Feeding ecology of the mangrove crab

Neosarmatium smithi (Crustacea: Decapoda: Sesarmidae)

R. L. Giddins¹, J. S. Lucas¹*, M. J. Neilson², ³ & G. N. Richards², ⁴

¹ Department of Zoology, James Cook University of North Queensland, Townsville, Queensland 4811, Australia
² Department of Chemistry, James Cook University of North Queensland, Townsville, Queensland 4811, Australia
³ Department of Biochemistry, La Trobe University, Bundoora, Victoria 3083, Australia
⁴ Wood Chemistry Laboratory, University of Montana, Missoula, Montana 59812, USA

ABSTRACT: Consumption experiments showed that the mangrove crab Neosarmatium smithi H. Milne-Edwards preferred litter of the mangrove Ceriops tagal decayed for some weeks. Determination of assimilation efficiencies, using a 72% sulfuric acid insoluble residue as indigestible marker, revealed that N. smithi assimilated little of the components of a 2 wk old litter diet, but that assimilation efficiencies (carbon, nitrogen, caloric content, organic matter) ranged from 30 to 60% when fed 4, 6 or 8 wk old litter. These values must be treated with caution because the validity of using a 72% sulfuric acid insoluble residue as an inert marker in determining assimilations in mangrove studies, as used here, is questioned. Due to the extremely high carbon-nitrogen ratios of the litter, these diets fall well below accepted nutritional standards. An additional source of nitrogen in the diet was determined by gut content analysis to be small crustaceans. N. smithi is not a direct link in the ecosystem food chain. Carrying the leaf litter into their burrows − where the leaves decay, are subsequently consumed, and faeces are produced − the crabs substantially reduce export of leaf litter by tidal transport and regenerate nutrients for other small invertebrates and the mangroves themselves.

INTRODUCTION

Detrital material formed from mangrove leaf litter is considered to be the basis of food webs within mangrove ecosystems (Mann 1975, Odum & Heald 1975). However, although certain macro-invertebrates have the ability to digest detrital matter directly (Adams & Angelovic 1970, Yingst 1976, Foulds & Mann 1978, Wainwright & Mann 1982, Klumpp & Nichols 1983), none of these organisms inhabits mangrove ecosystems. Several studies (Malley 1975, 1978, Beever et al. 1979, Wilson 1981, Mattson 1982) have shown significant consumption of mangrove leaf material by sesarmid crabs, but none assessed the ability of the organisms in question to actually assimilate the plant material.

Neosarmatium smithi, a large (50 mm carapace width) sesarmid crab, is the dominant crustacean in the Ceriops tagal australis zone of mangroves in the Townsville region (Australia). This zone is 3.1 m above chart datum, and is infrequently immersed. The crab emerges from its burrow at night to remove litter from the surface of the mud (Alexander pers. comm.). This study set out to determine at what stage of decomposition C. tagal leaves are consumed and which components of the leaf material are assimilated.

MATERIALS AND METHODS

Collection of leaf material. There were 3 collection sites in the region of Townsville, North Queensland, Australia (ca 19° S, 147° E). Neosarmatium smithi H. Milne-Edwards specimens and senescent Ceriops tagal leaves (defined as yellow and easily removed from trees) were collected at Cape Ferguson. Fresh leaves (mature, green) were collected from Ross River. Decayed leaves (black and fragmenting) were collected from Three Mile Creek. To produce detritus of known age, large collections of senescent leaves were made every 2 wk for a period of 8 wk. The collections were divided into 40 g subsamples and placed into 150 mm square nylon mesh bags (2 mm mesh). Groups of 3 bags were tied to mangrove trees in Ross River at
the mud surface. All bags experienced the same conditions and were able to float at high tide. The bags were recovered at the end of the tenth week to give a decomposition series designated S0 (0 weeks decomposition), S2, . . . , S10 leaves.

Representative sub-samples were removed, dry weighed (60 °C for 72 h) and ball-milled. These sub-samples were either chemically analysed immediately or stored at -20 °C until required. The remainder of each leaf sample was either kept at 5 °C if consumption experiments were to be carried out within 3 d, or kept frozen until use.

