, whereas the
other profiles were heterogeneous. Venoms from the south-west of
France (Haute-Pyrénées, Haute-Garonne, Hérault) have diverse
PLA2 profiles (Figure 9). Vaz venom 459 presents an additional
peak of AmI2 not found in Vaz venoms from the Hautes-Pyrénées.
The PLA2 SELDI profiles of Vaz venoms from Hérault were
complex and heterogeneous: AmI2 was found in venoms 504 and
505 but not in venoms 506 and 509. Venom SELDI profiles from
the south-east of France were homogenous (Figure 10). Their
complexity is in agreement with transcriptome data, which
indicate the presence of AmI1, AmI2, vasA, vasB, AtxA, AtxB,
and AtxC transcripts in the venom glands of some of the snakes
captured in this region [30]. The SELDI profiles suggest that
AmI2 is not present in venom 800. Interestingly, an additional
peak of molecular weight 13751 to 13759 Da was identified in
venoms 406, 458 and 514. SELDI experiments indicate that the
molecular weight of AtxB is 13765 Da and AtxC has a molecular
weight of 13724 Da (Figure 10). The molecular weights of AtxA
and AtxB predicted from their sequences differ only by two Da,
and therefore C-terminal sequencing is required to differentiate
between these two isoforms. Venom 406 was separated by SDS-
PAGE on a homogeneous 20% acrylamide gel; Figure 6B) and transferred onto a PVDF
membrane. The lowest of the two bands migrating at 14 kDa
was sequenced: a mixture of 2 N-terminal sequences was obtained
corresponding to AmI1 (main sequence, HLSQFGDMIN) and
vasA (NLFQFG) (Figure 6B).

The PLA2 profiles of isolated Vaz and Vaz PLA2 fractions
matched those of the complete venom samples; this confirms that
PLA2s are effectively bound onto the ProteinChip surface. The
first peak of Vaz venom probably contains a mixture of vasA and
AmI1 PLA2. Indeed, a pure vasA peak, as found in the Vaz
30 kDa PLA2 fraction, was thin (Figure 6A) whereas the peaks
detected in Vaz venom were thicker (Figure 4).

Next, we analyzed several individual venoms from snakes from
French departments and neighboring countries (Figures 7–11).
Venoms were grouped according to their geographical localization
and/or their subspecies. Vaz-type venoms from the Loire-
Atlantique (code no. 44) and Seine-et-Marne (code no. 77) were
characterized by the presence of two main peaks consistent with

### Table 2. Genome and transcriptome analysis of snake venom PLA2

| Label number | Administrative French department or country | Viper | Genome composition | Venomous gland composition (cDNA) |
|--------------|---------------------------------------------|-------|--------------------|----------------------------------|
|              |                                              |       | Aml1 | Aml2 | vasA | vasB | AtxA | AtxB | AtxC | AmlI | ART  | Aml1 | Aml2 | vasA | vasB | AtxA | AtxB | AtxC | AmlI |
| 406          | Alpes-de-Haute-Provence (04) Vaa            | yes   | yes  | yes  | yes  | yes  | no   | yes  | yes  | no   | yes  | yes  | yes  | yes  | no   | no   | no   | no   | no   | yes  |
| 513          | Alpes-de-Haute-Provence (04) Va?            | yes   | yes  | yes  | yes  | yes  | no   | yes  | yes  | no   | yes  | yes  | yes  | yes  | no   | no   | no   | no   | no   | yes  |
| 514          | Alpes-de-Haute-Provence (04) Va?            | yes   | yes  | yes  | yes  | yes  | no   | yes  | yes  | no   | yes  | yes  | yes  | yes  | no   | no   | no   | no   | no   | yes  |
| 603          | Alpes-Maritimes (06) Vaa                    | yes   | yes  | yes  | yes  | yes  | no   | yes  | yes  | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 459          | Haute-Garonne (31) Vaz                      | nd    | nd   | nd   | nd   | nd   | nd   | nd   | nd   | nd   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 564          | Hérault (34) Vaa                            | yes   | yes  | no   | no   | no   | no   | no   | no   | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 613          | Gironde (33)                                | no    | no   | yes  | yes  | no   | no   | yes  | yes  | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 303          | Loire-Atlantique (44) Vaa                   | yes   | yes  | yes  | yes  | no   | no   | no   | no   | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 452          | Puy-de-Dôme (63) Vaa                        | yes   | yes  | yes  | yes  | no   | no   | yes  | yes  | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 502          | Puy-de-Dôme (63) Vaa                        | yes   | yes  | yes  | yes  | no   | no   | yes  | yes  | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 601          | Puy-de-Dôme (63) Vaa                        | yes   | yes  | yes  | yes  | no   | no   | yes  | yes  | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 456          | Puy-de-Dôme (63) Vaz                        | no    | no   | yes  | yes  | no   | no   | no   | no   | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 508          | Hautes-Pyrénées (65) Vaz                   | yes   | yes  | no   | no   | no   | no   | no   | no   | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 651          | Haute-Savoie (74) Vaa                       | yes   | no   | no   | no   | no   | no   | yes  | yes  | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 620          | Vaucluse (84) Vaz                           | yes   | yes  | yes  | yes  | no   | no   | no   | no   | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 605          | Italy Vaf                                   | yes   | yes  | yes  | yes  | no   | no   | yes  | yes  | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 604          | Italy Vah                                   | yes   | yes  | no   | no   | nd   | nd   | yes  | yes  | nd   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 608          | Spain VII                                   | yes   | yes  | no   | no   | no   | no   | no   | no   | no   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  |
| 709          | Bulgaria Vannon                             | yes   | yes  | no   | no   | no   | no   | nd   | nd   | nd   | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  | yes  

