Marker concentration patterns of labelled leaf and stem particles in the rumen of cattle grazing bermuda grass (*Cynodon dactylon*) analysed by reference to a raft model*

D. P. Poppi†, W. C. Ellis‡, J. H. Matis§ and C. E. Lascano||

Texas A&M University College Station, Texas 77843, USA

(Received 11 January 2000 – Revised 18 September 2000 – Accepted 10 November 2000)

Large (>1600 μm), ingestively masticated particles of bermuda grass (*Cynodon dactylon* L. Pers.) leaf and stem labelled with $^{169}$Yb and $^{144}$Ce respectively were inserted into the rumen digesta raft of heifers grazing bermuda grass. The concentration of markers in digesta sampled from the raft and ventral rumen were monitored at regular intervals over approximately 144 h. The data from the two sampling sites were simultaneously fitted to two pool (raft and ventral rumen–reticulum) models with either reversible or sequential flow between the two pools. The sequential flow model fitted the data equally as well as the reversible flow model but the reversible flow model was used because of its greater application. The reversible flow model, hereafter called the raft model, had the following features: a relatively slow age-dependent transfer rate from the raft (means for a gamma 2 distributed rate parameter for leaf 0.0740 v. stem 0.0478 h$^{-1}$), a very slow first order reversible flow from the ventral rumen to the raft (mean for leaf and stem 0.010 h$^{-1}$) and a very rapid first order exit from the ventral rumen (mean of leaf and stem 0.44 h$^{-1}$). The raft was calculated to occupy approximately 0.82 total rumen DM of the raft and ventral rumen pools. Fitting a sequential two pool model or a single exponential model individually to values from each of the two sampling sites yielded similar parameter values for both sites and faster rate parameters for leaf as compared with stem, in agreement with the raft model. These results were interpreted as indicating that the raft forms a large relatively inert pool within the rumen. Particles generated within the raft have difficulty escaping but once into the ventral rumen pool they escape quickly with a low probability of return to the raft. It was concluded that the raft model gave a good interpretation of the data and emphasized escape from and movement within the raft as important components of the residence time of leaf and stem particles within the rumen digesta of cattle.

**Marker: Residence time: Rumen: Raft: Leaf and stem particles**

Passage of digesta from the rumen is clearly a major determinant of intake of fibre-containing diets (Blaxter et al. 1956; Balch & Campling, 1962; Ulyatt et al. 1967; Thornton & Minson, 1973). Measuring rate of passage in terms of mass/h or fractional outflow rate has been an important area of research to explain differences in intake. This has been done in a variety of ways, summarized by Faichney (1975, 1986) and Ellis et al. (1984a,b). The rationale has been to develop techniques that can be used to examine diet effects on digesta passage from the rumen and also to develop models which describe how digesta mixes within and exits from the rumen. The conceptual delineating of pools within the rumen and the quantitative values associated with their turnover enable identification of rate-limiting pathways and how these may interact with forage type and physiological state of the animal in determining intake by the animal.

Models of digesta flow through the rumen have considered rumen digesta to be a homogenous pool (Blaxter et al. 1956; Grovum & Williams 1973a,b, 1977).

**Abbreviations:** MRT, mean residence time; MRTR, mean residence time in the rumen raft compartment; MRTV, mean residence time in the ventral rumen compartment; VRR, ventral rumen–reticulum.

* Corresponding author: present address School of Land and Food Sciences, The University of Queensland, Q.4072, Australia, fax +61 73365 1177, email d.poppi@mailbox.uq.edu.au

‡ Present address: Animal Science Department, Texas A&M University, College Station, TX 77843, USA.

§ Present address: Department of Statistics, Texas A&M University College Station, TX 77843, USA.

|| Present address: CIAT, Apartado Aereo G713 Cali, Colombia, South America.
or one with a gradation of particle size pools often separated for convenience into large and small particle pools (Hungate, 1966; Baldwin et al. 1976; Poppi et al. 1981; Egan & Doyle, 1984; Faichney, 1986; Ellis et al. 1984b; Czerkawski, 1986; Ulyatt et al. 1986; Wagborn et al. 1986; Kennedy & Murphy, 1988). Sutherland (1988) has suggested an alternative concept to particle size pools to examine digesta movement through the rumen and to explain why only small particles escape. This is that particles separate into those having buoyant properties acquired via entrapped fermentation gases (newly ingested and especially large particles of stem) and those having sedimentation properties (have been largely digested and tend to be small particles). Flotation results in raft formation and entrapment of particles. Raft, in this context, refers to a pool of particles of varying size which float, but which by the nature of the constraints of the rumen are in contact with each other, cannot separate easily into distinct particles and form a pliable mass.

The concept of buoyant related pools was used to formulate a two-pool rumen digesta model comprising a raft pool (raft digesta low escape potential) and an escape pool (ventral rumen—reticulum (VRR) digesta, high escape potential) with an added feature that particles from the escape pool could return to the raft pool. This is not a feature in the sequential particle size pools in previous models (Baldwin et al. 1976; Poppi et al. 1981; Ellis et al. 1984b; Faichney 1986). Large masticated particles (>1600 μm) of leaf and stem were labelled with 169Yb and 144Ce, dosed into the rumen digesta raft and the concentration of these markers in the raft and ventral rumen was followed (Lascano, 1979; Ellis et al. 1982, 1985). The concentrations at the two sites were simultaneously fitted to determine rate parameters for flow between and out of two pools with low and high escape potential particles. This approach enabled a number of models to estimate kinetic parameters to be compared and also to prepare a means by which Sutherland’s model may be parameterized.

