Indonesian Journal of Agricultural Science 5(2) 2004: 45-62

CONTRIBUTION TO THE PHYLOGENY OF THE PANGASIIDAE BASED ON MITOCHONDRIAL 12S RDNA

L. Pouyaud^a, R. Gustiano^b, and G.G. Teugels^c

^aInstitut de Recherches pour le Developpement (IRD), GAMET, B.P. 5095, F-34033 Montpellier, France ^bResearch Institute for Freshwater Fisheries (RIFF), Jalan Sempur No. 1, Bogor 16151, Indonesia ^cAfrica Museum (MRAC), Ichthyology Department, B-3080 Tervuren and Katholieke Universiteit Leuven, Laboratory of Comparative Anatomy and Biodiversity, B-3000 Leuven, Belgium

ABSTRACT

Catfishes are generally one of the economically important groups of fresh and brackish water fishes in the world. In many countries, they form a significant part of inland fisheries, and several species have been introduced in fish culture. Judging from literature, the main constraint to cultivate wild species and to optimise the production of pangasiid catfishes is due to the poorly documented systematics of this family. In the present contribution, the phylogenetic relationships within Pangasiidae are studied to contribute to a better insight in their taxonomy and evolution. The genetic relatedness is inferred using mitochondrial 12S rDNA gene sequences. To resolve the phylogenetic position of Laides in this group of catfish, five genera of Asian and African Schilbeidae are also considered. The results showed that a species group (complex) could be clearly seen in the genetic tree. Pangasius is more derive than the other genera. By using approximate molecular clock/evolutionary calibration from mitochondrial gene, a new episode of speciation for the family marked explosive radiation about 5-8 million years ago (mya). This adaptive radiation extended until the Late Pleistocene. Regarding the relationships between the Pangasiidae and Schilbeidae, two families show an allopatric distribution with slight overlap. The Pangasiidae occur mainly in Southeast Asia, while the Schilbeidae are seen mainly on the Indian subcontinent (including Myanmar) and Africa. It confirms the separation between Schilbeidae and Pangasiidae occurred in the Early Miocene.

[*Keywords*: Pangasiidae, evolution, biogeography, taxonomy, morphology, mitochondrial DNA, cytochrome b gene, 12 S rDNA gene, genetic distance]

INTRODUCTION

Advances in DNA sequencing technology over the past decade have been phenomenal (Hillis *et al.* 1990; Meyer 1993; Simon *et al.* 1994). The power of the technique has ensured that it has become one of the most utilized molecular approaches for inferring phylogenic history (Hillis *et al.* 1990). The primary attraction of nucleic acid sequencing relates to the

fact that the characters (nucleotides) are the basic units of information encoded in organisms (the genome), and that the potential of informative data sets are immense.

Fish may contain more than 10^8 nucleotide pairs per haploid genome, although the number of independent characters that could be used in phylogenetic analysis is considerably lower. To use nucleotide sequence positions in the phylogenetic studies, orthologous sequences must be aligned. The number and size of orthologous sequences that can be aligned will differ depending on the level of comparison, but for most studies, systematically informative variation is essentially inexhaustible (Hillis *et al.* 1990).

Due to the lack of molecular data, considerable confusion has arisen in the systematics of pangasiid catfishes. This is illustrated by the genus *Laides* Jordan, 1919, placed either in the Pangasiidae (Roberts 1989) or in the Schilbeidae (Vidthayanon 1993). Following Mo (1991), the Pangasiidae are the sister group of the Schilbeidae.

In Mekong Delta, the aquaculture production of *Pangasius* exceeds significantly the production from capture of fisheries, showing the economically importance of their aquaculture in the global fisheries sector. Meanwhile, in Indonesia, although more than 10 pangasiid species have been listed, the only *Pangasius* cultured remained *Pangasianodon hypophthalmus*, introduced from Thailand. Of the 28 valid species, few have been reproduced successfully: *P. hypophthalmus* since 1966, and more recently several others including *Pangasianodon gigas*, *Pangasius bocourti*, and *Pangasius djambal* (Roberts and Vidthayanon 1991; Legendre *et al.* 2000).

In the present contribution, the phylogenetic relationships within the Pangasiidae are studied to contribute to a better insight in their taxonomy and evolution. For this purpose, the genetic relatedness 46

between 23 out of the 24 valid species of the genera *Pangasius* and *Helicophagus* will be inferred using mitochondrial 12S rDNA gene sequences. These data will be combined with results from previous studies based on cytochrome b/allozyme phylogenies (Pouyaud *et al.* 2000). To resolve the phylogenetic position of *Laides* in this group of catfishes, five genera of Asian and African Schilbeidae (*Clupisoma* Swainson, 1838; *Eutropiichthys* Bleeker, 1862; *Silonia* Swainson, 1838; *Pseudeutropius* Bleeker, 1862; and *Schilbe* Oken 1817) will also be considered.

MATERIALS AND METHODS

Origin and number of samples used are listed on Table 1. Fresh specimens collected from 1996 to 2000 were dissected on site and tissue samples (eye, muscle, and liver) were stored in liquid nitrogen for transfer to the laboratory. They were then stored at -20°C until analysis in the molecular genetic laboratory in Pasar Minggu, Jakarta, Indonesia.

Mitochondrial analysis consisted in sequencing part of the 12S rDNA gene. One to five individuals were analysed for 22 species of Pangasiidae and 7 species of Schilbeidae. Hundred milligrams of muscle were used to isolate total genomic DNA using a hexadecylmethyl-ammoniumbromide extraction process (Doyle and Doyle 1987).

Primers used for the amplification of the mitochondrial gene included 12S light strand 5'-TTACACATGCAAGTCTCCGC-3' and 12S heavy strand GTTACGACTTGCCTCCCTT-3' defined on the complete mitochondrial sequence of Cyprinus carpio (EMBL X61010; Chang et al. 1994). DNA amplification and sequencing were performed following the procedures developed in Pouyaud et al. (2000). To check sequence accuracy and to correct any ambiguity bases, both strands were sequenced using each one of the two initial PCR primers. Sequences of both strands were compared with each other and aligned using the sequence editor ESEE (version 3.1 s; Cabot and Beckenbach 1989). Comparison of absolute numbers of transitions and tranversions were calculated and plotted versus corrected genetic distances using Kimura's distance two-parameter method (Kimura 1980). Phylogenetic inference was based on the neighbour joining (Saitou and Nei 1987) method (NEIGHBOR program in PHYLIP; Felsenstein 1993) from Kimura's distances among species. The reliability of the topologies was assessed with bootstrapping on 1000 replicates (SEQBOOT and CONSENSE programs in PHYLIP; Felsenstein 1993).

L. Pouyaud et al.

RESULTS AND DISCUSSION

All material listed was sequenced (Table 1). For many taxa, the results showed single sequence (no intraspecific variation) except for *Pangasius polyuranodon* and *Pangasius kunyit* having different mitochondrial haplotype from distinct geographic populations. In the following section, genetic interrelationships of the Pangasiidae, genetic interrelationships between Pangasiidae and Schilbeidae, evolutionary process based on speciation peaks, systematic position of the genus *Laides*, and speciation process are discussed.

Genetic Interrelationships of the Pangasiidae

Seven hundred and thirty seven nucleotides encoding the internal part of the mitochondrial 12S rDNA gene were aligned for all pangasiid species (Table 2). In the 737-bp sequences, 123 (17%) sites were polymorphic and 72 of these (59%) were phylogenetically informative.

In a first step, the genetic differentiation within each species studied was assessed to confirm the validity of the species recognised by Roberts and Vidthayanon (1991), Pouyaud et al. (1999), Pouyaud and Teugels (2000), and Ng and Kottelat (2000). For this purpose, corrected pairwise Kimura's genetic distances were computed considering a population as the unit of analysis. A phylogenetic tree was obtained from the genetic distance matrix using the nearest neighbour joining method (data not shown). This dendrogram revealed that all populations analysed for a given species aggregated and constituted in all cases of a monophyletic group (bootstrap between 77% and 100%), which confirms the existence of 23 out of 24 species recognised or described previously. Except for P. polyuranodon and P. kunyit, genetic distances among conspecific populations were null or insignificant [Pangasius macronema: two nucleotidic substitutions between Chao Phraya (Thailand) and Mekong (Vietnam) populations; Pteropangasius micronemus: a unique substitution between Batang Rajang (Sibu, Malaysia) and Batang Hari (Sumatra, Indonesia) populations].