Legend: ART: Bov-b line retroposon; nd: not determined.

Venn diagrams illustrating the relationship of venom PLA2 encoding genes amplified by PCR using specific probes from liver DNA (genomics) and venomous gland cDNA (transcriptome) as described in the

Material and Methods section. ART: Bov-b line retroposon. nd: not determined.
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Aspis Venom Neurotoxicity

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correspond to AtxB (Figure 5D). This result is consistent with transcriptome data, which indicated that few AtxA transcripts are present in the venom glands of snake 406 [30]. SELDI profiles of Haute-Savoie Vaa venom are very similar to those of Vaa venom from Loire-Atlantique (Figure 11). AmI2 appears to be absent from the venoms of Vaf (Figure 11).

The spectra obtained were grouped into geographical clusters (cluster A: Loire-Atlantique; cluster B: Puy-de-Dôme; cluster C: Hautes-Pyrénées; cluster D: Hérault; cluster E: Haute-Garonne; cluster F: Alpes-Maritimes and Alpes-de-Haute-Provence; cluster G: Seine-et-Marne; cluster H: Vaf from Italy) and analyzed with CiphergenExpress software to study the relationships between the profiles. A list of 4 peaks with p-values < 0.05 were selected, which helped discriminating between the various venom samples: 13530 Da (AmI2), 13658 Da (AmI1 or vasA), 13753 Da (AtxA or B) and 13846 Da (vasB). A heat map was generated to see how well groups could be separated in a non-supervised way, based on the data set provided: it gave an interesting classification of the venoms (Figure 12). According to the dendrogram obtained, the venoms are separated into two major groups. The first group is distinguished by the down-regulation of the expression of AmI2 (13530 Da) and the over-expression of vasB (13854 Da). The second group is mainly characterized by the down-regulation of the expression of vasB. All venoms of Hautes-Pyrénées and Hérault are clustered into the first group, whereas all venoms from Loire-Atlantique, Seine-et-Marne and Haute-Savoie are found in the second group. Eight out of 11 venoms collected in Puy-de-Dôme are clustered in the first group whereas venoms from Alpes-Maritimes and Alpes-de-Haute-Provence are distributed in a homogeneous way in the two groups. A heterogeneity is also noted for Vaf venoms, which are distributed into the two groups (Figure 12). Venoms 401, 406, 458 and 514 were clearly characterized by the over-expression of a peak at an average molecular weight of 13753 Da. This is consistent with the molecular weight of AtxB since we have shown that the level of expression of AtxA is very low in the venom glands of snakes 406 and 514 [30] and only AtxB was detected in venom 406 by microsequencing (Figure 6B).
Immunome of snake venoms

Cross-reactions to Atx antibodies were previously shown to identify the presence of neurotoxins (Atx and/or vaspin) in the blood of patients presenting neurological symptoms following snake bite by vipers in “Alpes-Maritimes” and “Alpes-de-Haute-Provence” (south-east of France) [18]. The same approach was therefore used to identify snakes that express neurotoxins in their venom (Table 3). Vaa venoms from Loire-Atlantique, Seine-et-Marne and Haute-Savoie did not show any cross-reaction with anti-Atx antibodies consistent with the absence of neurological cases of envenomation in these areas (Table 3).

