Thermodynamic modeling of ruminal fermentations

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Abstract – The main weakness of current rumen models is their inability to explain the end-products of fermentation (volatile fatty acids, gas, and microbial matter) in a satisfactory way. The objective of this study was to improve this prediction based on the application of thermodynamic laws to microbial metabolism. Therefore, a dynamic model of rumen fermentation was developed. In this original approach, the reactions evolved with a decrease in Gibbs energy according to thermodynamic laws. Some simulated results obtained with this model were similar to experimental observations. The predicted post-prandial evolution of the volatile fatty acid profile was satisfactory. However, simulations of some other parameters (pH, redox potential) were less reliable and further studies should be conducted to represent these better. The results allowed some properties resulting from thermodynamic principles to be identified and strengthened the importance of this subject.


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NOMENCLATURE

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Name</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>[x]</td>
<td>concentration of species x</td>
<td>mol·L⁻¹</td>
</tr>
<tr>
<td>E</td>
<td>oxidation-reduction potential</td>
<td>V</td>
</tr>
<tr>
<td>F</td>
<td>Faraday constant (96 485.31 C·mol⁻¹)</td>
<td>C·mol⁻¹</td>
</tr>
<tr>
<td>G</td>
<td>extensive Gibbs energy of a system</td>
<td>kJ</td>
</tr>
<tr>
<td>ΔrG</td>
<td>reaction Gibbs energy for specified concentrations of species at specified T, P</td>
<td>kJ·mol⁻¹</td>
</tr>
<tr>
<td>ΔrG(x)</td>
<td>Gibbs energy of formation of species x at specified T, P</td>
<td>kJ·mol⁻¹</td>
</tr>
<tr>
<td>H</td>
<td>extensive enthalpy of a system</td>
<td>kJ</td>
</tr>
<tr>
<td>ΔrH⁰(x)</td>
<td>standard enthalpy of formation of species x at specified T, P</td>
<td>kJ·mol⁻¹</td>
</tr>
<tr>
<td>k₁, k₂</td>
<td>rate constant of the forward and reverse reaction</td>
<td>dimensionless</td>
</tr>
<tr>
<td>n</td>
<td>number of electrons involved in a cell reaction</td>
<td>dimensionless</td>
</tr>
<tr>
<td>P</td>
<td>pressure</td>
<td>atm</td>
</tr>
<tr>
<td>pH</td>
<td>– log[H⁺]</td>
<td>pH unit</td>
</tr>
<tr>
<td>R</td>
<td>gas constant (8.31451 J·K⁻¹·mol⁻¹)</td>
<td>J·K⁻¹·mol⁻¹</td>
</tr>
<tr>
<td>r(t)</td>
<td>rate of the reaction</td>
<td>mol·L⁻¹·min⁻¹</td>
</tr>
<tr>
<td>T</td>
<td>temperature</td>
<td>K</td>
</tr>
</tbody>
</table>

1. INTRODUCTION

During recent decades, farm animal models and feed evaluation systems have been developed in most industrialized countries. The role of the rumen in the nutritional response of the ruminant to its diet is essential. Models of whole rumen function have been proposed [2, 9, 19, 32] although these do not contain a mechanistic means to predict the variation of carbon (C) partitioning among microbes, gas and volatile fatty acids (VFA) [33]. The validation of the prediction of VFA profiles pointed out that simulations have not been entirely satisfactory [3, 11, 28]. Regarding the importance of these products [10], the current study intends to focus on this aspect. It was hypothesized that the failure of currently published models to predict C partitioning is due to their lack of explicit representation of thermodynamic principles.

Thermodynamic principles could be applied to determine which process can occur, in which direction and how strong the tendency is for the changes to take place. It is based on the principle that substances tend to react to achieve the lowest energy state. Kinetic laws rather describe the rates of the reactions. For a given bacterial species, the metabolic pathways and flows are mainly determined by the enzymatic environment, which is the outcome of the cell’s genetic material. Therefore, in a monoculture, the kinetic laws are more important than thermodynamic ones. In contrast, in the case of a highly diversified microflora, like in the rumen, numerous metabolic pathways are a priori possible. So, it can be assumed that successful species and metabolic pathways are those which optimize the thermodynamic yield. Consequently, in the rumen, thermodynamic laws could play a more important driving role than kinetic laws. Some global thermodynamic approaches of microbial ecosystems have already been developed. These are mainly based on the use of macrochemical equations expressing system inputs and outputs and take into account the principles of Gibbs energy [14, 38]. Kohn and Boston [17] developed a dynamic model of glucose fermentation based on thermodynamic limits. However, the thermodynamic efficiencies
were considered static at steady state. To progress in this area, a simple thermodynamic driven model of ruminal fermentations was developed with the objective to predict variation in C flow partitioning among VFA, fermentation gas and microbial biomass.

