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#### RESEARCH ARTICLE

## **Evolutionary process of a tetranucleotide microsatellite locus** in **Acipenseriformes**

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#### Abstract

The evolutionary dynamics of the tetra-nucleotide microsatellite locus Spl-106 were investigated at the repeat and flanking sequences in 137 individuals of 15 Acipenseriform species, giving 93 homologous sequences, which were detected in 11 out of 15 species. Twenty-three haplotypes of flanking sequences and three distinct types of repeats, type I, type II and type III, were found within these 93 sequences. The MS-Align phylogenetic method, newly applied to microsatellite sequences, permitted us to understand the repeat and flanking sequence evolution of Spl-106 locus. The flanking region of locus Spl-106 was highly conserved among the species of genera Acipenser, Huso and Scaphirhynchus, which diverged about 150 million years ago (Mya). The rate of flanking sequence divergence at the microsatellite locus Spl-106 in sturgeons is between 0.011% and 0.079% with an average at 0.028% per million years. Sequence alignment and phylogenetic trees produced by MS-Align showed that both the flanking and repeat regions can cluster the alleles of different species into Pacific and Atlantic lineages. Our results show a synchronous evolutionary pattern between the flanking and repeat regions. Moreover, the coexistence of different repeat types in the same species, even in the same individual, is probably due to two duplication events encompassing the locus Spl-106 that occurred during the divergence of Pacific lineage. The first occured before the diversification of Pacific species (121-96 Mya) and led to repeat types I and II. The second occurred more recently, just before the speciation of A. sinensis and A. dabryanus (69-10 Mya), and led to repeat type III. Sequences in the same species with different repeat types probably corresponds to paralogous loci. This study sheds a new light on the evolutionary mechanisms that shape the complex microsatellite loci involving different repeat types.

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#### Introduction

Microsatellites are tandemly repeated short sequences widely dispersed throughout most eukaryotic genomes. Because of their highly polymorphic allele size caused by changes in repeat number, microsatellites are very popular genetic markers in population genetics and other evolutionary studies (Goldstein and Schlötterer 1999). However, the evolution of microsatellites themselves needs to be investigated, because the molecular mechanisms that generate microsatellites is not

completely understood. There are several ways to examine the evolutionary process of microsatellites.

Theoretically, some mutational models have been proposed, i.e. (i) the infinite allele model (IAM; Kimura and Crow 1964), which assumes that microsatellite mutations may create an infinite number of repeated units and allelic states not present in the population; (ii) the stepwise mutation model (SMM; Kimura and Ohta 1978), in which microsatellite mutations have the same probability of gaining or losing one repeat unit; (iii) the generalized stepwise model (GSM) or two phase model (TPM; Di Rienzo *et al.* 1994), an

**Keywords.** microsatellite evolution; homologous sequences; sturgeon phylogeny; sequence alignment; tandem repeat alignment; MS-Align.

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extension of the SMM model considering the probability for a microsatellite mutation to involve more than one unit; (iv) the *K*-allele model (KAM; Crow and Kimura 1970), where there are *K* allelic states and equal probabilities to mutate towards any of the other *K*-1 alleles.

Another straightforward and conclusive way of investigation is by direct observation of microsatellites in pedigrees. For example, Messier *et al.* (1996), mapped microsatellite sequences onto a phylogeny, and revealed the 'birth' of two microsatellites. Similarly, Taylor *et al.* (1999a) characterized microsatellite 'death' by placing the interrupted microsatellite sequences into a phylogeny. Moreover, many other studies focussed on phylogenetic and comparative genomic evidence for microsatellite evolution (Li *et al.* 1996; Ortí *et al.* 1997; Zhu *et al.* 2000; Chirhart *et al.* 2005; Leclercq *et al.* 2010).

It is accepted that the high polymorphism of microsatellites is mainly caused by replication slippage, which leads to the allelic length variation (Levinson and Gutman 1987). Previous studies have demonstrated that the repetitive sequences also contain many mutational events other than allelic length changes, such as base substitution and insertion/deletions (reviewed in Estoup *et al.* 2002). In general, when microsatellites are applied as genetic markers, only the allelic size variations information based on the electromorphs are used to analyse the genetic structure of populations or phylogenetic relationships, while the basic sequence variation information caused by substitution or indels is neglected. The consequence is that homoplasy is frequent in most studies based on microsatellite length polymorphism.

Basic variation information in the repeat sequences is then useful for investigation of sequence evolutionary process and the mechanism of microsatellite evolution. One of the methods should be to organize logically the allele sequences (cladistic tree) and to compare it with species phylogeny. However, because of the insertion or deletion of entire repeated blocks of nucleotides, the classical sequences alignment methods are not suitable for repetitive structure.

By-product of repeated region sequences, the substitution rate in microsatellite flanking sequences is similar to that in intron sequences (Brohede and Ellegren 1999), and microsatellite flanking sequences can be specific to each locus and informative for phylogenetic relationship investigation among alleles, populations and species (Zardoya *et al.* 1996; Makova *et al.* 2000). Flanking region evolution can be regarded as a part of the repeated region evolution, being a cause or a consequence of some mutational events modifying the repeat structure.

To extract the basic variation information, other than length variation, in microsatellites alleles sequences, a new alignment method, the MS-Align program (Bérard and Rivals 2003), which was successfully applied on minisatellite sequences (Bonhomme *et al.* 2007), have been applied in this study on microsatellite allele sequences. With this method, both the length variation and basic sequence variations were considered in the analysis.

