**DIRS1**-like retrotransposons are widely distributed among Decapoda and are particularly present in hydrothermal vent organisms

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**Abstract**

**Background:** Transposable elements are major constituents of eukaryote genomes and have a great impact on genome structure and stability. Considering their mutational abilities, TEs can contribute to the genetic diversity and evolution of organisms. Knowledge of their distribution among several genomes is an essential condition to study their dynamics and to better understand their role in species evolution. **DIRS1**-like retrotransposons are a particular group of retrotransposons according to their mode of transposition that implies a tyrosine recombinase. To date, they have been described in a restricted number of species in comparison with the LTR retrotransposons. In this paper, we determine the distribution of **DIRS1**-like elements among 25 decapod species, 10 of them living in hydrothermal vents that correspond to particularly unstable environments.

**Results:** Using PCR approaches, we have identified 15 new **DIRS1**-like families in 15 diverse decapod species (shrimps, lobsters, crabs and galatheid crabs). Hydrothermal organisms show a particularly great diversity of **DIRS1**-like elements with 5 families characterized among Alvinocarididae shrimps and 3 in the galatheid crab *Munidopsis recta*. Phylogenetic analyses show that these elements are divergent toward the **DIRS1**-like families previously described in other crustaceans and arthropods and form a new clade called AIDIRS1. At larger scale, the distribution of **DIRS1**-like retrotransposons appears more or less patchy depending on the taxa considered. Indeed, a scattered distribution can be observed in the infraorder Brachyura whereas all the species tested in infraorders Caridea and Astacidea harbor some **DIRS1**-like elements.

**Conclusion:** Our results lead to nearly double both the number of **DIRS1**-like elements described to date, and the number of species known to harbor these ones. In this study, we provide the first degenerate primers designed to look specifically for **DIRS1**-like retrotransposons. They allowed for revealing for the first time a widespread distribution of these elements among a large phylum, here the order Decapoda. They also suggest some peculiar features of these retrotransposons in hydrothermal organisms where a great diversity of elements is already observed. Finally, this paper constitutes the first essential step which allows for considering further studies based on the dynamics of the **DIRS1**-like retrotransposons among several genomes.

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Background
Deep-sea vents are chemosynthetic environments which are considered as extreme as compared to usual life standards due to their physico-chemical characteristics. For instance, they show high levels of temperature, pressure, heavy metals and sulfide [1]. They are also highly variable, with physico-chemical shifts occurring over very short spatial and temporal scales [2]. Although other marine environments (e.g., the intertidal zone) are also variable, hydrothermal environments are particularly unstable due to (i) the intensity of variations observed in deep-sea vents, e.g., variation of temperature from 400 to 2°C over few centimeters [3], (ii) local random mix of vent fluids with surrounding waters [4] and (iii) the relatively short lifetime of vents [5]. Because such unstable environment may be difficult to live in, deep-sea vents are often considered as harsh and stressful. For example, the vent shrimps Rimicaris exoculata, which usually live between 15°C and 30°C, endure sudden changes of thermal conditions due to the convection of fluids and can survive to exposure to very high temperature vent emissions [6,7].

In terms of biome characteristics, hydrothermal ecosystems are also peculiar, as they show a much higher density of individuals compared with surrounding abyssal plains. For example, populations of R. exoculata can reach up to 2500 individuals per square meter [8]. On the other hand, in comparison with coastal environments, hydrothermal vents are associated to a small number of species [9,10] (less than 600 species described to date in [11]). These features, along with the variable nature of the hydrothermal environment, suggest that deep-sea vent organisms may present particular genetic characteristics in relation to their peculiar adaptive abilities. Studying transposable elements (TEs) constitute an interesting way to investigate the genomic bases of such adaptive capacities. Several studies have shown that environmental variations can promote genome plasticity through transcriptional activation and mobilization of TEs (especially retrotransposons), often in response to specific stimuli such as biotic stress (e.g., pathogens) and abiotic environmental changes (e.g., temperature) [12-15]. For these reasons, TEs have a large impact on genome structure and stability, contribute in particular to variations in genome size [16-19], and are therefore considered as one of the major sources of genetic variability in eukaryotes [20-24]. The order Decapoda is a great model to investigate the genomic nature of the adaptation of hydrothermal organisms and its possible relationship with TEs. First, decapod crustaceans (shrimps, lobsters, crabs, ...) are found in various environments, with about 50 species described in hydrothermal vents [11]. Second, they exhibit great variation in genome size, from 1.07 pg (1050 Mb) in the crab Carcinus maenas to 40.89 pg (40000 Mb) in the shrimp Sclerocrangon ferox [25,26], with several species (e.g., shrimps) showing particularly large genomes that are thus liable to harbor high TE contents. Given their abundance and diversity, decapods have been greatly underrepresented in studies of TEs with only few elements described to date [27-35] including a LTR retrotransposon, CalEa1, that we have recently characterized in the hydrothermal galatheid squat lobster Munidopsis recta [36].

LTR retrotransposons (Ty1/copia-like, Ty3/gypsy-like and Pao/Bel-like elements) and tyrosine recombinase encoding elements (DIRS-like, Ngaro, Viper) constitute two major groups of retrotransposons [37-39], a class of TEs specific to eukaryotes [40]. Several phylogenetic analyses have shown that the pol region of DIRS-like elements harbor reverse transcriptase (RT) and RNase H (RH) domains closely related to those of Ty3/gypsy-like LTR retrotransposons [41,42]. However, these elements differ fundamentally in structure from LTR retrotransposons as they are devoid of LTR, do not encode either Proteinase and Integrase and harbor a Methyltransferase domain (Fig. 1) [39,41]. Moreover, DIRS-like retrotransposons replicate through a particular mechanism that involves a tyrosine recombinase [43]. Within the DIRS-like superfamily [38], two distinct groups have been described: DIRS1-like and PAT-like elements. These two groups differ by the nature of their termini (inverted terminal and split direct repeats, respectively [44]) and by phylogenetic relationships [45].