2. MATERIALS AND METHODS

2.1. Application of thermodynamic principles in mathematical modeling

Thermodynamic principles state that spontaneous processes have a negative Gibbs energy change ($\Delta rG < 0$). In this situation, the system releases energy to its surroundings as the process occurs. This principle was the current working assumption and the magnitude of the Gibbs energy change was assumed to be one of the driving forces of biochemical reactions occurring in the rumen. Thermodynamic principles applied to biochemical reactions were integrated in the model. Let us consider a reversible reaction: $aA + bB \xleftrightarrow{\frac{k_1}{k_2}} cC + dD$, where the symbols A and B stand for the reactants of the reaction and the symbols C and D stand for the products of the reaction. The rate constant $k_1$ is for the forward reaction, $k_2$ is that of the reverse reaction. In writing biochemical equations, the correct stoichiometric coefficients (a, b, c, d) should be used to balance atoms (e.g., C, H, O) and charges if necessary. The overall rate ($r$) of the reaction can then be defined by:

$$r(t) = \frac{-1}{a} \frac{d[A]}{dt} = \frac{-1}{b} \frac{d[B]}{dt} = \frac{1}{c} \frac{d[C]}{dt} = \frac{1}{d} \frac{d[D]}{dt}. \quad (I)$$

Thus, biochemical reactions can be modeled with differential equations, describing the change in concentration of the various chemical species over time. The rate can also be written:

$$r(t) = k_1 \frac{[A]^a[B]^b}{K_{eq}} - k_2 \frac{[C]^c[D]^d}{K_{eq}}. \quad (II)$$

Equations (I) and (II) were the basic concepts for the construction of the current dynamic model of biochemical reactions.

Thermodynamic principles help to determine the rate constants for forward ($k_1$) and reverse ($k_2$) reactions. The change in Gibbs energy ($\Delta rG$, kJ-mol$^{-1}$) of the reaction is given by:

$$\Delta rG = \Delta rG^0 + RT \ln(K)$$

with

$$K = \frac{[C]^c[D]^d}{[A]^a[B]^b} \quad (III)$$

$\Delta rG^0$ represents the standard Gibbs energy change ($T = 298$ K, $[x] = 1$), $R$ the gas constant, $T$ the temperature in K, $[x]$ the activities of reactants and products expressed as a concentration in mol-L$^{-1}$ or as atmospheric pressure.

The thermodynamic laws may help to predict if a reaction would happen, whenever the Gibbs energy change is negative, the process is spontaneous. In a closed system, the Gibbs energy decreases to $\Delta rG = 0$, which corresponds to a dynamic equilibrium. When this thermodynamic equilibrium is reached, the equilibrium constant of the reaction ($K_{eq}$) can be specified from equations (II) and (III).

$$r(t) = 0 \rightarrow K_{eq} = \frac{k_1}{k_2} \quad (IV)$$

$$\Delta rG = 0 \rightarrow K_{eq} = e^{-\frac{\Delta rG^0}{RT}} \quad (V)$$

$K_{eq}$ is determined with the standard Gibbs energy change ($\Delta rG^0$) values, which can be calculated from tabulated thermodynamic data. Thus, in the model, the dynamic differential equations were written for each molecule involved in a reaction as below for A:

$$\frac{d[A]}{dt} = -ak_1 \frac{[A]^a[B]^b}{K_{eq}} + a \frac{k_1}{K_{eq}} \frac{[C]^c[D]^d}{K_{eq}}. \quad (VI)$$

These principles can be applied for oxidation-reduction reactions with electron...
(e⁻) and proton (H⁺) exchanges, which are common in the rumen: a Ox + ne⁻ + cH⁺ ↔ b Red, Red and Ox are respectively the reducing and the oxidizing agents, n is the number of electrons transferred in the half-reaction. The Nernst equation is also used to calculate the redox potential (E, volt) of the reaction:

\[ E = E^\circ + \frac{RT}{nF} \ln \frac{[Ox]^a[H^+]^c}{[Red]^b} \]  

(VII)

\[ E = E^\circ_{pH} + \frac{RT}{nF} \ln \frac{[Ox]^a}{[Red]^b} \]

with \( E^\circ_{pH} = E^\circ + c \times \frac{RT}{nF} \times \ln(10) \times pH \)

where \( E^\circ \) is the standard redox potential of the half-reaction (\( T = 298 \text{ K}, \; pH = 0 \)), \( E^\circ_{pH} \) is the standard redox potential for a given pH, F is the Faraday constant.

Two half-reactions should be coupled to achieve the electron balance. The electrons are then transferred from the reduction couple \((E_{\text{red}})\) to the oxidation couple \((E_{\text{ox}})\). The variation of Gibbs energy \((\Delta rG)\) is proportional to the redox potential of variation \((\Delta E)\) for the cell reaction:

\[ \Delta rG = -nF(E_{\text{ox}} - E_{\text{red}}) = -nF\Delta E. \]  

(VIII)

2.2. Model development

The model of ruminal fermentations was based on 23 compartments corresponding to the major biochemical molecules and another compartment representing microbial dry matter. The compartments are expressed in mol·L⁻¹ for solutions, in atm for gas and in g·L⁻¹ for microbial dry matter. Each of these is defined by a dynamic differential equation representing the difference between inflows and outflows in the compartment. The reactions are considered to be reversible and thermodynamic laws presented previously (IV) were used in the model. The Forrester diagram of the model
Thermodynamic modeling of ruminal fermentations is summarized in Figure 1. The equations, variables and parameters are listed in the annex. The model was developed using the Dynamo language [31]. The Euler integration method was used for the numerical integration of the 24 dynamic differential equations with a step time of 0.1 min. The simulation time was 4 days in order to reach a dynamic equilibrium. A simulation of day 4 is presented in the results. Feeding occurred at the beginning of each day.