The average Kimura's distances within species revealed significant genetic differentiation among populations of *P. polyuranodon* and *P. kunyit* located in vicariant biogeographic areas. *P. polyuranodon* is composed of three distinct genetic groups of populations, respectively from East Kalimantan (Samarinda), the rest of Indonesia (Sumatra, Central Kalimantan, West Kalimantan, Sarawak), and Indo-

Species identification ¹	n	Location (site, river, country)
Panaasius diambal Blooker 1846	1	Jambi Batang Hari Sumatra Indonesia
rangasius ajambai Bieekei, 1840	1	Parito, Cantral Pornao, Indonesia
Panageius rheaphilus Pouvaud & Teugels 2000	1	Tanjung Selor Kayan Bulungan East Borneo, Indonesia
Tungustus Theophitus Touyadd & Teugets, 2000	1	Tanjung Reden Berau Bulungan Fast Borneo, Indonesia
Pangasius bacourti Sauvage 1880	1	Mekong Vietnam
Pangasius nasutus Bleeker 1863	1	Musi Sumatra Indonesia
I angastas nasatas Dicelei, 1000	1	Palangkaraya, Kahayan, Central Borneo, Indonesia
	1	Kapuas. West Borneo. Indonesia
Pangasius conchophilus Roberts & Vidthayanon, 1991	1	Mekong, Vietnam
Pangasius larnaudii Bocourt, 1866	1	Chau Doc, Mekong, Vietnam
Pangasius sanitwongsei Smith, 1931	1	Chau Doc, Mekong, Vietnam
Pangasius pangasius Hamilton, 1822	1	Chandpur, Ganges, Bangladesh
Pangasius krempfi Fang & Chaux, 1949	1	Mekong, Vietnam
Pangasius kunyit Pouyaud et al., 1999	1	Sanga-sanga, Mahakam, East Borneo, Indonesia
	1	Pontianak, Kapuas, West Borneo, Indonesia
	1	Barito, Central Borneo, Indonesia
	1	Kinabatangan, Sabah, North Borneo, Malaysia
	1	Mekong, Vietnam
Pangasius polyuranodon Bleeker, 1852	1	Jambi, Batang Hari, Sumatra, Indonesia
	1	Barito, Central Borneo, Indonesia
	1	Kapuas, West Borneo, Indonesia
	1	Batang Rajang, Sarawak, North Borneo, Malaysia
	1	Mahakam, East Borneo, Indonesia
	1	Mekong, Vietnam
Pangasius macronema Bleeker, 1851	1	Mekong, Vietnam
Pangasius humeralis Roberts, 1989	1	Sintang, Kapuas, West Borneo, Indonesia
Pangasius lithostoma Roberts, 1989	1	Sintang, Kapuas, West Borneo, Indonesia
Pangasius kinabatanganensis Roberts & Vidthayanon, 1991	1	Sukau, Kinabatangan, Sabah, North Borneo, Malaysia
Pangasius nieuwenhuisii Popta, 1904	1	Mahakam, East Borneo, Indonesia
Pangasianodon hypophthalmus Sauvage 1878	1	Mekong, Vietnam
Pangasianodon gigas Chevey, 1930	1	Mekong, Thailand
Pteropangasius micronemus Bleeker, 1847	1	Batang Hari, Sumatra, Indonesia
	1	Indragiri, Sumatra, Indonesia
	1	Sihu Batang Baiang Sarawak Malawaia
Discongramme and an and a second seco	1	Choo Dhroup Theiland
Haliconhagus typus Blocker, 1858	1	Lambi Batang Hari Sumatra Indonesia
Helicophagus Ientorbynchus Ng & Kottelat 2000	1	Mekong Laos
Schilba mandihularis Günther 1867	1	Comoé Ivory Coast West Africa
Clunisoma garua Hamilton 1822	1	Chandpur Ganges Bangladesh
Laides sinensis Huang 1981	1	Chao Phraya Thailand
Silonia silondia Hamilton 1822	1	Chandpur Ganges Bangladesh
Eutropiichthys vacha Hamilton, 1822	1	Chandpur, Ganges, Bangladesh
Pseudeutropius brachypopterus Bleeker, 1858	1	Muara Bungo, Batang Hari, Sumatra, Indonesia
	1	Sintang, Kapuas, West Borneo, Indonesia

Table 1. Species identification, sampling origin, collection reference, sample size (n) analysed with mitochondrial DNA.

¹Species identification is based on Roberts and Vidthayanon (1991) and Pouyaud et al. (2000).

China (Vietnam and Thailand). The average genetic distances between populations from East Kalimantan and those from the rest of Indonesia or Indo-China (d = 0.0253) is equivalent to average distances between *Pangasius conchophilus* and *P. bocourti* or between *P. nasutus* and *P. larnaudii*. The genetic

differentiation between populations from the rest of Indonesia and Indo-China is the smallest (d = 0.0126), but corresponds to the genetic differentiation between *P. micronemus* and *P. pleurotaenia* (d = 0.014). Similarly, three genetic groups are found within the species *P. kunyit*, respectively from Indonesia

P. polyuranodon Rest												
of Indonesia	AGCCCAAGAC	GCCTTGCTAC GCCACACO	CAAGGGAATI	T CAGCAGTAAT	AAACATTAAG 0	CCATAAGTGT A	AACTTGACT T	AGTTAGGGC C	A-A-AAGGG TO	CGGTAAAT TC	GTGCCAGC CA	CCGCGGGTT
P. polyuranodon												
Indo-China	GG		· · · · ·	•	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	•		••••••
P. polyuranodon												
East Kalimantan	· · · ·	· · · · ·	· · · ·	•••••••••••••••••••••••••••••••••••••••	•	· · · ·	· · · ·	· · · ·	•	•	•	•
P. rheophilus	· · · ·	· · · · ·	· · · ·	•••••••••••••••••••••••••••••••••••••••	•	•	· · · ·	· · · ·	.T	•	•	•
P. djambal	•	· · · · · ·	· · · ·	•	· · · ·		· · · ·	A	· · · ·	•	•	
P. bocourti			· · · · ·	•	•	c	· · · ·	A	· · ·	•	•	
P. nasutus			· · · · ·	•	•	· · · ·			· · · ·	•	•	
P. conchophilus		· · · · · ·	· · · · ·	•	•	G	· · · ·	· · · ·	· · · ·	· · · ·	•	•
P. pangasius		T	· · · ·	•••••••••••••••••••••••••••••••••••••••	•	· · · ·	· · · ·	· · · ·	•	•	•	•
P. kunyit Indonesia		T	· · · ·	•	· · · ·	· · · ·	· · ·	· · · ·	· · · ·	•	•	••••••
P. kunyit Vietnam		T	· · · · ·	•	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	•	•
P. kunyit North Borneo		T	G	•		· · · ·	· · · ·	· · · ·	· · · ·	•	•	
P. sanitwongsei	••••••	•••••••••••••••••••••••••••••••••••••••	c.	••••••		A	· · ·	· · · ·	· · · ·		•	
P. larnaudii		· · · · · ·	T	•	· · · ·	· · · ·	· · · ·	Γ Τ	· · · ·	•	•	
P. humeralis		· · · · ·	· · · ·	•••••••••••••••••••••••••••••••••••••••	•	•	· · · ·	· · · ·	•	•	•	•
P. nieuwenhuisii		· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·	••••••		· · · ·	· · ·	· · · ·	· · · ·		•	
P. kinabatanganensis		· · · · ·	· · · ·	••••••	· · · ·	· · · ·	· · · ·	TA	· · · ·	•	· · · ·	
P. lithostoma		· · · · ·	· · · ·	•••••••••••••••••••••••••••••••••••••••	•	· · · ·	· · · ·	· · · ·	C.	•	•	•
P. macronema	· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·	T	••••••		· · · ·	· · · ·	· · · ·	· · · ·		· · ·	
P. micronemus		· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·	••••••	T	· · · · · · · · · · · · · · · · · · · ·	· · · · ·	· · · · ·	· · ·		•	
P. pleurotaenia		· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·	••••••	T		· · · ·	T	· · ·		· · ·	
P. hypophthalmus		· · · · · · · · · · · · · · · · · · ·	· · · ·	••••••	•	· · · ·	· · · ·	TA T	· · · ·		· · ·	
P. gigas		T	· · · · · · · · · · · · · · · · · · ·	••••••		c .	. g	TT	· · ·		•	
H. leptorhynchus	••••••	T	c.			•		c	• • • • • • • • • •		•	
H. waandersii		T	c.	•••••••••••••••••••••••••••••••••••••••	•	· · · ·	· · · ·	. A C	· · · ·	•	•	•
H. typus	•	G	· · · ·	•	•	. T	· · ·	· · · ·	•	•	•	•
S. mandibularis	Τ	· · · · ·	· · · ·	•••••••••••••••••••••••••••••••••••••••	T /	AGC .	c	Τ	•	•	•	•
L. sinensis		· · · · ·	· · · · · · · · · · · · · · · · · · ·	•••••••••••••••••••••••••••••••••••••••	•	· · · ·	· · · ·	AT T	.AT C.	•	•	•
C. garua		T	T	••••••	T	· · · ·	· · · ·	AT T	. AC	•	•	•
E. vacha	••••••	•••••••••••••••••••••••••••••••••••••••	c		T	c .		. A A T	.ATT C.		•	
S. silondia		· · · · ·	c	•	T	c	· · · ·	AT T	. ATT	•	•	•
L. hexanema		· · · · ·	TT	••••••	T	· · · ·	· · · ·	AT T	.AT	•	•	T
P. brachypopterus			T	•••••••••••••••••••••••••••••••••••••••	•	. T	•	T T	· · · ·	•	•	• • • • • • •

Table 2. Sequence alignment (12S rDNA gene) between Pangasiidae and Schilbeidae.

