Unexpected diversity of slow lorises (Nycticebus spp.) within the Javan pet trade: implications for slow loris taxonomy

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Abstract

Since the 1950s, Sundaland (Borneo, Java, Sumatra and their surrounding islands) was thought to be inhabited by a single slow loris species, the greater slow loris Nycticebus coucang. Early taxonomies as well as recent morphological and genetic studies, however, point to at least three species native to this region: N. coucang, N. menagensis, and N. javanicus. In the light of this taxonomy, all Sundaland slow lorises, previously considered Least Threatened, have been listed as Vulnerable or Endangered. Of particular concern is the fact that slow lorises are the most common protected primate species in the rampant Southeast Asian pet trade, resulting in their recent transferral to CITES Appendix I precluding all international commercial trade. Due to lack of knowledge regarding morphological differences between the three species, they are still managed as one, with potential serious affects to wild populations, as hard-release of individuals of unknown geographic origin is common. This paper examines morphological variability of 34 live slow lorises, all of which were rescued from the wildlife trade in Java, Indonesia. Morphometric data and diagnostic images were collected, various species descriptions were considered and statistical analyses were conducted and compared with other taxonomists’ classifications. A discriminant function analysis provided support for four distinct groupings: Nycticebus coucang and N. javanicus, as well as evidence for two new taxa that correspond closely to N. hilleri and N. ornatus. The morphological traits that varied significantly and the external characteristic trends described in this study that contributed to these groupings might provide a baseline to classify Nycticebus taxa. This information is pertinent for appropriate captive management and specific designation of rescued individuals and for designing proper in-situ and ex-situ conservation strategies.

Introduction

Until recently, the cryptic appearance and lifestyles of nocturnal primates concealed an abundance of species within taxonomic groups including Tarsiiformes, Lemuriformes, Galaginae, and Aoutidae (Bearder, 1999). Long-term and detailed studies of the behaviour, ecology, morphology and genetics of these groups revealed that the real number of species was highly underestimated and have led to a wide-spread acceptance of new speciose taxonomic arrangements, (Masters, 1988; Zimmermann et al., 1988; Martin, 1995; Bearder, et al., 1995; Hafen et al., 1998; Nietsch, 1999; Pastorini et al., 2003). Many of these new species, formerly thought to have a broad distribution, are now known to inhabit highly restricted ranges, increasing threat via habitat loss or other stochastic events (Ganzhorn et al., 1997).

It is not surprising, then, that similar trends might be found amongst the Lorisiinae, the lorises of Asia, in particular the slow lorises (Nycticebus) which range from Northern India to the Philippines (Nekaris and Bearder, 2007). Historically, taxonomists recognized at least nine species within the genus Nycticebus (Table 1). In 1953, these taxa were consolidated to a single species, N. coucang (Osman Hill, 1953). Later, Groves (1971, 1998) argued that N. pygmaeus was consistently different enough from N. coucang to be a species in its own right. Captive slow lorises from that point were managed as two distinct species, despite enormous variation in body size and different management needs (Fitch-Snyder and Schulze, 2000). In the last few years, an accumulating number of studies pointed to genetic, morphological and behavioural variability within the lorises (Duckworth, 1994; Ravosa, 1998; Groves, 1998; Supriatna and Hendras, 2000; Fitch-Snyder and Ehrlich, 2003; Roos, 2003; Miehs and Nekaris, 2005; Chen, et al., 2006; Groves and Maryanto, in press). Five species are now generally recognized and have been independently assessed

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in the IUCN Red List (Nekaris and Nijman, 2007), yet many researchers, zoological collections managers, and rescue centre workers still operate in accordance with the 1971 taxonomy (Supriatna et al., 2001; Molur et al., 2003; Schulze and Groves, 2004; Perez, et al., 2005; Shapiro, 2007).

One factor more than any other demands clarification of slow loris taxonomy. Slow lorises are repeatedly found to be the most common protected primate species represented in Southeast Asian animal markets (Malone et al., 2003; Harris, 2003; Webber and Nekaris, 2004; McGreal, 2007). The extent of this trade led to a successful proposal by Cambodia to transfer all members of the genus from CITES (Convention on International Trade in Endangered Species) II to I (Nekaris and Nijman, 2007). The illegal wildlife trade not only contributes to the loss of wild populations (Hunyh, 1998; Ratajszczak, 1998; Fitch-Snyder and Thanh, 2002), but also leads to arbitrary release of animals to forested areas without consideration of their geographic origin (Schulze and Groves, 2004). Confiscated animals of different species housed together may hybridise causing further complications (pers. obs.). Most animals, however, die due to lack of specialised knowledge for keeping slow lorises, or due to complications of health problems acquired whilst in transit or in animals markets (Streicher, 2004; den Haas and Sanchez, pers. comm.).