Feeding experiments. Consumption experiments were carried out at 25 °C ± 2 °C with a 12 h light/12 h dark photoperiod. The crabs were kept individually in ventilated plastic containers with ca 25 ml of seawater in the base to prevent desiccation. In an initial experiment, crabs were given mixtures of known weights of leaves of fresh, senescent, and decayed condition. After 24 h the leaf material was removed, sorted into the 3 different types, and weighed after surface drying (by blotting). Four controls, without crabs, were also run, and the specific consumption rate (SCR) data (g wwt of leaf g⁻¹ crab 24 h⁻¹) were corrected accordingly. Some leaf material was finely fragmented but not ingested and this could not be distinguished from faecal particles, and thus consumption is here defined as having been processed by the mouthparts. In a second experiment, crabs were fed each leaf type in the Cerops tagal decay series separately. In this case, SCR was expressed in terms of leaf dry weight (dwt). One half of a leaf was wet weighed and fed to the crabs, and the other half dry weighed. After 24 h the remains were removed, dried, and weighed to give SCR values. Again, controls without crabs were used. SCR was then expressed as relative consumption rate (RCR):

\[
\text{RCR} (L) = \frac{\text{SCR} (L_i)}{\text{SCR} (L_0) + \text{SCR} (L_2) + \ldots + \text{SCR} (L_{10})}
\]

where \(L_0\) = 0 wk litter, etc. This gives the SCR of 1 crab, on a particular diet, relative to the total of its SCRs for all diets. RCR is dimensionless, and as such crab weight has no effect on it.

Bacterial counts. Bacterial numbers on litter were determined as in Hobbie et al. (1977), except that homogenization was used in the place of ultrasonication. Ellery & Schleyer (1984) consider the latter method more effective, but they were studying sediments. Bacteria within the leaf tissue may not have been revealed with ultrasonication. The dilution water was kept bacteria-free by the addition of 'Tween 80' at a concentration of 50 mg l⁻¹. Two samples of each of the 5 litter types were taken. From each of these, 2 filter preparations were made, and 10 microscope frames were counted for each filter.

Fractionation of samples into water-soluble and insoluble components. Microbial and mud contamination of decay series leaves made it necessary to modify the fractionation procedure described by Neilson et al. (1986). Leaf samples (S2 to S10), with petioles removed, were cut into ca 1 cm² pieces, extracted with hot acetone (10 ml acetone g⁻¹ wwt sample) for 15 min and gently vacuum filtered through Terylene cloth, which allowed mud and microbial matter to pass through. Extraction was repeated, the mixture cooled, homogenized and filtered. The cloth was folded about the leaf material and then sonicated in cold acetone until most of the mud was removed. Although some mud ‘coating’ was visible on the larger meal fragments of S8 and S10, it was assumed that its contribution to the total dry weight would be minimal. Corresponding faecal samples (F2 to F10) were treated similarly but without the homogenization and sonication steps.

The acetone-insoluble residues were then extracted (24 h) under nitrogen in the dark, with ethylenediaminetetra-acetic acid (EDTA: 0.1M, pH 6, 25 °C; 40 ml EDTA g⁻¹ dwt sample). The skurry was cooled, centrifuged (4 °C, 10 000 × g) for 30 min, and the supernatant aspirated off and held at 4 °C. The extraction was repeated and the combined supernatants were exhaustively dialysed in the dark (4 °C, 48 h) against distilled water in pre-washed tubing, freeze-dried and weighed to yield the EDTA-soluble component.

The EDTA-insoluble fractions were washed thoroughly with water by centrifugation, acetone dried at the pump, vacuum dried (1 mm Hg, < 40 °C, P₂O₅, 12 h) and weighed.

Analyses. Caloric value of samples was determined using a Gallenkamp Ballistic Bomb Calorimeter. Ash Free Dry Weight (AFDW) was determined by ashing dried samples at 500 °C for 16 h (Rice & Tenore 1981). Carbon and nitrogen analyses were performed in a ‘Leco’ CHN600 Elemental Analyser. Proteinaceous nitrogen was defined as that nitrogen lost after boiling with potassium hydroxide (Odum et al. 1979).