In contrast to LTR retrotransposons that have been found in a wide continuous range of species [39,46,47], DIRS1-like elements have a more patchy distribution being yet reported only in a restricted number of phylogenically diverse eukaryote organisms (Table 1) [41,43,46,48-56]. Although most of the DIRS1-like retrotransposon identifications were based on in silico approaches [41,43], they have not been yet described in several well-studied phylum (e.g., mammals or plants), and are absent from model organisms such as Saccharomyces cerevisiae and Drosophila melanogaster. Moreover, only few different DIRS1-like families have been found per species [41] in contrast to the large variety of LTR retrotransposon families [38,57-59]. In the slime mold Dictyostelium discoideum, DIRS1-like retrotransposons have been shown to harbor an active heat-shock promoter in their inverted terminal repeats [60,61]. Considering the high level of thermal instability in vents, the presence of such promoters could favor TE activation in hydrothermal species.

Hence, studying DIRS1-like elements in Decapoda could allow both to better understand the role of TEs in the adaptation of species in hydrothermal ecosystems and to investigate the dynamics of these peculiar elements. We focused on ten hydrothermal decapods that represent the three major infraorders observed (6 caridean shrimps, 3 brachyuran crabs and 1 anomuran galatheid crab). We
have detected DIRS1-like retrotransposons in all vent shrimps and in M. recta as well as in 8 other non-vent decapods. To our knowledge, this paper constitutes the first study based on DIRS1-like element distribution both in a continuous range of species within a single taxon and in species living in various ecosystems. Fifteen new DIRS1-like families are at all described among 15 out of the 25 species tested. These unusual elements are thus surprisingly more widely distributed among decapods than expected.

Results
Fishing out DIRS1-like pol sequences in hydrothermal shrimps
In the course of looking for Ty3/gypsy-like LTR retrotransposons in three hydrothermal shrimps (Rimicaris exoculata, Chorocaris chacei and Mirocaris fortunata) using degenerate primers GD1 and GD2 corresponding to the “RMPFGL” and “LITTDAS” conserved motifs of the RT domain (Fig. 1), we PCR-amplified additional fragments in supplement of those that correspond to Ty3/gypsy-like elements (data not shown). Cloning and sequencing of these additional fragments revealed that they share high sequence identity with the pol region of DIRS1-like elements at the protein level. Detailed sequence analysis showed that the fragments observed originated from either GD1/GD1 amplifications (sequences RexAlvi1a from R. exoculata and CchAlvi1a from C. chacei, [GenBank:FJ707538-FJ707539]) or from GD1/GD2 amplification with annealing of degenerate primer GD2 at a non targeted site (sequence MfoAlvi3a from M. fortunata, [GenBank:FJ707541]). Fishing out of DIRS1-like elements using primers designed for Ty3/gypsy-like retrotransposons can be explained by the close phylogenic relationships between RT domains of Ty3/gypsy-like and DIRS1-like elements [43]. Although the regions in which the primers were designed are only partly conserved between Ty3/gypsy-like and DIRS1-like elements, partial conservation may be sufficient for them to anneal on DIRS1-like sequences, particularly because primers are degenerate. To further confirm the presence of DIRS1-like elements in these species, we designed a pair of degenerate primers, DD1 and DD2, to amplify the region comprised between the “LGFLIN” and “DLFAS” motifs of the DIRS1-like elements pol region (Fig. 1). We tested these primers in R. exoculata. Unfortunately, they did not allow for amplifying a single band, but rather led to the amplification of numerous fragments (smear). After long and tedious optimization of the PCR conditions, we managed to obtain a major amplicon of 870 bp, which we then cloned and sequenced. Sequence analysis revealed that it corresponds actually to a DD2/DD2 amplification product, which shares similarity with pol region of DIRS1-like elements only in the last 532 bp (RexAlvi2a, [GenBank:FJ707540]). Due to the technical difficulties to obtain specific amplicons, we did not use this primer pair for other analyses. To get insights on the genetic diversity of the DIRS1-like elements found, we aligned the partial sequences of the pol region that we obtained. Alignment of the 4 sequences allowed us to define 3 families. RexAlvi1a and CchAlvi1a sharing 89% nucleotide sequence identity, they were clustered into a single family that we called Alvi1 (see Additional file 1). On the other hand, RexAlvi2a and MfoAlvi3a shared too low sequence identity to either Alvi1 (< 53% and < 50% sequence identity, respectively) or to each other (71% sequence identity) to be clustered and were therefore considered as members of different families that we called Alvi2 and Alvi3.

Screening for Alvi elements in other decapods
To date, DIRS1-like retrotransposons have been identified and characterized in only a restricted number of species (Table 1). To better understand the evolutionary dynam-
ics of DIRS1-like elements, we investigated the diversity and the distribution of the Alvi families among several Decapoda crustacean genomes. We first analyzed the distribution of our 3 novel DIRS1-like families among several decapods, from both hydrothermal and non-hydrothermal ecosystems. For this, we screened for Alvi families among the 25 decapods species using different combinations of primers (Table 2) including specific primers of each element and Retro1+, a primer shared by all 3 elements (see Methods and Additional file 2). Although pairs of specific primers would allow for amplification of closely related elements, we expected that the association of Retro1+ with a single specific primer would be particularly useful to extract more divergent sequences suggesting some specificity troubles especially due to the use of a single specific primer.

Results, presented in Table 2, show that Alvi elements are only detected in hydrothermal shrimps. Two families (Alvi1 and Alvi2) are widely shared among these species while the two others (Alvi3 and Alvi4) are restricted to the species where they were firstly identified (M. fortunata and A. stactophila, respectively). Alvi2 elements could not be detected in M. fortunata. Using Alvi2 specific primers on this species we were only able to identify non-targeted elements of the Alvi3 family, because nucleotidic differences with the other 4 families were too spread out on the sequences. Hence, we finally analyzed the distribution of DIRS1-like elements in decapods using 4 families: Alvi1-4.