In Indonesia, trade of slow lorises is illegal, and possession of a loris by buyers or traders is punishable by law. None of the previous studies of slow lorises in Indonesian markets identified the animals at the species or subspecies level, classing them all as N. coucang. If Indonesian trade laws are enforced and animals are not transported between islands, one would hypothesise that animals in markets and in confiscations should come from the island on which they are traded or confiscated (but see Schulze and Groves, 2004). In this study, we had access to 34 slow lorises confiscated in the Javan animal trade, and predicted that all animals should be in accordance with N. javanicus. We provide a detailed analysis of variation within this group of confiscated slow lorises in order to address a number of issues relating to loris conservation. Are multiple taxa present in the Javan trade? If so, which ones? Are these taxa identifiable? What features can be used to discern Indonesian slow loris taxa in order to manage their rescue and rehabilitation?

**Methods**

Thirty-four live Nycticebus specimens of unknown origin, all rescued from the illegal wildlife trade in Java, Indonesia, were maintained at the Schmutzer Primate Centre, Jakarta, Indonesia. In May 2006, the animals were measured for 32 morphological and pelage-related characters using templates specifically designed for
Nycticebus spp. (Fitch-Snyder and Schulze, 2000) (Table 2). We used a large number of pelage-related characters for two reasons. Firstly, accumulating evidence suggests that the markings of nocturnal animals may play an important role in species recognition systems (Bearder, 1999; Bearder, et al., 2006). Secondly, because animals were not anaesthetised, visual assessment with the aid of a colour chart allowed for rapid collection of many characters with limited stress to the animal. All morphometric measurements were taken with digital callipers, and with a Pesola 1000 g spring scale. Hair samples were taken from the thoracic region of the dorsum at the root and measured for length with a digital microscope; the average of ten hairs for each individual was used in analysis. Diagnostic photos were taken of each individual. Later, measurements and images were collected of specimens from the Natural History Museum London, Zoological Museum Amsterdam and Naturalis Leiden for subsequent species comparison of ‘trade’ animals of unknown origin with museum specimens of known origin to ascribe the individuals to a known taxon.

SPSS 12.0 was used for all statistical analyses. Discriminant analyses were conducted to test the validity of grouping of animals based on their close similarity to museum specimens. Non-parametric statistical tests were used for group and pair-wise comparisons due to small sample size, with probability set at the 0.10 level (Zar, 1999).

### Results

Four morphotypes of slow loris were distinguished in the sample when comparing to museum specimens, principally using their characteristic facial masks (including inter-ocular stripe, preauricular hair pattern, circumocular patch shape, and shape and diffusion of crown) (Fig. 1): a greyish small Sumatran form (n=16), a large rufescent Sumatran form (n=9), a short-coated Javan form (n=6), and and a long-coated larger Javan form (n=3). Although more research needs to be conducted comparing these animals with type specimens, they correspond closely to the following taxa: *N. cf. coucang*, *N. cf. hilleri*, *N. cf. javanicus*, and *N. cf. ornatus*. For ease of discussion, the species names are referred to in the rest of this paper, but it should be noted that more research is required with a larger sample size to explore the taxonomic status of these species further.

| General | Pelage Characteristics |
|---------|-----------------------|
| 1. Sex  | 9. Colour of dorsal hair, general |
| 2. Weight | 10. Frosting on back present or absent |

### Table 2. Characters used in the current study; figures showing precise measurements and scores for facial pattern are available in Schulze et al. (2007).

| Pelage Characteristics | Pelage Characteristics |
|-----------------------|-----------------------|
| Face                  | Body                  |
| 3. Facial markings scored | 9. Colour of dorsal hair, general |
| 4. Colour of dark facial markings | 10. Frosting on back present or absent |
| 5. Colour of forehead | 11. Colour of ventral hair, superficially |
| 6. Colour of median facial stripe | 12. Ventral hair base colour |
| 7. Colour of preauricular hair | 13. Throat hair base colour |
| 8. Skin pigmentation | 14. Girth measurement of zone with dark dorsal hair, measured as a percent of girth circumference |
|                       | 15. Dorsal stripe enclosed by light lateral stripe, zone, light neck? |
|                       | 16. Dorsal stripe extends to the more caudal part of trunk? (long vs short) |
|                       | 17. Hair quality: woolly? Amount of stiff guard hair? Wavy or curly? |
|                       | 18. Hair length [mm] on back |