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Carbohydrate contents were determined by the phenol-sulfuric method (Dubois et al. 1956). For EDTA-soluble fractions, the method was applied as described in Neilson et al. (1986). EDTA-insoluble fractions were subjected to total acid hydrolysis based on modified Saeman conditions (Saeman et al. 1954) prior to assay. Samples (15 to 25 mg) were weighed into a centrifuge tube (10 ml), titrated with 72% sulfuric acid (100 µl) at 25 °C and kept in a desiccator with occasional stirring for 1 h. Water was added (2.5 ml) and the tube and contents tared prior to secondary hydrolysis under reflux for 3 h at 100 °C. The mix-
ture was cooled, weight adjusted with water (if necessary), centrifuged (1000 × g, 10 min) and aliquots taken for phenol-sulfuric assay. The insoluble residues produced on hydrolysis were washed free of acid (check by pH paper) with water, vacuum dried and weighed. These ‘crude lignin’ samples were ashed (600 °C, 4 h), weight adjusted, reported as acid-insoluble residue, and used to calculate assimilation efficiency values (see below).

Newly-recognized flavologlycan (FG) (Neilson et al. 1986) was present in the EDTA-soluble fraction isolated as described above. Total flavolan or condensed tannin content in the fraction was estimated colorimetrically by the vanillin-HCl method detailed in Broadhurst & Jones (1978) with catechin as a reference. Since the reactivity of the vanillin reagent with catechin is much greater than with flavolans (Goldstein & Swain 1963), these values indicate the relative amounts of total flavolan content.

Model leaching experiment. Intact senescent leaves (49 g, 21 g dwt) were placed in a 1 l flask and immersed in sterilized, artificial sea-water (800 ml) (Reichelt & Baumann 1973). Sodium azide (0.02%) was added to prevent any microbial activity. The flask was capped with aluminium foil and kept at 25 °C in the dark. After 2 and 4 wk, filtered (Millipore, 0.45 μm) aliquots of the ‘mother liquor’ were analysed for flavolan and total carbohydrate.

Assimilation efficiency (AE). AE was estimated by comparing the proportion of a dietary component in the food to that in the faeces relative to an indigestible marker substance (Klumpp & Nichols 1983). In this case, the insoluble residue after acid hydrolysis (see above). Klumpp & Nichols used a similar ‘lignin’ material as the marker (see below). Preliminary experiments showed that starving crabs may take more than a week to evacuate their guts, thus precluding the use of a simpler quantitative method for determining AE. Crabs were fed a particular diet for 2 d, the chambers were cleaned, and the crabs returned together with litter of the same type as before. Faecal matter was collected after a further 24 h and frozen for later analysis.

Table 1. Neosarmatium smithi. Consumption rates when offered a choice of fresh, senescent and decayed Ceriops tagal leaves

<table>
<thead>
<tr>
<th>Leaf type</th>
<th>Mean specific consumption rate (g wwt leaf g⁻¹ crab 24 h⁻¹)</th>
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</thead>
<tbody>
<tr>
<td>Fresh</td>
<td>0.004 (0.006)*</td>
</tr>
<tr>
<td>Senescent</td>
<td>0.011 (0.014)</td>
</tr>
<tr>
<td>Decayed</td>
<td>0.062 (0.055)</td>
</tr>
<tr>
<td>* ± 1 standard deviation</td>
<td></td>
</tr>
</tbody>
</table>

Gut content analysis. Five specimens were caught in the field for gut analysis. They were killed by destroying the major thoracic ganglion, preserved in 30% seawater formalin and dissected within 24 h. At a magnification of 40 ×, 10 frames were assessed from each sample. The proportion of the total volume of the gut made up by each food type was thus estimated.

RESULTS

Gut content analysis

Gut analysis of 5 Neosarmatium smithi showed that plant material was 90 ± 7% (mean ± 1SD) of the total volume of the gut, animal material was 8 ± 2%, and the remainder was unidentifiable. The vast majority of the animal material was crustacean in origin.