For all other decapods, no Alvi element could be detected even in other hydrothermal organisms or in the other closely related caridean shrimps [62] (Crangon crangon and Palaemon serratus). In the hydrothermal galatheid crab Munidopsis recta, although no PCR amplification fragment of the expected size was obtained, we PCR-amplified an additional fragment (545 bp) when using Retro1+. Sequencing of this fragment revealed that it originated from the Retro1+/Retro1+ amplification of a new DIRS1-like element, belonging to the retrotransposons previously reported even in partial sequences.

| Species             | Phylum       | Name         | Sequences       | Reference          |
|---------------------|--------------|--------------|-----------------|--------------------|
| Apis mellifera      | Insecta      | AmDIRS1      | [GenBank: AADG 02016821] | [48]              |
| Arbracia punctulata | Echinidea    | ApDIRS1      | [GenBank: BK004821] | [48]              |
| Cotesia congregata  | Brachyura    | CcBv31.6-31.7| [GenBank: A632329] | [51]              |
| Daphnia pulex       | Brachyura    | DpDIRS1-2    | [GenBank: ACB38665, ACB38666] | Unpublished |
| Dario rena          | Teleostei    | DrDIRS1-3    | [GenBank:AL590134, BK001258, BK001259] | [43,49] |
| Dictyosteleus discoideum | Ectoprocta | DIRS1        | [GenBank: M131319] | [52]              |
| Gopherus agassizii  | Sauropoda    | GaDIRS1      | [GenBank: AC147866] | [41]              |
| Lytechinus variegatus | Echinidea | LvDIRS1      | [GenBank: AC131505] | [49]              |
| Nasonia vitripennis | Insecta      | NvDIRS1      | [GenBank: NW_001820604] | Unpublished |
| Nemastomatina      | Cnidaria     | NvDIRS1      | [GenBank: X01627585]; Repbase | [53] |
| Okolophora dioica  | Urochordata  | OdDIRS1      | [GenBank: Y634219] | [54]              |
| Oncorhynchus mykiss | Teleostei    | OmDIRS1      | [GenBank: DQ156151] | Unpublished |
| Phycyces blakesleeanus | Zenophora | Ptl          | [GenBank: Z54337] | [55]              |
| Rhizopus oryza     | Zygomycota   | RoDIRS1      | Retrobase        | [49]              |
| Salmo salar        | Teleostei    | SsDIRS1      | [GenBank: DQ156149, DQ156150] | Unpublished |
| Strongylocentrus purpuratus | Echinidea | SpDIRS1-4    | [GenBank: BK005158]; Retrobase | [43] |
| Takifugu rubripes  | Teleostei    | FrDIRS1      | Repbase          | [56]              |
| Tetraodont nigriviridis | Teleostei | TnDIRS1      | [GenBank: AF442732] | [43] |
| Tribolium castaneum | Insecta      | TrDIRS1      | [GenBank: AY331876] | [48] |
| Xenopus laevis      | Amphibia     | XIDIRS1      | [GenBank: BGSS15156] | [43] |
| Xenopus tropicalis  | Amphibia     | XtDIRS1-2    | [GenBank: AC144974, AC145807] | [43] |

Notes: -1 Accession numbers deposited in DDBJ/EMBL/GenBank databases, in Repbase database [46] or in Retrobase database [50].
like sequence (Ymur1a, [GenBank:FJ707610]). It shared low sequence identity to Alvi1-5 families (< 50% sequence identity) and was therefore considered as member of a novel family that we called Ymur1.

Diversity and phylogenetic relationships of Alvi DIRS1-like retrotransposons

Screening for Alvi elements among decapods led us to obtain 36 partial sequences of the pol region (see Additional file 3). To gain power in our investigation of the diversity of these elements, we completed these data with additional sequences. Two complementary approaches were performed: extension of the sequences by PCR walking and PCR amplification of larger internal sequences using specific primers (see Methods and Additional file 4). Seventeen new sequences were thus obtained (see Additional file 3), enabling to well-cover the pol region of DIRS1-like elements at least from the "YLDD" RT motif to the end of the RH domain (Fig. 1).

All DNA sequences obtained were aligned using ClustalX program [63] and corresponding alignment was used to analyze the relationships between families. First, we checked the boundaries of the different families through the construction of a sequence identity matrix using the pairwise gap deletion method (see Additional file 1). A group of sequences was considered as a family if its higher intra-group divergence was lower than its inter-groups divergence, without overlap of the two distributions. The clustering obtained from sequence identity matrix validates the five families that we had previously defined (Alvi1 to Alvi5, Fig. 2). Among them, the Alvi1 copies are more diverse (from 63 to 95% sequence identity, mean: 76% sequence identity) than other families ones. The Alvi1 family is also more divergent toward the four other families (53% divergence in average between the Alvi1 copies and the others) whereas Alvi2-5 families appear to be more closely related as copies from these families differ one by one by only 35% of divergence in average.

Table 2: Alvi elements detected in decapods.

| Specimens and phylum | Pol region |  |  |  |  |  |  |  |  |  |  |  |
|----------------------|------------|---|---|---|---|---|---|---|---|---|---|---|
| Rimicaris exoculata  | hydrothermal shrimps | RexAlvi1 | RexAlvi1 | RexAlvi2 | RexAlvi2 | na | na | na | RexAlvi1 |
| Chorocaris chacei     | hydrothermal shrimps | CchAlvi1 | CchAlvi1 | CchAlvi2 | A | na | CchAlvi2 | na | A |
| Mirocaris fortunata   | hydrothermal shrimps | na | MfoAlvi1 | MfoAlvi3 | A | MfoAlvi3 | MfoAlvi3 | na | MfoAlvi3 |
| Alvinocaris markensis | hydrothermal shrimps | AmaAlvi1 | AmaAlvi1 | AmaAlvi2 | A | na | AmaAlvi5 | na | A |
| Alvinocaris municola  | hydrothermal shrimps | A | AmuAlvi1 | AmuAlvi2 | A | na | na | na | A |
| Alvinocaris stactophila | hydrothermal shrimps | A | AstAlvi1 | AstAlvi2 | A | na | AstAlvi4 | AstAlvi4 | AstAlvi4 |
| Segonzacia mesatlantica | hydrothermal crabs | na | na | na | na | na | na | na | na |
| Bythograea thermydron | hydrothermal crabs | na | na | na | na | na | na | na | na |
| Cyaneaglaea praedator | hydrothermal crabs | na | na | na | na | na | na | na | na |
| Munidopsis recta     | hydrothermal galatheid crabs | na | na | na | na | na | na | na | na |
| Palaemon serratus    | intertidal shrimps    | na | na | na | na | na | na | na | na |
| Crangon crangon      | intertidal shrimps    | na | na | na | na | na | na | na | na |
| Other decapods tested* | intertidal or seamount | na | na | na | - | na | na | - | na |