Vu venom 620 (Table 3) and Vu venoms from Alpes-Maritimes (data not shown) did not cross-react with Atx, showing that vaspin expression is low or inexistent. Venoms from Puy-de-Dôme had diverse cross-reactivities ranging from 0 to 2 depending on their area of capture. A higher heterogeneity was observed among venoms from Alpes-de-Haute-Provence with cross-reactivities ranging from 0 to 4. The Atx cross-reaction scores for snake venoms from Hautes-Pyrénées were between 1 and 3. Venoms from Hérault and Haute-Garonne were all highly reactive with anti-Atx antibodies. We also tested the cross-reactivities of Vaz venom samples collected in other departments of the south-west of France: Tarn (code no. 81), Aveyron (code no. 12), Pyrénées-Atlantique (code no. 64), Aude (code no. 11) and Pyrénées Orientales (code no. 66). They all presented maximum scores of cross-reactions (data not shown).

Venoms from Vll from Burgos (Spain) did not show any cross-reactivity with anti-Atx antibodies, consistent with the PLA2 composition of the Vll genome, which includes only AmI1 and AmI2 genes. Vaf venoms collected in Tuscany, Liguria and Calabria (Italy) showed Atx cross-reactivity scores of 1 to 3.

Interestingly, all venoms from SELDI group 1 but one (452) had Atx cross-reactivities ranging from 1 to 4 whereas 73% of venom from group 2 did not cross-react with Atx. This observation shows that the SELDI technology is definitely efficient to differentiate Vaz-type neurotoxic venoms from classical ones. Neurotoxic atypical venoms like those from the south-east of France are less easily identified unless they clearly over-express Atx, which can be easily identified in the heat map.

Figure 5. Isolation of PLA2 fractions from pooled Vaa and pooled Vaz venoms on Sephadex G100 Superfine column. (A) Pooled Vaa and (B) pooled Vaz were fractionated on Sephadex G100 superfine as described in the Material and Methods section. Fractions containing PLA2 activity are shown in red.

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Figure 6. PLA2 analysis by SELDI and SDS-PAGE. A) Vaa and Vaz PLA2 fractions isolated from pooled venoms as shown in Figure 5 were spotted on WCX2 ProteinChips. Normalised mass (m/z) for each peak (in Daltons (Da)) is demonstrated on the X-axis, while intensity (µA) is plotted on the Y-axis. B) Venoms 406 and 508 were separated by SDS-PAGE on a 20% homogeneous acrylamide gel under reducing conditions. N-terminal and internal sequences were determined as described in the Material and Methods section.

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Geographical distribution of neurotoxic snakes 
venoms and cases of envenomation

The unusual features of immunological cross-reactions observed in Puy-de-Dôme, Alpes-Maritimes and Alpes-de-Haute-Provence as well as the complexity of their classification according to their proteomic profile prompted us to compare the geographical distribution of snake venom data and cases of envenomation. The Vipera aspis vipers of Puy-de-Dôme whose venom belong to SELDI group 1 (CiphergenExpress) and cross-react with Atx are grouped in the south-west of the department (Figure 13). These venoms were classified in the same group as that of Vaz venoms from Hautes-Pyrénées. Interestingly, only classical features of Vipera aspis envenomations were observed for patients collected at Clermont-Ferrand hospital. However, it must be specified that the localization of the geographical sector where the bite took place is not specified by the epidemiologic investigation. The distribution was more complex in Alpes-Maritimes and Alpes-de-Haute-Provence (Figure 14). In Alpes-Maritimes, all snakes and all envenomation cases were neurotoxic as far as they were located at distance of the mediterranean coast. In Alpes-de-Haute-Provence, an equal number of neurotoxic and non-neurotoxic venoms and envenomations were reported. At Digne-les-Bains, a classical envenomation was observed and a non-neurotoxic viper was captured. At Colmars-les-Alpes, a neurotoxic case was reported whereas both neurotoxic and non-neurotoxic snakes were captured.
Cases of neurological envenomation by *Vipera aspis* snakes have regularly been reported since 1992 in the south-east of France (Alpes-Maritimes and Alpes-de-Haute-Provence). The presence of genes encoding PLA2 neurotoxins in the *Vaa* snake genome led us to conduct an epidemiological survey of snake bite in various regions of France. Neurological signs were reported after snake bite in three regions of France: Languedoc-Roussillon, Midi-Pyrénées and PACA. All cases involved ptosis, which appears as a characteristic sign of neurotoxic *Vipera aspis* envenomations. The time to between envenomation and the onset of symptoms was variable, but symptom onset was often late. Ptosis and ophtalmoplegia, two of most frequent symptoms, were shown to occur after $9.03 \pm 5.6$ hrs and $10 \pm 6.62$ hrs respectively.