Mechanical alteration of leaf litter by N. smithi

Fig. 1 illustrates several SEM views of the teeth of the gastric mill of Neosarmatium smithi. These show a series of heavily dentate ridges that interlock when the lateral teeth pass over the dorsal tooth (Fig. 1a, b, c). Also, a number of curved ‘spines’ (Fig. 1d) that meet together over the dorsal tooth are visible.

Consumption rates

When offered a choice of fresh, senescent, and decayed Ceriops tagal leaves (Table 1), Neosarmatium smithi exhibited a distinct preference for decayed material (1-way ANOVA, p < 0.0001).

A partial correlation of SCR with litter type, controlling for the effect of crab weight, revealed that the effect of leaf age on SCR was significant above any effect of crab weight (r = 0.41, p < 0.0001). The consumption rate was expressed as RCR to determine an accurate relationship with decomposition stage. This transformation was performed on all crabs fed on all types of litter. A plot of RCR against litter type (Fig. 2) shows that a maximum RCR is reached for the S6 litter (mean RCR at S6 litter = 21.1%).

Assimilation efficiencies for Ceriops tagal decay series

Table 2, 3 and 4 list the quantities of the various components of food and faeces, and Table 5 shows AE values for these components for all litter types except S0 leaves.
Fig. 1 Neosarmatium smithi. Electron micrographs of dorsal and lateral teeth of the gastric mill. (a) and (b) A lateral tooth showing other large ridges which interlock with those on the dorsal tooth. (c) Dorsal tooth seen ventrally, showing large ridges used in crushing plant fibres. (d) Large spines on a lateral tooth possibly used for keeping food in place. All scales 100 μm.

When both flavolan and soluble carbohydrate components were present (S2 and S4 litters) they were totally assimilated. The insoluble carbohydrates were assimilated from the S2 and S4 litters (56 and 35 % respectively), but negative AEs were recorded for the S6, S8 and S10 litters (−44, −115, and −63 % respectively). The AEs for the nitrogen, carbon, organic matter (AFDW), and energy components were either low or slightly negative for the S2 litter (−2, 1, −1, and −1 % respectively). For the S4, S6, and S8 litters, AE values ranged between 24 and 59 %. In the S10 litter all of these components were assimilated only in small amounts, AE values ranging from 4 to 10 %. AE for bacteria in the S10 litter was only 9 %.

The possibility that some of the carbohydrate content shown in Tables 2 and 3 was derived from bacterial polysaccharides is eliminated since these would have been removed during the isolation procedure.
DISCUSSION

Consumption of leaf litter by *Neosarmatium smithi*

Gut content analyses indicate that leaf litter is the major component in the diet of *Neosarmatium smithi*. Electron micrographs of the gastric mill detail graphically how well *N. smithi* is adapted to feeding on this material. A similar gastric mill is found in the mangrove tree crab *Aratus pisonii* which is also known to consume mangrove leaves (Warner 1977). As contrasting gastric mill ossicles are found in crabs that have alternative feeding habits (Warner 1977) this mill would seem to be specifically adapted to deal with vascular plant material.

| Table 2. Compositional analysis of *Ceriops tagal* litter types S0 to S10 |
|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|
| Sample                  | Nitrogen 1                | Carbon 1                 | AFDW 1                   | EDTA-soluble 2 carbohydrate | Flavolan 3 | Acid-insoluble 4 residue |
|                         | 0.2 (0.01)                | 36.7 (0.4)               | 77.0 (0.3)               | 1.8                       | 4.9           | 25.1                     | 15.0                     |
| S2                      | 0.4 (0.01)                | 38.4 (0.1)               | 71.2 (2.3)               | 1.4                       | 3.0           | 26.6                     | 30.7                     |
| S4                      | 0.4 (0.02)                | 39.7 (0.2)               | 73.4 (0.5)               | 1.3                       | 2.0           | 16.6                     | 27.6                     |
| S6                      | 0.4 (0.00)                | 35.2 (0.9)               | 65.2 (2.0)               | 0.7                       | 0.2           | 7.7                      | 26.4                     |
| S8                      | 0.4 (0.01)                | 38.2 (0.1)               | 69.4 (1.1)               | 0.5                       | 0.1           | 3.8                      | 17.9                     |
| S10                     | 0.6 (0.05)                | 39.7 (0.1)               | 70.1 (2.5)               | 0.0                       | 0.1           | 11.5                     | 37.1                     |