In black bold: Amplicon sequences corresponding to the element targeted. Underlined: PCR amplification products in species in which the element targeted was firstly identified. In italics: Amplicon sequences corresponding to another element than the one targeted. A: PCR amplification product of the expected size observed but not sequenced. na: No PCR amplification of expected size. -: Not tested. * see Methods for more details.

$ single specific primer associated to the Retro1+ primer.

In italics: 1 specific primer $ associated to the Retro1+ primer.

Single primer $ associated to the Retro1+ primer.
Second, we analyzed the relationships of the sequences among and between the 5 families using a Neighbor Joining phylogenetic analysis with pairwise gap deletion. The tree obtained (Fig. 3) confirmed that each Alvi family is well-supported with bootstrap values being greater than 94%. It also confirmed that the Alvi1 family highly diverges to the other 4 families, with strong bootstrap support (100%) of the separation into an Alvi2-5 clade, and an Alvi1 clade. Analysis of the Alvi1 subtree also revealed a lack of structuration as compared to the other families, this confirming the large sequence diversity observed with the matrix analysis. On the contrary, although relationships between families within the Alvi2-5 group cannot be resolved by our tree, relationships of sequences within each family are well resolved considering most of bootstrap values being greater than 70%. It is especially the case for the Alvi2 family which sequences were obtained from various hydrothermal shrimps. Phylogenetic relationships among this family also fit with those of the species [62] in which they were identified, with only few exceptions like the AmuAlvi2a sequence from A. muricola that clusters with Alvi2 sequences from A. markensis.

**Screening for DIRS1-like elements in decapods**

Capacity of detection of TEs using a PCR approach depends of the choice of the primers, as shown by the identification of some elements using specific/aspecific primer pairs and not using specific/specic pairs. Hence, absence of PCR signal could be due to the divergence of the elements rather than their absence from the species analyzed. To detect such more distant DIRS1-like retrotransposons, we designed degenerate primers using two types of primers: "highly" degenerate and "softly" degenerate (see Methods). Although both of these approaches have flows (low copy number elements could be missed using the "highly" degenerate primers and highly divergent elements could be missed with the "softly" ones), they are complementary and should, when used in parallel, raise the detection sensitivity. Six different combinations of primers were tested (Table 3), five using "highly" degenerate primers, and one using two "softly" ones (see Methods).

As expected from our previous PCR results, amplification fragments were obtained in hydrothermal shrimps whatever the degenerate primer combination used (one fragment, AmaAlvi2of, was sequenced as control in A. markensis, [GenBank:FJ707542]). We have shown that M. recta harbors DIRS1-like retrotransposons (Ymur1 element). Using "highly" degenerate primers, clear amplification fragments of the expected size were only obtained with the GD1/rcDD4 combination. Cloning and sequenc-
ing allow to obtain three additional related sequences (Ymur2a-c, [GenBank:FJ707543-FJ707545], > 75% sequence identity). Two others related sequences (Ymur3a-b, [GenBank:FJ707546–FJ707547], 93% sequence identity) were also obtained using the "softly" degenerate primer combination, one of them (Ymur3b) corresponding to an unexpected additional fragment about 1100 bp (GD1soft/GD1soft amplification). Sequence analysis revealed that the DIRS1-like sequences obtained in M. recta can be clustered into 3 distinct families (Ymur1-3, see Additional file 1) and phylogenetic analyses indicates that Ymur1-3 families positioned allowing to obtain three additional related sequences (Ymur2a-c, [GenBank:FJ707543-FJ707545], > 75% sequence identity). Two others related sequences (Ymur3a-b, [GenBank:FJ707546–FJ707547], 93% sequence identity) were also obtained using the "softly" degenerate primer combination, one of them (Ymur3b) corresponding to an unexpected additional fragment about 1100 bp (GD1soft/GD1soft amplification). Sequence analysis revealed that the DIRS1-like sequences obtained in M. recta can be clustered into 3 distinct families (Ymur1-3, see Additional file 1) and phylogenetic analyses indicates that Ymur1-3 families positioned themselves as a single well supported clade (bootstrap value of 100%) (Fig. 3).

In other decapods, the two methods gave different results (Table 3): while using "highly" degenerate primers combinations no amplification of the expected size was obtained, the use of GD1soft/rcDD4soft led to either trace amplification or amplification fragments of the expected size. Cloning and sequencing of all single bands observed allowed us to discover new DIRS1-like elements ([GenBank:FJ707548–FJ707560]) in 8 new species (Fig. 4 and Additional file 3): the two coastal shrimps (Crangon crangon and Palaemon serratus), the two coastal lobsters (Nephrops norvegicus and Homarus gammarus), 2 seamount galatheid squat lobsters (Agononida laurentae and Eumunida annulosa) and 2 coastal crabs (Xantho pilipes and Maja squinado). The sequences obtained in a given species, except X. pilipes, cluster into a single family (> 93% sequence identity, see Additional file 1). In the crab X. pilipes, the two sequences obtained define two distinct families (Ypili1a and Ypili2a, 52% sequence identity). Moreover, elements of the galatheid crabs E. annulosa and A. laurentae (Yannu1a-b and Yago1a, respectively) can be clustered into a single family we called Yannu1 (> 90% sequence identity). Likewise, elements of the two lobsters N. norvegicus and H. gammarus (Ynmo1a-b and Yhoga1a,
respectively) clustered into the same Ynno1 family (> 82% sequence identity). We also detect in the EST databases two DIRS1-like sequences ([GenBank:CN853875.1, FC556055.1]) of the lobster Homarus americanus, that share high similarities with Ynno1 family (80% sequence identity in average). Altogether, this study led us to discover 7 new DIRS1-like families (see Additional file 3). All of these families diverge to Alvi1-5 and Ymur1-3 families with < 56% sequence identity (see Additional file 1).

