# Diet and habitat of Siwalik primates *Indopithecus*, *Sivaladapis* and *Theropithecus*

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Received 14 July 2013, final version received 12 Nov. 2013, accepted 13 Nov. 2013

Patnaik, R., Cerling, T. E., Uno, K. T. & Fleagle, J. G. 2014: Diet and habitat of Siwalik primates *Indopithecus, Sivaladapis* and *Theropithecus. – Ann. Zool. Fennici* 51: 123–142.

We present dental enamel stable carbon and oxygen isotope data, histological analyses of daily cross striations and perikymata, and microwear data of Late Miocene primates *Indopithecus* and *Sivaladapis nagrii* and an early Pleistocene primate *Theropithecus delsoni*, known from the Indian Siwaliks. The results indicate that the Late Miocene giant ape *Indopithecus* had a C<sub>3</sub> diet, likely including nuts, seeds and fruits with hard cover, and most probably lived in a seasonal forest that experienced long dry and wet seasons. While the seasons changed as depicted by intra-annual variation in  $\delta^{18}$ O values, the C<sub>3</sub> component of diet was similar throughout the year, as indicated by consistent  $\delta^{13}$ C values. *Sivaladapis nagrii* dental enamel  $\delta^{13}$ C and  $\delta^{18}$ O values, striations and pit percentages indicate that this small primate (3 kg) was a C<sub>3</sub> mixed folivore-frugivore. *Theropithecus delsoni* dental enamel  $\delta^{13}$ C and  $\delta^{18}$ O values and microwear pattern indicate a diet comprising mainly C<sub>4</sub> grasses or sedges.

# Introduction

Indopithecus giganteus (or Gigantopithecus bilaspurensis) was an ape the size of a female gorilla (150+ kg, Fleagle 1999) that lived in the sub-Himalayan Siwaliks of India and Pakistan, in the Late Miocene from ca. 8.6 to 8.85 Ma ago (Pillans *et al.* 2005). The genus *Indopithecus* was erected by Von Koenigswald (1950) based on an unusually large isolated molar (GSI D-175) recovered from Alipur, now in Pakistan, which had been described as *Dryopithecus giganteus* by Pilgrim (1915). Subsequently, the

same specimen was allocated to *Dryopithecus* (*Sivapithecus*) *sivalensis* by Simons and Pilbeam (1965), who considered this specimen to be similar to *Sivapithecus*. The discovery of a mandibular specimen (CYP359/68) from Haritalyangar, India, and its similarity to those of the Chinese Pleistocene giant ape, *Gigantopithecus blacki*, led Simons and Chopra (1969) to propose a new species, *Gigantopithecus bilaspurensis*. However, Cameron (2001, 2003), based on his studies on functional morphology and phylogenetics, argued that the mandibular specimen should be reallocated to the species *Indopithecus gigan*.

*teus*. Szalay and Delson (1979) and Wolpoff (1999) were of the opinion that *Indopithecus* was a "thick enameled ground ape". Pillans *et al.* (2005) described an isolated tooth of *Indopithecus* (HDI-1) from Haritalyangar (Fig. 1A and B), India, which has been studied here for stable isotope, dental microstructure, and microwear analyses.

Sivaladapis was a small late-Miocene primate, weighing around 3 kg (Fleagle 1999) and belonging to the adapoid family Sivaladapidae. Early sivaladapids, including Hoanghonius, Rencunius, Guangxilemur and Wailekia appeared in the Middle Eocene of China and Thailand (Woo & Chow 1957, Gingerich et al. 1994, Ducrocq et al. 1995, Tong 1997, Qi & Beard 1998, Gebo et al. 1999, Tong et al. 1999), and sivaladapids have also been reported from the Eocene of Myanmar (e.g. Beard et al. 2007). In the Oligocene deposits, Guangxilemur occurs at Bugti Hills, Pakistan (Marivaux et al. 2002). During the Miocene, the sivaladapids were taxonomically and geographically very diverse with Sivaladapis and Indraloris from southern Asia, Sinoadapis from China, and Siamoadapis from Thailand (Lewis 1933, Tattersall 1968, Prasad 1970, Gingerich & Sahni 1979, 1984, Thomas & Verma 1979, Wu & Pan 1985, Pan & Wu 1986, Pan 1988, Sahni 1989, Barry et al. 2002, Pillans et al. 2005, Flynn & Morgan 2005, Chaimanee et al. 2008). By the end of the Miocene sivaladapids became extinct (Gingerich & Sahni 1984). The three sivaladapid specimens, Hari I-6, Hari I-7 (Pillans et al. 2005) and P11 (Gingerich & Sahni 1984) studied here come from Dhok Pathan Formation, Haritalyangar (Fig. 1A and B).

One mandibular fragment of the papionin cercopithecine, *Theropithecus delsoni* (A643), is known from the locality of Mirzapur, near Chandigarh (Fig. 1C) (Gupta & Sahni 1981, Delson 1993, Delson *et al.* 1993). Its age is tentatively considered ~1 Ma (Pinjor Formation) as the specimen does not come from a well dated section (Patnaik & Nanda 2010). The presence of *Theropithecus* in India holds significance as it is considered that *Homo erectus*, *Theropithecus*, *Megantereon whitei*, and *Hippopotamus* were among the first mammals to disperse out of Africa in the early Pleistocene (Martinez-Navarro 2004, Rook *et al.* 2004, Fleagle *et al.* 2010).