The biological model on which this method is applied in this study is the order Acipenseriformes which includes 25 extant sturgeons and two paddlefish species. The phylogenetic relationships of this group have been investigated based on the mitochondrial gene sequences (Krieger *et al.* 2008) and morphological evidence (Artyukhin 2006). A comprehensive tree has been reconstructed by combining eight presently available mitochondrial gene sequences (4406 bp) (Krieger *et al.* 2008). In addition, five nuclear genes, *18S* rRNA gene (Krieger and Fuerst 2002), *5S* rDNA gene (Robles *et al.* 2005), *Hind*III and *Pst*I satellite sequences (Robles *et al.* 2004) and RAPD markers (Comincini *et al.* 1998), have been examined for phylogenetic analysis, confirming most topology, but brought no new decisive information with them.

Having a look at the systematics, several discrepancies still exist between the phylogenetic trees based on molecular and morphological data:

- The most basal group is composed of the genera *Polyodon* and *Psephurus* (the American paddlefish and Chinese swordfish, respectively);
- Then comes the two species, *Acipenser sturio* and *A. oxyrhinchus*, the European and Atlantic sturgeons, sister species, considered as ancestral while showing classical sturgeon morphology;
- An intermediate group composed of three species of the genus *Scaphirhynchus* (pallid, shovelnose and Alabama sturgeons) breaks the monophyly of the *Acipenser* genus;
- Finally, most sturgeons belong to a monophyletic group, the 'sturgeons sensu stricto' or 'true sturgeons', split into two geographic lineages: the Atlantic and the Pacific sturgeons. This structure is contradicted by the nomenclature: included in the 'true sturgeons' are the genus *Huso* composed of two species, one species in each geographic lineage and the genus *Pseudoscaphirhynchus*, composed of three species.

The contradictions between phylogeny and taxonomy have still to be resolved, such as the monophyly (and so the relevance) of genus *Huso*, and the basal position of two Atlantic sturgeons (*A. sturio* and *A. oxyrinchus*), which is not highlighted in the nomenclature (logical necessity of a distinct genus for these two species).

In this study, the conservation of the flanking sequences and the high variation of repeat sequences in microsatellite locus *Spl-106* among sturgeon species provided us the opportunity to understand the sequence evolution of this locus across the Acipenseriform species. We attempted to use the sequence variation both at the flanking and repeat region.

Understanding the microsatellite sequence evolution of locus *Spl-106* as a model was the first objective. Checking if the nuclear new information provided here confirms published phylogenies based on mtDNA sequences, mainly Peng *et al.* (2007) and Krieger *et al.* (2008) was the second objective. The new information was also to be confronted with the systematics anomalies given above.

#### Materials and methods

#### Sturgeon samples

Fin or muscle samples were collected in sturgeon farms, sturgeon research institutes or laboratories in different countries and preserved in 95% ethanol, except for *Scaphirynchus platorynchus*, for which the original sequence of microsatellite locus *Spl-106* was downloaded from database (GenBank: AF276175, McQuown *et al.* 2000). For each species, the sample size, quality, origin, collection sites and providers are listed in table 1.

#### Molecular methods

DNA extraction, PCR programme, electrophoreses, gel extraction of PCR products, TA cloning and sequencing procedures are same as described in Shao *et al.* (2005). The primer sequences of *As-100*, 5'-GGGAGAAAACTGGGGTAAA-3' and 5'-CCAAAAGAAGAATGGTAGACGG-3' (Shao *et al.* 2002), were used in the present study for all samples. The IAquick Gel Extraction kit (Qiagen, Shanghai, China) was used to extract each of the PCR products with the size between 100 bp and 500 bp from the 4% intermediate melting temperature (75°C) MetaPhor<sup>®</sup> agarose

(FMC BioProducts, Rockland, ME, USA) gel which can separate 4 bp difference alleles. Since some sturgeon species are polyploidy species, such as A. sinensis, if some individuals have more than two bands with size between 100 bp and 500 bp, we extracted all of them. The TA cloning kit (Takara Biotech, DaLian, China) and DH5α or DE3 competent cells were used to clone the PCR products. Unilateral sequencing of inserted DNA was performed on an ABI PRISM 377 DNA sequencer (PE Applied Biosystems, Foster City, CA, USA). For each PCR product fragment, at least three clones were sequenced in order to obtain at least two identical sequences of each extracted PCR product from the gel. However, the nonidentical sequences with the same size would also be used in alignment for discovering the homoplasy sequences and evolutional process of microsatellites.

#### Data analysis

The flanking and repeat sequences were aligned separately because of their structure differences. The flanking sequences were aligned by using the Clustal X version 1.81 Program (Thompson *et al.* 1997) and checked by hand.

