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## Morphology of the Gastrointestinal Tract in Primates: Comparisons With Other Mammals in Relation to Diet

DAVID J. CHIVERS and C.M. HLADIK Sub-Department of Veterinary Anatomy, University of Cambridge, Tennis Court Road Sub-Department of Veterinary Anatomy, University of Cambridge, Tennis Court Road,
Cambridge, England, and C.N.R.S., Museum National d'Histoire Naturelle, Brunoy, France

ABSTRACT Three categories of dietary adaptation are recognized-faunivory, frugivory, and folivory-according to the distinctive structural and biochem ical features of animal matter, fruit, and leaves respectively, and the predominance of only one in the diets of most species.
Mammals subsisting mainly on animal matter have a simple stomach and colon and a long small intestine, whereas folivorous species have a complex stomach and/or an enlarged caecum and colon; mammals eating mostly fruit have an intermediate morphology, according to the nature of the fruit and their tendency to supplement this diet with either animal matter or leaves. The frugivorous group are mostly primates: 50 of the 78 mammalian species, and 117 of the 180 individuals included in this analysis are primates.
Coefficients of gut differentiation, the ratio of stomach and large intestine to small intestine (by area, weight, and volume), are low in faunivores and high in folivores; the continuous spread of coefficients reflects the different degrees of adaptation to these two dietary extremes,

Interspecific comparisons are developed by allowing for allometric factors. In faunivores, in which fermentation is minimal, the volume of stomach and large intestine is related to actual body size, whereas these chambers are more voluminous in larger frugivores and mid-gut fermenting folivores; fore-gut fermenters show a marked decrease in capacity with increasing body size. Surface areas for absorption are related to metabolic body size, directly so in frugivores; area for absorption is relatively less in larger faunivores and more in larger folivores, especially those with large stomachs.
Indices of gut specialization are derived from these regressions by nonlinear transformation, with references to the main functional features of capacity for fermentation and surface area for absorption.

These are directly comparable with the dietary index, derived from quantitative feeding data displayed on a three-dimensional graph, with all species within a crescentic path from $100 \%$ faunivory through $55-80 \%$ frugivory to $100 \%$ folivory, perhaps illustrating, at least for primates, the evolutionary path from primitive insectivorous forms through three major ecological grades.

Recent field studies of primates have produced major advances in our understanding of their feeding behaviour and diet (CluttonBrock, '77; Chivers and Herbert, '78; Hladik, 75). The introduction of quantitative assessments of diet in these field studies allows precise comparisons between food intake and the morphology of the gastro-intestinal tract.

Detailed descriptions of gut morphology in mammals (Mitchell, '05; Bolk et al., '39;

Grassé, '55; Hill, '58) have also lacked effective quantification. The first quantitative effort at comparing gut morphology with diet (Cuvier 1805) produced no obvious correlations. Magnan ('12), however, working with areas rather than lengths, demonstrated a correlation be tween structure and diet (in general terms), but data are not presented so that his conclu sions can be verified. Preliminary efforts at a more critical amalysis (Fooden, '64; Hladik,
'67) showed interesting relationships among primates, but data on diet were still inadequate.
Our aims in this paper are 1) to describe various features of gut morphology with greater precision and quantification, 2) to present data from our field and laboratory studies, 3) to account for allometric factors in the discussion of interspecific differences, and 4) to compare these data on morphology with what is now known about the feeding ecology what is now known about
The combination of data on primates with The combination of data on primates with
those on domestic and other mammals is useful, because it allows a group of closely-related pecies with considerable dietary flexibility to species with considerable dietary fexibility to be contrasted with others which have become
highly specialized for markedly different diets. highly specialized for markedly different diets. Wract is fairly homogeneous among the differdract is fairly homogeneous among the diflerell ders of mammals, there have been par in variousopments of different parts of reflect adaptations to different foods, which can be classified into three major groups, according to structure and biochemical composition, and the resulting digestive requirements:

1) "Animal matter," including invertebrates, fish, and other small vertebrates from the secondary production of the ecosystem, which provide sources of protein and fat that are easily digested and, therefore, require a relatively short and simple gut.
2) "Fruits," including unripe (e.g., flowers) and ripe (fleshy) parts, seeds, and tubers mostly the reproductive parts of plants which are foods containing short-chain sugars that are hydrolyzed rapidly in tracts of large intestinal area for rapid absorption and immediate use.
3) "Leaves," including young and mature leaves, grasses, stems, as well as barks and gums- the vegetative parts of plants-which are foods usually containing protein and longchain sugars that require fermentation in an enlarged stomach or large intestine.
According to the predominant items consumed, three categories of dietary adaptation may be recognized, and in this paper they are referred to hereafter as faunivore, frugivore and folivore respectively (recognition of insectivore, carnivore, and herbivore, with their taxonomic and other connotations, contributes little to this analysis). These categories represent a gradation, for a generalized mammal, from foods that are relatively difficult to col lect but easy to digest (prey), through those
available in limited quantity (fruit), to those that are widely abundant but relatively diffi cult to digest (leaves). Hence the need for marked differentiation of feeding strateg $y$ and gut morphology. A classification in terms of three dietary grades (Hladik, 78 a ), with appropriate subdivisions, allows greater flexibil propriate subdivisions, allows greater fexibilty, and seemes of greater admixture of the ionary stages of great

COMPARATIVE ANATOMY OF THE GASTRO-
INTESTINAL TRACT
The structure of the wall of the gastro-in testinal tract follows a pattern common to all vertebrates: the inner lining of mucous mem brane is separated by connective tissue from an outer cylinder of at least two layers of muscle. Variation in histological structure ef fects divisions into stomach, small intestine (duodenum, jejunum, ileum), and large intestine (caecum and colon). Brief reference wil be made to various configurations of the mu cosa and underlying connective tissue, which apparently assist digestion mechanically, by mixing or slowing the passage of food or by increasing the surface area for digestion and absorption, e.g., papillae, rugae (folds), haus trae (sacculations), villi,
In this section we shall try to identify those structures relating to each of the three main dietary adaptations by supplementing previous knowledge with new observations. The latter are made from relaxed guts immersed in water and positioned to show the main features clearly; a complete reconstruction impossible by photograph, is achieved by mov ing parts of the tract while drawing, and adjusting the dimensions of each region after dissection and measurement.

## Faunivores

The basic pattern of gut structure among faunivores consists of a simple globular stomach, tortuous small intestine, short conical caecum, and simple smooth-walled colon. This pattern is exhibited by primates feeding mainy on invertebrates, such as Arctocebus (Fig. 1), Loris, and Tarsius. In other mammals there may be structural specialization in one direc fion or another. The smallest mammalian gut known is found in the insectivorous bat, Rhinopome; its tract is only four-fifths of body length (Grassé, '55). Simplification of the gut is extreme in haemophagous bats, such as is exmodus, with the stomach as a blind endin tube a very short colon, and no caecum. Such
reductions are clearly specializations, rather a "muscular tooth" compensating for the lack than representing the primitive condition. Specializations of faunivores may also involve the stomach. Some ant-eating edentates and the gut is only seven times body length, but the stomach contains of oral teeth (Grassé, 55). A similar muscular specialization is found in pholidotes, such as the termite-eating pangolin, Manis (Fig. 1), supplemented by a keratinized area in the pylorus and by the presence of small stones.


Fig. 1. Gastro-intestinal tracts of faunivores, drawn by C.M.H., with accurate scaling of proportions, main flood vessels to show the disposition of mesenteries, and conventional shading of the different morphological in terms of morphology. The pangolin, Manis gigantea (specimen MR, juvenile) is presented below with an open stomach to show the muscular "tooth;" the arrow marks the junction of ileum and colon, determined from microscopic examination of the mucosal wall: the extreme length of the small intestine did not allow it to be
drawn completely unfolded, as in other drawings.

In cetaceans the stomach has three main compartments (Harrison et al., 70). The first and largest is covered by folds of thick, kerainized epithelium, the second by spirally ar ranged folds of thick, glandular epithelium making a direct channel along the lesser curvature), and the third, tubular compart ment has a simple pyloric mucosa and strong phincters at both ends, e.g., the porpoise Phocaena (Fig 2). The small intestine, start from a dilated duodenum, is very long ing from a dilated duodenum, is very long about 10 cm , identified und a very short colon conly 10 cm , identified under the dissecting microscope
In the Insectivora the simple stomach is followed by a very short small intestine and, followed by a very short small intestine and usually, no caecum-as illustrated by Potomogale (Fig. 3), which is adapted for feeding on freshwater fish and crustaceans. In Sorex the tract is only 2.6 times body length, and some faeces are reingested to permit a second opportunity for digestion. This phenomenon o eflection (Crowcroft, '52) helps to explain the reduction in gut size as a physiological/behavoral specialization. In one species of Tenrec which eats foods other than insects, the tract may be seven times body length. Tree shrews, Tupaia, which also supplement their invertebrate diet with fruit, have a slightly larger colon than other insectivores and a small caecum (Grassé, '55). Some rodents subsist on diet composed entirely of insects or other nimal prey In the African murid Lophuroys, for example, specialization to such a diet pos a chan in the nchud (Genest-Villard 1968): llard, 1968)
Other mammals, such as some feeding on ertebrates, show no obvious specialization. In fissipedes, for example, the stomach is simple, the small intestine four to six times body ength, the caecum small or absent, and the colon reduced, as shown in the Viverridae by the African linsang Poiana (Fig. 3) and the mongoose Atilax, and in the Felidae by the golden cat Profelis (Fig. 4). The shape, inter nal features, and relative sizes of fundus and pylorus vary slightly among such mammals, as described by Ellenberger and Baum ('21 and illustrated here by the domestic dog, Canis (Fig. 5).

## Frugivores

This group contains most primates, but none of them subsists entirely on fruit. All frugiores supplement their diets with varying amounts of insects and/or leaves, but have no
distinctive structural specialization in the gut, although its morphology may show considerable variation between species

Some Carnivora also have this mixed diet, but retain the structural features of faunivores, e.g., the palm civet Nandinia feeds heavily on fruit (Charles-Dominique, '78), but has no caecum and a reduced colon (Fig. 6).
Myoxid rodents also have no caecum (Grassé, '55), and their predation on birds, as a supplement to seeds and fruit, places them on the border between faunivores and frugivores. In the stomachs of cricetine rodents, the vores. fundus ar dimensions, in their dimensions, separated by a fold of varying shape (Carleton, in). In frugivorous bats the stomach is relatively complex, with a distinct cardial region, a long pyloric diverticulum folded back on itself, and a lateral "caecum"; the true caecum is present in several genera (Grassé, '55).
Among artiodactyls, the pigs have a stomach that is clearly divided into zones, and in some cases into compartments; they have an especially long small intestine, a large caecum, and a relatively complex colon, so that the whole tract is about 20 times body length. These elaborations relate to the inclusion of roots and other vegetative parts of plants in their diet.

Gut structure is more homogeneous among frugivorous primates (Figs. 7, 8, 9). The stomach is essentially simple and globular in structure (Hill, '58). Marmosets show some elongation of the fundus, whereas those of cebids are more specialized with a globular fundus, conical body, and cylindrical pylorus Aloust conical body, and cylindrical pylorus. Alouatby weight Hodik and Hladik '69), shows the by weight, Hadik and Hadik, 69), shows the greatest complexity, with a capacious globular sac, narrowing towards the bent tubular pylorus, which is guarded by strong pillars; rugae radiate from the carda and run longitudinally within the body. Ateles, which is one of the most frugivorous and swallows many stones, has an enlarged J-shaped stomach. Old World primates, other than colobine monkeys, have a single smooth-walled sac; among the apes it is more globular and man-like in gibbons, even more globular in gorillas, and more elongated in chimpanzees and orangutans (Hill, '58).
The duodenum is commonly C-shaped, in contrast to the elongated U-shape of other mammals; in some cebids and all catarrhines it is retroperitoneal. The caecum is large in frugivorous prosimians, short and wide in
marmosets, and hook-shaped in cebids; in catarrhines the base is globular, the body shor and capacious, and the apex blunt and conical, with a terminal vermiform appendix in hominoids (Hill '58)
The colon is simple and straight in cebids such as Saimiri; there is a transverse colon in Cebus and Aotus, and a right colon as well in Callicebus, Cacajao, and Pithecia. Further elongation (and folding) occurs in callitrichids, agothrix, and all catarrhines (Hill, '58 Taenia coli (reduction of longitudinal muscle
into bands) are lacking in Saimiri, Cebus, and most prosimians, but there may be one or two in Nycticebus, Perodicticus (Fig. 6), Lemur and callitrichids, and cebids and most catarrhines otherwise have three, except for gibbons with four (Hill, '58). The ansa coli loop in the transverse colon is common in prosimians (Fig. 6); this part of the colon is also long and dependent in apes. The capacious colon of gibbons (Fig. 10) is indicative of considerable leaf content in the diet and its potential for fermentation.