2.2.1. Model compartments

2.2.1.1. Soluble carbohydrates

The system simulated a daily “bolus” intake of cellulose and degradable starch, main carbohydrates ingested by the animal, of respectively 50 and 25 glucose-equivalent moles. Offner et al. [29] compared different rumen models on their way of predicting starch digestion. Both cellulose and starch were degraded into glucose, with assumed fractional rates of 0.08 and 0.3%·min⁻¹ respectively. Glucose was then converted to pyruvate by glycolysis, which is summarized by one reaction:

\[
C_6H_{12}O_6 + 2(ADP + Pi^2-) + 2NAD^+ \leftrightarrow 2C_3H_3O_3^- + 2ATP + 2(NADH+H^+) + 2H_2O. \quad (1)
\]

The pyruvate formed was converted to volatile fatty acids. Two pathways can lead to the formation of propionate. Pyruvate can be reduced to propionate by the lactate pathway (2, 3) or by the succinate pathway (4):

\[
C_3H_3O_3^- + NADH + H^+ \leftrightarrow C_3H_5O_3^- + NAD^+ \quad (2)
\]

\[
C_3H_5O_3^- + NADH + H^+ \leftrightarrow C_3H_5O_2^- + H_2O + NAD^+ \quad (3)
\]

\[
C_3H_3O_3^- + 2NADH + 3H^+ + ADP + Pi^2- \leftrightarrow C_3H_5O_2^- + 2H_2O + 2NAD^+ + ATP. \quad (4)
\]

The succinate pathway (4) was assumed to produce 1 ATP although there is no confirmation of this value. Pyruvate can also be oxidized into acetylCoA, which is a precursor for acetate and butyrate. Reactions for acetate (C₂) and butyrate (C₄) formation are the following:

\[
C_3H_3O_3^- + (ADP+Pi^2-) + NAD^+ \leftrightarrow C_2H_2O_2^- + CO_2 + ATP + NADH \quad (5)
\]

\[
2C_3H_3O_3^- + (ADP+Pi^2-) + 2H^+ \leftrightarrow C_4H_4O_2^- + 2CO_2 + ATP + H_2O. \quad (6)
\]

Pyruvate can also lead to the formation of minor VFA in C₅ or C₆, like valerate:

\[
2C_3H_3O_3^- + 3NADH + 4H^+ \leftrightarrow C_5H_9O_2^- + CO_2 + 2H_2O + 3NAD^+. \quad (7)
\]

VFA leave the system with a fractional outflow rate of 0.5%·min⁻¹ corresponding approximately to their transit and absorption from the rumen.

2.2.1.2. Gas

Carbon dioxide is formed during reactions (5), (6) and (7). NAD⁺ can be regenerated by methanogenesis or acetogenesis from CO₂. The two reactions are assumed to be associated with the formation of 1 and 0.25 mol of ATP respectively.

\[
CO_2 + 4NADH + 4H^+ + (ADP+Pi^2-) \leftrightarrow CH_4 + 3H_2O + 4NAD^+ + ATP \quad (8)
\]

\[
2CO_2 + 4NADH + 3H^+ + 0.25(ADP+Pi^2-) \leftrightarrow C_2H_3O_2^- + 2.25H_2O + 4NAD^+ + 0.25ATP. \quad (9)
\]

The dynamics of H production and utilization was modeled with NAD⁺/NADH as the main electron acceptor and donor. The model does not consider other electron acceptors and donors even though others are known to exist. By convention, NAD⁺/NADH were chosen to represent the electron acceptor and donor. Hydrogen can be formed from H⁺ by an oxidation-reduction reaction:

\[
H^+ + NADH \leftrightarrow H_2 + NAD^+. \quad (10)
\]

CO₂, CH₄ and H₂ activities were expressed in atm. When gas pressure in the rumen
was above 1 atm, gas was assumed to escape the rumen relative to the gas proportions [17]. The relation was then similar to a mass action law.

2.2.1.3. Regulatory components

Protons (H\(^+\)) are involved in several reactions:
- formation of carbon dioxide from bicarbonate ions HCO\(_3^-\) coming from saliva (inflow of 0.01 mol·L\(^{-1}\)·min\(^{-1}\)):

\[
\text{HCO}_3^- + \text{H}^+ \leftrightarrow \text{H}_2\text{CO}_3 \leftrightarrow \text{CO}_2 + \text{H}_2\text{O}.
\] \(^{(11)}\)

- formation of hydrogen by oxidation-reduction according to reaction (10),
- equilibrium between formed anions and corresponding acids:

\[
\text{A}^- + \text{H}^+ \leftrightarrow \text{AH}.
\] \(^{(12–16)}\)

The pH had an influence on all the reactions where it is involved in the determination of equilibrium constants.