**Table 1.** Specimen information of 15 sturgeon species studied.

Species	Abbr. of species name	Sample size	Quality (wild/W or farm/F)	Number of chromosomes <sup>a</sup>	Ploidy level <sup>c</sup>	Origin	Sample providers <sup>d</sup>
Scaphirynchus platorynchus	SP	1	W	120	2 <i>n</i>	USA	GenBank: AF276175
Pacific lineage Acipenser sinensis	AS	15	W	264±?	4 <i>n</i>	China	Ms Xiao H. Dr Wei Q. W.
A. dabryanus	AD	10	F (owi) <sup>b</sup>	?	4 <i>n</i>	China	Mr Zhou S. W.
A. schrenckii	AK	10	F (owi)	240	4 <i>n</i>	China	Mr Ma G. J.
Huso dauricus	HD	13	F (owi)	120	2n	China	Mr Ma G. J.
A. transmontanus Atlantic lineage	AT	10	W	226-288	4 <i>n</i>	USA	Dr Congiu
A. baerii	AB	9 10	F F	236-256	4 <i>n</i>	Russia Russia	Mr Ma G. J. Dr Congiu
A. naccarii	AN	10	6W, 4F	232-254	4n	Italy	Dr Congiu
A. fulvescens	AF	10	Ŵ	262±6	4n	USĂ	Dr Congiu
H. huso	HH	4	F	116-128	2n	Italy	Dr Congiu
A. gueldenstaedtii	AG	5 5 1	F F F	239-264	4 <i>n</i>	Russia Russia	Dr Congiu Mr Lan Z. Q. Anonymous
A. stellatus	AL	2	F	114-152	2n	Russia	Mr Lan Z. Q.
A. ruthenus	AR	10	F	114-122	2n	France	Mr Lan Z. Q.
A. sturio Psephurus gladius	Stu Pgl	11 1	W W	116±4 ?	2 <i>n</i>	France China	Dr Rochard E. KunShan,
Polyodon spathula	Psp	1	F	120	2 <i>n</i>	USA	Anonymous

<sup>&</sup>lt;sup>a</sup>The chromosome numbers were referred from <a href="http://web.unife.it/progetti/geneweb/sturgeon.html">http://web.unife.it/progetti/geneweb/sturgeon.html</a>, in which different chromosome numbers for each species from different literatures were collected from published literatures by Dr Fontana.

<sup>&</sup>lt;sup>b</sup>Owi means offsprings of wild individuals.

<sup>&</sup>lt;sup>c</sup>Ploidy levels of Acipenseriformes were referred from Peng et al. (2007).

<sup>&</sup>lt;sup>d</sup>Ms Xiao Hui, Institute of Chinese sturgeon, YiChang, P. R. China; Dr Wei Qiwei, Institute of Yangtze River Fisheries, JingZhou, P. R. China; Mr Lan ZeQiao, TianXia Sturgeon Fishery Ltd., YiChang, P. R. China; Dr Leonado Congiu, University of Ferrara, Italy; Mr Ma GuoJun, Institute of Amur River Fisheries, P. R. China; Mr Zhou ShiWu, Institute of Rare Aquatic Animals, YiBin, P. R. China; Dr Eric Rochard, Cemagref, France.

The maximum-likelihood phylogenetic tree of flanking sequences was constructed by the online software PhyML (Guindon and Gascuel 2003) through the bioinformatics platform Phylogeny. fr on the internet website http://phylogeny. lirmm.fr (Dereeper *et al.* 2008), using the default parameters.

The evolutionary relationships of the repeat region among sequences were constructed based on the alignment distances computed with the MS-Align program (Bérard and Rivals 2003). This program can extract all the information included in repeated DNA sequences, which is impossible to do using classical sequence alignment methods. Considering only the differences in the number of repeats is also inappropriate, since *Spl-106* is composed of distinct repeated units, called variants (mainly 'GAAA', 'TAAA' and 'TAGA'), most of which are 4-nucleotides long and A/T rich, and probably derived from each other by point mutations (Shao *et al.* 2005). Indeed, alleles of the same or similar length can be completely different in sequence.

To apply this method on tetra-nucleotide repeat sequences, we encoded the repeat unit with letters (see figure 4). To realistically reflect the similarity of distinct repeat units, we weighted the changes between variants (or units) in the alignment, e.g., changing the variant TAGA into TAAA or viceversa costs the weight of a nucleotide substitution in the alignment score. Computing an alignment score between all homologous sequences pairs yielded a matrix of pairwise distances between sequences (Bérard *et al.* 2006). We then fed this distance matrix into FastME (Desper and Gascuel 2002), a variation of the neighbour-joining (NJ) algorithm, to reconstruct a tree representing the relationships among sequences.

Another point is to estimate the robustness of the tree with respect to the alignment parameters and the level of confidence in its internal nodes. For this, we repeated the analysis with 19 sets of parameters and, each time, computed two mathematical measures: the variance accounted for (VAF), also known as percentage of explained variance, and the rate of elementary well-designed quartets (Re). The formula for the VAF and Re are detailed in Guenoche and Garreta (2000). The VAF quantifies the adequacy of representing the distances within a tree, while the Re gives a confidence value for each internal edge as the percentage of quartets supporting the split of this edge. Both measures are comprised in [0,1]. According to these two measures we chose the best tree among all 19 analyses. The complete protocol of this analysis has already been used for instance in Bonhomme et al. (2007), where it is more detailed.

To simply compare the repeat structures among sequences, the encoded (letters) sequences of the repeat region (including repeat-like (RL) region, see Results) were aligned by Clustal X version 1.81 program (Thompson *et al.* 1997) and checked by hand. Since the Clustal X program can only recognize the sequences composed of G, A, T and C, we used G, A and T to encode the three main repeat units (e.g., G = TAGA, A = TAAA, T = GAAA), and used C to encode all the other rare variable units. After alignment,

to avoid the confusion between encoding sequence and nucleotide sequence, and to make the encoding map readable, we replaced G, A and T with L, O and J, respectively, and replaced C with other letters for different variable units (figure 4).