| Measurements (mm) |
|-------------------|
| 19. Chest girth circumference | 27. Head and body length |
| 20. Tail length from hind edge of anus | 28. Maximum length of head |
| 21. Upper arm (humerus) length | 29. Muzzle length |
| 22. Forearm (radius) length | 30. Head breadth |
| 23. Thigh (femur) length | 31. Face breadth |
| 24. Leg (tibia) length | 32. Ear length |
| 25. Hand span |
| 26. Foot span |
When all taxa and all characters were included in a single discriminant function analysis, the percentage of correct classifications obtained was 100%, and groups were clearly discriminated on the basis of two functions (Fig. 2). The first function significantly explained 89.7% of the variance between the four taxa and was dependent on overall colour and facial pattern (Wilks’ Lambda: $\chi^2 = 174.5$, df = 90, $p<0.0001$). The second function significantly explained 7.5% of the variance and was dependent on overall size (Wilks’ Lambda: $\chi^2 = 85.4$, df = 58, $p<0.01$). A second analysis was run on the same data combining the Javan forms due to small sample size of $N. ornatus$. In this case, the percentage of correct classifications obtained was 100%, and groups were still clearly discriminated on the basis of two functions. The first function significantly explained 98.7% of the variance between the four taxa and was dependent on overall colour and facial pattern (Wilks’ Lambda: $\chi^2 = 175.7$, df = 62, $p<0.0001$). The second function significantly, dependent on size, explained 1.3% of the variance (Wilks’ Lambda: $\chi^2 = 53.6$, df = 30, $p<0.01$).

As facial pattern was the main character used to classify the animals based on museum specimens, a second analysis was run excluding this character, still grouping the Javan forms. The classification remained at 100% for the three groups (Fig. 3). The variance explained significantly by the first function, still relating to colour pattern, reduced to 90.5% (Wilks’ Lambda: $\chi^2 = 168.8$, df = 87, $p<0.0001$), whereas variation as explained by size rose to 7.8% (Wilks’ Lambda: $\chi^2 = 78.6$, df = 56, $p<0.03$).

Table 3 summarises the morphometric measures of each taxon, showing relatively (but not significantly) larger ($N. javanicus$ and $N. hilleri$) and relatively smaller ($N. ornatus$ and $N. coucang$) forms on each island. Table 5 highlights pelage characteristics that differ significantly amongst the groups. General size and colour patterns in 12 different characters differentiate these groups of lorises, despite the small sample size (Fig. 4).

Pairwise comparisons were then run amongst significant traits to examine differences between groups. The only ordinal character that distinguished the two Javan taxa was hair length (Mann-Whitney U: $U=0$, $p<0.03$). Sumatran taxa were distinguished by head breadth (Mann-Whitney U: $U = 36$, $p<0.05$) and brachial index (Mann-Whitney U: $U = 42$, $p<0.09$). They were further distinguished by several categorical traits:
facial mask \( (\chi^2 = 21.1, \text{df} = 1, p<0.0001) \), dorsal stripe length \( (\chi^2 = 6.8, \text{df} = 2, p<0.03) \) and preauricular hair colour \( (\chi^2 = 6.2, \text{df} = 3, p<0.10) \).

Finally, pairwise comparisons were run on these same traits grouping all Javan lorises and all Sumatran lorises. Eight ordinal characters distinguish these groups as well as six characters relating to pelage (Tables 3 and 4).

**Discussion**

Two species are clearly discernible in the Javan trade – *N. coucang* and *N. javanicus*. Indeed, nearly two thirds of the sample in this study was from Sumatra, where trade in slow lorises is known to be high (Shepherd *et al.*, 2004). Variability within these groups aside, the high proportion of Sumatran lorises in this sample was similar to that seen in the trade in orang-utans and gibbons on Java (Nijman 2005), suggesting that inter-island trade is poorly enforced. Slow lorises are wholly protected in Indonesia. Although fines and jail sentences should be imposed on those trading or owning lorises, and indeed other protected wildlife, effective law enforcement with respect to wildlife protection laws is all but non-existent in Indonesia (Nijman 2006).

Futhermore, Javan slow lorises are considered Endangered, and trade has been identified as a consider-
able threat. Non-systematic surveys by International Animal Rescue, an organisation that does discern between the Javan and Sumatran forms, noted a marked decrease of Javan forms in the markets over the last few years (den Haas and Sanchez, pers. comm.; Jaffe, 2005). As demand for slow lorises is still high, and numbers of Javan animals has decreased, trade could be a direct indicator of decrease of these animals in the wild. Similar trends have been identified in Vietnam and Cambodia, where lorises have become both increasingly difficult to observe in both the wild and the trade (Streicher and Nadler, 2003; Starr, pers. comm.).