Fig. 2. The compound stomach of the harbor porpoise, Phocaena phocaena (D35), is shown open and flattened in a dissecting tray. The oesophagus (center) leads right into the first compartment, which in turn opens into the glandular chamber (lower); the first sphincter, opening into the pyloric tube, is just visible
(lower center) leading up to the pyloric sphincter and thence into the duodenal diverticulum; the mucosal folds, which run the lengtt of the intestine, can be seen (upper left). Photo by D.J.C. and Department of Anatomy,
Cambridge University.


Fig. 3. Gastro-intestinal tract of Potamogale velox (MX) presented as in Figure 1, with the arrow marking the junction of small intestine and colon. In Poiana richardsoni (MS), to the right, the limit of the colon is clearly marked by a short caecum. Drawings by C.M.H.


Fig. 4. Gastro-intestinal tracts of Atilax paludinosus (MW) and: on the right: Profelis aurata (MZ). Drawings by C.M.H.


Fig. 5. Internal view of the stomach of the domestic dog, opened around the greater curvature, showing the oesophageal apening intact (at Anatomy, Cambridge University.

## Folivores

The long-chain $\beta$-linked carbohydrates predominant in the leaves, grasses, stems, barks, and gums consumed by these animals require and gums consumed by these animals require considerable degradation organisms. The most conspicuous adaptations are chambers for the bacterial fermentation of cellulose and for the absorption of tation of cellulose and for the absorption of volatile fatty acids and other metabolites, either in the stomach or in the large intestine. This dichotomy might mask further diversification as shown by the expansion of the right colon as well as, or instead of, the caecum, the presence or absence of caecotrophy, and variation in stomach structure

The large intestine is enlarged in those prosimians which feed on leaves or gums. In

Lepilemur a mechanism similar to refection (see above) allows efficient use of a diet very high in fiber content (Hladik and CharlesDominique, 74). This case of caecotrophy is Dominique, why the ill intestine is of the shortest why the (Fig 11) An the among manals. gated and and Euolicus. Since gums require fermentation for digestion they are classifled with folivores, along with Indri, which shows similar features (Fig. 11) and is a true folivore.
The rabbit provides the classic case of caecotrophy (Morot, 1882; Taylor, '40). In lagomorphs and myomorph rodents some faeces are reingested after fermentation in the capacious caecum, so that metabolites from the
herbivorous diet can be absorbed in the small ntestine.
The caecum is very coiled and elongated in specialized folivores, such as the "gliding" squirrel Anomalurus (Fig. 11)-even more so than in Lepilemur. The most complex large intestine is found in Dendrohyrax (Fig. 12), where the first caecum is followed by two more after about 20 cm of colon.
With enlargement of the colon in mammals the migration of the ileo-caeco-colic junction can be traced from the left cranial part of the abdominal cavity round to the right cauda aspect, so that the caecum comes to poin caudally rather than cranially (Hill, 58), In those species with a voluminous caecum, however, cranial rotation has occurred so that it comes to occupy the ventral part of the abdomen, as in the horse (Fig. 13). Perissodactlys and proboscids have large colic loops in addi-
tion to the huge sacculated caecum for the breakdown of their fibrous diet. As in other mammals which cope with this kind of diet the horse has a large area of keratinized epithelium in its stomach, which, however remains simple (Fig. 14). Carleton ('73) sug gests that the variable cornification of the stomach lining in different species of cricetine rodents might be correlated with the amount of cellulose in the diet
In contrast to perissodactyls, proboscids have a large folded stomach and a short small intestine of large internal area. Sirenians, such as the dugong, have a complex two-chambered stomach, with one part fulfilling the role of the duodenum; they also have a very wide caecum (Grassé, '55).
The most elaborate tracts are found in those folivores, usually subsisting almost entirely on grasses, with complex stomachs for bacte-


Fig. 6. Gastro-intestinal tracts of frugivores. On the left, from Perodicticus potto (FM), a frugivorous prosimian feeding Fig. 6. Gastro-intestinal tracts of an animal matter. On the right, the palm civet, Nandinia binotata (MY is a carnivore feeding mainly on fruit; cum, the junction of colon and small


Fig. 7 Gastro-intestinal tracts of frugivorous monkeys, Above, the mangebey Cercocebus albigena (FF, juvenile), and below, the guenon Cercopithecus cephus (FD). Drawings by C.M.H.
rial fermentation, as exemplified by the artiodactyl ruminants. Macropod marsupials some edentates, hippopotami, camels, and colobine monkeys show evolutionary conver gence with ruminants in their adaptations of stomach structure for folivory (Moir, '68). In these groups there is actually a continuum of diets from frugivore to folivore, as shown in the preceding section for pigs and peccaries whose stomachs show some similarity to
those of ruminants). Among the ruminants, for example, there are pure frugivores, such as Cephatophus and Hyemoschus (chevrotain), intermediate types such as the spotted deer Axis, and pure folivores, such as Neotragus, or pure herbivores, such as the buffalo Syncerus (G. Dubost, pers, comm.). These extremes of the continuum are the most specialized forms Macropod marsupials have a long tubular stomach sacculated along much of the greater


Fig. 8. The disposition of the gastro-intestinal tract within the abdomen of the long-tailed macaque, Macaca fascicularis (P21). Note small size of stomach (a
left), and loops of colon, with taenia coli (center). Photo by D.J.C.
rvature, with an oesophageal groove (rumnant feature linking oesophagus with omasm) (Grassé, '55). The stomach leads into a ong intestine with a wide caecum. Among livorous edentates, such as the sloth Bra-有 there is a keratinize spall "rumen" with two diverticula and an esophageal groove, and an "abomasum" with an expanded pyloric region with a very thick
muscular wall.
The hippopotamus has the oesophagus opening into a vestibule, into which open two unequal diverticula, and which leads into a third tubular chamber; all three chambers have stratified epithelium thrown into projecting folds with numerous papillae. There is a very long intestine, but no caecum. Camels have a stomach that is smooth and ovoid in


Fig. 9. Gastro-intestinal tract of the barbary macaque, Macaca sylvana (FO), showing a rather larger colon than occurs in other frugivorous pimates, which correlates with a diet including large amounts of plant matter. Drawing by C.MH.
hape, composed of two glandular sacs; the the sac, than in Asian colobines, where the omasum and abomasum are merged into a single tube
Colobine monkeys have a similarly large and complex stomach, with much distension and sacculation proximally and a U -shaped tube distally, sacculated along the proximal part of the greater curvature (Hill, '58). These sacculations are produced by the reduction of ongitudinal muscle into two or more bands taenia). The stomach of African colobines is more elongated, with the tube bent back on
sac is roughly spheroidal (Fig. 15, cf. Kuhn, 64). The colon is long and sacculated, and the caecum is of moderate size
The artiodactyl ruminants are well known for their four-chambered stomach (Comline et for their four-chambered stomach (Comline et
al., '68), which is dominated by the vast rumen, divided into dorsal and ventral sacs by muscular pillars, and covered by keratinized squamous epithelium with papillae of varying size and shape (Fig. 16). The oesophagous


Fig. 10. On the left, an internal view of the stomach of the siamang, Hylobates syndactylus (P27), opened around esser (to left) and greater curvatures and laid out in a dissecting tray; the oesophagus opens at the upper left, with a dark region of cardial glands, and the pyloric sphincter is at the lower left. On the right, an external view of the siamang's large sacculated colon, with taenia coli, partly distended with, and immersed in, water; the ileum is clamped by forceps
in the upper right, where the vermiform appendix projects down from the caecum. The lower end of the left (descending) colon is clamped in the lower right. The large volume relates to the large intake of leaves in its diet. Photos by the Department of Anatomy, Cambridge Unviersity
opens into a much smaller reticulum, which has a distinctive honeycomb pattern of ridges hexagonal in cow and sheep, pentagonal in goats) and is covered by small conical papillae. The rumen connects with the glandular part of the stomach through the small ovoid omasum, which is partitioned by many leaves of varying size for water absorption. The interhal surface of the glandular abomasum is thrown into folds throughout the fundic region (Fig. 16). The intestine is again very long, the caecum is relatively short, and the colon is ong and elaborately flexed and coiled.
Efforts at demonstrating homologies with the ruminant stomach of bovids have had
limited success. For example, the stomach of New World camelids has only three compartments, with the ventricular groove running from the first to the last; only the terminal fifth of the third, tubular compartment has true fundic and pyloric glands (Vallenas et al 71). While this chamber has mucosal pleat over much of the rest of its length, the firs two sac-like compartments have areas of large glandular saccules, which not only contain considerable amounts of ingesta, but are capable of frequent eversion. Thus, they seem more likely to contribute secretions to buffer stomach contents, rather than to absorb water. It is claimed that such structures aid


Fig. 11. Gastro-intestinal tracts of folivorous prosimians (left) and a rodent with extreme development of the caecum related to their specialized diets. The sportive lemur, Lepilemur leucopus (DO), upper left, has the shortest small intestine of all primates; it is the only genus in which caecotrophy occurs and the ileo-caeco-colic "plate" (arrowed) probably plays an important role in regulating this behavior (Charles-Dominique and Hladik, '71). The needle-clawed bush-baby, Galago chain carbohydrates, that also require fermentation. The flying squirrel, Anomalurus fraseri (MT) has a similar gut norphology related to a diet known to be mainly leaves. Drawings by C.M.H.
in a greater efficiency of digesting poor-quality egetation at high altitudes, where cattle and goats cannot graze.
Janis ('76) suggests that horses also have an advantage over cattle in their ability to a more fibrous diet of low protein content都 pidly, rough more ing more effient digestion of cellulose. In dis issing the evolutionary strategy of equids, then ecology, she conrasts their digestive system with that of rumnants, and refers to the greater extension of bivores that do not practice caecotrophy.

In conclusion, this review of the principa distinguishing features in the mammalian gastro-intestinal tract has emphasized the simple stomach and long small intestine of nammals known to subsist mainly on anima matter, and the elaboration of the stomach and/or small intectine in leaf or grass eatin forms, with frugivores showing an intermedite morphology (Table 1) Most mammalian eatures of gut mor logy, except for the eature gut more specialized, occur ang primates, wich forray dond whed nspecialized forms and which have not a ained the extreme adaptations found in other


Fig. 12. Gastro-intestinal tract of the tree hyrax, Dendrohyrax dorsalis (MU), showing the most complex arrangement of eaeca and colon, the exact functions of which are not yet known. Drawing by C.M.H.

QUANTITATIVE ANALYSIS OF GUT MORPHOLOGY Methods
Gastro-intestinal tracts were taken from 180 individuals of 78 mammalian species in England, France, Morocco, Gabon, Madagascar, Sri Lanka, Malaysia, and Panama. There are 117 primates of 48 species, 13 temperate mammals of 7 species ( 2 aquatic), and 24 tropical nonprimate mammals of 17 species. One-hundred forty-eight specimens were caught in their natural habitat by hunters during pest control operations or by local people for food; 29 animals, mostly primates, died in captivity, from illness or old age. In addition, 26 domestic mammals of six species were put down during routine marketing, research, or teaching operations:
Larger samples of certain species indicate the level of intraspecific variation, and the


Fig. 13. The large intestines of the domestic horse (D14) showing, above, external shape and large size of the caecum center) surrounded by dorsal and ventral loops of the primitive right colon, with sternal and diaphragmatic flexures (lower and the smaller size of the transverse and left colons, also with taenia coli (lower left corner). Below, the internal appearance of the caecum after opening and washing, before cutting the taenia coli, which increases the length from 80 to 240 cm . Photos


Fig. 14. Internal aspect of the stomach of the domestic horse (P14), opened around the greater curvature to show the large extent of folded, keratinized mucosa around the oesophagus (ef. Figure 2) and up into the saccus caecus (above), the dark, fundic mucosa (to left and right), and the pyloric mucosa (below and center). The sites of attachment of bot-fly Department of Anatomy, Cambridge University
reliability of small samples. Thirty-seven of the 78 species are represented by only one specimen, 15 species by only two, and 10 species by three individuals. There are four specimens of Galago (Euoticus) elegantulus, G. alleni, G. demidovii, Alouatta palliata, Cercopithecus cephus, C. nictitans, Presbytis obscura, and the domestic goat; five specimens of Arctocebus, Cheirogaleus, Miopithecus, Vulpes, and Dendrohyrax; six specimens of Presbytis melalophos and the domestic cat: and nine domestic dogs.