#### **Results**

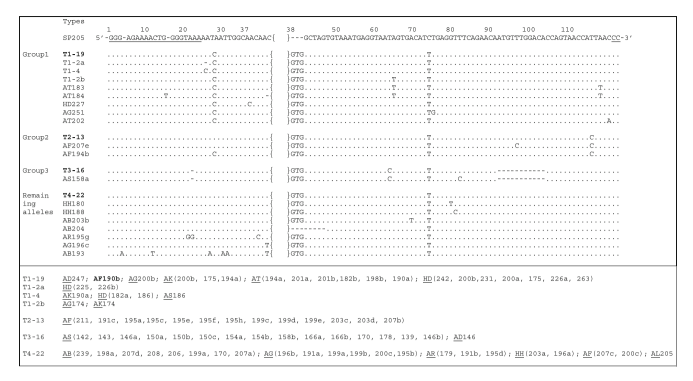
The primer set *As-100* successfully amplified in 13 out of the 15 species, failing in *A. naccarii* and *Psephurus gladius*, probably due to bad quality of samples. Among the 220 sequenced PCR products of the 13 species, 93 different homologous sequences were detected in 11 of 13 species. No homologous sequence of locus *Spl-106* was detected among the PCR products from two species, *Acipenser sturio* and *Polyodon spathula*.

Among 220 sequenced PCR products of the 13 species, 122 of them contained conserved flanking region of the target locus Spl-106 with flanking region identity level higher than 96%, 46 sequences from A. sinensis and A. dabryanus were less conserved (identity = 89%) due to the 11-bp deletion in the 3'- end (figure 1), while the remaining 52 sequences from different species contained flanking region with less than 79% identity to that of Spl-106. The flanking sequences of P. spathula and A. sturio have less than 75% identity and the repeat structures of these sequences were also different from that of Spl-106. As we did not observe any flanking sequence with a percentage of identity between 79% and 89%, we excluded from further analysis of all PCR products whose flanking regions exhibit a percentage of identity lower than 79%, as they probably originate from other loci than Spl-106. Finally, 93 different homologous sequences of locus Spl-106 were recorded from 11 species for further analysis.

#### Flanking region

Excluding the RL region, the length of the flanking region of locus *Spl-106* is 131–132 bp in most alleles, while it is 120 bp in alleles of *A. sinensis* and *A. dabryanus* because of the 11-bp deletion. No mutation occured in the 20 bp 3′-primer sites, so we did not show it in figure 1. Including the original sequence SP205 from *S. platorynchus* (Shao *et al.* 2002), 23 haplotypes of flanking sequences were observed among the 94 homologous sequences (figure 1). Four haplotypes gathered the most common types, accounting for 70 of the 94 sequences (i.e., T1-19, T2-13, T3-16 and T4-22). The remaining 19 rare haplotypes were observed from one (i.e., haplotypes names beginning with two specific letters) to four sequences (i.e., haplotypes T1-2a, T1-4 and T1-2b).

Compared to the original sequence SP205, two common mutation sites were observed among the 93 new sequences (positions 38–40 and position 69), and they are the only mutations observed in haplotype T4-22, from which all the other sequences evolved by a few point mutations. Below



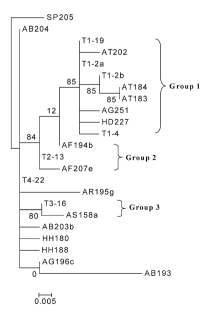
**Figure 1.** Sequence alignments of 23 haplotypes in the flanking region of locus *Spl-106*. The parentheses between position 37 and 38 indicate the repeat region. 5'-primer site is underlined on the left side, while 3'-end primer site is omitted on the right site of this map because no mutations occured in this primer region. Dots indicate identical sequences. Dashes indicates deletions. The dominant haplotypes were named T1 to T4 with the number of alleles behind them. For example, T4-22 means that the T4 haplotype present in 22 alleles, and the name of these alleles for each hyplotype are listed under the alignment map. Rare haplotypes, which is present in only one allele, were named directly with the code of corresponding allele.

the haplotype T4-22, in figure 1, there are seven different hyplotypes, which appear only one time each. Except T4-22 and these seven rare haplotypes; all the other haplotypes can be classified into three groups according to the group-specific mutations (figure 1). Each group includes a common haplotype, i.e. T1-19, T2-13 and T3-16, respectively. This classification is supported by the maximum-likelihood phylogenetic tree based on the flanking haplotypes sequences (high-bootstrap values for the splits leading to the groups; see figure 2).

It is noticeable that group 1 encompasses sequences of five species of Pacific lineage: A. sinensis, A. dabryanus, H. dauricus, A. schrenckii and A. transmontanus (but also four exceptions from Atlantic lineage species: the three alleles of one tetraploid A. gueldenstaedtii individual (AG251, AG200b and AG174) and one allele of A. fulvescens (AF190b)); group 2 can be recognized as species-specific haplotypes of A. fulvescens; group 3 gathers sequences only from the sister species pair A. sinensis – A. dabryanus of Pacific lineage (figure 1). The hyplotype T4-22 and the seven rare haplotypes belong to six sturgeon species of Atlantic lineage (A. gueldenstaedtii, A. baerii, A. stellatus, A. ruthenus, A. fulvescens and H. huso).