Materials and methods

Animals, forage and markers

The data were obtained by Lascano (1979) in a grazing study designed to determine effects of maturity and selective grazing of Coastal bermuda grass (Cynodon dactylon L. Pers.) upon forage intake and kinetics of particle size reduction in and escape from the reticulo—rumen (Ellis et al. 1984b, 1994). Four plots each of three different maturities (20, 33 and 42 d regrowth plots) were each sequentially grazed for 2 d by successively grazing animals (A, B and C grazers) on two occasions. An average grazing pressure of 22.5 kg pasture DM/100 kg live weight resulted. A 6 d measurement period followed a 14 d period of adjustment to each treatment (three maturities and three successive grazers) during two occasions (n 18).

The 8–10-month-old Brahman X Jersey heifers (mean 309 kg live weight) had previously established oesophageal and ruminal cannulas (Ellis et al. 1984b). Prior to each measurement period, samples of the digestively masticated forage were collected from each animal, freeze-dried and sieved through successive sieves of 3350, 1600 and 100 μm apertures. Negligible particles were retained by the 3350 μm aperture sieves. Particles retained by the 1600 μm aperture sieve were separated into leaf and stem, extracted for 1 h with a neutral detergent solution without the chelating agent, and thoroughly washed with water and acetone. Following drying at 60°C, 14–78 MBq 144CeCl3 or 169YbCl3 were sprinkled onto 60–170 g extracted stem and leaf fractions respectively and allowed to air dry for 24 h before dosing. Some marker may migrate from the particles after this procedure but this is likely to be small.

The 144Ce- and 169Yb-labelled fractions were dosed via a rumen cannula into the dorsal rumen strata of each animal during mid-morning when the animals were ruminating following their usual intense morning period of grazing. A cavity approximately 10 cm wide and 20 cm deep was excavated within the dorsal rumen digesta impinging onto the interior orifice of the 10 cm rumen cannulas. After dosing into the cavity, the excavated digesta was returned. Samples of dorsal rumen digesta were subsequently collected from the depository site of marked particles at 4, 8, 16, 20, 24, 28, 40, 48, 64, 72, 88, 96, 112, 120, 136, and 144 h after the dose.

Following collection of dorsal digesta, samples of the ventral rumen digesta were collected via a 2.54 cm internal diameter plastic pipe inserted through the dorsal rumen digesta to the floor of the ventral sac of the rumen. The pipe was stoppered, withdrawn and its initial digesta sample discarded. The pipe was reinserted to the floor of the ventral sac through the digesta tunnel remaining from the initial insertion. The tube was withdrawn 4–6 cm and digesta allowed to flow into the pipe via hydrostatic pressure, stoppered and the sample contained within the tube withdrawn.

Digesta samples were dried, ground, 1.0–1.5 g samples ashed and the 144Ce and 169Yb dissolved in 50 ml 3 M-HNO3. A 7 ml aliquot was assayed using a multichannel γ-spectrometer with a sodium iodide detector. A portion of the labelling solution was simultaneously assayed and used to compute decay-corrected count rate of each radioisotope per g digesta sample DM.

Model development

Sutherland’s concept (1988) relating to particle movement through the rumen can be represented simply as a two-pool model involving turnover of particles from raft and ventral rumen—reticulum pools. Based on the biological characteristics described by Sutherland (1988), the escape rate from the raft pool was assumed to increase as digestion of particles approached their potential for digestion. Such ‘age-dependent’ escape rates were modelled by assuming that the distribution of elapsed time during a single visit, i.e. ‘the retention times’, of particles in the raft pool is an integer gamma of 2 or greater (Matis, 1984; Pond et al. 1988). Under this assumption, the initial escape rate at t 0 was zero, and after infinite ageing, equals some constant that we denote as λ1. Models with gamma 2 and gamma 3, denoted G2 and G3, retention times were investigated. In the model with G2, the raft pool may be represented for mathematical convenience as two sequential, homogenous compartments, R1 and R2, as given in Fig. 1. In the model
with G3, the raft pool would be represented for subsequent mathematical analysis using three homogenous compartments. In either case, the sequence of homogenous compartments does not imply a physical partitioning of the raft pool; rather the sequence produces mathematically a combined effect, which yields the age-dependent distributed retention times that ingested particles spent in the raft pool prior to their escape.

Particles escaping the raft pool enter the VRR which, due to its rather uniform composition, is considered a homogenous compartment. Escape from the VRR compartment is by flow either to the omasum or return to the fermentatively spent, small particle compartment of the raft pool. Both are mass action processes and hence are represented with age-independent escape rates, $k_0$ and $k_2$ respectively, which imply exponentially distributed retention times ($G1$). Subsequent escape of the fermentatively spent, small particles recycled back into the raft compartment is via mass action, which yields an age-independent rate. This escape rate under the previous assumptions is $l_1$ and it is implemented into both the $G2$ and $G3$ models by recycling the particles into the second compartment, $R_2$, of the raft pool (Fig. 1). The two proposed reversible flow models are denoted as $G3 \Leftrightarrow G1$ and $G2 \Leftrightarrow G1$; where the mass action passage of the recycled particles is understood.

No assumption is made as to what are the escape characteristics of particles from the rumen, but Sutherland (1988) has noted that the ability to sediment is a necessary characteristic for particles to get into the appropriate digesta stream(s) to reach the omasal orifice and that this is related to the extent of digestion of the particle and its particle size.

**Model equations**

Solutions relating to the equations for the $G2 \Leftrightarrow G1$ model are given in the Appendix (p. 563). The equations for the $G3 \Leftrightarrow G1$ model are similar.

The equations used for $G2 \Leftrightarrow G1$ are:

$$
\dot{C}_{R1}(t) = -\lambda_1 C_{R1},
$$

$$
\dot{C}_{R2}(t) = -\lambda_1 C_{R2} + \lambda_1 C_{R1} + k_2 r C_v,
$$

$$
\dot{C}_v(t) = -(k_0 + k_2) C_v + \lambda_1 C_{R2}/r,
$$

$$
MRTR = (2k_0 + k_2)/(k_0\lambda_1),
$$

$$
MRTV = k_0^{-1},
$$

where $k_0$, $\lambda_1$, $k_2$ are rate parameters, $r$ is ratio of quantities (ventral rumen:raft); $C_{R1}$, $C_{R2}$, $C_v$ are concentrations at dosing and sampling sites in raft and in the ventral rumen with derivatives $\dot{C}_{R1}(t)$, $\dot{C}_{R2}(t)$ and $\dot{C}_v(t)$; and MRTR, MRTV are mean residence times in raft and ventral rumen compartments respectively.