Mammalian dental enamel contains periodic growth structures that preserve the history of development of the tooth. Each sub millimeter tooth enamel increment (perikymata) is secreted in 8 days in Sivapithecus indicus, and 9 days in Sivapithecus parvada (Mahoney et al. 2007). In Gigantopithecus blacki the periodicity has been found to be 11 days per perikymata (Dean & Schrenk 2003). Graecopithecus, a large-bodied Miocene ape from Greece, which has been considered phylogenetically closer to Indopithecus (Gelvin 1980, Cameron 2001), has a periodicity of 8 days (Smith et al. 2004). Among the extant apes, Hylobates lar which weighs around 6 kg, shows a periodicity of 4 days (Dirks & Bowman 2007); Pan troglodytes (33-45 kg) has a periodicity of 6 to 7 days (Smith et al. 2003); Gorilla (98-163 kg) has a periodicity of 7 to 10 days; Pongo (36–78 kg) has a periodicity of 8–11 days (Schwartz et al. 2001, Hu et al. 2011). Hence, it appears that there is a positive correlation between periodicity and body mass in hominoids (Reid et al. 2002, Hu et al. 2011).

Tooth enamel bioapatite of extant and fossil species preserves a record of carbon and oxygen isotopes incorporated through ingested food and water. In most tropical and many subtropical ecosystems, the carbon isotope composition ( $\delta^{13}$ C) of tooth enamel in modern herbivores distinguishes between diets dominated by C<sub>2</sub> plants (dicots including trees, shrubs, herbs, forbs; and cool season (high latitude or altitude) grasses or sedges), and those dominated by  $C_4$  plants (mostly monocots, such as warm-season grasses and sedges); mixed-feeding comprises a mixture of  $C_3$  and  $C_4$  plants (Lee-Thorp & Van der Merwe 1987, Bocherens et al. 1996, Cerling et al. 1997). Enamel is enriched in <sup>13</sup>C by 12‰ to 14‰ with respect to diet (Cerling & Harris 1999, Passey et al. 2005). For the post-Miocene, interpretations of C<sub>3</sub> vs. C<sub>4</sub> diets of extinct mammals are based upon certain cut-off values of  $\delta^{13}C$ in dental enamel. For instance, the  $\delta^{13}$ C values from dental enamel representing a pure C<sub>3</sub> diet of a modern mammal would range from approximately -20% to -8%, with the most negative values indicating a diet from the ground-level of closed canopy rainforest, and the higher values indicating a C<sub>3</sub>-diet derived from an open habitat, which could range from open forest or woodland



**Fig. 1.** (**A**) Location of the Himachal Pradesh reentrant [modified from Brozovic & Burbank (2000)]. (**B**) Magnetostratigraphic section at Haritalyangar (after Pillans *et al.* 2005). (**C**) Fossil localities in the vicinity of Chandigarh [modified from Sahni & Khan (1964)]. (**D**) Area around Haripur situated 100 km east of Chandigarh [modified from Kumar *et al.* (2002)].



**Fig. 2.** Monthly (January–December) rainfall amount and  $\delta^{18}$ O (SMOW) values illustrating the amount effect for (**A**) Hong Kong and (**B**) New Delhi, (modified from fig. 5 of Nelson 2005, reproduced with permission from Elsevier).

to grassland (Cerling *et al.* 1997, Cerling *et al.* 2004). On the other hand, a pure  $C_4$  diet would yield  $\delta^{13}C$  (enamel) values between +1‰ to +4‰. Prior to the late Miocene, most ecosystems on Earth were dominated by  $C_3$  plants so that the modern  $C_3$ -browsing/ $C_4$ -grazing distinction cannot be used for these older periods of time.

The oxygen isotope ratio of dental enamel depends upon body water, which is derived from drinking water, water in plants, and metabolically derived water (Kohn 1996, Kohn et al. 1996, Sponheimer & Lee-Thorp 1999). Differing proportions of these components, including intake of O<sub>2</sub> and loss of CO<sub>2</sub> and water vapour, determines the  $\delta^{18}$ O of body water which, in turn, determines the  $\delta^{18}$ O of tooth enamel (Luz & Kolodny 1985, Podlesak et al. 2008, 2012, O'Grady et al. 2010, 2012). Drinking water is generally derived from meteoric waters, which often show a seasonal cycle; in the monsoon region of Indo-Pakistan, summer precipitation is depleted in <sup>18</sup>O as compared with winter precipitation (e.g., Fig. 2) due to the "amount effect" (Dansgaard 1964, IAEA 1981). "Food water" is derived from leaves, which is significantly enriched in <sup>18</sup>O relative to its source water. Metabolic water is derived from the oxygen bound in foods (e.g., CH<sub>2</sub>O) which is released during respiration by reaction with atmospheric oxygen. The isotopic ratio of drinking water, and the relative proportions of the three water input sources, are both likely to change over seasonal cycles; thus, it is not possible to determine if <sup>18</sup>O-enriched tooth enamel is associated with "dry" or "wet" seasons without a priori assumptions or information. However, significant annual variations in body water are likely to indicate high seasonality in the isotope ratios or the relative proportions of these different water sources. Several studies have shown that intratooth variation in oxygen isotopic composition of both modern and fossil mammalian taxa provides information on intra-annual seasonal variation (Koch et al. 1989, Bryant et al. 1994, 1996, Fricke & O'Neil 1996, Fricke et al. 1998, Cerling & Sharp 1996, Sharp & Cerling 1998, Gadbury et al. 2000, Fox & Fisher 2001) but it is often difficult to extrapolate this directly to seasonal amplitude in the isotopic ratio of local precipitation.