According to divergence time of Acipenseriformes estimated by Peng *et al.* (2007) and the pairwise Jukes–Cantor distance of haplotypes calculated by MEGA 4.0 (Tamura

et al. 2007), we estimated the evolutionary rate of the flanking sequence (table 2). The average sequence divergence rate of the flanking region of this locus in Acipenseriformes is



**Figure 2.** Maximum-likelihood phylogenetic tree of the flanking sequences reconstructed by the program PhyML (Guindon and Gascuel 2003) using the default parameters.

 $0.278 \times 10^{-9}$  with a range from 0.106 to 0.789  $\times$   $10^{-9}$  per site per year.

#### Repeat-like region

Between the 5'-end flanking and the repeat regions, an interrupted RL poly (A) structure  $T(A)_n TAAA$  distinguishes the Pacific and Atlantic lineages, with n = 5 to 7 (mainly 7) in the sequences of Pacific species and n = 8 or 9 in Atlantic (figure 4). We included it as a part of repeat region because of its variable RL structure and of its special position. The RL region appears at both sides of the repeat region in several sequences from Pacific species, while it was absent in some others (see figure 4, Pacific group).

#### Repeat region

The pairwise divergence of the repeat regions of the 94 homologous sequences was estimated by the tandem repeat alignment program MS-Align (Bérard et al. 2006). This yielded a pairwise distance matrix, which served to reconstruct a NJ tree representing the relationships among homologous sequences. To understand the effect of the variable RL region on the tree topology, we inferred two trees, one based on repeat region with RL region (tree I in figure 3, VAF = 0.95, Re = 0.54 at the root) and other without RL region (tree II, VAF = 0.99, Re = 0.52 at the root). Both trees share a similar topology; only few sequences are placed differently. Except for the solely branched sequence SP205 in both trees and the sequence AB239 in tree I, all other sequences were basically clustered into two nodes (support value Re = 0.62 (up node) and 0.63 (down node) in tree I; Re = 0.77(up) and 0.81 (down) in tree II) due to the differences in the composition of repeat units and in number of copies of the three main repeat units (see figure 4). At the exception of six sequences (AR195g, AG199a, AB199a, HD175, AK175 and AT201b) placed within clade AS-AD (A. sinensis and A. dabryanus, tree I), further split clades are entirely Pacific or Atlantic. Although the sequences of the Atlantic group are not clustered in one well-supported clade in tree II, the division of Pacific and Atlantic groups are still obvious. Only one small Pacific/Atlantic mixed clade (Re=0.64) is observed in tree II. However, inside this mixed clade, the next nodes split individuals of Pacific and Atlantic species with higher support values as well.

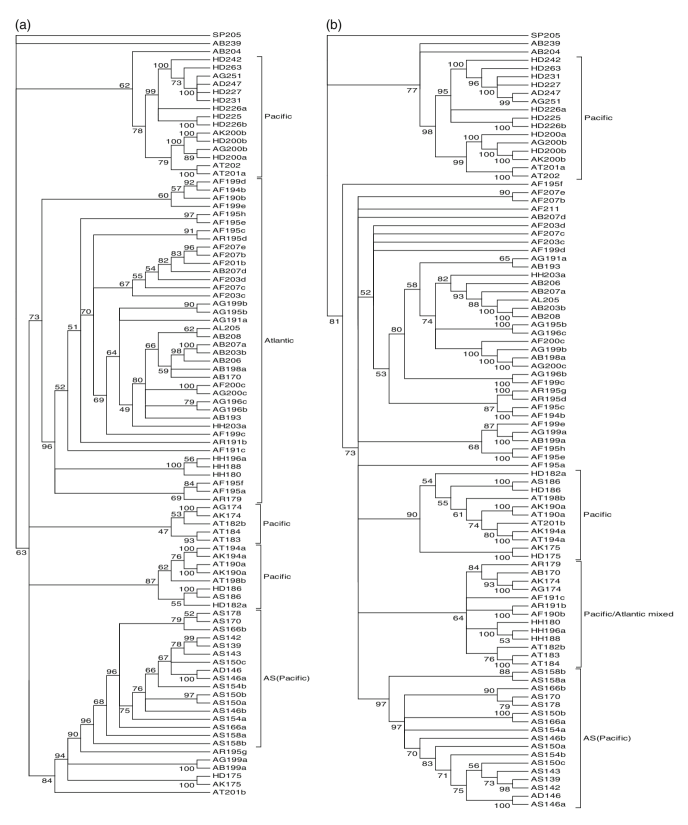
The repeat structures were compared between the sequences from species of Pacific and Atlantic lineages in figure 4. We found no identical repeat structure between sequences of Pacific and Atlantic lineages, while identical sequences were often observed among species within the lineage. The only exceptions are the four sequences from two species of Atlantic lineage: three AG alleles: AG251, AG200b and AG174, and one A. fulvescens allele: AF194b, which contain the same repeat structure or RL region as sequences of Pacific lineage species. Since the three AG alleles (AG251, AG200b and AG174), found in the same individual (given by an anonymous small fish farm, see table 1). unexpectedly belong to the Pacific lineage both at the flanking and repeat structure, the species of this individual is probably misidentified. Thus, we excluded these three alleles from further discussion.