: polyuranodon Rest											
of Indonesia	ATACGAAAGA	CCCTAGTTGA	TAGCTACGGC GTAAAGGGTG	GTTA-AG-GG AC	CATA-AA AA	ATAAGCTA A	AGATCCTCT /	AAGCCGTCAT /	CGCACTCCG	GGACACGAA ACCCCAACA	C GAAGTAGCT
. polyuranodon											
Indo-China				TGT	G		•		•	G	
. polyuranodon											
East Kalimantan				Γ	C	· · · ·	· · · ·		· · · ·		· · · ·
. rheophilus			A	Τ.Α	•	· · · ·	C		· · · ·	T G	
? djambal					A	G	CT	•	· · · ·		
? bocourti				· · · · ·	A	· · · ·	CT		· · · ·		
? nasutus			· · · · ·	T. A	AT	· · · ·	C		· · · ·		
?. conchophilus				A	A		C	•	· · · ·	T G	
?. pangasius			G	Τ	•		C	•	T	G	•
?. kunyit Indonesia	•				•		C	•	•	GG	•
. kunyit Vietnam					•		C	•		G	
. kunyit North Borneo					•		C	•	•	G G	•
. sanitwongsei				· · · ·		· · ·	C.		C.	T	
? larnaudii				A	•	T	C	•		GTT	
?. humeralis				T. A		· · · ·	•	T	· · · ·	G	
? nieuwenhuisii	· · · · · · · · · · · · · · · · · · ·			T. A		•		T			
. kinabatanganensis	· · · ·			Γ	•••••••••••••••••••••••••••••••••••••••		•		•	· · · · ·	· · · ·
? lithostoma		с.			G.		C		•	GT	
). macronema				Γ	G	· · ·	•	•	•	G	
. micronemus	· · · ·	C	AA	· · · · · · · · · · · · · · · · · · ·	A		C	· · · ·	•	T	· · · ·
?. pleurotaenia			A	A	A		•		· · · ·		
?. hypophthalmus		с	A	Γ			CT		· · · ·	AG TG	
? gigas			A	A	L	L	CT	•	· · · ·	AG TG	
I. leptorhynchus	•	C	c		•		C		•	GTT	•
I. waandersii		c.	c	· · · ·		T	C.		•	GTT	
I. typus		c			•		C	•		G	
. mandibularis			c	AT GT	c.ac.	· · ·	C.	•	T.		
. sinensis	•		c	A. AA.			C.T	T	T	AG GA	
garua			c	GA	A.AAAG.T		C	T	ATA 0	iGA	
. vacha		A	c	TA	••••••••••••••••••••••••••••••••••••••			гтт	.AT	CA	
. silondia			c	GA	••••C•AC•••		c	T	.AT	·····A····	
hexanema	•••••••••••••••••••••••••••••••••••••••	· · · ·	c	.GTGA .A			C.T	T	.AT	AA	· · · ·
brachypopterus			TAC	G.A	C.TT	T		T	C.	GT.T G.T.A	

P. polyuranodon Rest											
of Indonesia	TTAAATAAAA	TTAAACCTGA CCC	CACGAAA GCT	AAGAAAC AAACT	IGGGAT T	AGATACCCC	ACTATGCTTA	GCCTTAAACC	TAGATGTAAT	ATTACATATA	CATCCGCCCG
P. polyuranodon											
Indo-China	c	· · · ·	· · · ·	· · · ·	•		$\ldots T \ldots T$		· · · · · · · · · · · · · · · · · · ·	· · · ·	
P. polyuranodon											
East Kalimantan	C	.c	· · · ·	· · · ·	•	· · ·			A	GG	
P. rheophilus	$\dots \dots T$.ACTT.	· · · ·	· · · ·	•			c.	T.	C	
P. djambal		· · · · · · · · · · · · · · · · · · ·	•	•	•			c.	TA	TC.	
P. bocourti		· · · · · · · · · · · · · · · · · · ·	· · · ·	· · · ·	•			c.	TA	$T \ldots CG$	
P. nasutus		· · · · ·	· · · ·		•			c.	TC	cc.c.	
P. conchophilus			•		•			c	TA	cc.c.	
P. pangasius	$\dots \dots T$	· · · · ·	· · · ·	· · · ·	•			A	T.	cc.c.	
P. kunyit Indonesia	$\dots \dots \dots$	· · · · ·	· · · ·	· · · ·	•			c.	c.	cc.c.	
P. kunyit Vietnam	T	· · · ·	•	· · · ·	•			c.	C.	cc.c.	
P. kunyit North Borneo	$\dots \dots T$	· · · ·	· · · ·	· · · ·	•			c	T.	cc.c.	
P. sanitwongsei	c	· · · ·	· · · ·	· · · ·	•			A	TC	cc.	
P. larnaudii		· · · ·	•	· · · ·	•			c.	T.	cc.c.	
P. humeralis	C.G-T	T.	· · · ·	· · · ·	•			c.	c.	C	
P. nieuwenhuisii	G-T	AT.	· · · ·	· · · ·	•			c.	C.	C	
P. kinabatanganensis	9		•	· · · ·	•			с.	T.	TC.C.	
P. lithostoma	T	- CTT	· · · ·	· · · · · · · · · · · · · · · · · · ·	•			c.	TC	cc.c.	
P. macronema	C	· · · · ·	· · · ·	· · · ·	•						
P. micronemus	T	··· · · · · · · · · · · · · · · ·	•	•	•				TC	$T \dots A \dots A$	
P. pleurotaenia	$\ldots c \ldots T$	T	· · · ·	· · · ·	•				TC	$T\ldots C.A.$	
P. hypophthalmus	c	C	· · · ·	· · · ·	•				C.	TC.C.	T.
P. gigas	$\ldots C \ldots T$.GTC		· · · · · · · · · · · · · · · · · · ·					cc	$T\ldots C.C.$	•••••••••••••••••••••••••••••••••••••••
H. leptorhynchus	$\ldots \ldots T$	AGGC	· · · ·	· · · ·	•			c	T.	cc.c.	
H. waandersii	T	AGGC						c	T.	cc.c.	•••••••••••••••••••••••••••••••••••••••

ed
inu
nti
<u>)</u>
4
e
q
\mathbf{Ia}

TCCGCCCG GGTACTACGA GC-ACAGCTT

 T....

......

..... T.... ...c....

.....A.

.....T.

 $\ldots T \ldots \ldots$

 $\ldots \ldots \ldots T \ldots$

T.....C... TG.....T.

....CA..TC

.....c......

.

с.т.с.-...

.....c....

.....G...G.

P. brachypopterus

L. hexanema

S. silondia E. vacha C. garua

.....c....

..c.c.-...

...T.C.-...

 $c.c.c.\ldots \ .T\ldots \ldots$

.....c.....

.....T

.....C.... AAC.C.-...

....CA..T. $\dots \dots T$

....CA..CA

T..... TG.....

T.C..... TG.....T.

••••••

TG.....

Τ.....

....CA..TC

.....c.....

.....A. TG.....

T....C.C.

....CA..TC

.....c.....

...T-..T..... ...T.CT-.... A..........

....T

· · · ·

S. mandibularis

H. typus

L. sinensis

.....C...G.G...G.

.....c.....

......

· · · ·

C.T.C.....T.....

cc....ccA. Τ.....