1 ± 1 standard deviation
2 By phenol-sulfuric acid method using D-glucose as standard
3 By vanillin-hydrochloric acid method (catechin equivalents)
4 Residue remaining after total acid hydrolysis
5 S0 = leaves aged for 0 wk, etc.

| Table 3. *Neosarmatium smithi*. Compositional analysis of faecal matter produced when fed on litter types S2 to S10 |
|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|
| Sample                  | Nitrogen 1                | Carbon 1                 | AFDW 1                   | EDTA-soluble 2 carbohydrate | Flavolan 3 | Acid-insoluble 4 residue |
|                         | 0.4 (0.04)                | 37.4 (0.0)               | 70.4                     | Neg. 6                   | Neg.           | 11.5                     | 30.2                     |
| F4                      | 0.3 (0.03)                | 38.2 (0.0)               | 70.1                     | Neg.                      | Neg.           | 14.3                     | 36.7                     |
| F6                      | 0.4 (0.04)                | 36.0 (0.0)               | 60.4                     | Neg.                      | Neg.           | 18.2                     | 43.3                     |
| F8                      | 0.4 (0.05)                | 38.3 (0.0)               | 64.5                     | Neg.                      | Neg.           | 18.6                     | 40.8                     |
| F10                     | 0.6 (0.04)                | 39.2 (0.0)               | 66.9                     | Neg.                      | Neg.           | 18.7                     | 39.9                     |

1 ± 1 standard deviation
2 By phenol-sulfuric acid method using D-glucose as a standard
3 By vanillin-hydrochloric acid method (catechin equivalents)
4 Residue remaining after total acid hydrolysis
5 Negligible
**Table 4. Neosarmatium smithi. Caloric values and bacterial counts for food used and corresponding faecal matter produced when fed on different litter types**

<table>
<thead>
<tr>
<th>Litter type</th>
<th>Bacterial counts ((10^9 ; g^{-1} ; dwt))</th>
<th>Caloric value ((kJ ; g^{-1} ; AFDW))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Food</td>
<td>Faeces</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S0</td>
<td>Not determined</td>
<td>26.6 (0.9)</td>
</tr>
<tr>
<td>S2</td>
<td>–</td>
<td>29.8 (4.8)</td>
</tr>
<tr>
<td>S4</td>
<td>–</td>
<td>27.4 (1.0)</td>
</tr>
<tr>
<td>S6</td>
<td>–</td>
<td>32.4 (2.9)</td>
</tr>
<tr>
<td>S8</td>
<td>–</td>
<td>30.0 (2.1)</td>
</tr>
<tr>
<td>S10</td>
<td>36.6 (6.3)</td>
<td>35.5 (6.8)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>30.1 (5.1)</td>
</tr>
</tbody>
</table>

1 Not determined
2 ±1 standard deviation

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**Table 5. Neosarmatium smithi. Assimilation efficiencies of the various components of litter types S2 to S10**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Nitrogen</th>
<th>Carbon</th>
<th>AFDW</th>
<th>Bacteria</th>
<th>Energy</th>
<th>EDTA-soluble carbohydrate</th>
<th>Flavolan EDTA-insoluble carbohydrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>S2</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>–</td>
<td>–</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>S4</td>
<td>33</td>
<td>28</td>
<td>28</td>
<td>–</td>
<td>24</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>S6</td>
<td>34</td>
<td>38</td>
<td>44</td>
<td>–</td>
<td>39</td>
<td>Neg.</td>
<td>Neg.</td>
</tr>
<tr>
<td>S8</td>
<td>50</td>
<td>56</td>
<td>59</td>
<td>–</td>
<td>56</td>
<td>Neg.</td>
<td>Neg.</td>
</tr>
<tr>
<td>S10</td>
<td>4</td>
<td>7</td>
<td>10</td>
<td>9</td>
<td>5</td>
<td>Neg.</td>
<td>Neg.</td>
</tr>
</tbody>
</table>