All oxidation-reduction reactions (1–5 and 7–10) can be considered to be indirectly coupled with NAD\(^+\)/NADH. In a closed system, the dynamic equilibrium is reached when \(\Delta G = \Delta E = 0\). An open system is never at equilibrium, \(\Delta E \neq 0\), therefore the convergence of the redox potentials of the different reactions over time is a priori not possible to obtain. However, the average redox potential of the rumen (\(E_{\text{mean}}\)) was defined by the weighed average of the redox potentials of each couple (\(E_i\)). The weighting coefficients were proportional to the amount of the oxidized and reduced forms of each oxidation-reduction couple. For a given oxidation-reduction couple, reaction flows have been considered as an exponential function of the difference between \(E_{\text{mean}}\) and \(E_i\). This principle was previously applied in modeling [20]. The objective was to favor the convergence of the redox potentials to an average value.

Finally, ATP is produced by fermentation processes (1, 4, 5, 6, 8, 9) and is used for maintenance and growth of microbes.

2.2.1.4. Microbial population

The amount of microbial dry matter in the rumen is regulated by flows related to growth, lysis and transit. The flows for growth and lysis are expressed as functions of the amount of microbial dry matter and basal growth rate of microorganisms. The rates for growth and lysis are also largely dependent on the ATP/ADP ratio. Growth is defined as a positive exponential function of \(x\), where \(x = k \times (\frac{\text{ATP}}{\text{ADP}} - 1)\). This expression allowed favoring growth when energy status was positive. In contrast, lysis is defined as a negative exponential function of \(x\). The flow for transit simply follows a mass action law. ATP and NADH are required for microbial growth and maintenance; this utilization depends on the amount of microbial dry matter. Carbon needed for microbial growth is assumed to come from pyruvate. The outflow of pyruvate changes with the apparent microbial growth. Microbial DM is supposed to contain 25% carbon [14].

2.2.2. Model parameters – equilibrium constants

The equilibrium constant of each reaction was calculated from the thermodynamic values of all molecules involved. Table I gives the standard thermodynamic properties as they are found in thermodynamic tables and provides the standard formation properties for the standard state, which is the state in a hypothetical ideal solution with all activities set to 1. The standard enthalpy of formation (\(\Delta_f H^\circ\)) does not depend on temperature, contrary to the standard Gibbs energy (\(\Delta_f G^\circ\)). The Van’t Hoff equation linked the two parameters to calculate values of \(\Delta_f G^\circ\) for different temperatures (Tab. I).
Table I. Thermodynamic properties for compounds of ruminal metabolism: standard Gibbs energy and enthalpy of formation. $\Delta_f G^\circ$ and $\Delta_f H^\circ$ expressed in kJ·mol$^{-1}$, pH = 0.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$\Delta f G^\circ$ at 298 K</th>
<th>$\Delta f H^\circ$</th>
<th>$\Delta f G^\circ$ at 311 K$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose</td>
<td>-916.97</td>
<td>-1263.78</td>
<td>-901.84</td>
</tr>
<tr>
<td>Pyruvate</td>
<td>-474.50</td>
<td>-596.60</td>
<td>-469.17</td>
</tr>
<tr>
<td>Lactate</td>
<td>-516.72</td>
<td>-686.64</td>
<td>-509.31</td>
</tr>
<tr>
<td>Propionate</td>
<td>-360.00</td>
<td>-511.08</td>
<td>-353.41</td>
</tr>
<tr>
<td>Acetate</td>
<td>-369.60</td>
<td>-556.80</td>
<td>-360.18</td>
</tr>
<tr>
<td>Butyrate</td>
<td>-392.04</td>
<td>-535.55</td>
<td>-364.91</td>
</tr>
<tr>
<td>Valerate</td>
<td>-368.40</td>
<td>-566.80</td>
<td>-360.18</td>
</tr>
<tr>
<td>HCO$_3^-$ (aq)</td>
<td>-587.10</td>
<td>-692.29</td>
<td>-582.51</td>
</tr>
<tr>
<td>H$_2$O (L)</td>
<td>-237.19</td>
<td>-285.84</td>
<td>-235.07</td>
</tr>
<tr>
<td>CO$_2$ (g)</td>
<td>-394.50</td>
<td>-393.60</td>
<td>-394.54</td>
</tr>
<tr>
<td>CH$_4$ (g)</td>
<td>-50.79</td>
<td>-74.85</td>
<td>-49.74</td>
</tr>
<tr>
<td>H$_2$ (g)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>H$^+$</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>ADP$^{3-}$</td>
<td>-1711.55</td>
<td>-2005.24</td>
<td>-1698.74</td>
</tr>
<tr>
<td>ATP$^{4-}$</td>
<td>-2573.49</td>
<td>-2997.91</td>
<td>-2554.98</td>
</tr>
<tr>
<td>Pi$^{2-}$</td>
<td>-1094.10</td>
<td>-1294.00</td>
<td>-1085.38</td>
</tr>
<tr>
<td>NAD$^-$</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>NADH$^{2-}$</td>
<td>22.65</td>
<td>-31.94</td>
<td>25.03</td>
</tr>
</tbody>
</table>

$^a$ Values at rumen temperature $T = 311$ K were calculated using the Van’t Hoff equation: 

$$
\Delta f G^\circ(T_2) = \frac{T_2}{T_1} \left( \Delta f G^\circ(T_1) - \frac{T_2 - T_1}{T_1} \times \Delta f H^\circ \right).
$$

Knowing the standard Gibbs energy of formation for each compound, the standard change in Gibbs energy of a reaction was calculated with the following formula: 

$$
\Delta_r G^\circ = \Sigma \Delta f G^\circ \text{ products} - \Sigma \Delta f G^\circ \text{ reactants}.
$$

Table II presents the change in standard Gibbs energy for key reactions in the rumen. Equilibrium constants ($K_{eq}$) could then be determined with equation (V). The rate constants for the reverse reaction were defined arbitrarily for the reactions, so that flows were realistic. Unfortunately, no direct measurements of reverse rates were available.