The observed concentrations in the raft were ascending initially, which indicated that the sampling site was different from the dosing site (Fig. 2). For simplicity, the sampling site was assumed to be the $R_2$ compartment in the rumen raft pool. Therefore, the two regression functions fitted to the data were $C_{R2}(t)$ and $C_v(t)$, which were obtained numerically using computer software.

The statistical package PCNONLIN (CM Metzler and DL Weiner; PCNONLIN Nonlinear estimation program, 1986, Statistical Consultants, Inc., Lexington, KY, USA) was used. The marker concentration data from the dorsal and the ventral rumen were fitted simultaneously, which enabled estimation of the relative size of raft and VRR pools. The sensitivity of parameter estimates to the starting values was also examined. This is a robust procedure allowing parameter estimates not possible previously.

**Alternative models**

Instead of having reversible flow between the VRR and the
raft a sequential flow model may be assumed and the data fitted simultaneously using PCNONLIN (Statistical Consultants, Inc.). This can be designated the $G_n \rightarrow G_1$ model.

In many studies of marker kinetics in the rumen only, the descending portion of the concentration curve is examined and assumed to represent the slow turnover pool of major biological importance (Grosvum & Williams 1973; Cruickshank et al. 1990). This was also examined here using the Procedure NLIN of SAS (1986; Statistical Analysis Systems Inc. Cary, NC, USA) for fitting the $G_2 \rightarrow G_1$ model to data from each sample site independently as distinct from the simultaneous fitting procedure used earlier. The model representation as $G_n \rightarrow G_1$ is the same.

In addition, a simple procedure of linear regression with the starting value selected visually at a point away from the peak, where concentration was definitely descending, was also analysed. The rate parameter, $k$, was estimated by linear regression of the equation:

$$\ln A(t) = \ln A(0) - kt,$$

where $A(t)$ is the marker concentration at time $t$ and $A(0)$ is the marker concentration at time $t_0$. This can be represented as a $G_1$ model.

### Statistical procedures

The procedure GLM of SAS (Statistical Analysis Systems Inc.) was used to test effects of forage part (leaf or stem), order of grazer (A, B or C), period (six combinations of three maturities x two cycles) and some interactions upon the parameters estimated for individual data sets by the models. The ANOVA model is: response = part + grazer + period + (part x grazer) + (grazer x period) + error.

### Summary of models used

Eight models were fitted to the data. Four models were used when the data were fitted simultaneously to marker concentration data from the raft and ventral rumen. The simultaneous fitting procedure is novel, robust and enables estimates of relative pool sizes to be determined. The four models were a gamma 2 or gamma 3 order of age dependency with reversible flow, designated as $G_2 \rightarrow G_1$ or $G_3 \rightarrow G_1$, or without reversible flow, designated as $G_2 \rightarrow G_1$ or $G_3 \rightarrow G_1$. In addition, the $G_2 \rightarrow G_1$ sequential flow model was fitted independently to the concentration data from each sample site. Finally, the simple one-pool exponential model, designated $G_1$, was fitted only to the descending marker concentration data of each sample site.

### Results

#### Choice of model

The $G_2 \rightarrow G_1$ model consistently fitted all data better than the $G_3 \rightarrow G_1$ model as judged by the combined error mean square values and plot of actual vs. predicted values. However, there was no major difference, using these criteria, to separate the reversible flow models and the sequential flow models fitted simultaneously to the two sample sites. The reversible flow model was accepted purely on the basis that such a phenomenon as flow of particles from the ventral rumen to the raft is known to occur (Ehrlein, 1979; Deswyssen & Ehrlein, 1981; Sutherland, 1988) and that solutions to the model in this form would have more general application. For these reasons, results for the $G_3 \rightarrow G_1$ model and from the simultaneous fitting of site data in the sequential $G_3 \rightarrow G_1$ and $G_2 \rightarrow G_1$ models are not presented. Data for the $G_2 \rightarrow G_1$ and $G_1$ models fitted separately to each site are presented in Table 1.

#### Parameter estimates

The parameter estimates relating to the $G_2 \rightarrow G_1$ model are summarized in Table 1 where they are presented within treatments. In all cases the order of the rate constants was $k_0 > \lambda_1 > k_2$ with at least a 3-fold difference between these rate parameters, where $k_0$ is the rate constant for disappearance from the VRR, $\lambda_1$ for exit raft to VRR and $k_2$ for recycling to raft (Fig. 1). The VRR pool size:raft pool size ratio averaged 0·27 for leaf and 0·16 for stem. The order of the rate parameters is reflected in the mean residence time (MRT) in the various compartments shown in Table 2. The MRTR was much greater than MRTV (leaf 31·1 v. 4·6 h, stem 45·4 v. 4·5 h). There were large standard errors in estimating MRTV (e.g. 12·9 (SE 6·19) and 5·4 (SE 15·06) h) but the errors associated with MRTR and the total MRT in the rumen were much lower (e.g. MRTR 27·9 (SE 3·01) h, total MRT 40·8 (SE 5·36) h).

There were no significant differences due to order of grazer (A, B or C) in any parameter for either leaf or stem marked particles. Stem particles had a significantly longer MRT than leaf (31·1 vs. 4·6 h) and a larger mean residence time (45·4 vs. 27·9 h). The MRTR was much greater than MRTV (31·1 vs. 4·6 h, leaf 31·1 h) but not in the ventral rumen (45·4 vs. 27·9 h). There was a significant difference between the MRTR (45·4 h, leaf 31·1 h) but not in the ventral rumen (45·4 vs. 27·9 h) (Table 2).