Research on dental microwear patterns (percentage and width of pits and striations) of modern primates has been used to infer dietary preferences of their extinct counterparts (Grine & Kay 1988, Ungar 1996, Merceron et al. 2005). Several comparative studies using pit and striation ratios in modern apes such as Gorilla, Pan and Pongo and extinct apes/hominids (e.g., Sivapithecus, Gigantopithecus, Paranthropus, Australopithecus) have provided useful information about their diet (Teaford & Walker 1984, Daegling & Grine 1999, Nelson 2003, Ungar et al. 2008, Grine et al. 2009). It also permits understanding interspecific dietary differences and diet variation linked to seasonal and ecological factors (Teaford 1986, Teaford & Oyen 1989, Teaford & Robinson 1989, Teaford & Glander 1996). As such, it is the abrasive component associated with food that is responsible for the formation of pits and scratches. A low percentage of pits on wear facets is interpreted to indicate folivory, whereas a high percentage of pits indicates a fruit dominated diet. A high percentage of long, wide parallel oriented scratches indicate a grazing habit. Primates that eat a lot of hard food such as seeds, nuts and roots have very high number of pits, large pits and wide scratches (Ungar 1996, King et al. 1999). However, one should be careful while making such interpretations, as post-mortem changes prior to the final burial (taphonomic modifications) of the samples may lead to formation of similar scratches and pits (Ungar et al. 2006). To reconstruct dietary behaviour of small primates, body mass and shearing quotients (SQ) are also widely

used (Kay 1975, Kay & Covert 1984, Kirk & Simons 2001, Ramdarshan *et al.* 2010). Primates under 350 g obtain protein mainly from insects or other invertebrate fauna, while those above 600 g are usually more herbivorous. Molars of primates that eat insects and leaves have long and sharp crests, while those eating fruits have shorter crests and shallower basins.

# Material and methods

This study from the Indian Siwaliks includes the primates *Indopithecus* and *Sivaladapis nagrii* from the Miocene Dhok Pathan Formation, and *Theropithecus delsoni* from the Pleistocene. Paleosol carbonates from the Miocene localities near Haritalyangar were collected from well-developed paleosols using the criteria as described in Quade and Cerling (1995). Associated mammalian fauna collected from Dhok Pathan Formation exposed near Haritalyangar and Upper Siwaliks around Chandigarh region is also included in this study.

#### Stable isotope analyses

Stable isotope analyses were performed using both conventional  $H_3PO_4$  digestion and laser ablation methods. For conventional analysis of enamel, approximately 5 milligrams of powdered enamel was treated using conventional methods to remove carbonates (Cerling *et al.* 2011); paleosol carbonates required no pretreatment. Samples were reacted with 105%  $H_3PO_4$ at 90 °C and analyzed on an isotope ratio mass spectrometer (IRMS) with results reported in the permil notation:

$$\delta^{13}\mathrm{C} = (R_{\mathrm{sample}}/R_{\mathrm{standard}} - 1) \times 1000 \qquad (1)$$

where  $R_{\text{sample}}$  and  $R_{\text{standard}}$  are the <sup>13</sup>C/<sup>12</sup>C ratios for the sample and standard, respectively. An analogous expression is used to report <sup>18</sup>O/<sup>16</sup>O ratios. For both  $\delta^{13}$ C and  $\delta^{18}$ O the isotope standard is V-PDB (Vienna-Pee Dee Belemnite). Fossil isotope reference materials and an acid fractionation factor of 1.00725 were used to correct  $\delta^{18}$ O enamel values to 25 °C (*see* Passey *et al.* 2007).

**Fig. 3.** Laser ablation pits and perikymata on a silicon rubber mould of the *Indopithecus* (HDI-1) molar surface (A–G lateral and H–J cuspal enamel). There are 38 perikymata between samples A and G.

For selected samples, analyses were made using laser ablation (LA) to liberate CO<sub>2</sub> from bioapatite enamel, using the methods of Cerling and Sharp (1996) as modified by Passey and Cerling (2006) (Figs. 3 and 4A). Multiple analyses of a single tooth from the Siwaliks using both methods was used to normalize the LA-IRMS results to the conventional  $H_3PO_4$  results; the offset was 5.1% between the two methods for  $\delta^{18}O$ results; no correction was applied to the  $\delta^{13}C$ values because the offset between the laser and conventional results was similar to the analytical uncertainty (~0.1‰).

#### Growth periodicity studies

For determining growth periodicity of *Indopithecus* and *Theropithecus* a fractured piece of enamel was removed from the tooth, its transverse surface was cleaned with acetone, and was etched with 5% HCl for 100 seconds. In case of *Sivaladapis*, the removed enamel piece was embedded in polyester resin, its transverse surface was polished, cleaned with acetone and etched with 5% HCl for 80 seconds. The surface was coated with gold palladium and was examined under a scanning electron microscope (JSM JEOL, 6460) at the Geology Department, Panjab University to record the presence of both short- and long-period incremental growth lines in enamel. Short-period lines, or cross-striations,





**Fig. 4.** (A) *Sivaladapis nagrii*  $M_1M_2M_3$  (P11) with laser scan pits (scale 5 mm). (B) SEM image of labial view of *Sivaladapis nagrii*  $M_1$  (Hari I-6). (C) a portion of the  $M_1$  magnified to show perikymata.

represent daily increments of enamel deposition. Long-period lines, referred to as Retzius lines or striae of Retzius (manifested on the tooth enamel surface as 'perikymata'), represent periodic disruptions in enamel deposition that occur every six to twelve days in extant great apes and humans, although the periodicity is constant within any given individual (Dean 1987, 1998, 2000, FitzGerald 1998).

#### **Microwear studies**

For dental enamel microwear studies, a mould of the *Indopithecus* tooth was prepared using a RTV silicone rubber, followed by the preparation of a cast made by using a polyester resin (Araldite). The semi-transparent cast was surveyed under transmitted light of a Leica S8APO microscope to find areas that were unaltered taphonomically. The cast was then coated with gold palladium and was viewed under SEM. In case of Sivaladapis a P4 (Hari I-6) was coated with gold and viewed under SEM for microwear studies. Several images of Indopithecus, Sivaladapis and Theropithecus Phase II wear facets were taken and one from each tooth was selected to count/ measure pit percentage, pit width and length and scratch width. To check the reliability and consistency of the microwear pattern, two adjacent areas were scanned at a 500× magnification (see El-Zaatari et al. 2005 for details). The micrographs were saved as gray-scale image files at 200 dpi resolution and cropped to  $640 \times 480$ pixels. The counts and measurements were made by one observer (RP) using MICROWARE 4.0.2 (Ungar 2002). A comparison of these features was made with those reported in published works on both extant and extinct primates (Walker & Teaford 1989, Rafferty et al. 2002, Nelson 2003, El-Zaatari *et al.* 2005, Merceron *et al.* 2005, Ungar *et al.* 2006, Ramdarshan *et al.* 2010).