The clinical symptoms observed in “Alpes-Maritimes” and “Aveyron” were the most diverse, with nine different neurological signs. The diversity of symptoms observed was not related to the severity of envenomation, or to the age of the patients. Moreover, similar symptoms were observed after envenomation by venoms containing vaspin or ammodytoxin. This was true whether these toxins acted at the presynaptic part of the neuromuscular junction, like the monomeric ammodytoxin, or bound to the postsynaptic part, like heterodimeric vaspin.
We used several approaches to characterize the venom PLA₂ composition of snakes captured in the areas in which the epidemiological survey was performed. We analyzed the genes and transcripts encoding venom PLA₂s. We used SELDI technology to study the diversity of PLA₂ neurotoxins in various venom samples. Electrospray and MALDI-MS is a faster, more accurate approach than SDS-PAGE for the characterization of venom components [31,32]. SELDI-TOF-MS can be considered as an extension of the matrix-assisted laser desorption/ionization (MALDI)-TOF-MS method [33]. In the SELDI method, protein solutions are applied to the spots of ProteinChip Arrays, which have been derivatized with planar chromatographic chemistries. The proteins actively interact with the chromatographic array surface, and become sequestered as well as separated from salts and other sample contaminants by subsequent on-spot washing with appropriate buffer solutions. Prefractionation, a sample preparation prerequisite that complicates the MALDI analysis, often resulting in sample loss as well as artifactual qualitative and quantitative variances, is therefore not required for SELDI analyses. This is particularly interesting if one works with samples of which the quantities are reduced. In parallel, we evaluated venom neurotoxicity by analyzing cross-reactivity with anti-Atx antibodies. We have previously used this method to detect neurotoxins in the blood of patients bitten by vipers in the south-east of France who presented neurological symptoms [18]. Three snake populations were identified as worthy of particular attention based on their neurotoxic venom PLA₂ content.

Figure 9. SELDI profiles of venoms from the south-west of France. Peak molecular weights consistent with those of PLA₂s are indicated on the spectra.
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The genomic analysis of snakes captured in the south-east of France revealed considerable diversity in the genes encoding PLA₂. Genes encoding two types of neurotoxin, Atx isoforms and vaspin, were identified. These genes were not constantly expressed, as shown by transcriptome data. Consistent with this finding, venom SELDI profiles and neurotoxin production were highly variable and diverse. This finding is consistent with the substantial variability of symptoms in this region, in which both neurotoxic and classical envenomations are reported (Figure 14).

The genomes of Vaz snakes from Gironde and Hautes-Pyrénées were characterized by the absence of the AmI2 gene. Venom PLA₂ profiles were highly homogeneous. All contained neurotoxic PLA₂₈, regardless of where the snakes were captured. However, neurotoxin levels varied between venom samples, with the lowest level of variability for snakes from the center of the range of distribution and the highest level of variability for snakes collected in zones of intergradation between Vaz and Vaa snakes.

The genomes of Vaa snakes from Puy-de-Dôme were very similar to those of Vaa snakes from Pays-de-la-Loire, but one snake did not possess an AmI2 gene; interestingly, vaspin transcripts were detected in this snake only. SELDI profiles of venoms from Puy-de-Dôme revealed that PLA₂ composition was related to the

Figure 10. SELDI profiles of venoms from the south-east of France. Peak molecular weights consistent with those of PLA₂ are indicated on the spectra. doi:10.1371/journal.pone.0001194.g010
geographic location of the corresponding snake. Venoms from the west of the department were identical to *Vaz* venoms, and contained vaspin. Those from the center and east of the department displayed heterogeneous profiles, and no cross-reaction with anti-Atx antibodies was reported.