Numerous fermentation processes are oxidation-reduction processes. Each oxidation-reduction couple can be characterized by its redox potential. Table III presents the redox potentials of the half-reactions of the main oxidation-reduction equilibria occurring in the rumen. The redox potential in the rumen is around -0.35 V. This thermodynamic condition does not favor pyruvate, lactate or H$_2$ accumulation.

For the reactions of acid dissociation (12–16), the equilibrium constants are defined by $K_a = 10^{-pK_a} = \frac{[A^-][H^+]}{[AH]}$. The values of pKa for weak acids correspond to the pH at which the acid is half dissociated: lactic acid 3.86, acetic acid 4.75, butyric acid 4.82, valeric acid 4.84 and propionic acid 4.87. These values indicate that for usual conditions of rumen pH ($6 < \text{pH} < 6.5$), VFA are largely present under the dissociated form.

3. RESULTS

3.1. Thermodynamic validation

Simulated changes in Gibbs energy of the reactions ($\Delta_r G$) varied from 0
Table II. Change in standard Gibbs energy for key reactions in the rumen, expressed in kJ·mol⁻¹ for various conditions of pH and temperature.

<table>
<thead>
<tr>
<th>Reaction</th>
<th>ΔrG° at 298 K</th>
<th>ΔrG° at 311 K</th>
<th>ΔrG° (pH)</th>
<th>ΔrG° (pH = 7) at 311 K</th>
</tr>
</thead>
<tbody>
<tr>
<td>NAD &lt;&gt; NADH</td>
<td>22.65</td>
<td>25.03</td>
<td>ΔrG° + RTLn(10) × pH</td>
<td>66.7</td>
</tr>
<tr>
<td>ADP &lt;&gt; ATP</td>
<td>−5.03</td>
<td>−5.93</td>
<td>ΔrG° + RTLn(10) × pH</td>
<td>35.7</td>
</tr>
<tr>
<td>(1) Glu &lt;&gt; Pyr</td>
<td>1.7</td>
<td>3.2</td>
<td>ΔrG° − 2RTLn(10) × pH</td>
<td>−80.2</td>
</tr>
<tr>
<td>(2) Pyr &lt;&gt; Lac</td>
<td>−65.2</td>
<td>−64.9</td>
<td>ΔrG° + RTLn(10) × pH</td>
<td>−23.2</td>
</tr>
<tr>
<td>(3) Lac &lt;&gt; Pro</td>
<td>−104.2</td>
<td>−103.1</td>
<td>ΔrG° + RTLn(10) × pH</td>
<td>−61.4</td>
</tr>
<tr>
<td>(4) Pyr &lt;&gt; Pro</td>
<td>−173.0</td>
<td>−175.3</td>
<td>ΔrG° + 3RTLn(10) × pH</td>
<td>−50.3</td>
</tr>
<tr>
<td>(5) Pyr &lt;&gt; Ace</td>
<td>−35.7</td>
<td>−37.8</td>
<td>ΔrG°</td>
<td>−37.8</td>
</tr>
<tr>
<td>(6) Pyr &lt;&gt; But</td>
<td>−221.6</td>
<td>−217.1</td>
<td>ΔrG° + 2RTLn(10) × pH</td>
<td>−133.7</td>
</tr>
<tr>
<td>(7) Pyr &lt;&gt; Val</td>
<td>−356.6</td>
<td>−361.6</td>
<td>ΔrG° + 4RTLn(10) × pH</td>
<td>−194.8</td>
</tr>
<tr>
<td>(8) CO₂ &lt;&gt; CH₄</td>
<td>−226.3</td>
<td>−231.4</td>
<td>ΔrG° + 5RTLn(10) × pH</td>
<td>−23.0</td>
</tr>
<tr>
<td>(9) CO₂ &lt;&gt; Lac</td>
<td>−146.8</td>
<td>−147.2</td>
<td>ΔrG° + 3.25RTLn(10) × pH</td>
<td>−22.2</td>
</tr>
<tr>
<td>(10) H⁺ &lt;&gt; H₂</td>
<td>−25.0</td>
<td>−22.6</td>
<td>ΔrG° + RTLn(10) × pH</td>
<td>19.1</td>
</tr>
<tr>
<td>(11) HCO₃⁻ &lt;&gt; CO₂</td>
<td>−47.1</td>
<td>−44.5</td>
<td>ΔrG° + RTLn(10) × pH</td>
<td>−2.8</td>
</tr>
</tbody>
</table>

Table III. Redox potentials at pH = 0 and at pH = 7 and T = 311 K.