# Results

# Relationship between isotope sampling and growth structures

Stable isotopes measurements in enamel represent an attenuated signal because of maturation of enamel over some period of time (Passey & Cerling 2002). The temporal discussion in this section refers to the crown formation time of enamel, which does not include maturation time. The time interval sampled for isotope analysis includes crown formation time and maturation time; however, the true interval cannot be estimated without a good model for the enamel maturation process in these species.

The right M<sup>2</sup>, VPL/HDI (1) of *Indopithecus* from 8.85 Ma and Middle Siwaliks of Haritalyangar (Pillans *et al.* 2005) had 10 laser scans, each comprised of 5–8 ablation pits (Table 1 and Fig. 3). Laser ablation pits were of ~250  $\mu$ m diameter and the outcropping Retzius lines formed ~50–70  $\mu$ m thick perikymata. In total 38 perikymata were sampled that represent ~418 days (based on the periodicity of 11 cross-striations per perikymata; Fig. 5) for samples A–G (*see* Fig. 3). Perikymata in scans H–J are not distinct (*see* Fig. 3).

For stable isotope analysis of Sivaladapis nagrii, a mandible comprising M<sub>1</sub>, M<sub>2</sub> and M<sub>2</sub> (P11) was used (Fig. 4A). A total 6 laser scans (m1c, m1c2, m1r, m2, m3c and m3r) were obtained from the labial face of the three teeth, each scan comprised of 5-8 ablation pits (Fig. 4A). Sivaladapid teeth were too small to sample by traditional drilling methods, so the LA-GC-IRMS method was the only means by which the tooth could be sampled for isotope analysis. Each pit was  $\sim 250 \ \mu m$  in diameter. The outcropping Retzius lines formed ~40–50  $\mu$ m thick perikymata. Therefore, the six scans covered around 30 perikymata on the lateral enamel of the molars of Sivaladapis nagrii (Fig. 4B). A growth periodicity of two days (Fig. 6) was estimated using SEM. For such periodicity, isotope data from 6 laser scans of around 30 perikymata would represent approximately a two-month record of diet and climate. We, therefore, did not expect to find a seasonal variation in  $\delta^{18}$ O values in the Sivaladapis nagrii molars studied here.

**Table 1.** Stable isotope results from laser ablation.  ${}^{18}\varepsilon_{laser-acid} = -5.1\%$  for all  $\delta^{18}O$  data. No difference was found between laser and conventional  $\delta^{13}C$  data.

	Distance from	$\delta^{_{13}}C$	$\delta^{18} { m O}$		
	the base of the crown (mm)		laser measured	CO <sub>3</sub> equivalent	
Indopithecus					
HD1-A	0.3	-12.2	-10.6	-5.5	
HD1-B	0.6	-11.8	-9.5	-4.4	
HD1-C	0.9	-11.7	-7.6	-2.5	
HD1-D	1.2	-11.9	-7.2	-2.1	
HD1-E	1.5	-11.8	-8.0	-2.9	
HD1-F	1.8	-11.6	-8.2	-3.1	
HD1-G	2.4	-12.4	-10.3	-5.2	
HD1-H	3.0	-12.1	-9.3	-4.2	
HD1-I	3.6	-12.2	-10.1	-5.0	
HD1-J	4.2	-12.2	-8.7	-3.6	
Sivaladapis					
P11_m1c		-11.6	-10.0	-4.9	
P11_m1c2		-11.3	-9.4	-4.3	
P11_m1r		-12.4	-9.8	-4.7	
P11_m2		-12.7	-9.3	-4.2	
P11_m3c		-13.4	-9.6	-4.5	
P11_m3r		-14.0	-10.1	-5.0	

Fig. 5. (A) Enamel microstructure as seen in a crosssection on the Indopithecus tooth (HDI-1). (B) Portion inside the box in A magnified. Short-period lines, or cross-striations, representing daily increments of enamel deposition are shown by black arrows. Longperiod lines, referred to as Retzius lines or striae of Retzius are shown by white arrows. Arrows point towards the occlusal surface of the tooth.

A sample for stable isotope analyses from the Theropithecus molar enamel was obtained using a dental drill that covered around 30 perikymata. Theropithecus molar enamel microstructure shows a periodicity of 7 days (Fig. 7), therefore carbon isotope data from the drill pit covering 30 perikymata gives an average diet over a period of ca. 7 months.

#### Stable isotopes in Siwalik primates

The laser scans A-J on the Indopithecus tooth (Fig. 3) yielded min, max and mean  $\delta^{13}$ C values of -12.4‰, -11.6‰ and -12‰, respectively; and the min, max and mean  $\delta^{18}$ O values (adjusted to

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-2.1% and -3.9%, respectively (Tables 1 and 2). The laser scans (Fig. 4A) of the m1c, m1c2, m1r, m2, m3c and m3r on Sivaladapis nagrii teeth yielded min, max and mean  $\delta^{13}$ C values of -14.0%, -11.3% and -12.5%, respectively; and the min, max and mean  $\delta^{18}$ O values (enamel-CO<sub>2</sub> equivalent) of -5.0‰, -4.2‰ and -4.6‰, respectively (Table 2).

The  $\delta^{13}$ C and  $\delta^{18}$ O values for a single sample from Theropithecus were -1.9%, and -2.0%, respectively (Table 2). For the mean  $\delta^{13}$ C and  $\delta^{18}$ O values from laser or conventional sampling for Siwalik primates, associated mammalian fauna, and paleosol carbonates see Fig. 8.