Analysis of the descending portion of the marker concentration curve in the raft and ventral rumen (model $G_1$) yielded similar estimates for rate parameters, $k$, for both sites (leaf 0·0358 v. 0·0330, stem 0·0270 v. 0·0260).
Table 1. Rate parameters (λ1, k0 and k2) of leaf and stem particles in the conceptual pools of the G2=G1 model with the ratio DM in the ventral rumen–reticulum (VRR): DM in the raft pools (i.e. VRR/raft) and the faster (λ1) and slower (k0 and k) turnover rate parameters estimated independently for the rumen dorsal (raft) and ventral digesta (VRR) sampling sites by G2 → G1 and G1 models†

(Mean values with standard errors of the means for eighteen heifers)

<table>
<thead>
<tr>
<th>Plant part and grazer</th>
<th>G2=G1, raft+VRR</th>
<th>G2→G1</th>
<th>G1</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Raft</td>
<td>VRR</td>
<td>Raft</td>
</tr>
<tr>
<td>λ1 (h⁻¹)</td>
<td>0.074</td>
<td>0.071</td>
<td>0.079</td>
</tr>
<tr>
<td>k0 (h⁻¹)</td>
<td>0.004</td>
<td>0.005</td>
<td>0.006</td>
</tr>
<tr>
<td>k2 (h⁻¹)</td>
<td>0.014</td>
<td>0.014</td>
<td>0.025</td>
</tr>
<tr>
<td>VRR:raft</td>
<td>0.21</td>
<td>0.34</td>
<td>0.25</td>
</tr>
<tr>
<td>λ1 (h⁻¹)</td>
<td>0.244</td>
<td>0.211</td>
<td>0.409</td>
</tr>
<tr>
<td>k0 (h⁻¹)</td>
<td>0.054</td>
<td>0.047</td>
<td>0.048</td>
</tr>
<tr>
<td>k2 (h⁻¹)</td>
<td>0.007</td>
<td>0.015</td>
<td>0.014</td>
</tr>
<tr>
<td>λ1 (h⁻¹)</td>
<td>0.237</td>
<td>0.193</td>
<td>0.179</td>
</tr>
<tr>
<td>k0 (h⁻¹)</td>
<td>0.039</td>
<td>0.032</td>
<td>0.054</td>
</tr>
<tr>
<td>k2 (h⁻¹)</td>
<td>0.032</td>
<td>0.031</td>
<td>0.036</td>
</tr>
</tbody>
</table>

A, B, C, order of grazer; RSD, residual standard deviation.
Mean values were significantly different, mean leaf vs. mean stem variable: *P < 0.05.
† For details of procedures and models, see p. 554.
‡ 1, k0 and k2 are rate parameters for escape from raft, escape from VRR to omasum, and backflow from VRR to raft respectively.

0.0286 h⁻¹, raft vs. ventral rumen respectively). These values and their corresponding MRT (i.e. 1/k) were lower than observed for the slow turnover pool (raft) in the G2=G1 model. However, residence time of leaf particles was less than stem particles (mean 29.4 h vs. 37.4 h) in agreement with the order observed in the G2=G1 model. The rate parameter k estimated by the single pool exponential G1 model was significantly greater for all marked particles in the raft of grazer A than for grazer B or C. The fit of the descending concentration curve to the model equation, lnA(t) = lnA(0) – kt, was good in most cases with 78% of all regressions having r² > 0.8.

The results of the sequential G2 → G1 model fitted independently to each data set are also summarized in Table 1. Parameters were similar irrespective of whether they were estimated from marker concentration in the dorsal rumen digesta or the ventral rumen digesta. As with the raft model, the rate constants were larger for leaf particles than for stem particles and this is also seen in the residence time estimates which were larger for stem (Table 2).

Table 2. Compartmental residence time (h) of leaf and stem particles in the raft (MRTR), the ventral rumen–reticulum (MRTV) and the total mean residence time in the compartmental system (MRTT) as estimated by the G2=G1 model and similar estimates for dorsal digesta (2/λ1, 1/k2 and total mean residence in dorsal digesta (MRTD)) and ventral digesta (2/λ1, 1/k2 and total mean residence in ventral digesta (MRTV)) both estimated by the G2 → G1 model fitted independently to each of the two sites and the retention time in dorsal digesta (DD) and ventral digesta (VD) estimated by the simple G1 model fitted independently to data from each site†

(Mean values with standard errors of the means for eighteen heifers)

<table>
<thead>
<tr>
<th>Plant part and grazer</th>
<th>G2=G1, raft + VRR</th>
<th>G2→G1</th>
<th>G1</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Raft</td>
<td>VRR</td>
<td>Raft</td>
</tr>
<tr>
<td>MRTR</td>
<td>29.5</td>
<td>3.6</td>
<td>33.1</td>
</tr>
<tr>
<td>MRTV</td>
<td>33.5</td>
<td>6.1</td>
<td>39.6</td>
</tr>
<tr>
<td>MRTT</td>
<td>30.3</td>
<td>4.0</td>
<td>34.3</td>
</tr>
<tr>
<td>Mean</td>
<td>31.1</td>
<td>4.6</td>
<td>35.6</td>
</tr>
<tr>
<td>141Ce-labelled stem A</td>
<td>42.2</td>
<td>4.3</td>
<td>46.5</td>
</tr>
<tr>
<td>B</td>
<td>50.6</td>
<td>4.0</td>
<td>54.6</td>
</tr>
<tr>
<td>C</td>
<td>45.2</td>
<td>5.1</td>
<td>50.3</td>
</tr>
<tr>
<td>Mean</td>
<td>45.4</td>
<td>4.5</td>
<td>49.8</td>
</tr>
<tr>
<td>RSD</td>
<td>7.55</td>
<td>4.49</td>
<td>9.68</td>
</tr>
<tr>
<td>SEM</td>
<td>1.8*</td>
<td>1.1</td>
<td>2.3*</td>
</tr>
</tbody>
</table>

A, B, C, order of grazer; RSD, residual standard deviation.
Mean values were significantly different, mean leaf vs. mean stem variable; *P<0.05.
† For details of procedures and models, see p. 554.
‡ λ1 and k2 are rate parameters for escape from raft and backflow from VRR to raft respectively.