Our results raise two issues concerning the clinical expression of venom neurotoxicity and the origin of the two neurotoxic populations of *Vaa*.

A correlation was found between the expression of neurological symptoms in humans and the intensity of the cross-reaction of venoms with anti-Atx antibodies, which is correlated with the level of neurotoxin expression. Neurotoxic envenomations were reported only in the areas with the highest scores of Atx cross-reactivity. Atx cross-reactivity scores of 4 are frequent for snake venoms in the south-east of France, where neurotoxic envenomations have been regularly reported (Supplementary Table S2, Figure 14). No such score was ever reported for snakes from Puy-de-Dôme and Languedoc-Roussillon are probably hybrids between *Vaa* and *Vaz* snakes, as suggested by the following factors: 1) difficulties in identifying clearly their subspecies; 2) the heterogeneity of their PLA2 SELDI profiles, which demonstrated the expression of AmI2 in the venom of two snakes; 3) the composition of their venoms that present by SDS-PAGE bands specific of *Vaa* venoms as shown in Figure 15.

We will now consider the origin of the two neurotoxic populations of *Vaa*. Two data sets must be considered for the southeastern neurotoxic snake population: 1) the regular reports of neurotoxic envenomations in south-east France since 1992 suggesting the recent appearance of a new snake population and 2) the presence of genes encoding Atx isoform genes and Bov-B line retroposons in the genome of this neurotoxic snake population. We found published reports of envenomations with neurological signs in four departments of PACA: 1) Alpes-Maritimes [38], 2) Alpes-de-Haute-Provence [39,40], 3) Hautes-Alpes, and 4) Vaucluse [41]. The case from Alpes-de-Haute-Provence [40] was wrongly attributed to *Vipera ursinii*, which can clearly be excluded in light of recent observations [42]. These data indicate that the neurotoxic *Vipera aspis* population of the south-east of France did not appear in the nineties. Thus, regular report of neurotoxic snake bites in south-east France may be explained either by the careful survey performed by the Poison Center of Marseilles since the start of the 1990’s, which could have highlighted neurotoxic snake bites or to the modification of the neurotoxic snake population behaviour. Interestingly, the epidemiological survey has indicated that most of the neurotoxic snake bites have been reported in Hérault and Aveyron (as previously described by [37]). Venoms tested in these departments also presented the highest scores for Atx cross-reactivity. Thus, the amount of vaspin was shown to vary between snakes. Intergradation areas were sites of high levels of vaspin expression. Snakes captured in Haute-Garonne and Languedoc-Roussillon are probably hybrids between *Vaa* and *Vaz* snakes, as suggested by the following factors: 1) difficulties in identifying clearly their subspecies; 2) the heterogeneity of their PLA2 SELDI profiles, which demonstrated the expression of AmI2 in the venom of two snakes; 3) the composition of their venoms that present by SDS-PAGE bands specific of *Vaa* venoms as shown in Figure 15.

![Figure 11. SELDI profiles of venoms from Haute-Savoie (code no. 74) and Italy. Peak molecular weights consistent with those of PLA2s are indicated on the spectra.](doi:10.1371/journal.pone.0001194.g011)
bites occurred near dwellings or inside towns. The consecutive rarefaction of the preys to forest fires and to the drought that has prevailed for a few years in the south-east of France could explain this phenomenon.

We analyzed the occurrence of neurotoxic envenomations in neighboring countries. We also analyzed the genome, transcriptome and proteome characteristics of snake venoms in these countries. Neurological cases attributed to \textit{Vaf} snake bites were reported in Italy from 1989 to 1995 [43,44]. The symptoms described, and their timing did not significantly differ from our observations. The genome of the \textit{Vaf} captured in Piemont (Italy) contains vaspin genes, but no Atx genes. \textit{Vaatra} is present in Italy, at the border with France. This species is also present in Switzerland, but no case of neurotoxic envenomation associated with \textit{Vaatra} snake bite has ever been reported by the Swiss National Poison Center of Zurich (personal communication). These observations are consistent with our genomic analysis indicating that the \textit{Vaatra} genome contains no genes encoding neurotoxins. We suggest that the French neurotoxic snake population has resulted from hybridization between \textit{Vipera aspis} and \textit{Vipera ammodytes}. The following observations support this view: 1) natural hybrids between \textit{Vaf} and \textit{Vam rufoi} can be found in the north-east of Italy, where these two species cohabit, 2) a phyloge-