A 25kV

X100

100µm

0004

RAM

#### **Paleosol carbonates**

The Haritalyangar paleosol carbonate  $\delta^{13}$ C values ranged between -12.2% (min) and -8.0% (max) with mean = -10.8% (Table 3 and Fig. 8). Soil carbonate  $\delta^{18}$ O values range between -9.2% (min) and -5.8% (max) with mean = -7.7% (Table 3 and Fig. 8).

#### Stable isotopes of associated fauna

The older Dhok Pathan associated fauna includes bovids, cervids, equids, rhinocerotids, suids and tragulids. The  $\delta^{13}$ C values range between -13.2% (min) and -10.7% (max) with mean = -12.1%, and the  $\delta^{18}$ O between -0.3% (min) and -11.9% (max) with mean = -5.7% (Table 2 and Fig. 8). Associated fauna from younger deposits (1 to 3 Ma) include bovids, camelids, equids, giraffids, hippopotamids, and rhinocerotids with the  $\delta^{13}$ C values ranging from -2.9‰ (min) to +2.0% (max) with mean = -1.7%. Elephantids are represented by *Elephas* and *Stegodon*; *Elephas* has the  $\delta^{13}$ C values of -0.9‰ (min) and -0.1% (max) with mean = -0.4%, whereas Stegodon has min, max and mean  $\delta^{13}$ C values of -11.9‰, -7.3‰ and -9.4‰, respectively (Table 2 and Fig. 8).  $\delta^{18}$ O values of associated fauna from the younger deposits ranges from -7.6% (min) to +0.3% (max) with mean = -2.8%.

#### **Enamel microstructure**

Lateral enamel of *Indopithecus* shows Rhetzius lines of ~55  $\mu$ m thick with 11 prism cross-striations representing 11 days (Fig. 5). This gives an enamel secretion rate of ~5  $\mu$ m per day. In *Sivaladapis nagrii* the rhetzius lines are ~10  $\mu$ m thick having 2 cross striations, representing 2 days. The enamel secretion rate in *Sivaladapis nagrii* is ~5  $\mu$ m per day. Enamel microstructure of *Theropithecus delsoni* shows ~35  $\mu$ m thick Rhetzius lines with 7 prism cross-striations representing 7 days (Fig. 7), with an enamel secretion rate of ~5  $\mu$ m per day.

Dental enamel microwear on *Indopithecus* tooth (Fig. 9) reveals that the tooth surface is rather rough and heavily pitted with a high per-

Fig. 7. (A) Enamel microstructure as seen in a cross-section on the *Theropithecus delsoni*  $M_1(A643)$ . (B) Part of A magnified. Short-period lines, or cross-striations, representing daily increments of enamel deposition are shown by black arrows in B. Long-period lines, referred to as Retzius lines or striae of Retzius are shown by white arrows in A and B. Arrows point towards the occlusal surface of the tooth. Enamel surface is towards the top and enamel dentine junction is towards the bottom.

centage of pits (~58%). The average pit width is 5.57  $\mu$ m and the average pit length is 8.09  $\mu$ m. There is a moderate percentage (~42%) of mixed striations present. Striation width and length are on average 2.19 and 36.11  $\mu$ m, respectively. *Sivaladapis nagrii* dental enamel microwear on grinding lophs (Phase II facets) shows ~60% of small pits and ~40% of long parallel narrow to wide striations (Fig. 9). Pit width averages 1.88  $\mu$ m. The mean pit length is 2.72  $\mu$ m. Striation width and length are on average 1.22 and 64.25  $\mu$ m, respectively. Most of the striations were fine averaging ca. 1  $\mu$ m, whereas a few are as wide as 5  $\mu$ m. Few fine cross striations



are also present (Fig. 10). Theropithecus delsoni dental enamel exhibits  $\sim$ 34% of small pits and  $\sim$ 66% of fine to wide striations (Fig. 11D). Average pit width is 3.28  $\mu$ m and average pit length is 4.7  $\mu$ m. The average width and length of the striations are 1.3 and 22.48  $\mu$ m, respectively.

### Discussion

#### Indopithecus, Sivaladapis and Theropithecus diets and habitats

#### Indopithecus

The  $\delta^{13}$ C values for the *Indopithecus* reported

here range from -12.4‰ to -11.6‰, indicating a pure  $C_3$  diet similar to that of the ca. 9.2 Ma Sivapithecus from the Siwaliks as reported by Nelson (2007); thus, both Indopithecus and Sivapithecus had C<sub>2</sub>-dominated diets in the Dhok Pathan. A C<sub>3</sub>-dominated habitat is supported by the  $\delta^{13}$ C values obtained from associated Sivalhippus and other associated faunal remains, and from soil carbonates from equivalent strata exposed at a nearby site (Kursai 7; Table 2 and Fig. 8). However, because this period represented by the Dhok Pathan sediments is before the global expansion of  $C_4$  ecosystems (Cerling et al. 1997), the  $C_3-C_4$  distinction in diets and paleosols cannot be used to evaluate the degree of "openness" of the habitat at this time.

**Table 2.** Stable-isotope analysis results from mammalian taxa in the Haritalyangar region, Indian Siwaliks. Values in parentheses are averages for samples measured with the laser ablation system (Table 1) with  $\delta^{18}$ O given as the estimated equivalent value for the CO<sub>3</sub><sup>2-</sup> component of bioapatite.