Specimens were weighed intact, which was not always possible in the field, and their lengths were measured from bregma to ischium and from tip of nose to base of tail. The latter measure of length was not used in this
analysis, because of the distortion introduced by varying lengths of muzzle, especially when contrasting primates with other mammals. The guts of most specimens were examined and measured in the fresh state (or were preserved in a saturated saline and then preserved in a saturated saline and then washed water son later study in $10 \%$ formol saline, but meas-
urements under these conditions are affected adversely by contraction at the time of fixaadversely by contraction at the time of fixation. Many specimens were examined, drawn, displayed under water in a large dissecting tray. The dimensions of each region were then measured, for calculations of area and volume, and weighed after the removal of excess moisture.


Fig. 15. The gastro-intestinal tract of the dusky leaf monkey, Presbytis obscura (P19). Upper right, the disposition within the abdominal cavity; note the large size of the stomach occupying the upper half of the view, and the coils of colon below cef. Fig. 8). Upper left, the stomach (partially distended with water) displayed to show the large sac, the gastric tube (on the
right), and the pylorus (lower left). Below, the complete abdominal part of the tract, with a different aspect of the left, and the coils of small intestine, caecum (directed downwards), and colon leading around into the rectum stomach on o the right. Photos by D.J.C.


TABLE 1. Summary of adaptations of the gastro-intestinal tract in various orders of mammal, according to the predominance of either animal matter or fruit or leaves in their diet

| Dietary <br> Category | Mammalian Order | Features of Gastro-intestinal Tract |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | General | Stomach | Small Intestine | Caecum | Colon |
| Faunivores |  |  |  |  |  |  |
|  | Insectivora | $21 / 2-7 \times \mathrm{BL}^{1}$ | simple | very short | none |  |
|  | Chiroptera | - $4 / 5 \mathrm{BL}$ | blind-ending tube |  |  | very short |
|  | Primates |  | simple, globular | tortuous | short, conical | simple, smoothwalled |
|  | Carnivora |  | simple | 4-6×BL | small or absent | reduced |
|  | Edentat | $7 \times$ BL | "muscular tooth" |  | none |  |
|  | Pholidota |  | "muscular tooth," cornified, stones |  |  |  |
|  | Cetacea |  | 3 chambers, 2 sphincters | very long | none | very short |
|  | Rodentia |  | gastric gland changes |  |  |  |
| Frugivores |  |  |  |  |  |  |
|  | Chiroptera |  | rel. complex, distinct cardial region, long pyloric divertic., lat. 'caecum' |  | small or absent |  |
|  | Primates |  | simple, globular | C-shaped duodenum retroperitoneal | small-large | elongated, folded, taenia |
|  | Carnivora |  |  |  | none | reduced |
|  | Artiodactyla |  | zones, even chambers | very long | large | rel. complex |
|  | Rodentia |  | extensive cardial glands, separated from fundus by fold |  | none |  |
| Folivores |  |  |  |  |  |  |
|  | Marsupalia, macropod |  | long, tubular, sacculated, grooved | long | wide |  |
|  |  |  |  |  |  |  |
|  | lemurid colobine | caecotrophy | simple <br> sacs and tube | very short | elongated, coiled |  |
|  | Proboscidea |  | large, folded | short, capacious | huge, saccul. | large loops |
|  | Sirenia |  | 2-chambered |  | very wide |  |
|  | Hyracoidea |  |  |  | 3 caeca | most complex |
|  |  |  | cornified area |  | huge, saccul. | large loops |
|  | Artiodactyla |  |  |  |  |  |
|  | hippopotamid |  | 3 chambers | very long | none |  |
|  | NW camelid |  | 3 chambers |  |  |  |
|  | OW camelid bovid |  | 2 sacs, smooth, ovoid |  |  |  |
|  | bovid | most elaborate | 4 chambers, huge elaborate | long | large | long, folded and coiled |
|  | Edentate |  | cornified cardia, 'rumen', groove, 'abomasum' |  |  |  |
|  | Rodentia | caecotrophy | cornified variably |  | capacious $\mathbf{v}$. coiled, long |  |
|  | Lagomorpha | caecotrophy |  |  |  |  |

Techniques were standardized throughout between the two authors, on occasions when they worked together, so as to obtain comparable accuracy. Several hours were allowed to elapse after the death of the individual to permit complete relaxation of muscle tone in the gut wall. Measurements of length and breadth of stomach, small intestine, caecum, and colon were then made without stretching, after opening and flattening the gut wall after opening and fattenis the gut wall, usually under water in a dissect Beray (except for the larger specimens). Because different parts of the gut can be fully contracted or relaxed, simultaneously or sequentially, this seems to be the best compromise in functional terms for measuring, for comparative purposes, what is a very malleable system.
The surface areas of small and large intestines were calculated from lengths and a series of breadths; sometimes it was more appropriate to treat the caecum as a triangle rather than an elongated rectangle. The irregular shape of the stomach required summing the area of its parts, usually arranged to cover the different compartments or division into fundus and pylorus. The areas of such nontubular parts were also measured by cutting pieces of aluminum foil to the exact shape of the part(s) immersed under water, and then weighing for
calculation from the weight of unit area; this provided a means of checking the accuracy of ength and breadth measurements. Error re sulting from the different methods, or from repeated measurements, amounted to les than $5 \%$.
Little merit was placed on measuring vol umes by distending parts of the gut with water (even if the pressure could be controlled), if only because of the possible distortion of sub equent measurements of area. equer measmerative measurements were made in this way (Table 2) Usually, small and large intestines were considered as cyl inders for the calculation of volumes $(V)$ from their surface area $(A=b \times 1)$,

$$
\mathrm{V}=\pi\left(\frac{\mathrm{b}}{2 \pi}\right)^{2} 0.1=3.142\left(\frac{\mathrm{~b}}{6.283}\right)^{2} 0.1,
$$

and stomachs were treated as spheres,
$\mathrm{V}=\frac{4}{3} \pi\left(\sqrt{\frac{\mathrm{~A}}{4 \pi}}\right)^{3}=4.19\left(\sqrt{\frac{\mathrm{~A}}{12.57}}\right)^{3}$ (Table 3).
For stomachs, similar results were obtained in a few cases by calculating the volume from the greater curvature, assuming its length (L) to be the circumference of a sphere,

$$
\mathrm{V}=\frac{4}{3} \pi\left(\frac{\mathrm{~L}}{2 \pi}\right)^{3}=4.19\left(\frac{\mathrm{~L}}{6.283}\right)^{3} \text { (Table 3). }
$$

Volumes calculated from surface area are about $12 \%$ less on average than those from greater curvature or water-filling (Tables 2 and 3).

While all species can be compared against a common standard (calculating volumes from spheres of equivalent surface area), distortions occur in the case of species with a complex
stomach, where the whole clearly does not approximate a sphere. While some compartments resemble spheres (the ruminant reticulum, rumen and omasum, and the colobine presaccus and saccus), others approximate cylinders (the ruminant abomasum and the colobine gastric and pyloric tubes). Thus, volumes have been recalculated along these lines (Table 4), yielding values one-third less on average. Even the calculations of volumes of simple stomachs (whether from surface area of length of greater curvature) give variable

TABLE 3. Estimation of stomach volumes: considering the stomach as a sphere and calculating volume from (a) length of greater curvature ( $L$ ), equivalent to circumference and (b) surface area (A)

| Species | Volume, $\mathrm{cm}^{3}$ |  |  |
| :---: | :---: | :---: | :---: |
|  | Greater curvature | Surface area | latter ef. former $-\quad+$ |
| Prosimians |  |  |  |
| Arctocebus calabarensis | 9 | 5 | - |
| Avahi laniger | 12 | 19 | + |
|  | 23 | 39 | + |
|  | 46 | 62 | + |
| Cheirogaleus major | 9 | 10 | $=$ |
|  | 9 | 14 | + |
|  | 5 | 9 | + |
|  | 17 | 24 | + |
| Galago alleni | 17 | 8 | - |
|  | 17 | 9 | - |
|  | 4 | 6 | + |
| Galago demidovii | 2 | 2 | $=$ |
|  | 2 |  | - |
|  | 1 | , | = |
|  | 3 | 2 | - |
| Lepilemur mustelinus | 17 | 28 | + |
| Perodicticus potto | 2 | 3 | + |
| New World monkeys |  |  |  |
| Leontocehus midas | 4 | 4 | $=$ |
| Cebus griseus | 37 | 53 | + |
|  | 29 | 31 | + |
| Alouatta seniculus | 205 | 227 | - |
| Ateles paniscus | 264 | 212 | - |
| Old World monkeys |  |  |  |
| Miopithecus talapoin | 29 | 27 | - |
|  | 46 | 42 | - |
|  | 23 | 32 | + |
| Cercopithecus neglectus | 264 | 163 | - |
| Cercopithecus nietitans | 135 | 93 | - |
|  | 57 | 59 | + |
|  | 297 | 221 | - |
| Cercopithecus uethiops | 29 | 19 | - |
|  | 6 | 5 | - |
| Cercocebus albigena | 135 | 130 | - |
|  | 12 | 15 | + |
|  | 135 | 82 | - |
| Macaca syleanus | 29 | 20 | - |
| Papio papio | 9 | 8 | * |
| Papio sphinx | 9 | 7 | - |
| Ape |  |  |  |
| Pan troglodytes | 1079 | 965 | - |
|  | 135 | 72 | - |
| Pan gorilla | 116 | 88 | - |

results, apparently according to the degree of elongation of the stomach spindle.

## Results

The data collected, and resulting basal calculations, are displayed in tables $5-8$ for prosimians, New World monkeys, Old World monkeys (cercopithecine and colobine), apes, domestic mammals, and wild mammals (tem perate and tropical). Emphasis is placed on eralt animals taken from their natural habi adult ane considen fom their natural hab at, since considerable changes in gross di mensions may occur in captivity, even afte hor perids, e.g., fom a 38 reduction to an
 mestine in some cercopitheeires (Hadik, 67 Immature individuals are particularly susceptible to dietary effects on gut proportions, and their measurements should be treated with caution; the stomach and colon are relatively reduced in young folivores, and may be in creased in faunivores. Fixed specimens are also liable to distortion from the functional state.
Gut Differentiation. The sizes of stomach and large intestine relative to small intestine, in terms of surface area, weight, and volume, provide a simple quantitative index of gut differentiation, without regard to the size of the animal. These coefficients of gut differeniation vary considerably in the mammal studied, from tracts that are dominated by small intestine in faunivores to those that ar dominated by stomach and/or large intestine in folivores.

Considering surface areas, the coefficients of gut differentiation show considerable overlap between species of the three main dietary categories (Fig. 17). This overlap is explained to some extent by intermediate diets. The plots on a logarithmic scale represent species means, with the range of variation marked for those species with four or more specimens. While the values of the coefficients appear generally to represent structural adaptations indicative of the relative proportions of animal and plant matter in the diet, only those below 0.2 can be regarded as true faunivores and only those above 3.0 as exclusive folivores.
Interspecific variation may be appreciated more clearly by comparing coefficients within (and between) the various taxonomic or ecological groups of mammal (Fig. 18). In each of these groups dietary categories, as suggested by structure, are separated, albeit in different ways, with values for frugivores clustered around 1.0. This is especially marked among primates, with higher values reflecting a tract dominated by stomach and/or large intestine (for digesting leaves), and the few lower ones, among prosimians and ceboids, where the small intestine predominates (for digesting animal matter). The categories of "faunivore," "frugivore," and "folivore" are established according to structural discontinuities, and at this stage they can be no more than suggestive of diet. The overlap between them would seem to result from interspecific variation in the degree of admixture of animal, fruit, and leaf foods, which would be especially variable among frugivores.