 \ldots^{T}

P. polyuranodon Rest													
of Indonesia	AAACCCAAA	GGACTTGGCG GTG	TCTCAGA CC	CACCTAGA (3GAGCCTGTT	CTAGAACCGA	TAACCCCCGT	TAAACCTCAC	CACTTCTTGT	I-TTCCCCGC (CTATATACCG C	CGTCGCCAG CT	TACCCTGT
P. polyuranodon													
Indo-China			· · ·	•		•				· · · ·	· · · ·	· · · ·	
P. polyuranodon													
East Kalimantan		•	:	•		•			c.	· · · ·	· · · ·	· · · ·	
P. rheophilus			· · ·			•				· · · ·	· · · ·	T	
P. djambal			• • • • • • • • •	•		•	•			с	· · · ·	T	
P. bocourti			· · ·			•				· · · ·	· · · ·	T	
P. nasutus		•	:	•		•				· · · ·	•	T	
P. conchophilus		•	:	•		•	•••••••••••••••••••••••••••••••••••••••			· · · ·	•	T	
P. pangasius		•	:	•		•				· · · ·	•	T	
P. kunyit Indonesia	· · · ·	•••••••••••••••••••••••••••••••••••••••	· · ·	•		•	•••••••••••••••••••••••••••••••••••••••	· · · · · · · · · · · · · · · · · · ·	•••••••••••••••••••••••••••••••••••••••	· · · ·	•	T	
P. kunyit Vietnam		•••••••••••••••••••••••••••••••••••••••		•		•				· · · ·	•	T	
P. kunyit North Borneo		•••••••••••••••••••••••••••••••••••••••	· · ·	•							•	T	
P. sanitwongsei		•••••••••••••••••••••••••••••••••••••••	· · ·	•	· · · ·	•	· · · ·		· · · ·	· · · ·	•	T	
P. larnaudii	· · · ·	•••••••••••••••••••••••••••••••••••••••		•		•				· · · ·	•	T	
P. humeralis		•••••••••••••••••••••••••••••••••••••••	· · ·	•							•	T	
P. nieuwenhuisii		•••••••••••••••••••••••••••••••••••••••	· · ·	•	· · · ·	•	· · · ·		· · · ·	· · · ·	•	T	
P. kinabatanganensis		•	· · ·	•)	c	•		•	•	T	
P. lithostoma	Τ	· · · · · · · · · · · · · · · · · · ·	· · · ·	•		•••••••••••••••••••••••••••••••••••••••					• • • • • • • • • • • •	T	
P. macronema		•••••••••••••••••••••••••••••••••••••••	· · · ·	•		•••••••••••••••••••••••••••••••••••••••					•	· · · · · · · · · · · · · · · · · · ·	
P. micronemus		· · · · · · · · · · · · · · · · · · ·	· · · ·	•							•	T	
P. pleurotaenia		•••••••••••••••••••••••••••••••••••••••	· · ·	•					c.		•	T	
P. hypophthalmus		· · · · ·	•	•	· · · ·					•	· · · ·	T	
P. gigas	· · · ·	· · · · ·	· · ·	•		•••••••••••••••••••••••••••••••••••••••	· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·		••••••	· · · · · · · · · · · · · · · · · · · ·	T	· · · ·
H. leptorhynchus			· · · ·	•		••••••				T	•	T	
H. waandersii	• • • • • • • • • •	· · · · · · · · · · · · · · · · · · ·	· · ·	•		•••••	• • • • • • • • • •	••••••		T	• • • • • • • • • •	T	· · · ·
H. typus		· · · · · · · · · · · · · · · · · · ·	· · · ·	•		••••••		••••••			• • • • • • • • • • • • •	T	· · · ·
S. mandibularis		•••••••••••••••••••••••••••••••••••••••	· · ·	•					c.		•	T	A.
L. sinensis		•••••••••••••••••••••••••••••••••••••••	· · ·	•	· · · ·	•	· · · ·		.c	T	•	T	
C. garua	• • • • • • • • • • •	· · · · · · · · · · · · · · · · · · ·	· · · ·	•		••••••	• • • • • • • • • •	••••••	.c	.TT	· · · · · · · · · · · · · · · · · · · ·	T	· · · ·
E. vacha	• • • • • • • • • •	,		•		•••••	T	••••••	.c	.TT.	• • • • • • • • • •	T	· · · ·
S. silondia	• • • • • • • • • •	· · · · · · · · · · · · · · · · · · ·	· · · ·	•		•••••	T	••••••	.c	.TT.	• • • • • • • • • •	T	· · · ·
L. hexanema	••••••	· · · · · · · · · · · · · · · · · · ·	· · · ·	•		••••••		••••••	.c	T	• • • • • • • • • • • • •	T	
P. brachypopterus		•	:	•		•				. T	· · · ·	T	

-
Ð
<u>د</u>
_
ti
0
್
\sim
4
e
-
P
<u>_</u> 0

P. polyuranodon Rest												
of Indonesia	GAAGGCCT - A	GCAGTAAGCA A.	AATGGGCTC G(CCCAAAAAC G	TCAGGTCGA C	GTGTAGCGC A	CGAAGTGGG A	AGAAATGGG C	TACATTTTC 1	GCA-GC-AG A	ATACT-ACG GAC-GGCACC CTG	AAACA-A
P. polyuranodon												
Indo-China			•	•	•	•	•	•	· · · ·	A	$\ldots T \ldots A \ldots \ldots \ldots \ldots \ldots \ldots$	
P. polyuranodon												
East Kalimantan	T	AG	•	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	AA	T A	
P. rheophilus			C.	· · · ·		· · · ·	· · · ·	•		AGAG	$\ldots T \ldots A.T \ldots T \ldots T \ldots$	T
P. djambal	A	· · · · ·	•	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	AA	$\ldots T \ldots A \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots$	T.T.
P. bocourti			•	· · · ·		· · · ·	· · · ·	· · · ·	· · · ·	AA	T A	T.T.
P. nasutus		A	c.				•	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • •	AA	$\ldots T \ldots A \cdot T \ldots \ldots$	T
P. conchophilus	· · · · · · · · · · · · · · · · · · ·	· · · · ·	c.	· · · ·		· · · ·	· · · ·	· · · ·	· · · ·	AAT	$\ldots T \ldots A . T \ldots \ldots$	
P. pangasius				· · · ·	· · · ·	· · · ·	· · · ·	•		AA	$\ldots T \ldots \ldots A \ldots \ldots T \ldots \ldots T$	T
P. kunyit Indonesia			c.	· · · ·		· · · ·	· · · ·	· · · ·	· · · ·	AA	T A	
P. kunyit Vietnam		· · · · ·	C. A.	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	ATA	$\ldots T \ldots A \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots$	· · ·
P. kunyit North Borneo			c.	· · · ·	•	· · · ·	G	· · · ·	· · · ·	AA	T A	
P. sanitwongsei			.GA.	•	•	•	•	•	•	AA	$\ldots T \ldots A . T \ldots T \ldots T$	T
P. larnaudii		. T	c.	· · · ·	•	· · · ·		· · · ·	· · · ·	AA	T A.TTG.T	· · ·
P. humeralis	C	A	•	· · · ·	•	· · ·		•	· · · ·	AA	T A	
P. nieuwenhuisii	C	A	•	· · · ·	•	· · ·		•	· · · ·	AA	T A	
P. kinabatanganensis	T.	A	•	· · · ·	•	· · · ·		· · · ·	· · · ·	AA	$.c.T\ldotsA\ldots\ldotsT\ldots$	T
P. lithostoma	CT.	A	C. A.	· · · ·	•	· · ·	A	•	· · · ·	AA	$\ldots T \ldots A . T \ldots T A \ldots T A$	$\ldots T \ldots$
P. macronema			•	•	•	•	•	•	· · · ·	A	$\ldots T \ldots A \ldots \ldots \ldots \ldots \ldots \ldots$	
P. micronemus	$\ldots T \ldots \ldots$	AT	c.	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	AA	$\ldots T . T . T . \dots A . T . \dots \dots \dots \dots$	T.T.
P. pleurotaenia	T.A	AT	C.	•	•	•	•	•	•	AA	$\ldots T . T . T \dots A . T \dots \dots \dots$	T.T.
P. hypophthalmus				c	•	•	•	•	•	AAT	$\ldots T \ldots A . T \ldots \ldots \ldots \ldots \ldots \ldots \ldots$	
P. gigas	$\ldots T.C\ldots$	A	· · · · · · · · · · · · · · ·	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	AAT	$\ldots T \ldots A . T \ldots \ldots \ldots \ldots \ldots \ldots \ldots$	
H. leptorhynchus	· · · · · · · · · · · · · · · · · · ·	. T	TC.	•		· · ·	· · · ·	· · ·	· · · ·	AAT	$\ldots T \ldots A . T \ldots \ldots \ldots \ldots \ldots \ldots \ldots$	
H. waandersii	••••••	$. T \dots \dots$	Tc.	• • • • • • • • • •			•	•	•	AAT	$\ldots T \ldots A . T \ldots \ldots \ldots$	
H. typus	· · · ·	AT	c.	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	· · · ·	ATA	$\ldots T \ldots A . T \ldots \ldots \ldots \ldots \ldots \ldots \ldots$	T
S. mandibularis	\dots AT											
L. sinensis												
C. garua												
E. vacha	T.											
S. silondia												
L. hexanema												
P. brachypopterus												