Sample ID	$\delta^{_{13}}\mathrm{C}$	$\delta^{18} O$	Taxon	Locality	Formation	Age (Ma
Samples younger than 8 Ma						
M2	-2.9	-2.4	Bovid	Nadah	Pinjor	~1.8
M5	2.0	-1.9	Bovid	Nadah	Pinjor	~2.0
B281	0.0	-3.8	Bovid jaw	Mirzapur	Pinjor	~2.0
KB-1	-2.9	-4.5	Bovid	Kanthro	Tatrot	2.5
PUBMN1	-2.5	0.3	Camelus	Moginand	Tatrot	~2.5
DKE-1	-0.1	-5.7	Elephas	Devni Khadri	Tatrot	~2.5
F335	-0.3	-3.0	Elephas	Quaranwala	Pinjor	1.5
B594	-0.2	-4.0	Elephas	Quaranwala	Pinjor	1.8
B246	-0.9	-1.8	Elephas	Quaranwala	Pinjor	2.0
A586	-1.7	-3.7	Equus	Quaranwala	Boulder Congl.	~1.0
A600	-0.1	-0.1	Equus	Mirzapur	Pinjor	~2.0
161-G	-0.4	-0.8	Hemibos	Quaranwala	Pinjor	~2.0
F1140	1.9	-7.2	Hexaprotodon	Siswan	Tatrot	~2.5
DK1	0.1	-0.8	Hipparion	Devni Khadri	Tatrot	~2.5
A565	0.2	-4.5	Hipparion	Quaranwala	Pinjor	2.4
B30	0.7	-2.6	Rhinoceros	Quaranwala	Pinjor	~1.7
B-276	2.0	-2.0	Sivatherium	Quaranwala	Pinjor	~1.7
HPM1	-9.7	-3.8	Stegodon	Haripur Khol	Tatrot	2.5
DKS 1	-7.3	-7.6	Stegodon	Devni Khadri	Tatrot	~2.5
B-252	-11.2	-4.6	Stegodon	Quaranwala	Tatrot	3.0
A643	-1.9	-2.0	Theropithecus	Mirzapur	Boulder Congl.	~1.0
Samples older than 8 Ma						
HD1-C	-11.6	-2.7	Bovid	Hari Devi	Dhok Pathan	8.9
D1-1	-10.7	-1.2	Bovid	Dharamsala	Dhok Pathan	10.1
DP8	-13.2	-0.6	Cervid	Chob Ka Nala	Dhok Pathan	8.8
HT1-2	-11.5	-9.6	Dorcatherium	HT1	Dhok Pathan	9.1
K1D	-12.5	-3.6	Dorcatherium	Kursai-1	Dhok Pathan	9.2
HD1-1 average	(-12.0)	(-3.9)	Indopithecus	Hari Devi	Dhok Pathan	8.9
CHK-53	-11.0	-6.3	Propotamochoerus	Chob Ka Nala	Dhok Pathan	8.8
DP2	-13.1	-11.9	Rhinoceros	Kursai-1	Dhok Pathan	9.2
P11 average	(-12.6)	(-4.6)	Sivaladapis	HT1	Dhok Pathan	9.1
CK2	-12.1	-7.4	Sivalhippus	Chob Ka Nala	Dhok Pathan	8.8
K1S	-13.2	-7.3	Sivalhippus	Kursai-1	Dhok Pathan	9.2
HD 1–4	-12.6	-6.8	Sivalhippus	Hari Devi	Dhok Pathan	8.9



Table 3. Stable-isotope analysis results from paleosol carbonate nodules in the Dhok Pathan Formation, Haritalyangar.

Sample ID	$\delta^{13}{ m C}$	$\delta^{18} { m O}$	Locality	Age (Ma)
IN01-HP-301.1	-11.2	-7.6	Kursai-6	9.0
IN01-HP-301.2	-10.8	-8.1	Kursai-6	9.0
IN01-HP-317.1	-11.1	-7.5	Dangar-1	9.0
IN01-HP-317.2	-11.6	-7.5	Dangar-1	9.0
IN01-HP-318.1	-12.2	-6.7	Dangar-1	9.0
IN01-HP-318.2	-11.3	-7.8	Dangar-1	9.0
IN01-HP-319.1	-10.1	-8.0	Dangar-1	9.0
IN01-HP-319.2	-10.0	-9.2	Dangar-1	9.0
IN01-HP-320.1	-10.7	-8.1	Dangar-1	9.0
IN01-HP-320.2	-10.7	-7.9	Dangar-1	9.0
IN01-HP-325.1	-10.3	-8.2	Kursai-6	9.0
IN01-HP-325.2	-10.0	-8.3	Kursai-6	9.0
IN01-HP-326.1	-10.1	-7.5	Kursai-6	9.0
IN01-HP-326.2	-10.5	-7.2	Kursai-6	9.0
IN01-HP-327.1	-8.0	-5.8	Kursai-6	9.0
IN01-HP-328.1	-11.3	-7.2	Kursai-6	9.0
IN01-HP-328.2	-10.8	-7.6	Kursai-6	9.0
IN01-HP-330.1	-11.1	-7.4	Kursai-4	9.1
IN01-HP-330.2	-11.1	-7.2	Kursai-4	9.1
IN01-HP-302.1	-11.2	-7.7	K7	8.9
IN01-HP-302.2	-11.5	-8.3	K7	8.9
IN01-HP-303.1	-11.0	-7.7	Dharamsala	10.1
IN01-HP-303.2	-12.0	-7.5	Dharamsala	10.1
IN01-HP-304.1	-10.8	-7.2	Dharamsala	10.1
IN01-HP-304.2	-12.0	-7.1	Dharamsala	10.1
IN01-HP-306.2	-11.4	-8.8	Dharamsala	10.1
IN01-HP-307.1	-10.3	-8.9	Dharamsala	10.1
IN01-HP-307.2	-10.1	-9.2	Dharamsala	10.1

Fig. 9. (A) Indopithecus  $M^2$  (HDI-1) (scale 5 mm), and (B) scanning electron micrograph of the Phase II wear.

The isotope profile of *Indopithecus* (Fig. 12) shows an apparent annual variation in  $\delta^{18}$ O of ca. 3‰ with mean = -3.9‰; this range is attenuated from the primary "body water" range due to enamel maturation and sampling geometry (Passey & Cerling 2002). The actual range in  $\delta^{18}$ O of body water for *Indopithecus* was certainly higher than the measured 3‰, but without additional constraints on enamel maturation, we cannot provide quantitative estimates. *Sivapithecus* from Siwaliks of Pakistan had high  $\delta^{18}$ O values (mean  $\approx -2.6\%$ ), which was interpreted to mean that *Sivapithecus* obtained most of its water from enriched upper-canopy vegetation (Nelson 2007).