TABLE 4. Estimation of stomach volumes: complex stomachs, treated (a) as a

| Species | Volume, $\mathrm{cm}^{3}$ |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
|  | Single sphere | $\begin{gathered} \text { Spheres } \\ \text { \& } \\ \text { Cylinders } \end{gathered}$ | \% reduction of latter | n |
| Domestic mammals |  |  |  |  |
| Capra (goat) | 17,776 | 11,365 | 36 | 4 |
| Ovis (sheep) | 13,615 | 7,954 | 42 | 3 |
| Colobine monkeys |  |  |  |  |
| Presbytis cristata | 5,171 | 3,259 | 37 | 2 |
| Presbytis obscura | 4,124 | 2,767 | 33 | 4 |
| Presbytis melalophos | 3,917 | 2,313 | 41 | 5 |
| Presbytis rubicunda | 3,547 | 2,259 | 36 | 1 |
| Nasalis larvatus | 8,270 | 6,523 | 21 | 1 |
| Pygathrix nemaeus | 4,605 | 3,199 | 31 | 2 |
|  |  |  | $\overline{\mathrm{x}}=33$ |  |
| Colobus polykomos | 3,147 | 2,108 |  | 2 |
| Presbytis senex | 3,856 | 2,584 |  | 2 |
| Presbytis entellus | 5,532 | 3,706 |  | 2 |



COEFFICIENT of GUT DIFFERENTIATION
surface area of stomach + caecum + colon surface area of small intestine

Fig. 17. Coefficients of gut differentiation from surface area plotted in order of magnitude (smallest values below) on a logarithmic scale, indicating by arrows and stippling the three main morphological dietary categories and the overlap between them, which may or may not include species with intermediate diet. Range of values are denoted by horizontal lines for species with more than one specimen.



| Species | Ident. no. ${ }^{4}$ | Sex | Body length, cm | Body weight, gm | Surface area, $\mathrm{cm}^{2}$ |  |  |  | Weight, gm |  |  |  | Volume, $\mathrm{cm}^{2}$ |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  | Stom. | S.1. | Caec. | Colon | Stom. | S.I. | Caec. | Colon | Stom. | S.I. | Caec. | Colon |
| Cercopithecine monkeys |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Miopithecus, $t$ talapoin | P13 ${ }^{1}$ | M | 38 | 950 | 59 | 451 | 40 | 212 | 7 | 26 | 2 | 16 | 43 | 96 | 16 | 76 |
|  | P37 ${ }^{1}$ | F | 28 | 880 | 64 | 594 | 30 | 258 | 4 | 26 | 1 | 12 | 48 | 142 | 12 | 106 |
|  | AM | F | 30 |  | 59 | 157 | 26 | 214 |  |  |  |  | 43 | 14 | 18 | 81 |
|  | AO | F | 29 |  | 44 | 149 | 25 | 87 |  |  |  |  | 27 | 18 | 12 | 20 |
|  | B1 | M | 24 |  | 49 | 231 | 12 | 122 |  |  |  |  | 32 | 34 | 4 | 32 |
| Cercopithecus cephus | EM | F | 42 | 2650 | 187 | 784 | 76 | 371 |  |  |  |  | 240 | 218 | 46 | 192 |
|  | EY | M | 49 | 3800 | 70 | 446 | 45 | 325 | 13 | 15 | 5 | 30 | 55 | 71 | 18 | 129 |
|  | EZ | F | 41 | 2400 | 144 | 582 | 60 | 458 | 27 | 18 | 10 | 43 | 163 | 107 | 48 | 220 |
|  | FD | M | 54 | 4500 | 155 | 1008 | 56 | 877 | 37 | 81 | 17 | 86 | 181 | 321 | 83 | 572 |
| Cercopithecus neglectus | EO | F | 46 | 3850 | 206 | 710 | 83 | 827 | 61 | 49 | 12 | 132 | 278 | 260 | 69 | 605 |
|  | AR | F | 46 |  | 144 | 556 | 69 | 728 |  |  |  |  | 163 | 132 | 47 | 444 |
|  | P28 ${ }^{1}$ | M | 58 | 11340 | 289 | 1382 | 77 | 982 | 61 | 100 | 11 | 119 | 460 | 448 | 43 | 518 |
| Cercopithecus nictitans | AP | M | 57 |  | 99 | 760 | 87 | 654 |  |  |  |  | 93 | 195 | 75 | 340 |
|  | AS | F | 47 |  | 73 | 763 | 65 | 563 |  |  |  |  | 59 | 179 | 48 | 280 |
|  | AX2 | M | 58 |  | 177 | 798 | 95 | 753 |  |  |  |  | 221 | 214 | 80 | 410 |
|  | EL | M | 59 | 6500 | 120 | 927 | 80 | 816 | 61 | 56 | 13 | 88 | 124 | 225 | 57 | 441 |
| Cercopithecus aethiops | $\mathrm{AB}^{1}$ | M ${ }^{\text {P }}$ | 37 |  | 34 | 169 | 13 | 143 |  |  |  |  | 19 | 17 | 3 | 35 |
|  | $\mathrm{AC}^{1}$ | $\mathrm{F}^{3}$ | 40 |  | 14 | 234 | 31 | 105 |  |  |  |  | 5 | 42 | 15 | 25 |
| Cercocebus albigena | AN | F | 55 |  | 124 | 559 | 69 | 977 |  |  |  |  | 130 | 152 | 54 | 799 |
|  | $\mathrm{AX}$ | F | 51 |  | 91 | 657 | 67 | $580$ |  |  |  |  | 82 | $189$ | 51 | 268 |
|  | FK' | M | 47 | 3700 | 88 | 453 | 20 | 374 | 24 | 51 | 6 | 62 |  |  | 8 | 164 |
| Macaca sylvana | FN | F | 51 |  | 270 | 1187 | 72 | 1323 |  |  |  |  | 416 | 434 | 46 | 954 |
|  | FO | Fo | 38 |  | 180 | 795 | 59 | 826 |  |  |  |  | 226 | 222 | 31 | 503 |
| Macaca mulatta | P11 ${ }^{3}$ |  | 39 | 3350 | 77 | 618 | 81 | 377 | 15 | 56 | 13 | 32 | 64 | 160 | 47 | 123 |
| $\begin{aligned} & \text { Macaca } \\ & \text { sinica } \end{aligned}$ | DH | F | 42 |  | 226 | 587 | 62 | 772 |  |  |  |  | 320 | 185 | 51 | 558 |
|  | DJ ${ }^{1}$ | M | 47 |  | 122 | 638 | 25 | 310 |  |  |  |  | 127 | 191 | 9 | 125 |
| Macaca fascicularis | P29 | M | 37 | 2700 | 143 | 1011 | 28 | 693 | 15 | 22 | 3 | 31 | 160 | 346 | 8 | 364 |
|  | P45 | M | 40 | 3900 | 306 | 918 | 29 | 659 | 32 | 70 | 9 | 66 | 502 | 268 | 8 | 320 |
|  | P46 | F | 40 | 3050 | 178 | 1144 | 67 | 866 | 37 | 56 | 8 | 78 | 223 | 390 | 40 | 568 |


| Papio | FK | M | 72 | 16600 | 402 | 2854 | 64 | 1894 |  |  | 760 | 1596 | 47 | 1518 |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| sphinx | EP | F | 51 | 6100 | 250 | 1143 | 50 | 1228 | 76 | 97 | 13 | 163 | 372 | 409 | 33 | 1395 |  |  |
|  | FR | F | 63 | 12300 | 322 | 2435 | 70 | 2240 | 123 | 296 | 13 | 332 | 543 | 968 | 195 | 1782 |  |  |
|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  | 8 | 19 | 1 | 11 |


| Colobine monkeys |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Colobus polykomos | $A Q$ | F | 56 |  | 1021 | 556 | 26 | 549 |  |  |  |  | 2055 | 107 | 13 | 218 |
|  | FB | F | 58 | 6500 | 1056 | 925 | 15 | 630 | 154 | 149 | 5 | 72 | 2162 | 184 | 6 | 301 |
| Presbytis entellus | DF | M | 63 |  | 1585 | 1673 | 105 | 978 |  |  |  |  | 3974 | 633 | 80 | 845 |
|  | DL | M | 65 | 10000 | 1439 | 1167 | 140 | 760 |  |  |  |  | 3438 | 330 | 104 | 505 |
| Presbytis senex | DG | F | 48 |  | 938 | 740 | 139 | 548 |  |  |  |  | 1814 | 181 | 77 | 275 |
|  | DM | M | 59 |  | 1416 | 687 | 139 | 548 |  |  |  |  | 3357 | 162 | 96 | 336 |
| Presbytis cristata | P30 | F | 50 | 6850 | 1694 | 1929 | 90 | 966 | 265 | 60 | 9 | 99 | 4831 | 604 | 64 | 645 |
|  | P33 ${ }^{\text { }}$ | F | 53 | 5440 | 1175 | 1329 | 30 | 607 | 182 | 65 | 5 | 59 | 2209 | 385 | 12 | 285 |
| Presbytis obscura | P18 | M | 50 | 7960 | 1363 | 1953 | 64 | 670 | 180 | 77 | 8 | 66 | 3237 | 699 | 41 | 343 |
|  | P19 | $\mathrm{F}^{3}$ | 45 | 4230 | 956 | 1311 | 52 | 558 | 105 | 53 | 4 | 42 | 1788 | 397 | 27 | 264 |
|  | P26 | M | 53 | 7200 | 1351 | 1969 | 88 | 902 | 154 | 62 | 6 | 76 | 2974 | 734 | 51 | 588 |
|  | P32 ${ }^{1}$ | F | 56 | 6350 | 1282 | 1129 | 90 | 869 | 211 | 45 | 8 | 50 | 3139 | 327 | 64 | 546 |
| Presbytis melalophos | P14 | M | 51 | 6860 | 1020 | 1386 | 60 | 532 | 114 | 82 | 6 | 52 | 2072 | 336 | 34 | 156 |
|  | P16 | M ${ }^{3}$ | 44 | 5220 | 694 | 1075 | 23 | 369 | 109 | 76 | 3 | 44 | 1128 | 247 | 8 | 132 |
|  | P17 | F | 47 | 6410 | 1648 | 1796 | 60 | 552 | 145 | 107 | 6 | 63 | 3521 | 517 | 36 | 224 |
|  | P22 | M | 49 | 6510 | 1078 | 1389 | 42 | 695 | 90 | 50 | 4 | 42 | 2158 | 395 | 20 | 362 |
|  | P23 | F | 50 | 7340 | 1382 | 1695 | 68 | 848 | 124 | 71 | 8 | 50 | 3327 | 588 | 43 | 469 |
|  | P24 | F | 52 | 6880 | 1274 | 2021 | 38 | 612 | 150 | 60 | 7 | 71 | 2296 | 633 | 14 | 292 |
| Presbytis rubicunda | P38 ${ }^{1}$ | M | 56 | 6350 | 1125 | 1672 | 45 | 637 | 105 | 21 | 7 | 38 | 2259 | 505 | 20 | 329 |
| Nasalis larvatus | P25 ${ }^{1}$ | M | 64 | 15880 | 1978 | 3120 | 100 | 1234 | 357 | 153 | 6 | 82 | 6523 | 1127 | 66 | 655 |
| Pygathrix nemaeus | P34 ${ }^{1}$ | F | 53 | 4540 | 1243 | 1512 | 36 | 578 | 137 | 63 | 5 | 50 | 2960 | 444 | 12 | 261 |
|  | P30 ${ }^{1}$ | F | 60 | 3630 | 1431 | 1601 | 80 | 854 | 200 | 47 | 4 | 45 | 3442 | 531 | 42 | 433 |

TABLE 7. Measurements of body length and weight, and of surface area, weight, and volume of stomach, small intestine, caecum, and colon in apes, domestic mammals and temperate wild mammals