P. polyuranodon Rest									
of Indonesia	GTGCCCGAAG	GTGGATTTAG	TAGTAAAAAG	CAAATAGAGA	GTCCTTTTGA	ATTAGGCTCT	GAGACGCGCA	CACACCGCCC	GTCACTC
P. polyuranodon									
Indo-China	· · · ·	· · · ·	· · · · · · · · · · · · · · · · · · ·		· · · ·	· · · ·	· · · ·		
P. polyuranodon									
East Kalimantan	T								
P. rheophilus	T			C				· · · ·	
P. djambal				C		· · · ·		· · · ·	
P. bocourti				C					
P. nasutus	$A \dots T \dots$			C	T			· · · ·	
P. conchophilus	$A \dots T \dots$			· · · ·		· · · ·		· · · ·	
P. pangasius	T.T			C					
P. kunyit Indonesia	T			C				· · · ·	
P. kunyit Vietnam	T			C	· · · ·	· · · ·		· · · ·	
P. kunyit North Borneo	T			C	· · · ·	· · · ·		· · · ·	
P. sanitwongsei	T			C		•••••••••••••••••••••••••••••••••••••••		•••••••••••••••••••••••••••••••••••••••	
P. larnaudii	AT			C					
P. humeralis			C	rc					
P. nieuwenhuisii			c	c		· · · ·		· · · ·	
P. kinabatanganensis	T			T					
P. lithostoma	T			C		•••••••••••••••••••••••••••••••••••••••		• • • • • • • • •	
P. macronema									
P. micronemus	$A \dots T \dots$			· · · ·				· · · ·	
P. pleurotaenia	$A \dots T \dots$			•		•			
P. hypophthalmus	$A \dots T \dots$			T					
P. gigas	T			T					
H. leptorhynchus	$A \dots T \dots$			C		· · · ·		· · · ·	
H. waandersii	$A \dots T \dots$			C					
H. typus	$A \dots T \dots T$			C		· · · · · · · · · · · · · · · · · · ·		•••••••••••••••••••••••••••••••••••••••	

(Sumatra and Kalimantan), North Borneo (Sabah), and Indo-China (Vietnam). Within *P. kunyit*, the average genetic distances between populations from North Borneo and groups from Indonesia (d = 0.007) and Indo-China (d = 0.0097) or among the two latter (d =0.003) correspond respectively to those estimated between *P. macronema* and *P. polyuranodon* in Vietnam (d = 0.004), *P. djambal* and *P. bocourti* (d =0.0055), *Helicophagus waandersii* and *H. leptorhynchus* (d = 0.004).

Putative new *Pangasius* species inside *P. polyura-nodon* and *P. kunyit* will be examined in further studies, but for the present contribution, they will be considered as genetically distant populations. They will be respectively labelled as POL 1 (populations of *P. polyuranodon* from Indonesia except East Kalimantan), POL 2 (populations of East Kalimantan), POL 3 (populations of Indo-China: Mekong and Chao Phraya Rivers), KUN 1 (populations of *P. kunyit* from Indonesia), KUN 2 (populations of Indo-China), KUN 3 (populations of North Borneo).

Finally, in a second step, corrected Kimura's genetic distances (below diagonal) and absolute substitution/insertion/deletion number (above diagonal) are presented for each possible couple of species (Table 3). Pangasiid species are well differentiated with genetic distance comprised between 0.0042 and 0.0561. The most closely related species are *H. waandersii* and *H. leptorhynchus* (d = 0.0042), *P. djambal* and *P. bocourti* (d = 0.0055), *P. nieuwenhuisii* and *P. humeralis* (d = 0.0056), and the most differentiated are *P. gigas* and *P. rheophilus* (d = 0.0561).

A significant linear correlation between Kimura's distances and the number of transitions and transversions per site suggested that few multiple substitutions have occurred at most nucleotide positions and are far from being saturated. This indicated that interspecific genetic distances among pangasiids are likely within the range to be phylogenetically instructive (Lydeard and Roe 1997).

The nearest neighbour dendrogram was inferred from the corrected Kimura's genetic distance pairwise matrix among taxa (Fig. 1). Four monophyletic groups were validated by the bootstrapping tests: group 1 (bootstrap 84%) including *Pangasianodon hypophthalmus* and *P. gigas*; group 2 (bootstrap 92%) composed of *Pteropangasius micronemus* and *P. pleurotaenia*; group 3 (bootstrap 80%) containing *Helicophagus waandersii*, *H. leptorhynchus*, *H. typus*; and group 4 (bootstrap 100%) with *P. humeralis*, *P. nieuwenhuisii*, *P. polyuranodon*, *P. macronema*. Besides groups 3 and 4, seven highly differentiated species are branched in the internal part of the tree, namely P. lithostoma, P. nasutus, P. larnaudii, P. sanitwongsei, P. rheophilus, P. conchophilus, and P. kinabatanganensis. In this part of the tree, the poor resolution of branching order of most internal nodes does not allow resolution of the phylogeny, although some pairs of related species could be clearly evidenced, for example P. djambal and P. bocourti (bootstrap 100%), P. kunyit and P. pangasius (bootstrap 50%). This lack of resolution in the Pangasius (Pangasius) subgenus was also noted for allozymes and cytochrome b phylogenies (Pouyaud et al. 2000) and was interpreted to be an expected signature of an intense radiation in a short period, too brief to allow for the accumulation of synapomorphies (Johns and Avise 1998; Martin and Bermingham 1998). Overall, the topology of the 12S rDNA phylogeny is similar to those obtained with allozymes and cytochrome b (Pouyaud et al. 2000).

Genetic Interrelationships between Pangasiidae and Schilbeidae

Five hundred and twenty seven nucleotides encoding the first half of the 12S rDNA gene were aligned between seven schilbeids and all the pangasiids (Table 2). Hundred and thirthy one sites (25%) were polymorphic. Corrected Kimura's genetic distances (below diagonal) and transition/transversion ratios (above diagonal) were calculated between the schilbeids and five pangasiids (H. waandersii, Pangasius nasutus, P. polyuranodon, Pteropangasius pleurotaenia and Pangasianodon hypophthalmus) representing the hypothesised genera (Table 4). The nearest neighbour dendrogram was inferred from the corrected Kimura's genetic distance pairwise matrix among species (Fig. 2). The topology of the resulting tree separates Schilbeidae from Pangasiidae.

The same results were obtained when including all the pangasiid species, but the corresponding tree is not given. Schilbeidae are characterised by three synapomorphic substitutions (respectively in nucleotide positions 90, 275, and 277). However, the monophyly of Schilbeidae is not supported by high bootstrap value (approximately 50%) due to the existence of three distinct lineages separated by huge genetic divergence. As an example, the low transition/transversion ratios calculated for the 12S rDNA sequences of *Schilbe mandibularis* and *Pseudeutropius brachypopterus* confirm that nucleotidic substitutions are saturated. Nevertheless, a monophyletic group is supported by a strong bootstrap value (100%) in the schilbeids. This highly

substitution/insertion/deletion for each	
er of	
qunu	
absolute	
and	
diagonal)	
(below	
Pangasiidae	
between	
distances	
genetic	diagona
Kimura's	above (
Corrected	comparison
Table 3.	pairwise