Microwear study of the *Indopithecus* molar shown a high incidence of pits (58%) and a low percentage (42%) of mixed scratches; although

Fig. 10. (A) Sivaladapis nagrii  $P_4$  (Hari I-6) and (B) scanning electron micrograph of the Phase II wear facet.

most pits are small, a few of the pits are very wide and long (Fig. 9). Such a high pit percentage indicates frugivory, whereas presence of large pits similar to those found in extant baboons, such as *Papio*, may suggest reliance on fallback foods like hard seeds, nuts, barks and fruits with hard cover. Among the extinct apes, *Sivapithecus* comes closest to *Indopithecus* in having a large number of pits, some large pits and mixed scratches. The Phase II wear pattern of the Greek ape *Graecopithecus* (= *Ouranopithecus*) (see Merceron et al. 2005: fig. 3D) also shows striking resemblance to the texture seen in *Indopithecus*.

*Indopithecus* was a terrestrial giant ape that lived in a  $C_3$ -dominated habitat. Based on microwear and morphology, it is thought to have lived in a forested habitat. Its main food source might have been low-lying fruits. It might have also exploited other resources such as seeds, nuts, barks or even roots. Possible reliance on fallen







Fig. 11. Scanning electron micrograph of the Phase II wear facets of (A) *Sivalhippus* from Hari Devi site, (B and C) *Dorcatherium* molar from HT 1 site, and (D) *Theropithecus* (A643) molar from Mirzapur site.



**Fig. 12.**  $\delta^{13}$ C and  $\delta^{18}$ O of multiple ablation samples along the growth axes of M<sup>2</sup> of *Indopithecus* (HDI-1). **A–G** represents ~14 months.

fruits and roots might explain presence of abrasive silica in its diet, which in turn would have produced wide scratches. However, its occasional reliance on bamboos or other  $C_3$  grasses cannot be ruled out. In earlier works it was hypothesized that Indopithecus giganteus (Gigantopithecus bilaspurensis) occupied an open country grassland landscape, which experienced aridity and seasonality (Simons & Chopra 1969, Pilbeam 1970, Gaur & Vasishat 1989). The estimated enormous sizes of Indopithecus (150+ kg) and Gigantopithecus blacki (300 kg) suggest terrestrial locomotion in these giant apes (Fleagle 1999). Earlier microwear and phytolith studies on the Chinese specimens (G. blacki) are interpreted to indicate a diet of fruits and bamboos, and living in subtropical to tropical forest seasonal environments (Ciochon et al. 1990, Daegling & Grine 1994, Jablonski et al. 2000). A recent carbon isotope and enamel microwear study of G. blacki concluded that this giant ape had a pure C<sub>2</sub> diet and lived in a forested habitat; eight specimens of Gigantopithecus blacki yielded the  $\delta^{13}$ C values between -13% and -16.8% (Zhao et al. 2011, Zhao & Zhang 2012). These highly <sup>13</sup>C-depleted values for G. blacki have been interpreted to indicate a close canopy forest habitat (following Cerling et al. 2004).

#### Sivaladapis

The estimated body-weight of *Sivaladapis* (3 kg) and its molar morphology indicates an ability to shear and puncture suggest folivory; *Sivaladapis nagrii* is interpreted as being an arboreal primate (Gingerich & Sahni 1984). White (2006) interpreted *Sivaladapis* as having a mixed frugivorous-folivorous diet.

The  $\delta^{13}$ C values of *Sivaladapis nagrii* range between -11.3% and -14%, indicating a pure C<sub>2</sub> diet. The range of the  $\delta^{18}$ O values for associated fauna is very large (from ca. 0% to -12%); the intermediate  $\delta^{18}$ O value for *S. nagrii*, averaging -4.6‰, suggests that an evaporated source (e.g., leaves) was an important component of its water budget. This is seen in a  $\delta^{13}$ C vs.  $\delta^{18}$ O plot (see Fig. 8) of all the mammal enamel and soil carbonates coming from > 8 Ma. It is clear that overall the body water  $\delta^{18}$ O values of the mammals, particularly those of the primates and bovids, were more enriched than the soil carbonate water values (Tables 2 and 3). Indopithecus and *Sivaladapis* are relatively enriched in  $\delta^{18}$ O compared to most other fauna (Fig. 8).

The Phase II grinding wear facet microwear pattern (Fig. 10B) on *Sivaladapis nagrii* premolars and molars shows a high incidence of pits (~60%) and a low incidence of striations  $(\sim 40\%)$ ; most striations are fine and long, oriented parallel and close to each other, formed due to a sideways (labio-lingual) jaw movement. However, many striations are coarse; the presence of coarse striations could be due to feeding on leaves with large phytoliths or even  $C_2$  grass (Mac Naughton et al. 1985, Ungar 1994). This characteristic pattern was observed in the trailing linear edges of lower pre-molars and molars of all the ca. 15 individuals examined. An abrasive diet is also evident from the prevalent exposure of the dentine due to the erosion of the thin enamel, in most of the specimens observed. This pattern implies that Sivaladapis nagrii was not a specialised folivore: rather it ate both fruits and leaves (White 2006). The tragulid Dorcatherium lower first molar from the Sivaladapis site (HT1) also shows a microwear pattern (Fig. 11B and C) that result from intake of some sort of abrasive diet as well, besides feeding on leaves and fruits. Obligate folivores such as Gorilla show similar parallel-oriented striations and a very low incidence of pits, but the striations are usually finer.