The basic structures of the repeat region were diversified among species, even within species. Three main types of compound repeat structures were observed: type I: (TAGA)<sub>n</sub>  $(TAAA)_m$  with the repeat number n greater than m (n =12–23; m = 5-7); Type II:  $(TAGA)_n$   $(TAAA)_m$  with nless than or equal to m (n = 2-7; m = 5-12); type III:  $(TAAA)_m(GAAA)_n$  with m greater than n (m = 3-11; n =0-4). The difference between types I and II is mainly due to the different repeat number of the two basic repeat units, and the sequences with these two repeat types can have the same flanking haplotypes and RL region, for instance the sequences with flanking haplotype T1-19 (in figures 1 and 4). However, the sequences with type III repeats not only have very different basic repeat units, but also have an 11bp sequence deletion in the flanking region (haplotypes of group 3 in figure 1) and even no RL region at all (figure 4). The distribution of repeat types in species were clarified in

**Table 2.** Estimation of evolutionary rate of microsatellite-flanking sequences in sturgeons.

Pairwise comparisons <sup>a</sup>	Divergence time (My) Peng et al. (2007)	Jukes–Cantor distance	Evolutionary rate $(\times 10^{-9})$	
SP-T4	74.4 (150.8–86.4)	0.0079	0.106	
SP-T3	140.3 (150.8–10.5)	0.0176	0.125	
SP-T1	54.6 (150.8–96.2)	0.0159	0.292	
SP-T2	98.1 (150.8–52.7)	0.0159	0.162	
T4-T3	75.9 (86.4–10.5)	0.0085	0.112	
T4-T1	9.8 (86.4–96.2)	0.0077	0.789	
T4-T2	33.7 (86.4–52.7)	0.0077	0.229	
T3-T1	85.7 (10.5–96.2)	0.0171	0.200	
T3-T2	42.2 (10.5–52.7)	0.0171	0.406	
T1-T2	43.5 (96.2–52.7)	0.0155	0.357	
Average	, ,		0.278	

<sup>&</sup>lt;sup>a</sup>SP indicates SP205, T1, T2, T3 and T4 are the four most common flanking haplotypes.



**Figure 3.** Neighbour-joining trees based on repeat region. (A) Tree I based on repeat region with repeat-like (RL) region. (B) Tree II based on repeat region only. The pairwise molecular divergence among homologous was estimated by aligning their tandem repeat sequences using the MS-Align program (Bérard and Rivals 2003). The tree was inferred with FastME (Desper and Gascuel 2002) from the matrix of pairwise alignment distances. For each internal edge, the confidence value (between 0 and 1) and the rate of elementary well-designed quartets (Re) were computed (Guenoche and Garreta 2000). The abbreviations of the species names are given in table 1.

Pacific o	group	Atlantic group			
SP205	OKOLLLLLLLLL	SP205	OKOLLLLLLLLL		
	Type I		Type I		
HD263	OKOLLLLLLLLLLLLLLLLLLLLLN-OKOKOO	AB239	UKOLLLLLLLLLLLLLLLLLLO0000		
AG251	OKOLLLLLLLLLLLLLLLLLLLLNOOKOKOO	AB204	<u>UK</u> LLLLCLLDLLLLLLLLOX		
AD247	OKOLLLLLLLLLLLLLLLLLLLLNOOKOKOO		Type II		
HD231	OKOLLLLLLLLLLLLLLLLLNOOKOO	AF211	<u>UKO</u> LLLLLLL0000000000		
HD227	OKOLLLLLLLLLLLLLLLLLNOOKOO	AF207e	UKOLLLLLLL000000000		
HD242	OKOLLLLLLLLLLLLLLLLLLLLLOOKOO	AF207b	UKOLLLLYLL00000000		
HD225	OKOLLLLLLLLLLLLLLL-000OKOO	AF195f	UKOLLLLLL000000		
HD226b	OKOLLLLLLLLLLLLLLLL-000OKOO	AF195a	UKOOLLLLLLOOOOOO		
HD226a	OKOLLLLLLLLLLLLLOL-000OKOO	AB207d	UKOLLLLLL0000000000		
AG200b	LLLDLLLLLLL000K00	AF207c	UKOLLLLL00000000000		
HD200b	LLLDLLLLLLL000K00	AF203d	UKOLLLLL000000000		
AK200b	LLLDLLLLLLL000K00	AF199e	UKOLLLLL000000000		
HD200a	LDLLLLLLLL000K00	AG199a	UOOLLLLLOOOSOOOOO		
AT201a	OFLLLLDLLDLLLLOOKOO	AB199a	UOOLLLLL000S00000		
AT202	OKLLLLDLLDLLLLOOKOO	AF203c	UKOLLLL0000000000		
	Type II	HH203a	UKOLLLLOOOOOOOWO		
AT198b	OKOLLOLLOOOOOOJJ	AF199d	UKOLLLL0000000000		
AK175	U-OLLOLLOOOO	AF191c	UKOLLLL00000000		
HD175	U-OLLOLLOOOO	AF195h	LIOLLLL000000000		
AT201b	OFOLLOL-0000000000	AF195e	UKOLLLL000000000		
AK194a	OKOLLOL-00000000	AR195q	OUOLLLX000000000		
AT194a	OKOLLOL-00000000	AR195d	UKOLLLX00000000		
AK190a	OKOLLOL-0000000	AG196b	UILLL0000000000		
AT190a	OKOLLOL-0000000	AF199c	UKOLLL0000000000		
AS186	OKOLLOL-0000000	AF195c	UKOLLL000000000		
HD186	OKOLLOL-000000	AF194b	OKOLLL000000000		
HD182a	OKOLLOL-000000	AR191b	UKOLLL00000000		
AT182b	0K0LLL0000000	AF190b	UKOLLL00000000		
AG174	OKOLLL00000	AR179	UKOLLL000000		
AK174	OKOLLL00000	HH180	UKOLML000000		
AT183	0K0LLL0000U00	HH196a	UKOLML00000000		
AT184	0K0LLL0000U00	HH188	UKOLML0000000		
HIIOI	Type III	AB206	UIBLLOOOOOOOOVOWO		
AS158b		AB207a	UIBLLOOOOOOOUVOWO		
AS158a		AB207d	UIBLLOOOOOOO-UVOWO		
AS146b		AB203D AB208	UKOLL0000000-UV0W0		
AS154a		AL205	UIOLLOOOOOOO-UVOWO		
AS154a AS150a		AB198a	UIBLL00000000000		
AD146		AG200c	UIOLL00000000000		
AS146a		AG200C	UKOLL00000000000		
AS140a AS142		AF200c	UIOLL00000000000		
		AG195b	UKOLL00000000000		
AS143 AS139			<del></del>		
		AG196c	<u>nko</u> <u>rr0000000000</u>		
AS166a		AG191a	UKOLL000000000		
AS150b		AB170	UIBLL00000		
AS166b		AB193	<u>UEO</u> ZYOWOWOOOOO		
AS154b					
AS178					
AS170 AS150c					
O = TAAA	L= TAGA	X = TAAG	N = TAGGA		
K = AAAA	J = GAAA	Y = TAGC	M = TAAGA		
F = AAA	V = TATA	Z = CAGT	D = GA		
U = TAAAA		P = CAAA	B = TA		
_		S = TACA	D-IA		
I = AAAAA					