To study their distribution, two specific primers were designed for each newly discovered DIRS1-like family (sequences available upon request). They were used to screen all fifteen species in which we had observed the presence of DIRS1-like elements (Fig. 4) as well as closely related species (same Genus). For each family, no PCR signal was obtained except in species in which the families were firstly identified. This underlines that all elements previously obtained do not result from potential cross-contaminations. Nevertheless, DIRS1-like retrotransposons are widely distributed among decapod crustaceans, although this distribution appears to be heterogeneous according to the phyllum considered (Fig. 4). DIRS1-like elements are yet identified in all caridean shrimps and lobsters tested. In anomourans the distribution appears very patchy as DIRS1-like retrotransposons are characterized in 3 out of the 4 galatheid squats lobsters but not in porcelain crabs or hermit crab. Finally, the presence of these elements has been only detected in 2 out of the 8 brachyuran crabs tested with especially no detection in hydrothermal organisms.

Discussion

Methodological issues

DIRS1-like retrotransposons have yet been reported in a restricted number of eukaryote species (Table 1), in contrast to LTR retrotransposons that show a wide and almost continuous distribution. However, we found DIRS1-like elements in 15 out of the 25 decapod species studied, with 15 novel DIRS1-like families identified (see Additional file 3) and a particularly large diversity observed among hydrothermal shrimps (Fig. 4). To characterize transposable elements among different genomes, two approaches are usually performed: in silico approaches [57,58] and/or PCR-based procedures [27,28,64]. Most of previous studies aiming at identifying DIRS1-like retrotransposons were based on in silico approaches [41,43]. However, in the case of decapods, genomic sequences are scarce, with sequencing projects concerning only the creation of EST databases (e.g., Petrolisthes cinctipes, Joint Genome Institute). In particular, to our knowledge no large-scale genome sequencing project is ongoing. We therefore conducted several PCR-based approaches that enable to screen for partial DIRS1-like pol region simultaneously in a large number of species representing numerous decapod taxa (shrimps, crabs, galatheid crabs, porcelain crabs, hermit crabs and lobsters). Among the 73 sequences that we identified, 5 were obtained from PCR-amplified fragments which size differed from the one expected (see Additional file 3). Analyses of such additional fragments, which can be obtained for different species or using different primer combinations, provide an interesting way to identify new TEs. One issue with DIRS-like retrotransposons is that they are sometimes abusively grouped with Ty3/gypsy-like elements because their RT/RH domains are closely related. Distinguishing them from each other is however primordial, particularly if DIRS-like elements are in fact more frequent that previously supposed. For that purpose, it is important to check the partial pol sequence of each element for the presence of the MT domain (specific to DIRS-like elements) (Fig. 1) before assigning them to DIRS-like superfamily.

To classify our sequences into families, we used a nucleotide sequence identity matrix built using pairwise deletion method (see Additional file 1). This method presents the advantage to compute a distance for each pair of sequences, ignoring only the gaps that are involved in the comparison and therefore including the largest amount of information for the whole dataset. This method appears as the most appropriate to analyze our sequences, as our dataset is constituted of several PCR-based sequences that only partially overlap, and because many of our sequences harbor small internal indels. Finally, this approach appears efficient as it enables to clearly define the different families that we found (Fig. 2). Recently, in an attempt to find criteria for sequence-based TE classification for the annotation of TEs in large scale genome sequencing projects, the "80-80-80" rule has been proposed [38]. This rule states that two TEs belong to the same family if they share 80% (or more) sequence identity over segments longer than 80 bp representing at least 80% of the coding domains, internal domains, or terminal repeat regions. However, as noticed by the authors, this rule is more efficient on terminal repeat regions and non-coding regions than on coding regions like pol. This property was confirmed with our dataset, thus leading to the conclusion that, although certainly efficient for large-scale genome sequence analysis of TEs, this rule is not adapted to our PCR-based analysis.

Distribution of Alvi families within Alvinocarididae genomes

Screenings for DIRS1-like elements in decapod genomes indicates that the "highly" degenerate primers have a lower detection sensitivity than the "softly" degenerate ones. Oligonucleotide sequences of "softly" degenerate primers correspond to a subset of those present in the "highly" degenerate ones (See Additional file 5). Nevertheless, the combination of "softly" degenerate primers
allowed for identifying DIRS1-like elements in at least 15 diverse decapods whereas combinations of "highly" degenerate ones detect elements in only the 6 Alvinocarididae and M. recta (Table 3). Despite their weaker detection sensitivity, "highly" degenerate primers led to PCR amplifications in the six hydrothermal shrimps and Ymur1-3 elements were identified in Munidopsis recta. Support for individual clades was evaluated using non-parametric bootstrapping obtained from 1000 bootstrap replicates. Only bootstrap nodes value over 70% are indicated. Distances were calculated with Tamura 3 parameter model plus gamma distribution’s correction for nucleotides.