The analyses performed in this study clearly distinguished Sumatran lorises from Javan lorises. *Nycticebus javanicus* was first recognised as a distinct taxon in 2000 (Supriatna and Hendras, 2000). A subsequent molecular study relied on only a single specimen from Java, and this taxon could not be discerned from this small sample size (Chen et al., 2006). In a more detailed cranometric study, Groves and Maryanto (in press) decided that Javan lorises, based on a sample of 25 skulls, were consistently distinct enough, especially when combined with pelage characters, to merit separation as a distinct species. This study provides further evidence, using a set of characters used to distinguish other nocturnal mammals (Masters and Spencer, 1989), that *N. javanicus* warrants distinction as a species.

Earlier studies also pointed to great variation within *Nycticebus* from Sumatra and Borneo (Groves, 1971; McPhee and Jacobs, 1986; Ravosa, 1998). Indeed, in 1904, Lydekker (p. 346) noted “the marked distinction between the grey Malay phase and the rufous Sumatran phase of the species (Fig. 5).” The small sample size in this study clearly distinguished two Sumatran forms, in accordance with *N. coucang* and *N. hilleri*. Two Javan forms were also distinguished, but mainly on hair length, perhaps indicating altitudinal differences. These differences were enough, however, for them earlier to be distinguished as *N. javanicus* and *N. ornatus* (Osman Hill, 1953). In the case of Sumatran species, the red form fetches higher prices in markets and is said to be more rare (Dwi, pers. comm.).
sis, Sumatran *N. coucang* and *N. javanicus* have been isolated on their respected islands since the Holocene or postglacial period, and within their islands are also isolated due to river and mountain boundaries (van den Bergh *et al.*, 2001). Therefore, the results shown here and the suggestion that these taxa are distinctive on the species or subspecies level are not surprising. Also, the two Javan forms found within this sample coincide with the notion of altitudinal variation; a common factor classifying Indonesian primate subspecies (Brandon-Jones *et al.*, 2004).

Ravosa (1998), looking at craniometric variables, found high variability within this group, but suggested that the differences within *coucang* were clinal. What he and other authors studying *Nycticebus* skeletal morphology did not consider is that in many other nocturnal primate species, mate recognition systems depend on vocal and olfactory signals as well as contrasting facial patterns (Mayr, 1963; Bearder, *et al.*, 1995). Thus animals that are almost identical skeletally are often in fact distinct species (Masters and Bragg, 2000). Mayr (1942) argued that a taxon is a valid subspecies if it can be distinguished on 75% of its traits; from this sample, such is the case with *N. c. hilleri*. Separation of this taxon from *N. coucang coucang* might help to explain considerable variability and low percent discrimination of individuals into this taxon within Sumatran *N. coucang* observed in previous studies (Ravosa, 1998; Groves and Maryanto, in press). Further morphological, behavioural and genetic studies are required to understand variability within *Nycticebus* keeping this taxonomic arrangement in mind.

Whether or not taxonomic arrangements of the Sundas slow lorises should be altered, identification of the morphotypes is essential for rescue and release programmes. Trade is so great that numbers of lorises coming into rescue centres exceed their capacity (den Haas and Sanchez, pers. comm.). Reduced populations in the wild, combined with mounting evidence for both national and international trade, have resulted in transferral of the genus *Nycticebus* to CITES I – the first transferral of a primate species since 1989 (Nekaris...
Specialist needs of these species also mean that they perish quickly in captivity (Fitch-Snyder and Schulze, 2000); thus oftentimes immediate re-release is considered as preferable (Wiek, 2007). This means that in countless cases, animals are reintroduced without regard to their health status, their behavioural needs, or their geographic origin (Schulze and Groves, 2004; Streicher, 2004). In the single re-introduction programme conducted ethically and systematically, one third of the released animals perished despite following all IUCN protocol (Streicher and Nadler, 2003). The results for hard released animals are probably far direr.

These cases illustrate that it is essential that rescue centres and confiscation authorities are able to identify animals in order to release them to an appropriate locality (Streicher and Nadler, 2003; Schulze and Groves, 2004). As cryptic species, physical differences amongst the slow lorises are subtle. We hope that information provided in this study will be useful in developing further identification keys for individuals confiscating or rehabilitating lorises and for better understanding speciation amongst Nycticebus.

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