Species	pol.1	pol.2	pol.3	rhe	dja	boc	nas	con	pan	kun.1	kun.2	kun.3	san
P. polyuranodon Rest of Indonesia (1)		10	20	30	22	20	26	24	21	17	19	21	25
P. polyuranodon Indo-China (2)	0.0126		22	29	25	23	26	24	22	20	22	22	28
P. polyuranodon East Borneo (3)	0.0253	0.0253		38	33	31	31	33	31	27	29	10	35
P. rheophilus	0.0413	0.0384	0.0502		31	29	23	18	20	22	23	23	27
P. djambal	0.0268	0.0326	0.0428	0.0384		4	22	23	23	19	21	22	23
P. bocourti	0.0240	0.0297	0.0199	0.0155	0.0055		20	21	21	17	19	20	21
P. nasutus	0.0340	0.0354	0.0411	0.0283	0.0268	0.0239		11	17	15	17	18	20
P. conchophilus	0.0311	0.0325	0.0442	0.0240	0.0282	0.0253	0.0182		19	15	17	16	23
P. pangasius	0.0326	0.0297	0.0413	0.0268	0.0282	0.0254	0.0211	0.0239		10	12	11	19
P. kunyit Indonesia (1)	0.0239	0.0268	0.0355	0.0297	0.0225	0.0197	0.0182	0.0182	0.0140		2	5	19
P. kunyit Vietnam (2)	0.0268	0.0296	0.0384	0.0311	0.0253	0.0225	0.0211	0.0211	0.0168	0.0028		L	19
P. kunyit North Borneo (3)	0.0297	0.0296	0.0413	0.0311	0.0268	0.0239	0.0225	0.0197	0.0154	0.0070	0.0097		22
P. sanitwongsei	0.0341	0.0370	0.0458	0.0356	0.0269	0.0240	0.0268	0.0312	0.0253	0.0254	0.0254	0.0297	
P. larnaudii	0.0369	0.0398	0.0472	0.0312	0.0340	0.0311	0.0251	0.0254	0.0268	0.0182	0.0211	0.0225	0.0297
P. humeralis	0.0297	0.0268	0.0398	0.0326	0.0355	0.0326	0.0297	0.0297	0.0311	0.0251	0.0282	0.0296	0.0428
P. nieuwenhuisii	0.0297	0.0268	0.0370	0.0326	0.0355	0.0326	0.0297	0.0297	0.0311	0.0253	0.0282	0.0296	0.0428
P. kinabatanganensis	0.0311	0.0326	0.0384	0.0341	0.0282	0.0254	0.0254	0.0297	0.0253	0.0253	0.0282	0.0296	0.0326
P. lithostoma	0.0503	0.0533	0.0563	0.0460	0.0415	0.0386	0.0298	0.0386	0.0370	0.0298	0.0298	0.0341	0.0342
P. macronema	0.0084	0.0042	0.0210	0.0340	0.0282	0.0254	0.0111	0.0282	0.0254	0.0225	0.0253	0.0253	0.0326
P. micronemus	0.0385	0.0428	0.0444	0.0371	0.0370	0.0341	0.0255	0.0269	0.0370	0.0326	0.0355	0.0370	0.0371
P. pleurotaenia	0.0199	0.0442	0.0458	0.0415	0.0384	0.0355	0.0269	0.0312	0.0384	0.0341	0.0370	0.0384	0.0400
P. hypophthalmus	0.0443	0.0428	0.0517	0.0458	0.0428	0.0399	0.0384	0.0370	0.0427	0.0325	0.0354	0.0397	0.0414
P. gigas	0.0547	0.0576	0.0592	0.0561	0.0472	0.0443	0.0458	0.0458	0.0516	0.0384	0.0413	0.0457	0.0503
H. leptorhynchus	0.0502	0.0532	0.0562	0.0487	0.0458	0.0428	0.0370	0.0370	0.0384	0.0268	0.0268	0.0311	0.0414
H. waandersii	0.0517	0.0546	0.0577	0.0501	0.0472	0.0443	0.0355	0.0385	0.0398	0.0282	0.0282	0.0326	0.0429
H. typus	0.0370	0.0369	0.0443	0.0313	0.0327	0.0298	0.0226	0.0255	0.0254	0.0211	0.0211	0.0239	0.0356
	din anti-	ir a -		190000	-			-					

pol = Pangasius polyuranodon, the = P. theophilus, dja = P. djambal, boc = P. bocourti, nas = P. nasulus, con = P. conchophilus, pan = P. pangasius, kun = P. kunyit, san = P. sanitwongsei, lar = P. larnaudii, hum = P. humeralis, nie = P. nieuwenhuisii, kin = P. kinabatanganensis, lit = P. lithostoma, mac = P. macronema, mic = P. micronemus, ple = P. pleurotaenia, hyp = P. hypophthalmus, gig = P. gigas, lep = Helicophagus leptorhynchus, waa = H. waandersii, typ = H. typus

Species	lar	hum	nie	kin	lit	mac	mic	pie	hyp	sig Bi	lep	waa	typ
P. polyuranodon Indonesia (1)	27	22	22	24	38	7	32	32	32	38	35	36	27
P. polyuranodon Indochina (2)	30	21	21	26	38	3	34	33	32	41	38	39	28
P. polyuranodon East Borneo (3)	36	31	29	31	43	16	36	36	39	43	41	42	34
P. rheophilus	24	25	25	27	35	26	32	34	34	39	35	36	24
P. djambal	28	29	29	25	34	22	30	30	33	36	35	36	27
P. bocourti	26	27	27	23	32	20	28	28	32	34	33	34	25
P. nasutus	19	24	24	22	26	23	23	23	30	34	28	27	19
P. conchophilus	19	24	24	25	31	21	24	26	28	33	28	27	21
P. pangasius	20	23	23	20	28	19	31	33	31	35	27	28	19
P. kunyit Indonesia (1)	14	19	18	20	23	17	27	28	24	27	19	20	16
P. kunyit Vietnam (2)	16	21	21	22	23	19	30	30	26	29	19	20	16
P. kunyit North Borneo (3)	17	22	22	23	26	19	31	31	29	32	22	23	18
P. sanitwongsei	21	32	32	26	27	25	32	33	39	36	30	31	27
P. larnaudii		29	29	25	30	27	32	30	29	33	23	26	23
P. humeralis	0.0383		4	20	35	17	34	31	34	35	43	45	26
P. nieuwenhuisii	0.0384	0.0056		20	35	18	35	33	36	37	36	38	27
P. kinabatanganensis	0.0311	0.0268	0.0268		32	23	35	31	31	37	37	38	26
P. lithostoma	0.0385	0.0473	0.0474	0.0429		38	35	35	37	36	33	34	27
P. macronema	0.0355	0.0255	0.0225	0.0283	0.0488		32	31	29	38	35	36	25
P. micronemus	0.0371	0.0414	0.0414	0.0400	0.0401	0.0384		11	33	34	35	35	23
P. pleurotaenia	0.0356	0.0385	0.0414	0.0356	0.0415	0.0398	0.0140		32	28	36	38	24
P. hypophthalmus	0.0384	0.0457	0.0486	0.0398	0.0502	0.0384	0.0399	0.0413		23	33	34	28
P. gigas	0.0458	0.0487	0.0517	0.0502	0.0503	0.0532	0.0415	0.0341	0.0312		37	38	34
H. leptorhynchus	0.0312	0.0531	0.0532	0.0502	0.0429	0.0487	0.0430	0.0474	0.0458	0.0531		3	22
H. waandersii	0.0326	0.0546	0.0546	0.0516	0.0444	0.0501	0.0444	0.0488	0.0472	0.0545	0.0042		23
H. typus	0.0312	0.0355	0.0355	0.0326	0.0327	0.0326	0.0269	0.0298	0.0398	0.0472	0.0298	0.0312	
Pol = Pangasius polyuranodon, the = P. the	eophilus, dja	= P. djamba	$ll, boc = P. b_l$	ocourti, nas :	= P. nasutus	con = P. con	nchophilus,]	pan = P. pang	asius, kun =	P. kunyit, sar	I = P. sanitwo	ngsei, lar = H	larnaudii,
hum = P. <i>humeralis</i> , nie = P. <i>nieuwenhuisi</i>	ii, kin = P. ki	nabatangane	ensis, $lit = P$.	lithostoma,	mac = P. ma	<i>icronema</i> , m	ic = P. micro	nemus, ple =	P. pleurotae	nia, hyp = P .	hypophthaln	uus, gig = P. g	igas, lep =
Helicophagus leptorhynchus, waa = H . wa	aandersii, ty	p = H. typus	8										

Table 3. (continued).



Fig. 1. The nearest neighbour dendrogam for 12S rDNA gene within Pangasiidae. Numbers next to node show percentage group occurrence found with the 1000 bootstrapped trees.

differentiated group, characterised by three synapomorphic substitutions (positions 245, 482, 488) and assembling *Laides* with *Clupisoma*, *Eutropiichthys* and *Silonia* clearly demonstrates that the genus *Laides* cannot be included in the Pangasiidae. Moreover, when considering the average genetic distances within Pangasiidae as a reference (Table 4), Kimura's genetic distances computed within Schilbeidae suggest the possible occurrence of several distinct families. Actually, equivalent genetic divergences are found between the two lineage of Asian schilbeids (0.0863 < d_{Kim} < 0.1040 between *P. brachypopterus* and others) or between both lineage of Asian schilbeids and lineage of African schilbeids (d_{Kim} = 0.1028 and 0.0980 < d_{Kim} < 0.1167, respectively).