Downloaded from https://www.cambridge.org/core. IP address: 54.191.40.80, on 05 Apr 2017 at 14:57:19, subject to the Cambridge Core terms of use, available at https://www.cambridge.org/core/terms. https://doi.org/10.1079/BJN2001317
The values for the slow turnover pool in all models are not strictly comparable since they involve different assumptions concerning functional pool size which affect their numerical value as will be discussed subsequently.

**Discussion**

The question of which compartmental model to use to describe marker excretion or concentration in various parts of the digestive tract and in faeces has been debated frequently (Faichney & Boston, 1983; Dhanoa et al. 1985; France et al. 1985; Faichney, 1986; Spillane & Minson, 1986; Huhtanen & Kukkonen, 1995; Aharoni et al. 1999). The search for new models continues only in the desire to define more accurately the biological system in operation. New model formulation alone would be of no use unless it alters the way in which the system in question is perceived. It should be clearly recognized that the construction of models of digesta flow are done to identify the biological pathways involved and to provide the mathematical means to quantify transfer of digesta, the latter being important in the examination of diet effects on intake and digestion.

**Models, assumptions and linkage of sampling site data**

The three types of models used in the current report have different assumptions and utilize the same data in different ways to provide estimates of residence time. The simplest model, the G1 model, assumes that a single exponential distribution of residence times describes the mass action, diluting effect of particle turnover from the sampling site. Imperfect mixing and effects other than mass action are disregarded by fitting the G1 model only to the descending segment of the data. This is the most commonly used model by biologists in examining the turnover of a marker in a single pool.

In contrast to the G1 model, the G2 → G1 sequential two compartment model assumes that the distribution of residence times at the sampling site is the result of two sequential compartments. These two sequential compartments are distinguished by different quantitative and qualitative attributes of the residence time distributions assumed for each individual compartment, i.e. gamma 2 age-dependent (G2) and exponential (G1). When fitted to data, the G2 distribution of residence times is associated with the faster turnover compartment while the G1 distribution is associated with the slower turnover compartment (Pond et al. 1988). In the G2 → G1 model, sampling is assumed to be from the G1 compartment and the G2 compartment represents a preceding mixing compartment. If the assumptions are correct for both the G1 and G2 → G1 models, then the mean residence time for the slower residence time distributions, \(1/k\) and \(1/k_2\) respectively, should be equal since they represent turnover at the sampling site. That \(1/k\) does not consistently approximate \(1/k_2\) (Table 2) may indicate as incorrect the assumption of a homogenous exponential distribution of residence time in the descending segment of data as estimated by the G1 model. By assuming a ‘non-mechanistic’, age-dependent compartment to describe the distribution of shorter residence times within the G2 compartment, the G2 → G1 model should provide unconfounded estimates of residence time distributions for the longer residence time distributions associated with the age-independent, mass action diluting (exponential, G1) processes.

Ellis et al. (1984b) and Wylie et al. (2000) suggested that the shorter residence time distribution represented by the age-dependent G2 compartment of the G2 → G1 model (\(2/k\)) is the result of several processes affecting the movement (mixing) of particles within the ruminal digesta. Such imperfect mixing results in differences in concentration of marker between dosing and sampling site. In the present study, the dosing and sampling site were the same, i.e. the ruminal cannula. However, ascending values exist in the data indicating that dose and sampling sites within the digesta were obviously not the same even though the two were anatomically the same (Fig. 2). The results suggest that the marked dosed particles migrated away from the dosed site as a bolus and then dispersed back to the dose site which subsequently served as the sample site. In the present data, the mean compartmental residence time associated with these age-dependent processes (\(2/k\)) ranged from 7.5 to 16.1 h, did not differ due to sampling site nor to marked plant part and averaged 11.3 h (Table 2).

Without specifying mechanisms, the age-dependent compartment of the G2 → G1 model accounts for residence time involving mixing between dosed and sampled sites and the exponentially distributed, age-independent compartment accounts for mass action effected turnover at the sampled site. Thus, the mean residence time associated with marker emergence at (G2) and diluting turnover from (G1), the sampling site, exceeds that estimated from the G1 turnover model. The mean residence time estimated by the G2 → G1 system (MRT in the dorsal rumen and MRTV) differed according to the digesta site sampled and plant part marked (Table 2). Differences in mean compartmental residence time in the G2 → G1 model system were primarily due to differences in mean compartmental residence time associated with turnover at the sampled site (1/k_2). The compartmental mean residence time due to turnover, \(1/k_2\), was slower from the VRR site for \(169\)Yb-labelled leaf (25.1 v. 10.1 h), similar for \(144\)Ce-labelled stem (45.3 v. 44.3 h), and slower for \(144\)Ce-labelled stem than \(169\)Yb-labelled leaf from both the VRR (44.3 v. 25.1 h) and the raft (45.3 v. 20.1 h).

In contrast to the G1 and G2 → G1 models, which were fitted to each individual sampling site, the G2=G1 model (or raft model) was simultaneously fitted to data from both sampling sites which is a new procedure. The mean compartmental residence time for the G2=G1 compartmental system (total MRT) was intermediate to that estimated by fitting the G2 → G1 to each individually sampled site (MRT in the dorsal rumen and MRTV, Table 2). Differences between MRT in the dorsal rumen and MRTV appeared primarily due to differences between sampled sites in the exponentially distributed residence time (1/k_2). These differences contributed to the dominance of mean compartmental residence time estimated by the G2=G1 system being apportioned to the non-exponentially distributed, age-dependent raft compartment (MRTR, Table 2). Thus, the interaction of two pools that differed due to their modelled turnover appears to be the cause of
the age-dependent turnover from the raft as modelled by the G2→G1 system and portrayed in Fig. 1.