| Species | Ident. no. ${ }^{1}$ | Sex | Bodylength, length, cm | Body weight, gm | Surface area, $\mathrm{cm}^{2}$ |  |  |  | Weight, gm |  |  |  | Volume, $\mathrm{cm}^{3}$ |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  | Stom. | S.I. | Caec. | Colon | Stom. | S.1. | Caec. | Colon | Stom. | S.I. | Caec. | Colon |
| Apes |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Hylobates | PO5 ${ }^{2}$ | M | 40 | 5000 | 204 | 453 | 15 | 383 | 58 | 81 | 4 | 74 | 274 | 154 | 3 | 259 |
| lar | PO6 ${ }^{\text {a }}$ | F | 40 | 5400 | 104 | 268 | 6 | 403 | 53 | 35 | 2 | 68 | 100 | 68 | 2 | 267 |
| Hylobates pileatus | P41 ${ }^{1}$ | F | 47 | 7260 | 304 | 592 | 77 | 1128 | 56 | 77 | 17 | 144 | 499 | 596 | 34 | 920 |
| Hylobates syndactylus | P27 ${ }^{1}$ | F | 52 | 11340 | 457 | 2278 | 75 | 1557 | 146 | 150 | 10 | 230 | 919 | 1007 | 34 | 1891 |
|  | P40 ${ }^{1}$ | M | 56 | 7250 | 140 | 1708 | 81 | 954 | 55 | 183 | 22 | 184 | 156 | 697 | 58 | 883 |
| Pongo pygmaeus | P42 ${ }^{1}$ | M | 61 | 8620 | 256 | 1263 | 70 | 978 | 71 | 124 | 17 | 157 | 385 | 461 | 56 | 1071 |
|  | P35 ${ }^{1}$ | M | 95 |  | 880 | 6564 | 155 | 5774 | 331 | 568 | 22 | 980 | 2457 | 4046 | 55 | 7800 |
| Pan troglodytes | $\mathrm{AD}^{\prime}$ | F | 83 |  | 472 | 1700 | 162 | 1812 |  |  |  |  | 965 | 815 | 91 | 1451 |
|  | EN | M | 72 | 34000 | 690 | 3761 | 286 | 2925 |  |  |  |  | 1705 | 1967 | 407 | 4335 |
| Pan gorilla | EQ | M | 84 | 51000 | 1087 | 4018 | 590 | 4813 |  |  |  |  | 3370 | 1897 | 955 | 7006 |
| Domestic mammals |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Felis (cat) | D09 | M | 45 | 4000 | 144 | 345 | 8 | 125 | 24 | 104 | 1 | 17 | 163 | 60 | 2 | 48 |
|  | D12 | F | 40 | 2450 | 104 | 249 | 7 | 87 | 20 | 67 | 1 | 14 | 100 | 36 | 1 | 25 |
|  | D13 | M | 43 | 2730 | 120 | 372 | 9 | 123 | 16 | 84 | 1 | 17 | 124 | 71 | 2 | 40 |
|  | D18 | F | 42 | 2450 | 106 | 348 | 12 | 148 | 17 | 42 | 1 | 11 | 102 | 75 | 3 | 62 |
|  | D19 | F | 42 | 2700 | 132 | 374 | 11 | 130 | 21 | 43 | 1 | 11 | 143 | 86 | 3 | 52 |
|  | D20 | M | 46 | 4340 | 117 | 291 | 8 | 111 | 24 | 60 | 1 | 13 | 119 | 41 | 2 | 36 |
| Canis (dog) | D03 | M | 78 | 13500 | 300 | 1030 | 40 | 225 | 133 | 263 | 8 | 42 | 490 | 238 | 16 | 90 |
|  | D04 | M | 59 | 7250 | 215 | 585 | 30 | 125 | 46 | 125 | 5 | 30 | 297 | 102 | 8 | 43 |
|  | D05 | F | 60 | 10680 | 426 | 992 | 37 | 208 | 109 | 198 | 8 | 42 | 826 | 253 | 11 | 75 |
|  | D06 | F | 58 | 9400 | 196 | 562 | 25 | 75 | 52 | 130 | 4 | 26 | 258 | 136 | 10 | 20 |
|  | D21 | F | 81 | 12550 | 344 | 1445 | 40 | 192 | 153 | 327 | 6 | 53 | 599 | 483 | 16 | 61 |
|  | D24 | F | 48 | 4750 | 162 | 776 | 20 | 135 | 48 | 170 | 7 | 24 | 194 | 196 | 5 | 54 |


| Sus | D27 | M | 54 | 7650 | 381 | 1057 | 30 | 170 | 107 | 285 | 14 | 35 | 701 | 294 | 10 | 58 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| (pig) | D29 | M | 55 | 7450 | 193 | 656 | 30 | 132 | 71 | 178 | 5 | 23 | 253 | 193 | 10 | 45 |
|  | D30 | M | 55 | 7950 | 216 | 696 | 30 | 99 | 68 | 163 | 5 | 21 | 297 | 166 | 10 | 35 |
|  | D15 | M | 98 | 47980 | 160 | 14425 | 440 | 4702 | 333 | 1144 | 70 | 685 | 1597 | 9119 | 700 | 3710 |
|  | D41 |  | 126 | 65320 | 917 | 9968 | 432 | 6246 | 464 | 1010 | 78 | 984 | 2610 | 4828 | 619 | 5855 |
|  | D42 |  | 123 | 60780 | 792 | 14036 | 630 | 6824 | 332 | 1327 | 128 | 1106 | 2097 | 7420 | 902 | 6042 |
| Equus (horse) | D14 | M | 157 | 202730 | 728 | 10991 | 9240 | 27993 | 675 | 3204 | 1395 | 5450 | 1847 | 6207 | 28296 | 50551 |
| Capra (goat) | D01 | M | 127 | 84950 | 31297 | 8967 | 300 | 5131 | 2250 | 710 | 67 | 585 | 16220 | 2924 | 286 | 2452 |
|  | D02 | M | 145 | 94220 | 36475 | 11948 | 437 | 6047 | 3210 | 950 | 98 | 878 | 20878 | 4277 | 282 | 3339 |
|  | D33 | $\mathrm{M}^{3}$ | 84 | 21900 | 15029 | 6102 | 275 | 1830 | 686 | 607 | 27 | 186 | 6317 | 1747 | 273 | 592 |
|  | D34 | $\mathrm{M}^{3}$ | 82 | 23850 | 13195 | 4901 | 187 | 1601 | 671 | 579 | 33 | 241 | 5086 | 1208 | 164 | 467 |
| Ovis (sheep) | D43 |  | 104 | 40820 | 14110 | 15780 | 490 | 3642 | 1062 | 925 | 61 | 384 | 7989 | 6414 | 530 | 1587 |
|  | D44 |  | 99 | 47170 | 13760 | 10591 | 403 | 2496 | 1093 | 868 | 58 | 393 | 7568 | 3496 | 461 | 533 |
|  | D45 |  | 99 | 38100 | 11702 | 10299 | 150 | 2066 | 959 | 798 | 57 | 368 | 8040 | 3523 | 119 | 701 |
| Wild mammals (temperate) |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Oryctolagus cuniculus | MB |  | 43 | 2600 | 157 | 958 | 529 | 431 |  |  |  |  | 185 | 214 | 353 | 117 |
| Sciurus vulgaris | D47 | M | 24 | 600 | 26 | 307 | 54 | 136 | 5 | 9 | 3 | 4 | 13 | 43 | 29 | 30 |
| Mustela nivalis | D38 |  | 18 | 140 | 30 | 131 | 0 | 12 |  |  |  |  | 16 | 19 | 0 | 2 |
| Mustela erminea | D39 |  | 24 | 30 | 42 | 121 | 0 | 19 | 2 | 5 | 0 | 1 | 26 | 13 | 0 | 2 |
| Vulpes vulpes | D37 | M | 64 | 8000 | 406 | 929 | 48 | 229 | 78 | 106 | 4 | 25 | 768 | 329 | 23 | 91 |
|  | D46 | F | 65 | 5500 | 308 | 993 | 75 | 336 | 50 | 122 | 5 | 21 | 508 | 372 | 30 | 204 |
|  | D48 | M | 52 | 6250 | 320 | 992 | 56 | 232 | 68 | 130 | 7 | 27 | 538 | 329 | 18 | 100 |
|  | D49 | F | 58 | 5750 | 352 | 854 | 48 | 279 | 57 | 104 | 5 | 28 | 621 | 302 | 15 | 151 |
|  | D50 | M | 56 | 6200 | 259 | 929 | 54 | 236 | 60 | 105 | 5 | 26 | 392 | 340 | 19 | 117 |
| Phocaena phocaena | D35 | M | 130 | 29030 | 1151 | $1121311213$ |  | 11213 | 462 | 1125 |  |  | 3672 | 6831 |  |  |
|  | D36 | F |  |  | 1501 | $21309$ | $21309$ | 21309 | 636 | 1928 |  |  | 5471 | 18733 |  |  |
|  | D40 |  | 168 | 50450 | 1501 | 12455 | 0 | 53 | 885 | 1333 | 0 | 9 | 5471 | 9257 | 0 | 25 |
| Tursiops truncatus | D51 |  | 230 | $\begin{array}{r} 450000 \\ \mathrm{ca} \end{array}$ | 1588 | 25540 | 0 | 389 | 200 | 430 | 10 | 10 | 3701 | 20913 | 0 | 208 |

[^0]| Species | Ident. no. ${ }^{\text {. }}$ | Sex | Body length. cm | Body weight, gm | Surface area, $\mathrm{cm}^{2}$ |  |  |  | Weight, gm |  |  |  | Volume, $\mathrm{cm}^{2}$ |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  | Stom. | S.L | Caec. | Colon | Stom. | S.I. | Caec. | Colon | Stom. | S.I. | Caec. | Colon |
| Wild mammals (tropical) |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Atilax paludinosus | MW | F | 45 | 2380 | 70 | 575 | 9 | 74 | 31 | 86 | 3 | 19 | 55 | 124 | 4 | 21 |
| Potamogale velox | MX |  | 23 | 740 | 19 | 368 | 0 | 39 | 4 | 22 | 0 | 4 | 8 | 67 | 0 | 11 |
| Manis tricuspis | ME |  | 28 | 1550 | 189 | 517 | 0 | 134 |  |  |  |  | 245 | 85 | 0 | 55 |
| Manis gigantea | MR | M | 23 | 1950 | 39 | 615 | 0 | 50 | 12 | 34 | 0 | 11 | 23 | 73 | 0 | 10 |
| Epixerus ebii | $\begin{aligned} & \text { MG } \\ & \text { MI } \end{aligned}$ | $\begin{aligned} & \mathrm{M} \\ & \mathrm{~F} \end{aligned}$ | $\begin{aligned} & 24 \\ & 23 \end{aligned}$ | $\begin{aligned} & 580 \\ & 540 \end{aligned}$ | 30 48 | 201 216 | 21 21 | $52$ | $4$ | 6 | 2 | 4 | 16 | 22 | 5 | 6 |
| Heliosciurus rufobrachium | MH | M | 20 | 290 | 51 | 190 | 48 | 68 63 | 6 | 6 4 | $\begin{aligned} & 1 \\ & 3 \end{aligned}$ | $\begin{aligned} & 4 \\ & 3 \end{aligned}$ | $\begin{aligned} & 31 \\ & 34 \end{aligned}$ | $\begin{aligned} & 26 \\ & 25 \end{aligned}$ | $\begin{array}{r} 6 \\ 11 \end{array}$ | $\begin{array}{r} 12 \\ 8 \end{array}$ |
| Funisciurus anerythrus | MK |  |  |  | 16 | 161 | 15 | 30 | 3 | 5 | 1 | 2 | 6 | 18 |  |  |
| Funisciurus pyrrhopus | ML |  |  |  | 18 | 268 | 16 | 48 | 7 | 10 | 1 | 3 | 7 | 25 | 3 | 7 |
| Anornalurus fraseri | MT ${ }^{2}$ |  | 24 |  | 10 | 216 | 90 | 93 | 5 | 9 | 7 | 5 | 3 | 34 | 22 | 11 |
| Eidolon helvum | MF | F | 14 | 190 |  |  |  |  | 2 | 4 | 0 | 2 |  |  |  |  |
| Dendrohyrax dorsalis | MC | ${ }^{3}$ | 40 | 1405 | 114 | 311 | 350 | 335 |  |  |  |  | 81 | 48 | 196 | 74 |
|  | MO | F | 39 | 2323 | 140 | 456 | 482 | 572 | 54 | 25 | 61 | 43 | 110 | 73 | 331 | 169 |
|  | MP |  | 38 | 2066 | 120 | 391 | 393 | 549 | 44 | 29 | 48 | 33 | 86 | 53 |  | $\begin{aligned} & 169 \\ & 129 \end{aligned}$ |
|  | $\begin{aligned} & \text { MU } \\ & \text { My } \end{aligned}$ | F | 31 44 | 1415 | 143 | 315 588 | 275 | 304 | 25 | 17 | 30 | 12 | 114 | 55 |  | 129 80 |
|  | MV | M | 44 | 2720 | 126 | 588 | 362 | 594 | 54 | 40 | 60 | 44 | 94 | 131 |  | 156 |
| Nandinia binotata | MD | F | 42 | 2400 | 143 | 481 | 0 | 150 |  |  |  |  |  |  |  |  |
|  | MY | M | 45 | 2250 | 105 | 420 | 0 | 98 | 34 | 65 | 0 | 23 | 101 | 142 100 | 0 | 53 27 |
|  | NA | M | 43 | 2440 | 60 | 209 | 0 | 73 | 22 | 38 | 0 | 16 | 43 | 37 | 0 | $\begin{aligned} & 27 \\ & 15 \end{aligned}$ |
| Bradypus tridactylus | MA |  | 46 | $510$ | 601 | 411 | 0 | 262 |  |  |  |  | 986 | 97 | 0 | 88 |
| Poiana richardsoni | MS |  | 29 |  | 36 | 184 | 1 | 27 | 11 | 17 | 3 | 4 | 20 | 28 | 1 | 6 |
| Genetta servalina | MQ |  | 50 | 1480 | 63 | 274 | 2 | 48 | 19 | 54 | 5 | 15 | 47 | 42 | 3 | 15 |
| Profelis aurata | MZ | $\mathrm{F}^{3}$ | 57 | 5230 | 134 | 404 | 2 | 92 | 58 | 130 | 1 | 26 | 120 | 88 | 1 | 32 |
| Panthera pardus | MM | M | 84 | 15700 | 421 | 1580 | 22 | 342 | 131 | 392 | 4 | 61 | 813 | 593 | 10 | 163 |

${ }^{*}$ Nos. AA, BA, etc. - C.M.H.; Nos. P01, D01, etc. - D.J.C.