1 Not determined
2 Negligible

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**Consumption rates**

The preference shown for aged litter by *Neosarmatium smithi* in both experiments suggests that leaves taken into their burrows are stored before consumption. This was seen by Kneipp & Alexander (unpubl.) when they kept crabs in artificial burrows that permitted observation of their behaviour: the crabs placed S0 leaves on the walls of the burrow instead of eating them. The immediate cause for this behaviour is probably the more acceptable 'taste' of aged leaves. Rapid leaching of flavologlycan from S0 leaves occurs, as shown in a model experiment. After 2 and 4 wk, 26 and 58% of the total flavolan (vanillin/HCl) had leached from the leaves. Concurrently, 16 and 27% of the available soluble carbohydrates (including low molecular weight constituents) were solubilized. Most likely, the leached FG has astringent properties because flavolans (condensed tannins) are astringent (Goldstein & Swain 1963, Bate-Smith 1973). Consequently, plant tissues are rendered unpalatable (Cope & Burns 1971, Burns et al. 1972) or the flavolans present will immobilize enzymes, thus impeding microbial degradation of the tissues (Bate-Smith 1973, Vallis & Jones 1973, Wong 1973, Daiber 1975). It is quite conceivable that in this case the FG concentration in the leaves is rapidly reduced by leaching and then the micro-organisms grow on the leaves and/or *N. smithi* consumes the leaves. This postulate is consistent with the findings of Ford (1978). In a comparative study on the influence of chemical composition on the digestibility of several pasture legumes, he found that *Desmodium intortum* had a high polyphenol concentration (20%) and an associated low relative digestibility. He showed that *D. intortum* digestibility was significantly increased when the polyphenols were extracted prior to incubation. The interaction between FG and protein needs to be investigated using specific enzyme preparations and pertinent microbial populations.

In addition to this, an advantage of delayed consumption is that the carbon/nitrogen ratio is reduced. The C/N ratio of S0 leaves (183:1) is well above the maximum 17:1 value prescribed by Russell-Hunter (1970) as being necessary for a nutritional food. Also, Tenore (1977, 1981, 1983a, b) has determined that nitrogen is an important factor governing the availability of detritus to a consumer. As the relative nitrogen content of detritus increases with decomposition (de la Cruz & Gabriel 1974, Gosselink & Kirby 1974), by allowing decomposition the crab will reduce the high C/N ratio problem and increase nutrient availability.

The effect of crab weight on SCR was to be expected as smaller organisms usually have a higher specific metabolic rate (Gordon 1977). The lesser effect with S0 leaves is probably due to the generally lower consumption rates suppressing this effect.

The RCR reaches 91% of the asymptote in the S6 litter and increases extremely slowly after this point, indicating that factors affecting the consumption rate are negligible after 6 wk decomposition. As near maximum consumption rate is reached at this point, the litter is probably consumed at or before this stage. However, with the S6 type leaves having a C/N ratio of
100:1, they are still far above the 17:1 maximum suggested by Russell-Hunter (1970), and hence there must be another source of nitrogen. Gut content analysis shows that the detritus diet is supplemented by predation on small invertebrates.

Assimilation efficiencies

The markers used here and in the study by Klumpp & Nichols (1983) are essentially the same; they defined ‘lignin’ as the residue from hydrolysis in 72 % sulfuric acid, after the initial removal of protein (van Soest 1963). In this study there was no protein removal. However, as the protein content of these litters is never greater than 3.1 %, and is mostly bacterial, the result should not be significantly affected.