It should be noted that the mean compartmental residence time of particles in the conceptual raft pool, MRTR, exceeds that estimated by any of the models applied to individual sampling sites. This is a consequence of the raft pool being defined as a pool of related compartments, R1 and R2 (Fig. 1), whose turnover via the particle flow paths sampled is age-dependent. Thus, the raft pool is defined by function of digesta attributes affecting flow properties of particles and not necessarily identifiable in terms of static, physical properties of rumen digesta such as ‘raft’, ventral digesta, large particles or small particles. In the present data, it appears that the ‘raft’ or age-dependent flow pool dominates the residence time of particles in the rumen digesta if the two sampling sites employed in the current study represent the flow path of all particles through the rumen digesta. Similar conclusions were found by Huhtanen & Kukkonen (1995).

The age-dependent mean residence time for flow between dosing and sampling sites as estimated by the G2→G1 model, 2λ1, appears to be relatively unaffected by the variables studied and averaged 11·3 h. If this is taken as an estimate of mean residence time due to slow and imperfect mixing within all digesta, then the mean residence time for age-dependent turnover from the total ‘raft’ pool can be estimated by deducting 11·3 h from MRTR for the ‘raft’. This yields 19·8 and 34·1 h for age-independent, mass action estimated by deducting 11·3 h from MRTR for the ‘raft’. These concepts were developed by Sutherland (1988) using lucerne whilst the variables estimated in this study are derived from tropical grass leaf and stem particles.

The actual values were estimated with varying accuracy and this is reflected also in the standard errors associated with MRT in the raft and VRR which were large for the VRR. Estimates of \( k_0 \) and \( k_2 \) were somewhat sensitive to initial starting values used in PCNONLIN (Statistical Consultants, Inc.) but the order of parameter estimates \( k_0 > \lambda_1 > k_2 \) never changes, irrespective of the order of the initial starting values, being at least 3-fold different. It may be concluded that little reliance can be placed on some of the numerical estimates of the parameters but the order of these estimates is reliable. The order is important because that outlines the mechanism for digesta flow through the rumen and clearly identifies turnover of particles from the raft as the rate-limiting component. The order is also important because this sequence of rate constants (slow raft, fast VRR) differs from the sequence derived in particle size models (fast large particle, slow small particle) (Poppi et al. 1981; Dixon et al. 1983; Faichney, 1986) and earlier models (Blaxter et al. 1956). Some caution should be exercised in these interpretations because estimated pool size has a major influence on the estimates of residence times and rate constants.

The implications of the particle size model were that size was a factor constraining escape and that reduction in size was inversely correlated with the probability for escape (Poppi et al. 1980, 1985). Sutherland’s (1988) model, however, has the concept that fermentability is a constraining factor, a factor persisting throughout particle size reduction. In support of this is the observation that most digestion of bermuda grass fibre (approximately 0·9) occurs within the rumen (McCarthy et al. 1987) a feature which must occur if flotation and sedimentation are important factors influencing escape potential of particles. Both models imply escape as the basic problem but the current
examination of Sutherland’s model specifies escape from the raft as a rate-limiting component, whereas particle size models are not specific other than identifying escape of small particles as the rate-limiting component. Thus, raft structure, consistency and properties influencing particle movement through the raft require investigation (Faichney, 1986; Bailoni et al. 1998). Vega & Poppi (1997) have examined this by labelling faecal particles (fermentatively spent) or feed particles (large fermentative capacity) in the size range 0.5–1.12 mm. These ‘small’ particles as compared with the >1.6 mm used in the present study escaped at similar rates irrespective of source (feed or faeces), but the retention time was markedly influenced by the rumen conditions resulting from the type of diet which was consumed (concentrate, pelleted lucerne, pangola hay or lucerne hay). Together, these results suggest that features of the raft (size, dispersion) markedly influence retention time rather than the extent of fermentation of a particular particle.

The MRT of stem particles in the raft was longer than leaf particles and this agrees with the suggestion by Sutherland (1988) that stem particles have better buoynancy characteristics and maintain these longer than leaf characteristics. Masticated leaf and stem particles contained within nylon bags have similar digestion rates (Poppi et al. 1981) but disintegration of stem particles by digestion and detrition is slow whereas a significant proportion of leaf particles can disintegrate by this pathway (McLeod & Minson, 1988) and these properties influence the buoynancy characteristics of stem.

The values for the proportion of the total rumen DM as raft appear high, as Dixon et al. (1983) with two steers recorded a mean value 0.54. However, their animals apparently were not fed ad libitum and Robinson et al. (1987) have shown that the raft size varies in relation to level of feeding where raft digesta weight as a proportion of total digesta weight varied from 0.42 to 0.96 as intake in cows increased from 6 to 24 kg/d. The raft in three steers offered bermuda grass hay ad libitum varied from 0.7–0.95 total rumen DM as determined by manual emptying (DP Poppi and WC Ellis, unpublished results). This estimate is comparable with that estimated by the two marker methods used here for animals grazing bermuda grass (leaf 0.75–0.83, stem 0.84–0.89). The difference in raft proportion as estimated by leaf and stem markers may simply relate to the fact that there is no physical boundary to the two pools and the pools are defined as groups of particles having similar characteristics. This may be different for leaf and stem particles. In any case the raft effectively occupies most of the rumen which is greater than might be envisaged from observations on animals with restricted levels of intake (Evans et al. 1973; Dixon et al. 1983).

What needs to be considered is whether the non-raft pool, designated VRR here, is quantitatively important when it apparently accounts for a low proportion of the total MRT of marker in the rumen (0.13 leaf, 0.19 stem) and thus the raft is the primary pool of interest.