In complement to degenerate primers, PCR screening using specific primers is a powerful approach to study the distribution of previously characterized elements, as well as for identifying novel members of closely related families. Indeed, screening for Alvi1-3 families at a small phylogenetic scale (i.e. within the family Alvinocarididae) led to the detection of the Alvi4 and Alvi5 families. Since all hydrothermal shrimps harbor Alvi elements, these ones likely result from the activity of DIRS1-like retrotransposons that were already present in the genome of the common ancestor of all Alvinocarididae species. Alvi elements may thus belong to relatively ancient lineages of DIRS1-like retrotransposons. They could be as old as several million years, since molecular estimates of divergence show that the vent-endemic caridean shrimp species radiated 6.7 to 11.7 million years ago [62]. This is especially true for the Alvi1 and Alvi2 families that are shared among Alvinocarididae.

Our phylogenetic analysis (Fig. 3) also shows that Alvi2 to Alvi5 families cluster into a single clade, suggesting that these families diverged from a common ancestor. The Alvi3 and Alvi4 families are each restricted to a single species and may therefore correspond to lineage-specific evolution. In A. stactophila, the presence of both Alvi2 and Alvi4 families suggests that this last could result from the differentiation of a subset of the ancestral family. In M. fortunata, which is the most distant species among vent shrimps [62], we observe the presence of Alvi3 instead of Alvi2 (Fig. 4). Two alternative evolutionary scenarios could explain this observation. If elements of the Alvi3 family were present in the Alvinocarididae ancestor, the distribution observed would result from different lineage-specific burst-like events of different Alvi families, as observed for LTR retrotransposons in rice-related species [65], leading to the amplification of Alvi3 in M. fortunata and this of Alvi2 in the common ancestor of other related vent shrimps. On the other hand, if the Alvi3 family was absent from the Alvinocarididae ancestor, it likely results from a lineage-specific evolution of the ancestral Alvi2 family in the M. fortunata lineage. In addition, R. exoculata and C. chacei, which are phylogenetically very close [62], harbor the same distribution pattern for DIRS1-like families (Fig. 4) with close corresponding sequences (e.g. average of 89% sequence identity within Alvi1 family). All these observations suggest that the wide distribution of DIRS1-like retrotransposons among Alvinocarididae is closely related to the phylogeny of species and that the emergence of new elements is linked to the independent evolutionary history of Alvi families within each host genomes. Although we cannot privilege one evolutionary scenario, all possible scenarios are in accordance with a simple vertical transmission of the different Alvi elements.

**ADIRS1: A new clade of DIRS1-like retrotransposons**

Previous analyses of phylogenetic relationships among the DIRS1-like retrotransposons [41,48] highlighted two...
clades: the DrDIRS1 clade that groups the DIRS1-like elements identified in teleost fishes, and the TcDIRS1 clade that pools together elements from insects and sea urchins. To determine the relatedness of our elements with other DIRS1-like retrotransposons, we performed a new phylogenetic analysis using the Maximum Likelihood method. The core dataset included Alvi1-5 and Ymur3 families, the only families for which we got sufficiently long sequences, as well as the recently identified elements of another crustacean species, Daphnia pulex, and those of the cnidarian Nematostella vectensis, which were never reported in phyl-ogenetic analyses to date. The resulting tree (Fig. 5) is in accordance to those previously described. The two DpDIRS1-2 elements from D. pulex cluster with those of other arthropods (insects) in the TcDIRS1 clade. In contrast, DIRS1-like elements of decapods are phylogenetically divergent from these, with the Alvi1-5 and Ymur3 families forming a distinct clade among the DIRS1-like superfamily that we called AlDIRS1-clade.

These results indicate the presence of amino acid signatures for decapod elements, in particular a Tryptophane (W) instead of a Phenylalanine (F) amino acid in the RT domain at position 44 on the alignment (see Additional file 6). Interestingly, this signature is also shared by the sequences that we obtained in other decapods. Therefore, we propose that the decapod DIRS1-like retrotransposons may belong to the AlDIRS1 clade, although further studies on the phylogenetic relationships of these elements are needed to confirm this. For that purpose, two conditions should be fulfilled: to obtain additional sequences to build better amino acids consensus sequences, and to enlarge the sequences obtained for the pol region to improve the number of informative sites.

**Wide distribution of DIRS1-like elements among decapods**

Our analysis of the distribution of the DIRS1-like retrotransposons (Fig. 4), the first as such large scale (within an order) gives clues to better understand the peculiar scattered distribution of these elements among eukaryote genomes. Three species (M. recta, A. stactophila and A. markensis) harbor at least 3 different DIRS1-like families. Such results are similar to those observed in other well-studied species such as the sea urchin Stronglylocentrotus purpuratus (4 families, [43]) and the zebrafish Danio rerio (3 families, [49]). In particular, we show that DIRS1-like elements can be much more than just a feature of isolate species. When focusing on a given taxa (order Decapoda)
Species distribution of *DIRS1*-like retrotransposons (Fig. 4) appears more or less patchy, depending on the infraorder considered (as defined in [66]). Three different distribution patterns are observed: (i) presence of the elements in a large range of species (the 8 caridean and the 3 astacidean species). (ii) presence of the elements in a subpart of the infraorder (presence in 3 out of 4 galatheid crabs and absence of detection in hermit crab or porcelain crabs). (iii) presence of the elements in a restricted number of species (2 out of 8 crabs). So, Decapoda are the first example revealing that *DIRS1*-like element may be frequent, widely distributed and so probably ancient within an order.