COEFFICIENT of GUT DIFFERENTIATION
weight of stomach + caecum + colon weight of small intestine

Fig. 19. Coefficients of gut differentiation from weight in each taxonomic group of primates, in other temperate and tropical mammals, and in domestic mammals, indicating the extent and overlap of the main morphological dietary categories
by vertical broken lines.

GUT MORPHOLOGY AND DIET IN MAMMAIS


COEFFICIENT of GUT DIFFERENTIATION volume of stomach + caecum + colon volume of small intestine

Fig. 20. Coefficients of gut differentiation from volume in each taxonomic group of primates, in other temperate and tropical mammals, and in domestic mammals, indicating the extent and overlap of the main morphological dietary categories. The distortion of values for colobine monkeys and domestic ruminants, resulting from the standard treatment of stomachs as spherical, is corrected as outlined in table 4 and plotted as a small closed circle to the left of the standardized value; in the
case of primates it leads to a closer reatationt case of primates it leads to a closer relationship with other primates, but it disrupts the sequential pattern among domestic

While surface areas are taken as critical for pinpointing differences in digestive and absorptive functions between species, weights of each region provide an indication of the amount of muscle, and thus of physical activity in that region. In each mammalian group such data as are available provide a clear contrast between those faunivorous mammals with a relatively heavy small intestine, and those folivores with a heavy stomach and/or large intestine (Fig. 19). Comparing groups, however, we find different values for dietary boundaries (except for folivores), so that the overall spread of faunivores overlaps slightly with that of folivores, thereby obscuring frugivores as a group.
Volumes are most significant with reference to the capacity of those parts of the tract concerned with fermentation; it is presumed that the larger the volume, the more fermenation can take place. The extensive overlap of coefficients between dietary categories, refecting wide variation in the relative volume different parts of the fut, may make this differer the least accurate indicator of diet Fig 201. Thi would be becouse complexitie Fig. 20.. This wou paring directly the volumes of mainly ferparing directly the volumes of mainly fer-
menting and mainly absorbing regions in this way.
Initially, stomachs were treated as spheres and volumes were calculated accordingly from surface areas, as described above, thereby standardizing interspecific comparison. Corrections according to the shape of each chamber in complex stomachs gives a more accurate figure and a more precise indication of the dietary adaptation of the species concerned Discrepancies between different methods of calculating stomach volume are small compared with the differences between species.
In a fermenting chamber the corresponding area for absorption should vary according to the two-thirds power of the volume. Since absorption is an important function irrespective of diet, there should be a compromise between large volume and reduced surface between la fermenting chambers of the more area in the fermens of chasses and the more different species the presence of sacculations, folds, papillae, villi, and microvilli change olds, papilla, ing aur measurements we the relationship. In our measurne w could account only for the larger features, e.g. in ruminants such as the goat, papillae in crease the surface area of the rumen eightfold, "leaves" quadruple the area of the omasum, and folds double the area of the abomas-
um (Fig. 16). Such features also affect the weight of the organ. Similarly, villi increase the surface area of the mucosa of the small intestine by similar proportions, although there is considerable interspecific variation Hladik, 1967); at this level of analysis no correction is thought necessary, but further studies are in progress. Nevertheless, thes crude areal measures seem to provide the best indicators of dietary adaptation.
Allometric Relationships. To compare the dimensions of the gastro-intestinal tract and its component parts between species, an allometric correction must be introduced in relation to body weight or some other measure of body size. In the field it was easier to measure accurately the length rather than the weight of the specimen; in any case, the latter is more susceptible to changes in individual condition. The value of the constant of proportionality is determined by the shape of the body (McMahon, '73), which is fairly homogeneous among primates, and not very different in the other terrestrial mammals, Furthermore the use of values derived from length are not use of values derived for length are not distorted weight, such as fat, which are irrelevant to ${ }^{7}$ allom
The volumes of potential fermenting chambers (the sum of values for stomach, caecum, and colon) in each species are displayed on a logarithmic scale in relation to body size (taken as the cube of body length, as a measure related directly to weight) (Fig. 21). The regression of these volumes ( $V$ ) on body size ( $\mathrm{L}^{3}$ ) in 73 species, using the means for those with more than one specimen, is represented by the equation:
$\log V=1.02 \log L^{3}-2.69(r$
$=0.83, \mathrm{p}<0.001)$
The large scatter, reflected by the low value of $r$, is not surprising since species differing markedly in diet are grouped together. The slope of the regression equation is close to 1.0 , which means that the volume for fermentation which at first approximation to be proporappeal to ber (78), when proportional to body size. Parra (oren when in ing the size ofermenting chamions in nants and nonrumants, obtained a slope of 1.1 and suggested that the relative capacity of fermenting chambers increased with body size.

The relationship between the volume of fer-
menting chambers and body size can be illustrated by cubic models of animals of length L . and volume $\mathrm{L}^{3}$ (Fig. 24). The fermenting chambers are required to fill this volume with nutrients each day; this will be achieved if the size of the fermenting chamber is proportional to the size of the animal. However, this intake of nutrients relates to metabolic weight, which only increases to the threequarter power of body weight (Schmidt-Nielsen, '72), that is, $\mathrm{L}^{2.25}$ in our model. Furthermore, Tucker ('70) calculated that the energetic costs of movement in terrestrial mammals are relatively smaller in larger forms. In the arboreal environment of most primates, however, costs might be much higher for larger species, because they are theoretically proporcional to actual body weight (or $L^{3}$ ) for all vertical movements. Thus the correcting factor for gut volume to account for eco-physiological needs in relation to body size should lie between $L^{2.25}$ and $L^{3}$, slopes of 0.75 and 1.00 respectively in the $\log / \log$ graph.

If animals with similar diets are considered together, the wide scatter of plots (Fig. 21) is resolved into meaningful components (Fig. 22). Thus, reliable data on the volumes of potential fermenting chambers from mature wild mammals are regressed linearly against body size in four groups:

1) in 30 faunivores of 14 species, from 11 to 230 cm body length,
$\log \mathrm{V}=0.95( \pm 0.11) \log \mathrm{L}^{3}-2.52(\mathrm{r}=0.95$,

$$
\mathrm{p}<0.001)
$$

2) in 50 frugivores of 22 species, from 17 to 126 cm body length,
$\log \mathrm{V}=1.13( \pm 0.12) \log \mathrm{L}^{3}-2.94(\mathrm{r}=0.92$,

$$
\mathrm{p}<0.001) ;
$$

3) in 13 folivores of seven species, with large caecum and colon, from 17 to 157 cm ,
$\log \mathrm{V}=1.20( \pm 0.08) \log \mathrm{L}^{3}-2.95(\mathrm{r}=0.99$, $\mathrm{p}<0.001$ ); and
4) in 14 folivores of six species, with large stomachs, from $44-145 \mathrm{~cm}$ body length,
$\log \mathrm{V}=0.78( \pm 0.09) \log \mathrm{L}^{3}-0.69(\mathrm{r}=0.97$, $\mathrm{p}<0.001$ ).

For each of these equations the $95 \%$ confidence intervals were calculated according to standard practice (Moore et al., '72); those for the
different dietary groups are clearly separated (Fig. 22). The differences between the slopes in the equations for folivores with large colon and faunivores are also significant ( $\mathrm{p}<0.05$ )
In faunivores, where fermentation is mini mal, the volumes of these parts of the gut are related mostly to actual body weight (ex pressed as $\mathrm{L}^{3}$ ) $\operatorname{In}$ frugivores and folivores with large caecum and colon the potential ferment ing chambers are relatively more voluminous in larger animals, the wider scatter among rugivores seems to reflect the inclusion of species consuming significant amounts of either animal or plant matter in addition to fruit. The folivores with large stomachs are remarkable in that the sizes of potential fermenting chambers are relatively much small er in larger animals; compared with the other three groups, this represents a more efficien adaptation to metabolic needs, since the slope of regression is close to the theoretical 0.75 . Although there is a close correspondence between fermenting volume and body size across the two types of folivore, as shown by he crossing and lack of much divergence of the rossing and lack (Fig 22), the sig he respective regressions (Fig. 22), the sig mant difference in slopes reflects the diferive alratric relationships of the two distinc in stog of gidgut fermentation in stomach and caecum/colon respectively) Structurally this is a matter of "sacs" (spheres versus "tubes" (cylinders). The "sac" is ideal or fermentation, but it must be kept relative y small in the larger fore-gut fermenters, so as to provide adequate area for absorption: hey have a relatively larger small intestine to complete this process (see below). In conrast, the larger mid-gut fermenters require an extended, thus more voluminous, "tube" or adequate absorption; area and volume do not diverge so rapidly with increasing size of cylinder, as they do with spheres.
This leads directly to a consideration of absorptive activity within the gastro-intestinal tract, which is concerned with surface area rather than volume. It is our initial assumption, based on evidence from man, rhesus monkey, and dog (Bell et al., '63; Kayser, '63) that the small intestine is the most important region for absorbing the products of digestion Thus, to seek allometric relationships, the values for the surface area in each species are plotted against body size on a logarithmic scale (Fig. 23). The linear regression of these surface areas (A) in relation to body size ( $L^{3}$ ), using means for those of the 76 species for which there is more than one specimen, is


Fig. 21. Volumes of potential fermenting chambers plotted against body size (from body length, cm ) for each species, using mean values where there is more than one specimen.


Fig. 22. The relationship between the volume of potentinl fermenting chambers and body size in faunivores, frugivores, folivores with large stomachs (stom.), and folivores with large caecum and colon (cae.col.), in the
form of regressions derived from individual data, with the shaded areas demarcating the $95 \%$ confidence limits for the slopes.
represented by the equation:
$\log A=0.76 \log L^{3}-0.96(r=0.93, p<$ $0.001)$.

The scatter is less than that for volume (Fig. 21). The slope of 0.76 would mean that the area for absorption is proportional to metabolic rather than overall body size (see below for discussion of confidence intervals)

Returning to the cubic model of animals of varying length L and volume $\mathrm{L}^{3}$ (Fig. 24), varying length $L$ and volume $L^{3}$ (Fig. ${ }^{24 \text { ), }}$,,$~=1$ ationships between surface area and abrelationships between surface area and ab-
sorption can be described. If flow across the sorption can be described. If flow across the mucosa occurs at a constant rate, because o similar histology, the surface area of gut required to fill the animal at a rate of $1,000 \mathrm{~cm}^{3}$ day $/ \mathrm{cm}^{2}$ will be $0.001,1$, and $1,000 \mathrm{~cm}^{2}$ respectively for the three animals. Because animals of different sizes have different basal
metabolic rates (BMR), however, the volumes to be filled are proportional to the three-quarter power of body weight- $\mathrm{L}^{2.25}$ in our modeland the volume of nutrients absorbed in 1 day will be such that areas of $0.0056,1$, and 178 $\mathrm{cm}^{2}$ are required in the three animals respec $\mathrm{cm}^{2}$ are required in the three animals respec tively. It can be seen that there is a point beyond which further increase in body size is ment for vast intestinal area

Allowance must also be made for activity, which as mentioned above costs relatively more in terms of energy in a larger animal, despite an improved output. Thus the flow across the gut wall depends on a surface area of gut that relates to a value of body size somewhere between $\mathrm{L}^{3}$ and $\mathrm{L}^{2.25}$. Although the somewhere between $L^{3}$ and $L^{2.2 .}$. Although the correct factor to account for metabolic plus energetic costs with regard to absorptive area seems to fall in the same range as for fer-
$\mathrm{cm}^{2}$


Fig. 23. Surface areas of the main absorbing region (small intestine) plotted against body size (from body length, cm ) for each species, using mean values where there is more than one specimen.
menting volume, the nature of the relationship is different.
The relationship between the surface area The relationship between the surface area by regressing data for mature wild individuals by regressing data for mature wild individuals
in four groups (Fig. 25):

1) in 31 faunivores of 14 species,
$\log \mathrm{A}=0.65( \pm 0.07) \log \mathrm{L}^{3}-0.54(r=0.96$,

$$
\mathrm{p}<0.001 \text { ); }
$$

2) in 51 frugivores of 23 species,
$\log \mathrm{A}=0.75( \pm 0.08) \log \mathrm{L}^{3}-0.85(\mathrm{r}=0.92$, $\mathrm{p}<0.001$ );
3) in 14 mid-gut fermenting folivores of 8 species,
$\log \mathrm{A}=0.86( \pm 0.15) \log \mathrm{L}^{3}-1.46(\mathrm{r}=0.95$

$$
\mathrm{p}<0.001 \text { ); and }
$$

4) in 14 fore-gut fermenting folivores of 6 species,

$$
\begin{gathered}
\log \mathrm{A}=1.16( \pm 0.22) \log \mathrm{L}^{3}-3.09(\mathrm{r}=0.94 \\
\mathrm{p}<0.001)
\end{gathered}
$$

$$
\mathrm{p}<0.001 \text { ). }
$$

Thus, different patterns emerge; only frugi-
vores follow the expectation of absorptive area
being directly proportional to metabolic body
$\mathrm{p}<0.001$ ).
Thus, different patterns emerge; only frugi-
vores follow the expectation of absorptive area
being directly proportional to metabolic body being directly proportional to metabolic body


Fig. 24. Cubic models of body lengths 1,10 , and 100 cm , and theoretical weights . 001,1 , and 1000 kg . The areas necessary to fill these models in unit time by constant flow (shown by openings in top) are . 001,1 , and $1000 \mathrm{~cm}^{2}$. These are equivalent to the absorptive intestinal area. The actual volumes are $.001,1$, and 1000
liters, but the "metaholic volumes" that actually have to be filled, with middle-sized model as reference, are 0056,1 , and 178 liters (shown by broken lines on the smallest and largest models). Thus, to maintain the constancy of the organism, and to fill the different volumes in the same time, the absorptive intestinal areas must be 0056,1 , and $178 \mathrm{~cm}^{2}$.
size (in log values), but they have the widest scatter (for the reasons mentioned previously). Faunivores have relatively less small intestine when larger, whereas the larger the folivore, especially the fore-gut fermenters, the larger the small intestine relative to body size.