Evolutionary Process Based on Speciation Peaks

The distribution pattern of pairwise Kimura-corrected genetic distance on the 12S rDNA gene was studied for all Pangasiidae and Schilbeidae species (Fig. 3). Three distinct modes were observed, corresponding respectively to interfamily, intergenus, and intragenus assignation of pairwise comparisons. The first mode, distributed between 0.0575 and 0.1025, corresponds to the most ancient radiation between Schilbeidae and Pangasiidae. The second mode, distributed between the values of 0.0175 and 0.065, corresponds to the radiation between the genera of Pangasiidae and among some genera of the Schilbeidae, except

Pseudeutropius and *Schilbe*. The radiation of *Pseudeutropius* and *Schilbe* is more ancient (between 0.0875 and 0.1175) and overlaps with the interfamily radiation peak (right part) suggesting again that both genera probably belong to distinct families. Finally, the third peak, between 0.001 and 0.05, corresponds especially to the radiation within the different genera of both families. The right part of this peak, largely superposed with the intergenus mode between 0.03 and 0.05 mainly concerns an earlier radiation within *Pteropangasius*, *Helicophagus*, *Pangasianodon*, and *Clupisoma*. Except for *P. lithostoma*, *P. polyuranodon*, and *P. kinabatanganensis*, the radiation within *Pangasius* is more recent (between 0.002 and 0.03).

Systematic Position of the Genus Laides

Molecular analyses clearly showed that Pangasiidae and Schilbeidae are in two distinct families. They also reveal in the Schilbeidae, the existence of different lineages characterised by a large genetic divergence. The genus *Laides* forms a monophyletic assemblage with the genera *Eutropiichthys*, *Clupisoma* and *Silonia*, and this definitely indicates that it belongs to the Schilbeidae. The average genetic distances estimated among species in this group are equivalent to those observed in Pangasiidae and suggest that both distant taxa *Schilbe* and *Pseudeutropius* may belong to distinct families. The important genetic divergence noted in Schilbeidae is also enforced by their important morphological diversity and their

Species	pol	nas	ple	hyp	waa	man	sin.	gar	vac	sil	hex	bra
Pangasius (Pangasius) polyuranodon		9/3	10/6	16/4	17/7	20/16	25/9	25/10	25/15	26/10	25/11	24/10
Pangasius (Pangasius) nasutus	0.0236		11/5	16/2	14/5	20/13	25/6	26/8	28/12	26/7	28/8	20/10
Pangasius (Pteropangasius) pleurotaenia	0.0317	0.0296		17/3	21/7	21/15	25/10	25/12	24/16	22/11	26/12	23/14
Pangasius (Pangasianodon) hypophthalmus	0.0397	0.0377	0.0396		20/4	25/16	33/7	31/9	29/13	30/8	32/9	24/10
Helicophagus waandersii	0.0480	0.0377	0.0562	0.0499		29/19	38/10	27/14	30/18	29/13	32/14	22/12
Schilbe mandibularis	0.0739	0.0717	0.0716	0.0848	0.1005		30/17	31/19	32/23	28/17	35/17	28/20
Clupisoma sinensis	0.0693	0.0627	0.0693	0.0820	0.0867	0.0980		18/2	23/6	17/1	12/2	32/10
Clupisoma garua	0.0736	0.0692	0.0737	0.0821	0.0846	0.1049	0.0395		18/8	16/3	16/4	32/12
Eutropiichthys vacha	0.0827	0.0824	0.0805	0.0870	0.1006	0.1167	0.0583	0.0522		17/5	23/6	36/14
Silonia silondia	0.0735	0.0668	0.0650	0.0777	0.0866	0.0936	0.0355	0.0376	0.0437	1	18/1	34/9
Laides hexanema	0.0736	0.0733	0.0757	0.0842	0.0954	0.1114	0.0274	0.0396	0.0582	0.0375		34/10
Pseudotropius brachypopterus	0.0692	0.0629	0.0737	0.0715	0.0714	0.1028	0.0863	0.0905	0.1040	0.0880	0.0904	
nol = Paneasius (Paneasius) nalvuranadon na	s = Pangasi	ius (Pangas	ius) nasutu	s. $nle = P_{c}$	in pasius (P	teronangasi	us) pleurota	enia. hvn =	= Pangasius	(Pangasia	ouvh (nobor	phthalmus

pecies of Schilbeidae (below diagonal) and absolute number of substitution (transition/	
nd 7 s	
of Pangasiidae a	
5 species	agonal).
between	elow di
distances	oarison (b
genetic e	se comp
imura's	h pairwi
cted K	for eacl
. Corre	rsion) 1
ıble 4	ansvei

pol = rangastus (rangastus) potyuranoaon, nas = rangastus (rangastus) nasutus, pie = rangastus (rteropangastus) peurotaema, nyp = rangastanoaon) nypoprinatuus, waa = Helicophagus waandersii, man = Schilbe mandibularis, sin = Clupisoma sinensis, gar = Clupisoma garua, vac = Eutropiichthys vacha, sil = Silonia silondia, hex = Laides hexanema, bra = Pseudotropius brachypopterus.



Fig. 2. The nearest neighbour dendrogam from partial 12S rDNA gene of Pangasiidae and Schilbeidae. Numbers next to node show the percentage group occurrence found with the 1000 bootstrapped trees.



Fig. 3. Distribution of corrected Kimura's genetic distances at all positions respectively for intragenus, intergenus, and interfamily pairwise comparisons.

wide distribution through Asia and Africa, which suggests a long evolutionary process since the emergence of this family.

As demonstrated above, Pangasiidae and Schilbeidae probably stem from a common ancestor. At present, Schilbeidae mainly occurs in Africa and on the Indian subcontinent, while Pangasiidae are mainly found in Southeast Asia. The molecular phylogenies obtained from these groups always revealed important genetic differentiation. Referring to the abundant literature concerning approximate molecular clock/evolutionary time calibrations available on mitochondrial genes, an estimation of time divergence between both families or within them can be given below.

It is accepted that the overall rate of nucleotide substitutions for the complete mitochondrial genome may be 0.5-2% per million years depending on the various selective constraints occurring on the different gene families (Naylor et al. 1995; Zardoya and Meyer 1996; Kocher and Carlton 1997). The small ribosomal subunit (12S) gene is characterised by the lowest rate of evolution in the mitochondrial DNA (between 0.5 and 1% per million years) due to severe structural or/and functional constraints rather than a depressed mutation rate (Simon et al. 1994). By contrast, cytochrome b evolves faster (between 1 and 2% per million years) due to silent substitutions mainly occurring in third position of the codon (Meyer 1993). Pouyaud et al. (2000) estimated an average time of divergence between the Schilbeidae and Pangasiidae around 20 million years by using a rate of cytochrome b evolution proposed for marine fishes (1-1.2% sequence divergence per million years) (Bermingham et al. 1997). An equivalent estimation between 14 and 20 million years ago (mya) is estimated with 12s rDNA when using a rate of 0.5% per million years. Therefore, we consider that the separation between Schilbeidae and Pangasiidae occurred in the Early Miocene during the opening of the Red Sea and achievement of tectonic uplifts in Northern Thailand.

By using the same rate of evolution as above for 12S rDNA, the average time of divergence between the four pangasiids genera is \approx 7-11 mya. This estimation corresponds to the Middle and Late Miocene. More recently (\approx 5-8 mya), a new episode of speciation marked by an explosive radiation is observed in the genus Pangasius. Similar dating was estimated with other molecular markers (\approx 8-10 mya with cytochrome b and \approx 9 mya with allozymes) by Pouyaud et al. (2000). This adaptative radiation extended until the Late Pleistocene (1-4 mya, referring to the split between *Pteropangasius* species, *H*. waandersii and H. leptorhynchus, P. humeralis and P. nieuwenhuisii, P. polyuranodon and P. macronema, P. bocourti and P. djambal or different groups of P. kunyit).

Speciation Process

The divergence time assessed with molecular tools (Fig. 4) agrees with the hypothesis proposed by Vidthayanon (1993), mentioning that the dispersal of Pangasiidae cannot be merely explained by recent connections of river systems, but may be traced back to the early Tertiary or even earlier times when the present-day continental and insular sections constituted a land mass known as the "Indonesian-

Cathaysian land". Under these conditions, cumulative fluctuations of sea water levels coupled with important tectonic movements during millions of years undoubtedly have fashioned the pangasiids leading to the notable morphological and taxonomic diversity known at present (De Beaufort 1951; Taki 1978; Rainboth 1991).

The large river basins in Asian continent (the Mekong and Chao Phraya) and the island of western Indonesia (the Kapuas and Mahakam in Kalimantan), which possess the highest species diversity, have certainly played a major role in terms of refuge zones during the past (Roberts 1989; Kottelat 1995) when marine transgression occurred. Refugee zones represent sites with a peculiar flow regime and vegetation linking to the tropical forest. This resulted in a vicariant speciation process. A good example illustrating this scenario is the comparison of pangasiid fauna between continental Asia and the Indo-Malay Archipelago revealing the existence of related species with allopatric distribution. This is respectively true for H. leptorhynchus and H. waandersii, P. conchophilus and P. nasutus, P. bocourti and P. diambal or between population groups within P. kunyit and P. polyuranodon.