<table>
<thead>
<tr>
<th>Reaction</th>
<th>E⁺ (volt)</th>
<th>E⁻ (volt)</th>
<th>[Ox] / [Red] for Ez = −0.35 V</th>
</tr>
</thead>
<tbody>
<tr>
<td>(7) 2Pyruvate⁻ + 7H⁺ + 6e⁻ ↔ Valerate⁻ + CO₂ + 2H₂O</td>
<td>0.495</td>
<td>−0.009</td>
<td>6.98 × 10⁻³⁴</td>
</tr>
<tr>
<td>(3) Lactate⁻ + 2H⁺ + 2e⁻ ↔ Propionate⁻ + H₂O</td>
<td>0.410</td>
<td>−0.022</td>
<td>2.34 × 10⁻¹ⁱ</td>
</tr>
<tr>
<td>(4) Pyruvate⁻ + 4H⁺ + 4e⁻ ↔ Propionate⁻ + H₂O</td>
<td>0.309</td>
<td>−0.123</td>
<td>1.93 × 10⁻¹⁵</td>
</tr>
<tr>
<td>(2) Pyruvate⁻ + 2H⁺ + 2e⁻ ↔ Lactate⁻</td>
<td>0.208</td>
<td>−0.224</td>
<td>8.25 × 10⁻⁵</td>
</tr>
<tr>
<td>(8) CO₂ + 8H⁺ + 8e⁻ ↔ CH₄ + 2H₂O</td>
<td>0.161</td>
<td>−0.271</td>
<td>5.73 × 10⁻¹¹</td>
</tr>
<tr>
<td>(9) 2CO₂ + 7H⁺ + 8e⁻ ↔ Acetate⁻ + 2H₂O</td>
<td>0.059</td>
<td>−0.319</td>
<td>9.57 × 10⁻⁵</td>
</tr>
<tr>
<td>NAD⁺ + H⁺ + 2e⁻ ↔ NADH</td>
<td>−0.130</td>
<td>−0.346</td>
<td>7.42 × 10⁻¹</td>
</tr>
<tr>
<td>(10) 2H⁺ + 2e⁻ ↔ H₂</td>
<td>0</td>
<td>−0.432</td>
<td>4.55 × 10²</td>
</tr>
<tr>
<td>(5) Acetate⁻ + CO₂ + 2H⁺ + 2e⁻ ↔ Pyruvate⁻ + H₂O</td>
<td>−0.284</td>
<td>−0.716</td>
<td>7.28 × 10⁻¹¹</td>
</tr>
<tr>
<td>(1) 2Pyruvate⁻ + 6H⁺ + 4e⁻ ↔ Glucose</td>
<td>−0.095</td>
<td>−0.743</td>
<td>2.98 × 10²⁵</td>
</tr>
</tbody>
</table>

The redox potentials indicate that the reactions proceed in the spontaneous thermodynamic direction.

The system can either maximize the production of highly energetic molecules or minimize entropy (S) production depending on the energetic situation. There is a clear relationship between Gibbs energy and entropy production (G = H – TS). In case of energetic deficiency, the system tends to minimize the loss of Gibbs energy and so increase entropy. The system then favors the production of acetate. Sauvant et al. [34] reported that in this case, ATP and NADH production per carbon of VFA is greater. In contrast, in the most favorable situations, there is an adaptation to the energy and molecular hydrogen excess brought by the feeds; the system favors the production of longer chain VFA (propionate and minor VFA), leading to more molecular formation. Therefore, this situation is associated with lower entropy due to higher order (less molecules formed).

3.2. Study of glucose fermentation and VFA profile

The simulations of glucose fermentation showed, for thermodynamic reasons, that
glucose offered a rather strong resistance to microbial fermentation (Fig. 3). This was consistent with the presence of significant amounts of soluble carbohydrate observed in the rumen [21] or in vitro [4]. This could partly be explained by the low concentrations of ADP and NAD in the rumen. These two molecules are absolutely necessary for glycolysis.

Regarding the fluxes and partitions of VFA, the model reflected fairly well the dynamics and stoichiometry of the rumen (Fig. 4). Feeding was followed by an increase of propionate at the expense of acetate and butyrate. A post-prandial increase of the propionate to acetate and butyrate ratio is usually observed. However, this phenomenon was exacerbated compared to experimental observations. The acetate to propionate ratio (A/P) ranged normally between 2 and 5. Simulated values of A/P could reach an unrealistic value of 15.
during transient periods, especially before feeding. The A/P ratio was influenced by various factors, which are more or less linked to energy availability of the substrates. A literature review on starch digestion in the rumen indicated that the A/P ratio decreases with increasing amounts of starch digestible in the rumen [27].

3.3. Study of regulatory parameters

Predicted values for pH were fairly realistic, ranging from 6.4 to 6.7; however, the prediction of pH changes during the day was not satisfactory because it was opposite to experimental observations (Fig. 5). Indeed, after feeding, the increase in VFA
production leads normally to a rapid drop in pH followed later by an increase in pH. A delay probably occurred in the model where H⁺ ions were involved in acidobasic equilibrium after the formation of anions by the various metabolic pathways. This aspect challenges a next version of the model to correct for this delay.