Correlations within each group, however, Correlations within each group, however, are less close than in the analysis of volumes, and calculation of the $95 \%$ confidence intervals produces extensive overlap between the different regression lines. The most obvious explanation is that it is erroneous to assume that absorption of nutrients occurs only in the small intestine, especially in folivores and frugivores with a significant intake of leaves.

There is a mounting body of evidence that the large intestine in particular does much more than regulate water and electrolytes (Sinesh chekov, '65; Giesecke, '69; Kay and Pfeffer 69; Parra, '78).
Accordingly, after testing different combinations of areal proportions, so as to account a some extent for the absorption that occur o some extent for the absorption that occur in stomach, caecum, and colon, the best fit in regression was obtained by adding half the mombed area of these regions to the area of small intestine. Since the actual efficiency o absorption has not been measured in most secies, we have to rely on this arbitrary
$\log \mathrm{A}^{\prime}=0.86( \pm 0.15) \log \mathrm{L}^{3}-1.46(\mathrm{r}=$ $0.95, \mathrm{p}<0.001$ ); and
4) in 14 fore-gut fermenting folivores,
$\log \mathrm{A}^{\prime}=1.19( \pm 0.17) \log \mathrm{L}^{T}-2.97(\mathrm{r}=$

$$
0.96, \mathrm{p}<0.001)
$$

These regressions now have a better fit (higher value of $r^{2}$ ). Since values for the two folivore regression was applied to the total set of data. Thus, for 38 folivores of 16 species,
$\log \mathrm{A}^{\prime}=0.88( \pm 0.06) \log \mathrm{L}^{3}-1.17(\mathrm{r}=0.97$,

$$
\mathrm{p}<0.001)
$$

In the three dietary groups characterized by these equations (Fig. 26), the $95 \%$ confidence intervals have been calculated; they show no overlap except for animals of very small body weight, because of the convergence of the three lines. This latter feature suggests that smaller animals show similar structural adaptations for absorption, irrespective of diet.
The maximum and minimum values of the slopes of these regressions have been compared in terms of $95 \%$ confidence intervals. They differ significantly between folivores and faunivores, but frugivores differ from these two extremes at only the $85 \%$ limit. This lack of high significance is not surprising, since the frugivore sample comprises species with rather different diets based on fruit. Both frugivores and folivores fall within the limits derived for metabolic plus energetic requirements (slopes between 0.75 and 1.00 ). Faunivores, on the other hand, with a slope of 0.66 (less than 0.50 for a small set of highly specialized fau..50 for a small set of highly specialized faunivores), in their larger forms fall below the value expected for absorptive intestinal area. These large faunivores usually catch very vide ample vide ample rich food spasmodically. Such food is digested during long periods of rest, and a smaller intestine is adequate, because of the extra time available for absorption. Conversely, small faunivores, relying mainly on invertebrates, have access to a more regular supply of food, which corresponds to the eco-physiological patterns of the other dietary groups.

The significant difference in the slopes of regression for each dietary category do not allow the use of a single allometric factor. Such a factor would have been invaluable in interspecific comparisons independent of body size, as was attempted above with indices of gut differentiation. The results of the relation
between body size and the potential area for absorption (Fig. 26) show that this area must be divided by $\mathrm{L}^{1.07}$ in faunivores, by $\mathrm{L}^{2,37}$ in frugivores, and by $\mathrm{L}^{2.64}$ in folivores, in order to eliminate allometric factors and validate the comparison between species. The factor $\mathrm{L}^{2}$, used in earlier studies to compare gut surface areas (Hladik, 67), was a good approximation (apart from being below the theoretical range from $L^{225}$ to $L^{3}$, but it can now be seen as valid only for faunivores and some frugivores, The ultimate aim in our studies of gut morphology has always been to seek correlations with diet. The results presented in this tions with diet. The results presented in this section advance considerably our understanding of the morphological features relevant to how each different dietary groups, and show how each achieves the necessary compromise between adequate volume for fermentation and adequate surface area for absorption. Of the approaches adopted, however, the first did not yield conclusive ratios of gut differentiation, probably through ignoring allometric factors, and the second, accounting for such factors, did not yield a single morphological index directly comparable with diet. So far we have been considering diet from the morphological viewpoint, and before concluding the search for a single morphological index, some features of diet and feeding behavior need first to be stressed.

DIET IN RELATIION TO GUT MORPHOLOGY
The diets of most species, especially primates, are composed of varying combinations of each major category of food-animal mat ter, fruit, and leaves. This is well known from field studies and partly explains the scatter of morphological plots in the preceding analysis, mostly among frugivores while recogmizing the special significance of the gross dietary the special signifance of the gross dietary categories to which each species can usually be assigned, particularly the most specialized forms, we have tried to avoid any implication that a classification into faunivores, frugi vores, and folivores reflects exclusive diets.
In comparing the diets of wild mammals even among closely related forms, we immediately encounter problems resulting from different methods of both observation and analysis. Observations may be made continuously or sampled at intervals of varying duration, over periods that vary from a few hours to several days. Food intake may be assessed in gross terms or in fine detail, either from direct observation or even from analyzing stomach contents or faeces. The difficulties are compounded when such variable data are subject-


Fig. 26. The relationship between the potential area for absorption (surface area of small intestine and half the combined areas of stomach, caecum, and colon) and body size (from body length, cm ) in faunivores, frugivores and folivores two types combined), in the form of regressions derived from individual data. The correcting factors ( L ), accounting for he allometric relations in each group, are markedly different, because of the variations in slope. The stippled area
ed to different kinds of analysis, according to the relatively narrow question to which the researcher may be addressing him/herself (Hladik and Chivers, '78)
It need not matter if different methods of observation are used, so long as their reliability can be assessed to yield results that are truly comparable. In seeking correlations with gut morphology it is amounts of different foods ingested, rather than the time spent feeding on each, which are of paramount im portance. It is relatively easy, in sampling the behavior of wild primates, to record accurately the details of feeding bouts in terms of time Chivers, ${ }^{\text {7 }} 74$; Struhsaker, '78; MacKinnon and MacKinnon, '78), but it is much more difficult to measure the amounts ingested in terms of fresh (or dry) weight over a reasonable length of time (Hladik and Hladik '69, '72; Hladik 73: Iwamoto ' 74,78 . Raemaekers, '77) and to analyze food composition (Hladik et al., ' 71 Hladik, '77a,b; Goodall, '77). Data based on the analysis of stomach contents pose special problems, but may yield the kind of measures required (Gautier-Hion, '78).
While the amounts of leaves and other vegetative plant parts ingested can be deduced with sufficient accuracy from feeding times, ruits may be underestimated as much as fivefold, and insects may be overestimated as much as 15 times, compared with the actual amount by weight (Hladik, '77a,b). While such distortions may fortuitously cancel each other ut (Raemackers, '77), data based on time are bviously inappropriate for our purposes. mix ing data based on time and weight should be ing data with circumspection w

保
Different models have been proposed to repesent the diets of wild primates, so as to account for the average intake of an individual nd its variation over time (e.g., Hladik and Hladik, '69, '72; Suzuki, '65; Kay, '73; Mac Kinnon and MacKinnon, '78). The most effec tive method for representing a system con

- taining three variables - animal matter, fruit and leaves - is a tri-rectangular projection (or three-dimensional graph).
This approach involves plotting values for the three major categories of food in the diet within a three-dimensional system of converging axes (Fig. 27, right upper). Since the three variables are not independent (their sum is always $100 \%$ ), the projection of any combined dietary value will fall within the triangle AFL, where point A represents a diet of $100 \%$ animal matter, point F a diet of $100 \%$ fruit and point L a diet wholly of leaves. In this
triangle (Fig. 27), any point close to A represents a diet rich in animal matter, close to F rich in fruit, and close to $L$ rich in leaves. Because of the construction of the triangle AFL, the dietary values (in terms of per cent of animals, fruit, and leaves) are plotted more conveniently along the perpendicular axes Ox and $O y$. If $\mathrm{OL}=+100$ and $\mathrm{OA}=-100$, then $\mathrm{x}=(\%$ leaves $)-(\%$ animals $)$, and $\mathrm{y}=\sqrt{3}(\%$ fruit), for any point within the triangle.
In addition to pinpointing an average diet In addition ( Fig 27) , the graph indicate or range of variation through the year. While these ranges may overlap, even yong sym these ang it may not be the sam time of year, but, more importantly, of year, but, more importan, his simarity of gross categories obscures important diferences in food choice (species and its diferent parts), and thus in biochemical composition. identify and this analifis is sufficent to dentify and quantify significant differences between species, even those which have been regarded previously as similar, e.g., the omnivorous, more properly frugivorous, primates such as Cebus, Mucaca, and Pan. Although these three primates eat mostly fruit, the overlap represents but a small part of their dietary ranges; for the means ( $x=-5$, $+15,+26$ respectively), accounting for differ ences in the supplement of plant and animal matter, provide good diagnostic dietary indices.

To amplify the interspecific comparison, the mean diets of as many as possible of the mammalian species studied morphologically are plotted in Figure 28. This inevitably means adapting data based on time measures, but all values are derived from studies lasting more than 1 year (see, for example, CluttonBrock, '77). The absence of plots near the baseline of the triangle is conspicuous, and is not a consequence of inadequate sampling of species. No mammal mixes large quantities of animal matter and leaves without including fruit in its diet. Since, as has been shown previously, faunivory and folivory represent contrasting, and incompatible, adaptations, the quantity of fruit in such a mixed diet should always be considerable. From the demonstration by Hladik et al. ('72) that fruit is an adequate source of carbohydrate for the energetic requirements of most primates but inadequate in protein, Kay ('73) argues that those primates securing this protein from insects are necessarily much smaller than those obtaining it from leaves,
Thus, as exemplified by primates, there are


Fig. 27. Annual means and ranges of the diet of nine primate species from Panama, Gabon, and Sri Lanka idata from Hladik and Hladik, '69, '72. Hladik, '73) represented within a triangle Isee text for explanation of its derivation from tri rectangular projection, and its applications). The composition of the mean annual diet, in terms of proportions of animal matter, fruit, and leaves, is pinpointed by an arrow; most data, collected over 24 -hour periods at all times of year, are included in the shaded area (only daily records which differ markedly from the previous or following days are excluded,
because of the distortion they would introduce into the small sample)
optimum body sizes corresponding to the different feeding strategies:
Body size

Biomass
$\mathrm{Kg} / \mathrm{km}^{2}$ (Hladik, ${ }^{78}$ a)
Faunivores kg (Kay, '73)

Frugivores
5

Folivores
$20 \%$ dependence); for reasons given above, no species exceeds $20 \%$ for both animal matter and leaves.