**Model assumptions**

Two model assumptions require clarification: (1) the nature of outflow from the VRR; (2) mixing and age dependency in changes in marker concentration in the raft. Outflow from the ventral rumen has been assigned a rate parameter, k₀, and in many other passage rate studies similar values have been calculated for the whole rumen (e.g. Poppi et al. 1981). Such a representation implies that k₀ is a diet characteristic and the objective in application of these models and in simulations is to use these values (determined experimentally) in a wider context. If outflow (mass/time) is a function of physiological state or energy deficit of the animal (Weston, 1996) and perhaps raft characteristics, then the rate parameter, k, calculated is a useful numerical tool within the limits of the experiment. However, it may be limited in its wider application because its calculation is based on a pool size. What is required to be measured in relation to intake regulation is mass transport per time (a function of reticular contraction rate and mass transported per contraction) and the variation inherent in that rather than the variation in k. The representation of digestion rate as a rate parameter is more meaningful biologically as it is independent of pool size, is a characteristic of the substrate and there are no limits on mass digested per unit time. Most biologists would agree that newly ingested particles require time to be colonized by bacteria and that the rate parameters applying to digestion and passage would change with time before reaching some plateau value. Most studies ignore this but the stochastic approach used by Matis (1972, 1984, 1987) and Pond et al. (1988) have incorporated this. The gamma rate functions used, where the rate parameter increases with time at varying rates (different order of gamma function) up to a plateau, have been particularly useful (Pond et al. 1988). This has been used in the present study.

The question remains as to why some groups have needed to incorporate this approach whilst others have been able to ignore it. It appears that it is largely due to the method of marker introduction into the rumen and to the slow mixing (dispersion) of marker occurring between site of introduction and sampling. Those studies where marker is injected into a number of sites in the rumen minimize imperfect mixing as a component of marker dilution in the rumen or any pool with the rumen. The present study described here had large particles dosed into one discrete site in the raft. Sampling at a different site to the dosing site would lead to an ascending concentration curve as the marker dispersed throughout the pool to attain an equilibrium value. Thereafter, any further changes would only be due to disappearance of marker from the rumen and hence a descending concentration curve. This was obtained in the present study. In contrast, sampling at the bolus site would lead to an initial rapid dilution as the marker mixed and another dilution rate relating to disappearance. Employment of both dosing procedures (number of sites, single site) is in error if the aim is to examine the disappearance of a meal and both represent the extremities of the real situation.

Consumption of a labelled meal would represent the situation more realistically and where this was used (Dixon et al. 1983) exponential disappearance curves adequately explained marker disappearance. Thus, the mixing component, a feature in the present data, was not important where
Fig. 3. Theoretical concentration curves of marker in the raft after injection of labelled bolus into raft illustrating mass action, age-independent turnover from the dosed site, i.e. marker sampling site close to dispersing bolus (— — —), age-dependent turnover from the dosed site and/or marker sampling site separated from dispersing marker bolus (— — —). Note the slow mixing and separation of dosing and sampling sites results in initially ascending concentrations of marker concentration as marker dispersion to the sampling site achieves equilibrium. Also note that curve (— — —) expressing marker concentration at the sampling site would become curve (— — —) when marker concentration is expressed as a function of total rumen digesta and the slow initial disappearance represents age-dependent turnover, i.e. particle disappearance rate is initially low and increases with time to a plateau rate parameter as distinct from dispersion functions.

The importance of this is whether ignoring age dependency introduces error in estimating MRT of the overall meal in the rumen. In the data of Dixon et al. (1983) it would appear not to be great, but it is not known how universal this is, particularly as their animals were fed at restricted levels of feeding which would minimize age-dependent effects. Imperfect mixing may well be a feature of very full rumens as occurs under ad libitum feeding and marker injection into a number of sites may hide a real biological phenomenon. Sutherland’s concept (1988) of particle movement through the rumen also highlights a very real age-dependent process of particles losing buoyancy. Thus, caution is required in interpreting current marker studies and the desire of researchers to fit single exponential curves to this data may mask real biological processes which contribute to residence time in the rumen.

The stochastic approach of Matis (1984) offers one versatile way of incorporating this whilst Dhanoa et al. (1985) offer another based on multiple exponential pools. This possibility is exemplified by examining the rate constants and their inverse, the MRT, associated with the descending portion of the marker concentration curve (G1 model) (Tables 1 and 2). Sampling sites (raft or ventral rumen) did not influence the values obtained but the total MRT in the rumen was greater than these values by approximately 20% (leaf) and 35% (stem). The current data set do, however, exaggerate this phenomenon by the method of dosing. The important point is that an accurate estimate of MRT is required and the MRT derived from a simple one pool exponential model may be inappropriate in some circumstances.

**Model validation**

Whether the raft model is an appropriate way to examine marker kinetics in the rumen is not known particularly in the separation of raft and VRR pools. Of concern is the model output indicating the size and inertness of the raft and the rapid turnover of the ventral rumen with little mixing between the two pools. Particle size models have not identified these characteristics and have assumed that particles of a particular size can move throughout the rumen.

The issue of which model to fit has implications beyond marker kinetics in the rumen. The particle size model identifies disappearance of small particles as the rate-limiting step to increasing passage rate and this can be manipulated by increasing the area of the reticulum (increasing reticulum contraction rate and particles carried per contraction). The current model identifies the raft and particle entrapment within the raft as the rate-limiting component and to manipulate intake in these circumstances pool size needs to increase. This occurs in lactation, where the animal is faced with the need to increase intake to meet increasing nutrient demand.

It was concluded that the raft and the VRR appear to form two distinct pools within the rumen with particles entering the raft having escape rates that increase with time (age dependent) but which are still quantitatively slow in comparison to turnover in the VRR.

**Acknowledgements**

The financial assistance of BARD for this project is gratefully acknowledged. The assistance of Sandra Whitmore and M.J. Wylie is appreciated.

**References**


Ehrlein HJ (1979) Motility of the Forestomachs in Ruminants. Gottingen: Institut Wiss, Film.