The diurnal variation of the energy and the oxidation-reduction status in the rumen was satisfactorily predicted. The kinetics of ATP/ADP and NADH/NAD ratios well represented energy intake by feeding followed by energy utilization by microorganisms. The ATP/ADP ratio ranged between 0.1 and 5. The sensitivity of the VFA molar proportions to the initial ratio was considerable as illustrated in Figure 6 with different initial ratios of 0.25, 1 and 2. Typical rumen conditions correspond to a ratio of 0.25. Acetate production was favored at the beginning when the energy status of the ecosystem was low. This could be explained because ATP production per mole of carbon as VFA was more important in the case of acetate formation than for the other VFA. With the simulations, the redox potential ranged from –0.4 to –0.3 mV, which is fairly realistic. The lowest redox potential was observed before feeding; which is similar to experimental observations [5, 22]. Similar to what was observed for the ATP/ADP ratio, the sensitivity of the VFA profile to initial NADH/NAD ratio was considerable. The NADH/NAD ratio had a large influence on VFA partitioning. When the ratio was below 1, butyrate production was favored whereas acetate production decreased.

3.4. Study of methanogenesis and acetogenesis partition

Simulated fluxes for acetogenesis and methanogenesis (Fig. 7) were low compared to observed data. Acetogenesis always remained lower than methanogenesis, similar to what is observed in the rumen. Moreover, these two fluxes varied in the same direction and are closely linked with variations in NADH and NAD. Regarding gas production, model predictions were unsatisfactory because predicted methane production was too low. An inappropriate representation for the gas outflow rates is likely the cause of this. However, little data is available concerning the kinetics of gas production.
4. DISCUSSION

4.1. Potential and limits of the model

Until now, attempts to consider thermodynamic laws in rumen modeling have been scarce and not exhaustive [17]. The approach taken in this study is therefore original, integrating the major thermodynamic laws to model the fermentation processes in a mechanistic way. This approach was a first attempt to describe the biochemical equilibrium and to determine pH and redox potential. The ability of thermodynamic models to explain some ruminal digestive phenomena is highlighted. The results of the simulations were satisfactory and seemed realistic. The model was constructed based on sound biochemical information and integrated the major known regulations by the ratios of ATP/ADP and NADH/NAD+. These cofactors are involved in numerous metabolic pathways and may interact with the intensity of rumen fermentation. This current study expands on that proposed by Sauvant et al. [34]. In addition to the before mentioned cofactors, rumen pH and redox potential had a major impact on the equilibrium of fermentation.

Some aspects of the model were not satisfactory. The convergence of the redox potentials of the different reactions over time was not reached even for simulation lengths of ten days. Indeed, the range of rumen redox potential could be extensive [5, 24] and this parameter might influence metabolic activities [7] and therefore the concentrations of the different oxidation-reduction molecules. However, it is likely that the range of values provided by the model does not influence the global microbial activities [23]. The lack of convergence can be explained in open systems, like the rumen, which exchange matter with the surrounding area and that likely never reach thermodynamic equilibrium [39]. This result underlines that, in a context of an open system, it is necessary to cautiously apply the thermodynamic laws described for closed systems. Proton and electron transfers are closely linked. The proton motive force created by

Figure 7. Simulated fluxes of acetogenesis and methanogenesis.
the accumulation of H\(^+\) on one side of a cell membrane and not the other can also be responsible for the lack of convergence of the redox potentials [6]. This aspect of H\(^+\) metabolism should be better represented. The simulated pH differed from expectations and this could probably be due to a poor representation of acid dissociation reactions or of buffer-acid reactions, this could also be the outcome of an insufficient consideration of the role of bicarbonate [16]. Moreover, it would be interesting to modulate VFA absorption relative to pH. Pitt and Pell [30] proposed another approach to model rumen pH fluctuations from the ruminal buffer index (saliva production) and the deviation in rumen acidity (from VFA and lactate production).

Finally, it is important to point out that, by construction, the model was non linear. The size and the turnover of the compartments varied to a great extent. These two properties made the model very sensitive and difficult to parameterize.

### 4.2. Proposals for further developments

At this stage, it seems important to integrate a minimum of other information in the model by identifying characteristics that are not determined directly by thermodynamic laws. There is a need to include additional constraints in the model, and inclusion of kinetic laws appears to be the most relevant. Kinetic laws would help to define parameters like the rates of product formation (for instance, these reaction rates could depend on the influence of enzymes or co-factors on the reactions). In terms of fluxes of elements, mass action laws were only considered in the model. This study also highlighted the interest of experimental studies on the influence of change in concentrations of co-factors. However, little experimental data is available on this subject. The ATP/ADP and NADH/NAD ratios had a main influence on fermentation pathways, therefore, it seems necessary to better quantify the microbial specific requirements for ATP and NADH [13, 14]. More in vitro studies on the intracellular NAD/NADH ratio could bring valuable information [15]. This study also raises some questions for modeling microbial dynamics [12, 26]. When several microbial groups are involved, responses of population ecology are important to deal with the dynamics of populations of living organisms. Rumen bacteria can be classified in several homogeneous groups according to metabolic criteria, end-products of fermentation, sensitivity to change in redox potential or pH [35]. These parameters could be taken into account when building models of microbial populations. Ideally, a global model combining these sub-models could be developed with both thermodynamic laws (as the first assumption for a multi-species system) and the known interactions among different microbial species (responses of population ecology). Another issue raised by this study concerns the distinction of several rumen compartments. Indeed, fermentation takes place inside of microbial cells. The conditions there could be entirely different from the extracellular ruminal fluid conditions (e.g., pH may be quite different). Our approach made no difference between intracellular and extracellular levels. Though there are few quantitative data on the relations among different rumen compartments, it would be interesting to better characterize and possibly to distinguish a soluble fraction, a gaseous fraction, a contact area between dietary particles and microorganisms, and finally the rumen wall [8]. Various aspects of the model require further investigation, like the evaluation of the model sensitivity to some assumptions (compartment initial concentrations, reverse reaction rates); however, the results show that the integration of thermodynamic laws in rumen models would certainly help to better predict fermentation pathways.
5. CONCLUSION