Returning to Figure 27, it is not possible from the data available to predict a central minimum $y$-value, the lowest proportion of fruit taken by a frugivore. It is likely to be quite high, since the data are clearly distributed along a crescentic path from A to L via the vicinity of $F$, with the greatest range of $y$ values around the zero $x$-values. This reprevalues around the zero $x$-values. This represents, at least for primates, the evolutionary (Ripley, '79) through three ecological grades (Hladik, 78 b ).
The geometrical arrangement of the data, The geometrical arrangement of the data,


Fig. 28. Mean dietary characteristics of the 34 species included in this study (tables 5-8) for which quantitative data on diet are available. Locations within the triangle represent diets as shown in Figure 27
cators of diet, varying from - 100 (pure fau nivores) to +100 (pure folivores). The final task is to relate these values to the dimensions of the gastro-intestinal tract.
Considering first the potential volumes for Comentation, the data for faunivores and for ivores (excepting those with large stomachs) have been linearly regressed separately against body size (Fig. 22). These regression are used to represent the structural limits for coping with the two extremes of diet (Fig. 29) Most primates fall within these limits, but the concentration of frugivorous species, represented by the third regression line, closer to the upper line, reflects their greater similarity to folivores than to faunivores, In order to derive a morphological index for comparison with the dietary one, we need a scale of negative and positive values to reflect their direction and degree of adaptation to eithe faunivory or folivory respectively. To account for the asymmetry in the relative positions of the regression lines, it is more appropriate to
take the line of the regression for frugivores as the zero, rather than a line midway between the regressions for folivores and faunivores.
Thus the morphological location of any species can be described as the distance D above or below this zero line, by the following conditional formula
if $\mathrm{V}>\mathrm{V}_{\mathrm{t}}, \mathrm{D}=+\frac{\mathrm{V}-\mathrm{V}_{\mathrm{t}}}{\mathrm{V}_{1}-\mathrm{V}_{\mathrm{f}}}$
if $\mathrm{V}<\mathrm{V}_{\mathrm{t}}, \mathrm{D}=-\frac{\mathrm{V}_{\mathrm{t}}-\mathrm{V}^{2}}{\mathrm{~V}_{\mathrm{f}}-\mathrm{V}_{\mathrm{a}}}$
where $\mathrm{V}, \mathrm{V}_{\mathrm{a}}, \mathrm{V}_{\mathrm{f}}$ and $\mathrm{V}_{1}$ represent the potential volumes for fermentation in the subject, and in faunivore, frugivore, and folivore of the same size, respectively; these latter are calculated from the regression equations relating to Figure 22, where body size is derived from the length L , the distance between bregma and ischial callosity. Since body weight $W$ is
the more widely used measure of body size the regression equations were recalculated according to the approximation, $\mathrm{W}=\frac{\mathrm{L}^{3}}{30}$ ( L in $\mathrm{cm}, \mathrm{W}$ in $\mathrm{gm}^{1}$ ). which fits most of our specimens, yielding the same results:
for faunivores,
$\log \mathrm{V}=0.95( \pm 0.11) \log W-1.07 ;$
for frugivores,
$\log \mathrm{V}=1.13( \pm 0.12) \log \mathrm{W}-1.25 ;$
and for folivores,
$\log \mathrm{V}=1.20( \pm 0.08) \log \mathrm{W}-1.18$.

As shown in Figure 29, the asymmetry creates a discrepancy in the size of units above and below the zero line, since the extremes represented by the outer lines have values of -1 and +1 . To attain a homogeneous distribution of the units, and thus the desired index of gut specialization, the distances D are transformed mathematically into the values TR The conditional formula used (Fig 29)

is based on antilogarithms; although appear ing very complex, on the small programmable calculator, now in wide use, it is almost as easy to use as the ordinary logarithm ${ }^{2}$.
This nonlinear transformation not only renders comparable scores above and below the zero line, but yields values that never exceed -100 or +100 , however large the value of D Fig. 29). The conditional formula is designed to produce values of -90 and +90 on the regression lines for animalivores and folivores, respectively. Because of the scatter around each regression line, it is unreasonable to assume that the upper one accounts for

- A regression of all data available in Tables $5-8$ gives a relation, $W=0.041 \times 1^{2=-}\left(r=0.98\right.$ or $W=\frac{L^{3}}{24.4}$. The approximation $W$ $\frac{i^{3}}{30}$ in preferred, since it yields a better fit for most wild arboreal 30
primater.

2On the HP 25 calculator the program is $\mathrm{g} \times \geqslant 0 \mathrm{GTO} 071 \mathrm{CH}$ TO $1 x \geq y$ ABS Ifx $y$ GTO $36 x \geq y$ STO $x \geq y-1 x \geq$ $1{ }^{2} \mathrm{ETO} 44-1 \times \geq y-\mathrm{E} 10$ RCL $3 \times \geq y-$ RCL $1 \times 1$ STO $1 \times=y$ GTO 00 . After storing the values 1,10 and 100 in repister 1. 2, and 3 respetively, any value
into $\mathrm{TR}_{\text {t. }}$ by pressing the RS key.
$D>1$
$\mathrm{IR}_{(\mathrm{D})}=100-10^{[1-(\mathrm{D}-1)}$
$0 \leqslant 0 \leqslant 1$
$T^{R}(D)=D\left(100-10^{[D-(D-1)]}\right)-10\left(10^{D}-1\right)(D-1)$
$0>0 \geqslant-1$
$T_{(D)}=(-D)\left(10^{[1-(|D|-1)]}-100\right)+10\left(10^{|D|}+1\right)(D+1)$
$T_{(D)}=10^{[1-(|D|-1)]}-100$
diets including more than $90 \%$ of leaves, and the lower one more than $90 \%$ of animal mat ter.
Such a method of nonlinear interpolation between two diverging regression lines, can be used as an approximation of the percentage "tendency" for any biological character varying between two opposite poles. In this case we are indicating the extent to which each species is tending toward one dietary extreme or the other, having resolved the special allometric problems encountered. These indices of gut specialization, derived from potential fermenting volumes for all species except those with complex stomachs, are very similar to the dietary indices (see below).
The same method is applied to the potential absorbing area (including half the area of stomach and large intestine together with the area of small intestine) in all species. The regressions of these areas ( A ) are those of Figure 26. The asymmetry of the regression for frugivores in relation to the other two is even more marked; thus the conditional formula is used to locate each species, according to body size, at distance D above or below this zero line
if $\mathrm{A}^{\prime}>\mathrm{A}^{\prime}, \mathrm{D}=+\frac{\mathrm{A}^{\prime}-\mathrm{A}_{\mathrm{f}}^{\prime}}{\mathrm{A}_{\mathrm{t}}^{\prime}-\mathrm{A}^{\prime}}$
if $\mathrm{A}^{\prime}<\mathrm{A}_{\mathrm{f}}^{\prime}, \mathrm{D}=-\frac{\mathrm{A}_{\mathrm{f}}^{\prime}-\mathrm{A}^{\prime}}{\mathrm{A}_{\mathrm{f}}^{\prime}-\mathrm{A}_{\mathrm{a}}^{\prime}}$
where $\mathrm{A}^{\prime}, \mathrm{A}_{\mathrm{a}}^{\prime}, \mathrm{A}^{\prime}$, and $\mathrm{A}^{\prime}$, represent the potential absorbing areas in the subject, and in faunivore, frugivore, and folivore of the same size, respectively. As in the preceding case, the regression equations were recalculated with references to body weight, rather than length:
for faunivores,
$\log \mathrm{A}^{\prime}=0.66( \pm 0.06) \log \mathrm{W}-0.49$;
for frugivores,
$\log \mathrm{A}^{\prime}=0.79( \pm 0.09) \log \mathrm{W}-0.33 ;$
and for folivores,
$\log \mathrm{A}^{\prime}=0.86( \pm 0.15) \log \mathrm{W}-0.15$.

The asymmetry again creates a discrepancy in the size of units above and below the zero line, so the transformation $\mathrm{TR}_{(\mathrm{D})}$ is performed, as shown in Figure 29. This yields indices of gut specialization, in terms of area, which are also very similar to the dietary indices

In these operations we are locating each species in relation to all others, according to the gross dietary classification derived previously Thus, it is not surprising with the large samples, that there is good correspondence between thorphere The problem of scatter is not easy to indices The proble scalteultie in easy to resolve, mecause the dieulies moasuring such a malleable morphological system, and of variation within specie
The major advance resulting from this approach is that, having resolved the complex allometric problems, mammalian species fol lowing different adaptive strategies may be compared quantitatively. It is possible to delimit the "ecophysiological tendency" for each species, especially for primates, whose body sizes fall in the central part of the range investigated. For example, among the so called "omnivorous" primates, which feed on variable amounts of fruit, insects, and leaves, the significant differences in diet which have been demonstrated recently follow those between the various indices:

|  |  |  |  |
| :---: | :---: | :---: | :---: |
|  | by area, A' | by volume, V | $\times$ - |
| Miopithecus talapoin | -85 |  | - 40 estimate, Fig. 28 |
| Ceropithecus cephus | -27 | - 2 | - 10 estimate, Fig. 28 |
| Papio sphinx | + 9 | 0 |  |
| Cebus capucinus | $+11$ |  | - 5 |
| Macaca sinica | $+16$ | + 22 | $+15$ |
| Alouatta palliata | $+40$ | + 31 | + 40 |
| Macaca sylvana | $+45$ | +85 |  |
| Presbytis melalophos | +82 |  | folivore |

[^1]The similarities between indices for each species are pleasing, considering that morphol ogical indices refer to single individuals, and the dietary indices are derived separately from data of variable quality. The success of this approach depends on a) obtaining adequate gut samples that are truly representative of the population from which dietary data are obtained, b) obtaining adequate dietary data, in terms of weight of each food ingested and c) defining in which species morphological adaptations may confer greater dietary flexibility, e.g., variations across the geographica range and feeding behavior in captivity.
Reference has been made to the convergence of the regression lines for potential absorbing area, A' (Fig. 26), which renders this model napplicable to species smaller than 27 cm nappleargth In such cases the model based on ody length. potential fermenting volumes, V Figs. 22, 29 is quite satisfactory for small species. Figures or folvores with are not included in Figure 29. To derive indices of gut specialization (GSI) for such forms, either the A' model should be used, or the V model revised using the regression line for olivores with large stomachs (Fig. 22).
The specialized seed-eaters have not been included in these models; if primates eat seeds they usually do so in small quantities and when they are unripe. Certain frugivorous squirrels of Gabon also consume some insect. Emmons, '75) and Epixerus ebii, for example see Table 8), has a GSI from $V$ of -16 . Gut secialization indices of other rain-forest squirrels have been calculated from data collected recently in Malaysia (Payne, '79). The small Sundasciurus tenuis, of length 13 cm , which eats mostly bark, sap, and seeds, and ome insects, has an index of +100 . The fruiteating species Callosciurus notatus and $C$ revosti have indices of +97 and -22 respec fively, with the latter eating considerably fively, wit fort in contrast the seederably more soft frui, in contrast, heed-eating Ratura bich in man Seeds rich in protern and rat seem to need pocessing more like animal matter than the vegetative parts of plants. An unusual primate, Cercopithecus neglectus, whose diet is known to include large quantities of seed Gautier-Hion, '78), also has low indices. Thus further research is needed on the composition, consumption, and assimilation of seeds ingested by mammals.
In the search for a full physiological explanation of the relationships between morphology and diet, that we have quantified, two
main lines of research should be followed There needs to be more extensive analysis and quantification of the biochemical compositio of foods, in relation to diet, and of the fin structure and cellular populations of the gas tro-intestinal mucosa, in relation to morphol ogy. At the same time, the quantity and com parability of the kinds of morphological and dietary data discussed herein must be im proved, if we are to understand the real significance of "sacs" and "tubes."

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Adresse de C.M. Hladik en 2013 :

Claude Marcel HLADIK
Directeur de recherche émérite
Eco-Anthropologie et Ethnobiologie
Muséum National d'Histoire Naturelle
4 avenue du Petit Château
91800 Brunoy (France)
cmhladik@mnhn.fr
[http://www.ecoanthropologie.cnrs.fr/IMG/pdf_Site-WEB-Hladik-2013.pdf](http://www.ecoanthropologie.cnrs.fr/IMG/pdf_Site-WEB-Hladik-2013.pdf)


[^0]:    Captive. ${ }^{\text {FFixed }}$ "Immature
    'Nos. AA, BA, etc. C.M.H.; nos. P01, D01, ete. = D.J.C.

[^1]:    Fig. 29. Method for comparing the potential volume of fermentation in the gastro-intestinal tracts of various primates with contrasting diets. The log of volume of stomach, caecum, and colon) is plotted against the log of body weight (taking $\frac{\mathrm{L}^{3}}{30}$ as a good approximation for body weight), so as to eliminate allometric parameters. The resulting regressions from a large sample of faunivores and folivores (Fig. 22) are used again here, and considered as showing the structural limits for coping with diets containing $90 \%$ of animal matter and $90 \%$ of leaves, respectively. Most primates fall within these limits, and the distance D from the regression line for frugivores can be regarded as an indicator of the morphological tendency towards faunivory or folivory. The conditional formula presented allows a transformation of D into the index $\mathrm{TR}_{\mathrm{p}}$, which, in most cases, is the same number as the dietary index, $x$, as defined from the tri-rectangular projection (Fig. 27).