As a consequence of human activities and heavy exploitation, many species in this family are presently being threatened with extinction. The World Conservation Union (IUCN) already lists some species as endangered, e.g. *Pangasianodon gigas* Chevey, 1930 and *Pangasius sanitwongsei* Smith 1931. Thus, the extraordinary diversity and distribution pattern of the Pangasiidae based on a long evolutionary process is now being rapidly modified under human threats. The basic information derived from this study is very important in understanding the relationships, biology and hence the study of the aquaculture potential of species, the improvement of their seed production and growth performance.

CONCLUSION

The present study recognised four genera, i.e. *Pteropangasius* Fowler, 1937; *Helicophagus* Bleeker, 1858; *Pangasianodon* Chevey, 1930; and *Pangasius* Valenciennes, 1840. *Pangasius* is more derive than other genera. The oldest genus may already have existed when the Asian mainland was still connected to the islands in the southern part, about 20 million years ago (mya), during the Miocene glaciation.

The separation between the Schilbeidae and Pangasiidae occurred in the Early Miocene. Diver-



Fig. 4. Evolution within the Pangasiidae as assessed with molecular tools. Speciation has occurred in the Late Miocene for the four pangasiid genera followed by explosive radiation within in the genera in the Early Pliocene and extended to the Late Miocene. Geologic time scale adapted from Ingersoll and Ernst (1987).

gence between the four pangasiids genera dates to about 7-11 mya. A new episode of speciation (about 5-8 mya), marked by an explosive radiation, is observed in the genus *Pangasius*. This adaptive radiation extended until the Late Pliocene (1-4 mya).

ACKNOWLEDGEMENT

The authors thank S. Lenormand, J. Slembrouck, A. Pariselle, M. Legendre, Sudarto, W. Hadie, D. Sadili, A. Hari Kristanto, O. Komarudin, Mr. Maskur, N.V. Thuong, and L.T. Hung for their help in collecting fish samples. Special thanks are due to F. Volckaert who discussed and assisted the paper. This paper forms part of the INCO.DC PROJECT "Catfish Asia" financed by the European Union (contract IC 18-CT 96-0043).

REFERENCES

- Bermingham, E., S.S. McCafferty, and A.P. Martin. 1997. Fish biogeography and molecular clocks: Perspectives from the Panamanian Isthmus. *In* T.D. Kocher and C.A. Stepien (Eds.) Molecular Systematics of Fishes. Academic Press, San Diego, USA.
- Cabot, E.L. and A.T. Beckenbach. 1989. Simultaneous editing of multiple nucleic acid and protein sequences with ESEE. Cabios 5: 233-234.
- Chang Y.S., F.L. Huang, and T.B. Lo. 1994. The complete nucleotide sequence and gene organization of carp (*Cyprinus* carpio) mitochondrial genome. J. Mol. Evol. 38: 138-155.
- De Beaufort, L.F. 1951. Zoogeography of the land and inland waters. Sidwick and Jackson, London, UK. 208 pp.
- Doyle J.J. and J.L. Doyle. 1987. A rapid DNA isolation procedure for small quantities of fresh leaf tissue. Phytochem. Bull. 19: 11-15.
- Felsenstein, J. 1993. PHYLIP (Phylogeny Inference Package) Version 3.5c. University of Washington, Seattle: USA.
- Hillis, D.M., A. Larson, S.K. Davis, and E.A. Zimmer. 1990. Nucleic acids III: Sequencing. p. 318-370. *In* D.M. Hillis and C. Moritz (Eds.). Molecular Systematics. Sinauer Assoc. Inc. Publ., Sunderland, Mass., USA.
- Ingersoll, R.V. and W.G. Ernst. 1987. Cenozoic basin of coastal California. Prentice-Hall, Engelwood, Cliffs NJ, USA.
- Johns, G.C. and J.C. Avise. 1998. Tests for ancient species flocks based on molecular phylogenetic appraisals of *Sebastes* rockfishes and other marine fishes. Evolution 52: 1135-1146.
- Kimura, M. 1980. A simple method for estimating evolutionary rate of base substitutions through comparative studies of nucleotide sequence. J. Mol. Evol. 16: 111-120.
- Kocher, T.D. and K.L. Carlton. 1997. Base substitution in fish mitochondrial DNA: Patterns and Rates. *In* T.D. Kocher and C.A. Stepien (Eds.). Molecular Systematics of Fishes. Academic Press, San Diego, USA.
- Kottelat, M. 1995. The fishes of the Mahakam River, East Borneo: An example of the limitations of zoogeographic

analyses and the need for extensive fish surveys in Indonesia. Trop. Biodiv. 2: 401-426.

- Legendre, M., L. Pouyaud, J. Slembrouck, R. Gustiano, A.H. Kristanto, J. Subagja, O. Komarudin, Sudarto, and Maskur. 2000. *Pangasius djambal*: A new candidate species for fish culture in Indonesia. Indon. Agric. Res. Dev. J. 22: 1-14.
- Lydeard, C. and K.J. Roe. 1997. The phylogenetic utility of the mitochondrial cytochrome b gene for inferring relationships among Actinopterygian Fishes. *In* T.D. Kocher and C.A. Stepien (Eds.). Molecular Systematics of Fishes. Academic Press, San Diego, USA.
- Martin, A.P. and E. Bermingham. 1998. Systematics and evolution of lower Central American Cichlids inferred from analysis of cytochrome b gene sequences. Mol. Phylogenet. Evol. 9: 192-203.
- Meyer, A. 1993. Evolution of mitochondrial DNA in fishes. In Hochachka and Mommsen. p. 1-36. Biochemistry and Molecular Biology of Fishes. Vol. II, Elsevier Science Publ.
- Mo, T. 1991. Anatomy and systematics of Bagridae (Teleostei), and siluroid phylogeny. Theses Zool. 17: 1-216.
- Naylor, G.J., T.M. Collins, and W.M. Brown. 1995. Hydrophobicity and phylogeny. Nature 373: 565-566.
- Ng, H.H and M. Kottelat. 2000. *Helicophagus leptorhyncus*, a new species of molluscivorous catfish from Indochina (Teleostei, Pangasiidae). Raffles Bull. Zool. 48: 55-58.
- Pouyaud, L., G.G. Teugels, and M. Legendre. 1999. Description of a new pangasiid catfish from South-East Asia (Siluriformes). Cybium 23: 247-258.
- Pouyaud, L. and G.G. Teugels. 2000. Description of a new pangasiid catfish from East Kalimantan, Indonesia. Ichthyol. Explor. Freshwater 11: 193-200.
- Pouyaud, L., G.G. Teugels, R. Gustiano, and M. Legendre. 2000. Contribution to phylogeny of pangasiid catfishes based on allozymes and mitochondrial DNA. J. Fish Biol. 56: 1509-1538.
- Rainboth, W.J. 1991. Cyprinids of Southeast Asia. p. 156-210.
 In I.J. Winfield and J.S. Nelson (Eds.). Cyprinid Fishes: systematics, biology and exploitation. Fish and Fisheries, Ser.
 Chapman & Hall, London, UK.
- Roberts, T.R. 1989. The freshwater fishes of Western Borneo (Kalimantan Barat, Indonesia). Mem. Cal. Sci. 14: 1-210.
- Roberts, T.R. and C. Vidthayanon. 1991. Systematic revision of the Asian catfish family Pangasiidae, with biological observation and description of three new species. Proc. Acad. Nal. Sci. Philad. 143: 97-144.
- Saitou, N. and M. Nei. 1987. The neighbor-joining method: a new method for reconstructing phylogenetic trees. Mol. Biol. Evol. 4: 406-525.
- Simon, C., F. Frati, A. Beckenbach, B. Crespi, H. Liu, and P. Flook. 1994. Evolution, weighting, and phylogenetic utility of mitochondrial gene sequences and a compilation of conserved polymerase chain reaction primers. Ann. Entomol. Soc. Am. 87: 651-701.
- Taki, Y. 1978. Geographic distribution of primary freshwater fishes in four principal areas of Southeast Asia. Tonan Ajia Kenkyukyu (Southeast Asia Studies) 13: 200-214.
- Vidthayanon, C. 1993. Taxonomic revision of the catfish family Pangasiidae. Ph.D. Thesis. Tokyo University of Fisheries. 203 pp.
- Zardoya, R. and A. Meyer. 1996. Phylogenetic performance of mitochondrial protein-coding genes in resolving relationships among vertebrates. Mol. Biol. Evol. 13: 933-942.