Table 3. Evaluation of neurotoxicity of venoms collected in France by measuring cross-reactivity to AtxA antibodies.

| Label number | Administrative department | Viper identification | Cross-reaction with AtxA |
|--------------|--------------------------|----------------------|-------------------------|
| 402, 403, 404| Alpes-de-Haute-Provence (04) | Vaa                  | 0                       |
| 405          | Alpes-de-Haute-Provence (04) | Va ?                 | 0                       |
| 651          | Haute-Savoie (74)          | Vaatra               | 0                       |
| 712, 713,714,715,716 | Haute-Savoie (74) | Vaa                  | 0                       |
| 564, 602     | Hérault (34)              | Vaa «garrigue»        | 0                       |
| 301, 303, 304, 312 | Loire-Atlantique (44) | Vaa                  | 0                       |
| 452, 453, 454, 455, 501, 502, 503, 551, 601, 561 | Puy-de-Dôme (63) | Vaa                  | 0                       |
| 778, 779, 783, 784 | Seine-et-Marne (77) | Vaa                  | 0                       |
| 620          | Vaucluse (84)             | Vu                   | 0                       |
| 800          | Alpes-Maritimes (06)      | Vaa                  | 1                       |
| 508, 519     | Hautes-Pyrénées (65)      | Vaz                  | 1                       |
| 552, 559     | Puy-de-Dôme (63)          | Vaa                  | 1                       |
| 510, 512, 521, 553, 554 | Puy-de-Dôme (63) | Vaa ?                | 1                       |
| 513          | Alpes-de-Haute-Provence (04) | Vaa ?                | 2                       |
| 613          | Gironde (33)              | Vaz                  | 2                       |
| 516, 528     | Hautes-Pyrénées (65)      | Vaz                  | 2                       |
| 451          | Puy-de-Dôme (63)          | Va?                  | 2                       |
| 456, 560     | Puy-de-Dôme (63)          | Vaa                  | 2                       |
| 527          | Hautes-Pyrénées (65)      | Vaz                  | 3                       |
| 401, 406, 458| Alpes-de-Haute-Provence (04) | Vaa                  | 4                       |
| 514          | Alpes-de-Haute-Provence (04) | Va?                  | 4                       |
| 603          | Alpes-Maritimes (06)      | Vaa                  | 4                       |
| 565          | Gironde (33)              | Vaz                  | 4                       |
| 459          | Haute-Garonne (31)        | Vaz                  | 4                       |
| 504, 505, 506, 509 | Hérault (34) | Vaa ?                | 4                       |

Cross-reactions to AtxA antibodies were tested by ELISA as described in the Material and Methods section. Optical densities measured at dilutions of anti-AtxA IgG of 1/16,000 were used for cross-reactivity gradation: level 0: OD≤0.1, level 1: 0.1<OD≤0.2, level 2: 0.2<OD≤0.3, level 3: 0.3<OD≤0.4, level 4: OD>0.4.

Figure 12. Heatmap obtained with clusters with p-values<0.05. Each column in the heatmap corresponds to a spectrum, each row to a cluster (or a potential biomarker), and each cell to a peak, with the colour indicating intensity of expression (green: low expression, red: high expression). Related spectra are sorted and grouped according to similarity, resulting in the dendrogram. The length of a branch is inversely proportional to the relatedness of the spectra. Average normalised mass (m/z) of the 4 clusters with p-values<0.05 are indicated on the right of the heatmap.