Poppi DP, Hendricksen RE & Minson DJ (1985) The relative resistance to escape of leaf and stem particles from the rumen of

validity of the critical size theory for particles leaving the 

Poppi DP, Minson DJ & Ternouth JH (1981) Studies of cattle and 
sheep eating leaf and stem fractions of grasses. III. The 
retention time in the rumen of large feed particles. *Australian 
Journal of Agricultural Research* 32, 123–127.

on Modelling the Passage of 
Spillane KT & Minson DJ (1986) The relationship between 
and lower gut of sheep, the volume of fluid in the rumen and 
condition as factors affecting passage of liquid and digesta 
to the clearance of digesta from the reticulorumen. In 
[S. Sutherland TM (1988) Particle separation in the forestomachs of 

Sutherland TM (1988) Particle separation in the forestomachs of 

Thornton RF & Minson DJ (1973) The relationship between 
apparent retention time in the rumen, voluntary intake, and 
apparent digestibility of legume and grass diets in sheep. 

Ulyatt MJ, Blaxter KL & McDonald I (1967) The relations 
between the apparent digestibility of roughages in the rumen 
and lower gut of sheep, the volume of fluid in the rumen and 

Ulyatt MJ, Dellow DW, John A, Reid CSW & Waghorn GC 
(1986) Contribution by chewing eating and rumination to 
the clearance of digesta from the reticulorumen. In *Control of 
Digestion and Metabolism in Ruminants*, pp. 498–515 [LP 
Mulligan, WL Grovum and A Dobson, editors]. Englewood 
Cliffs, NJ: Reston.

Vega A & de Poppi DP (1997) Extent of digestion and rumen 
condition as factors affecting passage of liquid and digesta 

Weston RH (1996) Some aspects of constraint to forage 
consumption by ruminants. *Australian Journal of Agricultural Research* 47, 175–197.

Waghorn GC, Reid CSW, Ulyatt MJ & John A (1986) Feed 
commiption, particle composition and distribution between the 
four compartments of the stomach in sheep fed chaffed lucerne 
hay at two feeding frequencies and intake levels. *Journal of 

Wylie MJ, Ellis WC & Matis JHB Bailey EM James WD Beever DE 
segments of the digestive tracts of cattle. *British Journal of 
Nutrition* 83, 295–306.

---

**Appendix**

**Solution of model equations from gamma 2 rate functions**

Definitions:

- \( X_{R1}(t) \), \( X_{R2}(t) \) is the marker in dosing and sampling 
sites of raft pool respectively;
- \( X_v(t) \) is the marker in VRR pool;
- \( V_R, V_v \) is the quantity of DM in raft and VRR pools 
respectively (assumed constant over time);
- \( C_{R1}(t), C_{R2}(t) \) is the concentration of marker in dosing 
and sampling sites of raft respectively;
- \( C_v(t) \) is the concentration of marker in VRR pool;
- \( X_i(t), C_i(t) \) is the derivatives of \( X_i(t) \) and \( C_i(t) \);
- \( \lambda_1, k_0, k_2 \) are the rate parameters for escape from raft, 
escape from VRR to omasum, and backflow from VRR to raft respectively;
- \( r \) is the \( V_v/V_R \), which is the ratio of quantities 
(VRR:raft);
- \( K \) is the coefficient matrix;
- MRTR, MRTV is the MRT in raft and VRR respectively;
- MRTT is the MRT in total system.

Equations:

\[
\begin{align*}
\dot{X}_{R1}(t) &= -\lambda_1 X_{R1}(t), \\
\dot{X}_{R2}(t) &= -\lambda_1 X_{R2}(t) + \lambda_1 X_{R1}(t) + k_2 X_v(t), \\
\dot{X}_v(t) &= -(k_0 + k_2) X_v(t) + \lambda_1 X_{R2}(t), \\
\dot{C}_{R1}(t) &= X_{R1}(t)/V_R = -\lambda_1 C_{R1}(t), \\
\dot{C}_{R2}(t) &= X_{R2}(t)/V_R = -\lambda_1 C_{R2}(t) + \lambda_1 C_{R1}(t) + k_2 r C_v(t), \\
\dot{C}_v(t) &= X_v(t)/V_v = -(k_0 + k_2) C_v + \lambda_1 C_{R2}(t)/r.
\end{align*}
\]

These linear differential equations yield non-linear 
regression models. The parameter estimation for such 
models requires specialized statistical software (Bates & 
Watts, 1988). This study used PCNONLIN (Statistical 
Consultants Inc.) which utilizes numerical integration to 
solve the differential equations.

Residence time denotes the elapsed time accumulated by 
a particle during all of its visits to a site, i.e. residence time 
is the sum of one or more retention times. Matis *et al.* (1983) show that the mean residence times are given by 
\( K^{-1} \), where \( K \) is the matrix of rate coefficients. For 
this model, one has:

\[
K = \begin{bmatrix}
\lambda_1 & 0 & 0 \\
-\lambda_1 & -\lambda_1 & k_2 \\
0 & -k_0 & -k_2
\end{bmatrix}
\]

and

\[
-K^{-1} = \begin{bmatrix}
a_1 & 0 & 0 \\
a_2 & a_2 & a_3 \\
a_4 & a_4 & a_4
\end{bmatrix},
\]

where

\[
a_1 = \lambda_1^{-1},
\]

\[
a_2 = (k_0 + k_2)/(k_0 \lambda_1),
\]

\[
a_3 = k_2/(k_0 \lambda_1),
\]

and

\[
a_4 = k_0^{-1}.
\]

Hence

\[
\begin{align*}
\text{MRTR} &= a_1 + a_2 = (2k_0 + k_2)/(k_0 \lambda_1), \\
\text{MRTV} &= a_4 = k_0^{-1}, \\
\text{MRTT} &= \text{MRTR} + \text{MRTV} = (2k_0 + \lambda_1 + k_2)/(k_0 \lambda_1).
\end{align*}
\]