Species distribution of *DIRS1*-like retrotransposons (Fig. 4) thus appears related to species phylogeny. Some families are widely distributed among closely related species (e.g., Alvi1-2 and Ynno1) whereas some others (e.g., Ypili1-2 and Yannu1) seem to be only scarcely shared, even among species that belong to the same Genus. Considering the environmental conditions of species, *DIRS1*-like retrotransposons are not restricted to hydrothermal species. However, their higher detection sensitivity in vent organisms (Alvinocarididae shrimps and the galatheid crab *M. recta*) suggests some peculiar features that could be related to a greatest element diversity and/or a higher number of copies. To better understand the distribution of *DIRS1*-like elements among Decapoda, the next step will be to extend our study to other infraorders (e.g. Achelata (spiny lobsters) or Dendrobranchiata (large prawn)), even to other crustaceans. Such an analysis will also allow for analyzing the vertical transmission of *DIRS1*-like elements suggested by our results. Moreover, we are currently estimating the number of copies of *DIRS1*-like retrotransposons. This will provide crucial information (i) to point out possible particularities of vent species in comparison with other decapods and (ii) to understand the dynamics of these elements within Decapoda. We also wonder whether such widespread distribution and diversity of *DIRS1*-like retrotransposons are peculiar features of Decapoda, or if similar features can be observed in other clades, for instance one known to harbor *DIRS1*-like elements (Table 1). If the widespread distribution of *DIRS1*-like retrotransposons that we observe within a taxon is a usual pattern, then the presence of one element in a given species would indicate that *DIRS1*-like retrotransposons are likely to be found in phylogenetically closely related species.

**Conclusion**

Because of the small number of *DIRS1*-like elements described to date, little is known about the distribution and the evolutionary dynamics of these retrotransposons, which differ to other TEs by their peculiar tyrosine recombinase -based replication model [41]. This paper presents the first study using PCR approaches to specifically search for *DIRS1*-like retrotransposons in several related species. Fifteen new *DIRS1*-like families were identified in 15 decapod species which nearly doubles both the number of *DIRS1*-like elements described to date, and the number of species known to harbor such elements. Our study reveals a larger diversity of *DIRS1*-like retrotransposons as we distinguish a third clade (ADIRS1-clade) in addition to the TcDIRS1 and DrDIRS1 clades previously described. Our results provide a new vision of the distribution of the *DIRS1*-like superfamily, and reveals that these elements are not as scattered as previously thought. It is the first study showing that these retrotransposons could have a widespread distribution within a large order (Decapoda) as well as a great diversity within a family (Alvinocarididae). These observations open the way to further studies on the evolutionary dynamics of these elements.

**Methods**

**Biological materials**

Twenty-five species of Decapoda were sampled in 3 major types of environment: deep-sea hydrothermal vents, the intertidal zone and seamounts. Vent animals were sampled during different cruises. Specimens from the South East Pacific Rise (the crabs *Bythogriaea thermophila* and...
Cyanagrea praedator, and the galatheid squat lobster Munidopsis recta) were collected during the French cruise "BIOSEPEEDO" (March–May 2004) using the French "Nautile" deep-submergence vehicle operating from the N.O. "L’Atalante". Samples from the LAU Back-Arc Basin (the caridean shrimps Alvinocaris muricola and Alvinocaris stactophila) were collected by the DSV JASON II during the American cruise TUI07MV (June 2005). At last, specimens from the Mid-Atlantic Ridge (the caridean shrimps Alvinocaris markensis, Mirocaris fortunata, Romicarid exculata, Chorocaris chacei, and the crab Segonzacia mesatlantica) were sampled on two vent fields (Lucky Strike or Rainbow), during the French cruise "MoMARETO" (August 2006) with the suction sampler of the ROV "Victor 6000" operating from the R/V "Pourquoi pas ?".

For coastal decapods, 12 species (the caridean shrimps Palaemon serratus, Crangon crangon, the brachyuran crabs Carcinus maenas, Maja squinado, Necora puber, Xantho incisus, Xantho pilipes, the lobsters Homarus gammarus, Nephrops norvegicus, the porcelain crabs Porcellana platycheles, Pisidia longicornis, and the hermit crab Dardanus calidus) were collected in French Brittany.

Finally, three specimens of galatheid squat lobsters from seamounts (Agononida laurentae, Eumunida annulosa, and Eumunida stenomaculata) were collected south of New Caledonia on Norfolk seamounts during the prospecting campaigns Norfolk 1 (2001, IRD Nouméa) and Norfolk 2 (2003, MUSORSTOM).

For all samples, living specimens were fixed immediately after collection, in liquid nitrogen for vent species or in 70% ethanol for the other species. They were then stored at -80°C or 4°C, respectively.

DNA from one individual per species was isolated using the CTAB method [67]. Dry DNA pellets were suspended in TE Buffer (10 mM Tris and 0.1 mM EDTA).

Detecting DIRS1-like retrotransposons using degenerate primers
To look for retrotransposon pol fragments, we performed PCRs using several degenerate primer pairs (Fig. 1). Two primers (GD1 and GD2) were designed to amplify, in three hydrothermal shrimp species (Chorocaris chacei, Mirocaris fortunata and Romicarid exculata), conserved motifs of Ty3/gypsy-like retrotransposons: 'RMPFGL' and 'LTTDAS', which are also shared with DIRS1-like elements (see Additional file 5). Four other primers (DD1, DD2, DD4, rcDD4 (reverse complement of DD4)) were designed to amplify conserved motifs specific to the pol region of DIRS1-like retrotransposons: 'LGFLIN', 'DLFAS' and 'ASKSGWG'. Another primer, DD6, was designed on the 'IRLHCD' motif that was conserved among the elements firstly identified in this study (Alv1-3). Detection sensitivity of DIRS1-like RT fragments was also improved by using, two "softly" degenerate primers related to GD1 (GD1soft) and rcDD4 (rcDD4soft). Degeneration level of these primers corresponds to 384 different sequences possible instead of the 1024 or 8192 sequences for GD1 and rcDD4, respectively. Finally, we also defined a non degenerate primer (Retro1+) conserved among DIRS1-like elements of Alvinocarididae and that spans the highly well-conserved YLDD motif that is present in both LTR retrotransposons and DIRS-like elements. All primer sequences are available in supplementary data (See Additional file 2). PCR amplifications of all primer combinations tested were performed as described in [36] and amplification products were separated on 1% agarose gels. Bands with the expected molecular weight were excised from the agarose gels, purified with SpinX column (Costar) and cloned in pGEM-T vector according to the manufacturer’s recommendations (Promega, Madison, WI, USA). One to three clones were sequenced using the ABI prism automated sequencer (Genome Express) and the partial pol nucleotide sequences have been submitted to the GenBank database ([GenBank:FJ707538–H707569]).