The existence of a low AE for bacteria, some negative AEs for the S6, S8, and S10 litters, and the inconsistency of results when comparing the S10 litter to the S4, S6, and S8 litters suggests that the use of an acid-insoluble residue as a marker may not be accurate. The method employed here assumed that the marker used was inert in acid and was not assimilated at all by Neosarmatium smithi. Since up to 50 % of the dry weight of the newly senescent leaves of Ceriops tagal consists of polyphenols (M. Neilson unpubl.), it would be reasonable to expect that a portion of these polyphenols is acid-soluble, and this would lessen the significance of using ‘lignin’ as an inert internal marker in mangrove studies, since it is assumed that such a marker is inert in the gut and in acid. It is also probable that errors will arise as the polyphenols which generate the acid-insoluble ‘lignin’ vary during decomposition, leaching and passage through the digestive system. This factor may be especially important for the polyphenols which are bonded to polysaccharides as flavoglucans. Some aromatic products from acid hydrolysis of FG are acid-soluble and some are acid-insoluble. The ratio may vary due to the chemical nature of the flavolan moiety. More accurate efficiencies would have been determined by feeding the crabs, which had cleared their guts previously, with a known weight of litter, and collecting a known weight of corresponding faeces. However, as reported in the ‘Materials and Methods’, this was not possible.

Previous studies have shown that detritivores can assimilate a significant proportion of the non-living plant fraction of detritus (e.g. Adams & Angelovic 1970, Foulds & Mann 1978, Wainwright & Mann 1982). Klumpp & Nichols’ (1983) study – which documented that the portunid crab Nectocarcinus integrifons is able to assimilate up to 60 % of the components of the seagrass Posidonia australis – is, however, now questioned due to possible inaccuracies of the method involved, as determined here. (In this publication cellulose and hemicellulose were incorrectly defined in the footnote of their Table III: cellulose and hemicellulose terms must be interchanged.)

The total assimilation of the flavolan fraction is not under question here as there were only negligible amounts left in the faeces. Since this fraction is soluble, the water in the experimental container was also analysed for flavolans, but it contained none. The 100 % assimilated flavolan refers to the FG component only.

The markers used here and in the study by Klumpp & Nichols (1983) are essentially the same; they defined that can be separated according to their solubility in aqueous and non-aqueous solvents (Ribereau-Gayon 1972). With their high polyphenolic concentrations, this property may easily be confused in original and aged mangrove leaves. A check for absolute methanol-soluble (Ribereau-Gayon 1972) flavonoid material in the EDTA-insoluble residues from S2, F2, S8, and F8 showed that 5.8 % of the S2 residue dry weight (i.e. 3.9 % of the total leaf dw) consisted of this material, with <1 % being found in the remainder. Hence, EDTA (or water) extraction readily removed FG (flavolan bonded to polysaccharides) leaving low molecular weight flavolans soluble in absolute methanol. The flavolans would be original plant tissue components or, in the case of aged leaves, produced by various degradative processes. Detailed categorisation of the complex flavolans obviously present in these plant tissues was beyond the scope of this work. For the S2 diet, the amount of absolute methanol-soluble material in the EDTA-insoluble fraction was reduced by 85 % after passage through the gut, suggesting apparent assimilation by Neosarmatium smithi. This indicates that N. smithi has a mechanism for eliminating some ‘tannins’. It is possible that, although the flavolan fraction can be dealt with to some extent, it may still have a deleterious effect on assimilation. This could be by the flavolans inhibiting digestive enzymes. Thus there would still be lower assimilation efficiencies when flavolans were present before consumption.

Ecological implications

No implications can be inferred from AEs due to possible inaccuracies of the method employed. The removal of litter from the mud surface by Neosarmatium smithi is important in that this retains the nutrients present in the leaves within the mangrove ecosystem, allowing circulation back to the mangroves themselves. Faecal material of N. smithi is of considerable ecological significance. Faecal pellets have long been known to be important as sites of microbial activity and as food sources for many detritivores (e.g. Hargrave 1970). The considerable proportion of car-
bohydrates left in faecal matter by *N. smithi* (ca 18%) will probably be useful to other detritivores as food. The removal of the flavon fraction will greatly increase the availability of the detritus to other detritivores. These 'other' detritivores will mainly have to exist in the burrows of *N. smithi*. The crabs spend so little time out of their burrows that the great bulk of their faecal material will be deposited within the burrow. Habitation in the burrows will be restricted to small crustaceans and polychaetes.

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**LITERATURE CITED**


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