**Figure 4.** Encoding map of repeat and repeat-like (RL) region. The RL regions are underlined. The four exceptional sequences of Atlantic species, three AG and one AF are in bold font (see text). The different repeat types are divided and indicated by types I, II and III.

figure 4. Almost all sequences from Atlantic species contain repeat type II, except for two sequences from A. baerii, which shows repeat type I. However, in Pacific group column, each species displays two repeat types. A. schrenckii, Huso dauricus and A. transmontanus contain repeat types I and II, and the sequence quantity of each repeat type is similar. A. dabryanus and A. sinensis mainly include repeat type III with two exceptional sequences, AD247 and AS186, which have repeat type I and type II, respectively. It should be mentioned that in one individual of the diploid species H. dauricus, we found four alleles (it was a surprise, probably due to paralogy following a local duplication), HD242, 200, 186 and 182a, in which the two larger alleles have repeat type I and flanking haplotype T1-4, while the two smaller ones have repeat type II and flanking haplotype T1-19 (figures 1 and 4). Moreover, in A. sinensis and A. dabryanus, AS186 (in type II) and AD247 (in type I) differ from the other alleles of the same species, both in flanking sequence and repeat structure (see figures 1 and 4). Allele AS186 coexists with the other four alleles (in type III) in one individual of A. sinensis, which has been reported as a tetraploidy species.

#### Discussion

The flanking region of locus *Spl-106*, which is originally isolated from *S. platorynchus* by McQuown *et al.* (2000), was highly conserved among the species of genera *Acipenser*, *Huso* and *Scaphirhynchus*, which diverged about 150 million years ago (Mya) (Peng *et al.* 2007). The failed amplification of the homologous sequences in *A. sturio* and *P. spathula* is probably due to mutations occurring at the primers sites.

Long time of microsatellite loci stability have been observed in some other species, e.g., over 300 My in turtles (FitzSimmons *et al.* 1995) and over 450 My in fish species (Rico *et al.* 1996). Flanking region stability on long time diverged taxa is necessity to study microsatellite evolution and to resolve the phylogenetic relationships of lineages. However, only a distorted sight is expected since we did not analyse the allele sequences at different steps of their evolution, but rather the present sequences which were influenced by very ancient speciation and splitting events.

#### Flanking region evolution

Concerning the flanking region of microsatellites, the rate of sequence divergence has been estimated in some fish groups: from 0.02% to 0.05% per My among diverse fish species (Rico *et al.* 1996), and between 0.14%–0.83% in sunfish (Neff *et al.* 1999). Our estimation on the microsatellite locus *Spl-106* in sturgeons is between 0.011% and 0.079% with an average at 0.028% per My, which is similar to that estimated by Rico *et al.* (1996). However, it is slightly lower than that of satellite DNA in sturgeons (0.07% per My for *Hind*III and 0.11% per My for *Pst*I; Robles *et al.* 2004).

The slow rate of molecular evolution in Acipenseriformes had been demonstrated in satellite loci (*Hind*III and *Pst*I)

and mitochondrial loci (De la Herrán et al. 2001: Krieger and Fuerst 2002; Robles et al. 2004). Also, this general observation is now supported by our data set of microsatellite sequences. According to Neff et al. (1999), the rate of sequence divergence at the microsatellite flanking regions may be dependent on the locus itself. However, according to several observations, the slow rate of sequence divergence seems to be the general rule of microsatellite evolution in this ancient fish group, Acipenseriformes: (i) among the 108 loci screened in S. platorynchus, 65%–80% can be successfully amplified on Acipenser species (McQuown et al. 2000); (ii) the microsatellite loci originally screened in the American paddlefish P. spathula can be crossamplified on most species of the family Acipenseridae (Heist et al. 2002). This apparently easy cross-priming should be the mark of a general slow mutation rate.