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Table 3. Evaluation of neurotoxicity of venoms collected in France by measuring cross-reactivity to AtxA antibodies.
Figure 13. Snake venom characteristics and type of grade 2 and 3 envenomations observed in Puy-de-Dôme (code no. 63). Envenomation cases are presented in circles and snake venoms in rectangles. Classical venoms or envenomations cases are in white and neurotoxic venoms and envenomations are reported in black. 
doi:10.1371/journal.pone.0001194.g013

Figure 14. Snake venom characteristics and type of grade 2 and 3 envenomations observed in Alpes-de-Haute-Provence (code no. 04) and Alpes-Maritimes (code no. 06). Envenomation cases are presented in circles and snake venoms in rectangles. Classical venoms or envenomations cases are in white and neurotoxic venoms and envenomations are reported in black. The white line indicates the border between the two departs. 
doi:10.1371/journal.pone.0001194.g014
Figure 15. Hybrid SDS-PAGE venom profiles from snakes captured in Hérault (code no. 34) and Haute-Garonne (code no. 31). Venoms 504, 505, and 509 from Hérault, 459 from Haute-Garonne, Vaa-type venoms 508, 516, 519, 527, 528 from Hautes-Pyrénées and Vaa-type venom 303 from Loire-Atlantique were separated by a 20% SDS-PAGE under denaturing and reducing conditions. Molecular weight markers are shown on the left. Arrows indicate protein bands specifically found in Vaa venom.

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netic analysis of *V. aspis* snakes has shown that “neurotoxic” Vaa collected in south-eastern France belong to a phylogenetic group separate from that containing the other French specimens of this subspecies [27]. According to epidemiological data, the neurotoxic snake population spans the territory of at least four departments. Given the low dispersion rate of these snakes, the hybridization event was probably not recent and may have occurred in refuge areas during the last period of glaciation. The Southern Alps (Italy) served as refuge areas for *V. aspis* and *V. ammodytes* snakes during several ice ages [45,46]. Moreover, analysis of the phylogeography of *V. aspis* has suggested the existence of refuge areas in south-eastern France [47]. This is consistent with the presence of Atx genes and Bov-b line retroposons in the genomes of Vaf captured in the south of Italy, and of Vammon from Bulgaria.

The origin of the neurotoxic population of Puy-de-Dôme appears different. All neurotoxic individuals are clustered in the south-west part of the department, near the tributaries of the Dordogne River, which flow into the Gironde. These snakes may have originated from the Vaz population of Gironde. They may have colonized northern areas from the ice refuges of the south-west of France, by moving along the rivers, which constitute thermal corridors. Their PLA2 profile is consistent with this possibility, because their genomes, like those of Vaz snakes, contain no AmI2 gene. Our analysis of PLA2 SELDI profiles suggests that the area of distribution of this potentially neurotoxic snake population to the east may be limited towards the west by the Massif Central Mountains. More studies must be performed on snake venoms from the east part of the department to define the range of this neurotoxic population.

Fry et al. recently reconsidered the very concept of “non-venomous” snakes after studying the transcriptomes of the venom gland of non-front-fanged snakes: these transcriptomes contain sequences encoding several toxin types [48,49]. Similarly, our study shows that we have to reconsider *Vipera aspis aspis* as a “cryptoneurotoxic” species. Its genome contains an arsenal of PLA2 neurotoxins ready to be expressed under stimuli that still need to be identified. Our work demonstrates that a multidisciplinary approach can fully characterize the diversity in snake venom composition and identify its medical and public health consequences. Thus, by combining epidemiological data concerning snake bites with genomic, transcriptomic, proteomic and immunological analyses of the major toxic components of venoms, we were able to provide a clear description of the potential neurotoxicity of *Vipera aspis* bites in France. These observations as a whole illustrate the biodiversity of venoms within the genus *Vipera*. The geographical variations in the composition of *Vipera aspis* venom have major consequences for the preparation of neutralizing antivenoms [50,51]. Indeed, it strengthens the arguments in support of the use of polyvalent antivenoms able to neutralize the venom of *Vipera aspis*, *Vipera berus* and *Vipera ammodytes* in the treatment of viper envenomations in France.

**SUPPORTING INFORMATION**

Table S1

Found at: doi:10.1371/journal.pone.0001194.s001 (0.17 MB DOC)

Table S2

Found at: doi:10.1371/journal.pone.0001194.s002 (0.24 MB DOC)

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**Author Contributions**

Conceived and designed the experiments: VC. Performed the experiments: EF VJ IG Jd. Analyzed the data: EF Ld VJ SJ Jd. Contributed reagents/ materials/analysis tools: Ld SJ AT. Wrote the paper: VC. Other: Wrote the first draft of the paper: VC.