Extending the pol sequences
Sequences obtained with degenerate primers allowed to define several new DIRS1-like retrotransposons. In hydrothermal shrimps, a PCR walking approach (See Additional file 4) was then performed to extend pol sequences from each initial fragment, in order to study the diversity of these elements and to analyze their phylogenetic relationships. PCR amplification was performed using a specific primer (sequence available upon request) designed within the fragment in combination with a degenerate primer pointing outward of either the 5’ or the 3’ edge of the element (see Additional file 3 for details). PCR amplifications were performed as presented above. For each walking step, one to three clones were sequenced and each individual sequence (which corresponds to part of a single genomic copy), was deposited in GenBank ([GenBank:FJ707561–H707569]). Each new sequence was automatically validated as an extension of the initial fragment using the Cap contig assembly program included in the BioEdit software [68] with a minimum overlap between the two sequences of 50 bp, and a minimum DNA identity of 95%.

Analyzing the distribution of the DIRS1-like retrotransposons identified
Any DIRS1-like element that was newly identified in any species was searched in other decapods studied. Primers (sequence available upon request) were designed to be as specific as possible of each element, i.e. common to all sequences available for this element but unshared with the others. These specific primers were then used to PCR-screen all other species to look for well-conserved ele-
To investigate the distribution of DIRS1-like retrotransposons at a small phylogenetic scale, we focused on the Alvi elements identified within the family Alvinocarididae (hydrothermal shrimps). For that purpose, two kinds of PCR amplifications were performed, the first one using different combinations of specific primers, and the second using a specific primer in combination with the Retro1+ primer. Two or three specific primers (sequences available upon request) were designed for the Alvi1-4 families. Different combinations of primers were tested for each family (See Additional file 2) on either all 25 decapod species or on a selected subset of 12 species (i.e., all hydrothermal species and the two non-hydrothermal caridean shrimps).

**Sequence analysis**

Sequence similarity searches were performed using BLAST at the National Center for Biotechnology Information [http://blast.ncbi.nlm.nih.gov/Blast.cgi](http://blast.ncbi.nlm.nih.gov/Blast.cgi). Multiple DNA and protein sequence alignments were constructed using ClustalX [63] and manually curated using BioEdit [68]. DNA alignments were used to estimate pairwise distances and to build phenetic trees, using the pairwise deletion of gaps option of the MEGA4.1 software [69]. Pairwise distance calculation matrix was computed using p-distance model. Clustering of sequences into families was then performed in a Neighbor Joining tree [70] constructed using the Tamura 3 parameter model [71].

Representative amino acids consensus sequences of several elements were constructed by identifying the most common amino acid for each position of the alignment. Amino acid multiple alignment was checked using Gblocks [72] and ambiguously aligned sites were removed. Phylogenetic analyses were then conducted using Maximum Likelihood method [73] included in the Topali2 software [74] where the best-fit model (WAG model [75] with gamma distribution) was selected. For all phylogenetic analyses, support for individual clades was evaluated using non-parametric bootstrapping [76] using 1000 or 100 bootstrap replicates for nucleic acids and amino acids sequences, respectively.

**Abbreviations**

bp: base pair; DNA: DesoxyriboNucleic Acid; EST: Expressed Sequence Tag; ICR: Internal Complementary Region; indel: insertion/deletion; ITR: Inversed Terminal Repeat; LINE: Long Interspersed Element; LTR: Long Terminal Repeat; Mb: Megabase; MT: MethylTransferase; ORF: Open Reading Frame; PCR: Polymerase Chain Reaction; RH: RnaseH; RT: Reverse Transcripase; SINE: Short Interspersed Element; TE: Transposable Element; YR: Tyrosine Recombinase.
Authors’ contributions
MP carried out molecular and phylogenetic analyses and participated in the design of the study. EB conceived and coordinated the study, and participated in molecular analyses. The two authors wrote and approved the final manuscript.

Additional material

Additional file 1
Sequence identity matrix including all the nucleotide sequences acquired along the study. Sequence identity matrix performed using the pairwise deletion of gaps option included in the Mega 4.1 software. ID: Identity. The intra-family values are indicated in colour shaded boxes. Click here for file
[http://www.biomedcentral.com/content/supplementary/1471-2148-9-86-S1.xls]

Additional file 2
Retro1+ and degenerate primer sequences and specific primer combinations used for Alvi family screenings. Click here for file
[http://www.biomedcentral.com/content/supplementary/1471-2148-9-86-S2.xls]

Additional file 3
Summarization of the 73 sequences acquired in this study. Corresponding attributes of each sequence acquired are given: TE family, length, species, PCR methodology and primers used for the identification and GenBank accession number. Click here for file
[http://www.biomedcentral.com/content/supplementary/1471-2148-9-86-S3.xls]

Additional file 4
TE Walking methodology. Summarization of the methodology used to extend the transposable element pol sequences. Click here for file
[http://www.biomedcentral.com/content/supplementary/1471-2148-9-86-S4.pdf]

Additional file 5
Retro1+ primer and degenerate primer design. Correspondence between non specific primers used in this study and DIRS-like elements previously described. Mismatches are indicated in red letter. Click here for file
[http://www.biomedcentral.com/content/supplementary/1471-2148-9-86-S5.xls]

Additional file 6
Amino acids sequences alignment of DIRS1-like retrotransposons. Analysis including consensus sequences of the Alvi1-5 and Ymur3 families and well-described DIRS1-like retrotransposons from the YLDD motif of the RT domain to the ADALSR motif of the RH domain. Alignment performed using ClustalX program and corrected by Gblocks software. Amino acids signatures of all DIRS1-like retrotransposons are indicated in yellow. Amino acids signatures of DIRS1-like retrotransposons identified in hydrothermal decapods are indicated in blue. Click here for file
[http://www.biomedcentral.com/content/supplementary/1471-2148-9-86-S6.xls]

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