#### Repeat region evolution and paralogy

According to the diversified repeat structures, the evolutionary process of *Spl-106* is more complicated in the Pacific than in the Atlantic lineage. In the Atlantic lineage, slippage duplication is the main mechanism in the evolutionary process, and the repeat number changes among the two basic repeat units contribute a lot to the size homoplasy. However, in Pacific group, each species contains generally two types of repeat structures corresponding to two evolutionary routes.

It is improbable that the different repeat types found in the same species could be due to stepwise mutation model. The same flanking sequence with different repeat motifs should be an evidence of paralogy (Karhu et al. 2000). Here, several point mutations and indels also occured on the flanking sequences, which indicate that the two paralogs have been evolving independently for a long time. According to the distribution of the three repeat types on species (figure 4), two duplication events encompassing the locus Spl-106 probably occurred during the divergence of Pacific lineage: the first occured before the diversification of Pacific species (121-96 Mya) and led to repeat types I and II; the second occurred more recently, just before the speciation of A. sinensis and A. dabryanus (69–10 Mya) and led to repeat type III. Therefore, the sequences in the same species with different repeat types likely are paralogous loci. Paralogs are genes that derive from a single gene that was duplicated within a genome. This could also explain the phenomenon that four alleles, among which two contain repeat type I and the other two contain repeat type II, were present in one individual of the diploid species H. dauricus, and that five alleles, among which one has repeat type II, while the other four alleles have repeat type III, were detected in one individual of the tetraploid species A. sinensis.

#### Evolutionary relationships of flanking and repeat regions

Our results show a synchronous evolutionary pattern between flanking and repeat regions. Homologous sequences with different repeat types may have the same flanking sequences, but homologous sequences with different flanking haplotypes normally have different repeat structures. Especially, in the sequences of clade AS-AD (group 3; figure 2), the large variation due to 11-bp deletion in the flanking region is consistent with the great change in repeat structure (see AS and AD alleles in figure 4). Microsatellite loci are as vulnerable as the rest of the genome to point mutations, whose effect is to divide long repeat stretches into smaller subunits, and hence decrease the rate at which slippage occurs (Bell and Jurka 1997; Kruglyak et al. 1998). Without the effects of the slippage mutation, the rates of base substitution or indels should be similar between the flanking and repeat regions. Moreover, both flanking and repeat regions (including the RL region) support the splitting of Pacific and Atlantic lineages, and the homologous sequences evolved independently after this split.

#### Adequacy with sturgeon phylogeny

The phylogenetic relationships of Acipenseriformes have been extensively studied by morphological characters and mitochondrial genes; however, many issues remain unresolved (Artyukhin 2006; Peng *et al.* 2007; Krieger *et al.* 2008). Several nuclear genes, e.g. *Sox* genes, 5*S* rDNA and 18*S* rDNA, have also been used, and did not contradict the consensus phylogeny (Krieger and Fuerst 2002; Hett and Ludwig 2005). For instance, the *PstI* satellite sequence supported the observed Atlantic/Pacific split and *HindIII* satellite sequence supported the basal position of both *A. sturio* and *A. oxyrinchus* (Robles *et al.* 2004).

In our data set, the flanking regions of microsatellite locus Spl-106 are highly conserved at the subfamily level: among the genera Scaphirhynchus, Huso and most species of genus Acipenser, except for A. sturio, in which no homologous sequence of locus Spl-106 was detected by the same primer set. It confirms that A. sturio diverged earlier than Scaphirhynchus in the phylogeny of Acipenceriformes. The failed amplifications in A. naccarii and Psephurus gladius were probably due to the quality of the DNA extracted from limited sample material. We could not conclude that this locus was absent from A. naccarii, which is very close to A. gueldenstaedtii and A. baerii in all sturgeon phylogenies. However, it is possible that this locus does not exist in the paddlefish, Psephurus gladius, since we also did not find it in the other paddlefish, Polyodon spathula, allowing us to estimate the birth of this locus at about 185 Mya, during the early Jurassic (Peng et al. 2007).

The Atlantic and Pacific clades were previously suggested by Ludwig *et al.* (2001) using cytochrome-*b* DNA sequences and confirmed by Peng *et al.* (2007). In our results, both the flanking and the repeat regions follow the two geographical lineages. Our sequences are also influenced by the close relationships between (i) *A. sinensis* and *A. dabryanus*; (ii) *A. schrenckii*, *A. transmontanus* and *H. dauricus* (i.e. the Pacific clade); (iii) *A. fulvescens*, *A. ruthenus*, *A. baerri*,

A. gueldenstaedtii and A. stellatus, which have been supported by the other molecular evidence (Krieger et al. 2008). Our results indicate that the evolution of a given microsatellite locus is directly driven by the speciation history revealed by mtDNA sequences, but does not follow the morphological and taxonomic considerations (Artyukhin 2006). The two Huso species clearly belong to the genus Acipenser and are separately assigned to Pacific and Atlantic lineages. Although the flanking sequence is not variable enough for reflecting the phylogenetic relationships among all sturgeon species, we still found several specific diagnostic sites from the closely related species pair, A. sinensis and A. dabryanus (group 3 in figures 1 and 2), and the lake sturgeon A. fulvescens (group 2 in figures 1 and 2).

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