Soil carbonates and associated fauna from this time interval in the Dhok Pathan show a  $C_3$ -dominated ecosystem; as with *Indopithecus*, however, the relatively open nature of the habitat cannot be evaluated using stable isotopes because these fossils are older than the global  $C_4$  expansion.

Sivaladapids became extinct in the late Miocene and Sivaladapis nagrii was the last Siwalik species to disappear. Its last known occurrence is at the site HT-1 of Haritalyangar dated to 9.1 Ma (Pillans et al. 2005, Patnaik 2013). Late Miocene climate change and biotic exchanges have been hypothesised to have influenced the primate diversity in Asia (Barry et al. 2002, Zheng et al. 2004, Patnaik et al. 2005). A contemporary of *Sivaladapis*, the Miocene ape Sivapithecus was thought to have disappeared because of the spread of grasslands and deterioration of rainforest habitat during the late Miocene (Patnaik et al. 2005, Nelson 2007). It has also been suggested that sivaladapids might have lost their ground due to competition from invading cercopithecoids that arrived from Africa in the late Miocene (Gingerich & Sahni 1979, 1984, Kirk & Simons 2001).

#### Theropithecus

The *Theropithecus delsoni*  $\delta^{13}$ C and  $\delta^{18}$ O values of -1.9%, and -2.0%, respectively, are indicative of a C<sub>4</sub> diet. The extant gelada is a graminivore and C<sub>3</sub> grazer, and is limited to high altitudes in Ethiopia above the altitudinal C<sub>4</sub>–C<sub>3</sub>-transition zone (ca. 2700 m; Tieszen *et al.* 1979).  $\delta^{13}$ C values of Pleistocene *Theropithecus oswaldi* from eastern and southern Africa indicate a diet composed primarily of C<sub>4</sub> plants (Lee-Thorp *et al.* 1989, Lee-Thorp & van der Merwe 1993, Van der Merwe *et al.* 2003, Cerling *et al.* 2013).

Overall, the stable isotope values of other mammalian fossils collected from Mirzapur and surrounding areas indicate presence of grasslands during the Pleistocene time (Table 2). For instance, Equus, Hipparion, Hemibos, Sivatherium, Camelus and Rhinoceros have the  $\delta^{13}$ C and  $\delta^{18}$ O values indicating C<sub>4</sub> grazing under highly seasonal conditions. The Hexaprotodon  $\delta^{13}$ C and  $\delta^{18}$ O values indicate it was a highly water dependent  $C_4$ -grazer. Elephantids have both grazing and browsing to mixed-diet members: *Elephas* was a nearly pure C<sub>4</sub>-grazer based on  $\delta^{13}$ C values; the range of the  $\delta^{13}$ C values for Stegodon, -7.3‰ to -11.7‰, permits only a minor fraction of C4 plants in the diet and it is likely, therefore, that Stegodon was predominantly a C<sub>3</sub> browser.

*Theropithecus* microwear having a high incidence of long wide scratches and few pits also suggests grazing (Fig. 11D). The small mammalian assemblage from Pinjor Formation in general indicate presence of seasonal monsoonal climate (Patnaik 2003).

# Conclusions

Laser ablation stable carbon and oxygen isotope analysis was carried out on the lateral enamel surface of an isolated tooth of *Indopithecus*, a Late Miocene Siwalik giant ape. An SEM-based counting of growth lines (perikymata) sampled, and the number of daily cross striations (11) each perikymata contained, permitted us to resolve the individual's diet and body water over a ca. 1-year period (Fig. 12). The *Indopithecus* enamel  $\delta^{13}$ C values range between -12.4% and -11.6% (mean = -12%), which indicate a pure C<sub>3</sub> diet. In view of the attenuation of the isotope signal due to maturation of enamel (Passey & Cerling 2002), an apparent intra-annual variation in  $\delta^{18}$ O of ca. 3% may suggest a significant seasonality, comparable to present day monsoons (*see* Fig. 2). A dental enamel microwear study of *Indopithecus* molars show a high incidence (~60%) of pits similar to those found in dominantly frugivorous apes such as *Sivapithecus*. The presence of large pits and wide scratches on the molar of this giant ape, however suggest diet that included hard objects such as seeds, nuts, roots and fruits with hard cover.

The small primate *Sivaladapis nagrii* dental enamel  $\delta^{13}$ C and  $\delta^{18}$ O values (means being -12‰ and -4.6‰, respectively) and microwear showing both pits and striations suggest a C<sub>3</sub> diet comprised of leaves and fruits. However, the presence of coarse striations may indicate a diet of abrasive nature, not meant for its thin enamel.

All the taxa from ca. 8.8 to 10 Ma had  $C_3$ -dominated diets and paleosols from this time interval also indicate  $C_3$ -dominated ecosystems.

Stable carbon isotope and dental microwear data of *Theropithecus delsoni* reflect a primarily  $C_4$  grass diet; in the Plio-Pleistocene, the associated ungulate fauna in the northwest Indian Siwaliks had strongly  $C_4$ -dominated diet except *Stegodon*, which had a  $C_3$ -dominated diet.

#### Acknowledgements

We thank Dr. Anu Kaakinen (the editor) and Drs. Juha Karhu and Alistair Evans (the reviewers) for their valuable suggestions on the manuscript. We thank Dr. Yuri Kimura for helping us with photography of the sivaladapid specimen. RP learned the techniques of dental microwear and microstructure at the Max-Plank Institute of Evolutionary Anthropology, Leipzig and is grateful to J. J. Hublin for the opportunity. He would like extend his gratitude to Tanya Smith, Sireen El Zaatari, A. Olejniczak and M. Skinner for their kind help. JGF and RP thank Wenner-Gren Foundation grant Gr.ICRG-43 for financial support. Ashok Sahni generously provided some primate specimens for the present study. RP and TEC would like to thank David Cameron and Australian Research Council for funding collection of mammalian and soil carbonate samples from Haritalyangar. RP received financial support from DST SR/S4/ES-171/2005 and PURSE grants.

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