The model favors reactions following thermodynamic principles. Reaction rates are determined by their change in Gibbs energy. The reactions are supposed to be reversible and evolve along a decrease in Gibbs energy. Tables of standard Gibbs energy for the different components, after adjustment to rumen temperature, allow determining the various necessary constants. Reverse reaction rates were determined arbitrarily, which is a strong limit of this model. The model simulations adequately represented the post feeding evolution of the main VFA concentration. Thermodynamic laws were therefore helpful to understand fermentation pathways. The model was developed to be included in whole rumen models, or to be completed by incorporating new reactions. However, difficulties arise during model development because the system is open and gathers a large network of metabolic reactions. This makes the model very sensitive. Some results were not entirely satisfactory, especially for pH and redox potential. As represented, the thermodynamic laws were not sufficient to explain all observed variation. Other driving forces from kinetics and population ecology should also be taken into account, especially to modulate forward and reverse reaction rates.

REFERENCES


[16] Kohn R.A., Dunlap T.F., Calculation of the buffering capacity of bicarbonate in the
Thermodynamic modeling of ruminal fermentations


Appendix: Parameter values and list of equations of the thermodynamic model.

1. PARAMETERS

The compartments and their initial values, chosen to represent typical rumen conditions:

<table>
<thead>
<tr>
<th>Compartment</th>
<th>Initial Value</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>ST</td>
<td>0</td>
<td>Starch</td>
</tr>
<tr>
<td>CW</td>
<td>0.05</td>
<td>Cell wall</td>
</tr>
<tr>
<td>GLU</td>
<td>1E-4</td>
<td>Glucose</td>
</tr>
<tr>
<td>PYR</td>
<td>1E-8</td>
<td>Pyruvate</td>
</tr>
<tr>
<td>LAC</td>
<td>1E-3</td>
<td>Lactate</td>
</tr>
<tr>
<td>ACP</td>
<td>4E-4</td>
<td>Propionate</td>
</tr>
</tbody>
</table>

k = 1 Reverse reaction rates (min\(^{-1}\))
C = 10 Exponential factor used to favor convergence of redox potential
KA = 1E2 Acid formation rate (min\(^{-1}\))
KA1 = 0.0001 VFA outflow rate (min\(^{-1}\))
KA2 = 0.005 Outflow rates of corresponding anions (min\(^{-1}\))
KCW = 0.0008 Cellulose degradation rate (min\(^{-1}\))
KOUTCW = 0.0001 Cellulose outflow rate (min\(^{-1}\))
KST = 0.003 Starch degradation rate (min\(^{-1}\))
KOUTST = 0.0001 Starch outflow rate (min\(^{-1}\))
KM = 0.0001 Microorganism transit rate (min\(^{-1}\))
KMU = 0.075 Microorganism growth factor
MUBAS = 0.001 Microorganism basal growth rate (min\(^{-1}\))

CONSTANTS

<table>
<thead>
<tr>
<th>Constant</th>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>F</td>
<td>96.487</td>
<td>Faraday constant</td>
</tr>
<tr>
<td>R</td>
<td>0.00831451</td>
<td>Gas constant</td>
</tr>
<tr>
<td>T</td>
<td>311</td>
<td>Temperature in Kelvin</td>
</tr>
</tbody>
</table>

\[ \Delta G_{\text{ATP}} = -9 \] Standard Gibbs energy change for ATP formation at pH = 0

Redox potentials values in standard conditions\((pH = 0, T = 298 \text{ K})\)

<table>
<thead>
<tr>
<th>Redox Potential</th>
<th>Value</th>
<th>(pK_a) values of the acids</th>
</tr>
</thead>
<tbody>
<tr>
<td>EvOV</td>
<td>0.512</td>
<td>pK(_{HCO_3^-}) = 7.74</td>
</tr>
<tr>
<td>EvOL</td>
<td>0.418</td>
<td>pK(_{Lac}) = 3.86</td>
</tr>
<tr>
<td>EvOP</td>
<td>0.229</td>
<td>pK(_{Prop}) = 4.87</td>
</tr>
<tr>
<td>EvO2CO2ch</td>
<td>0.170</td>
<td>pK(_{Acet}) = 4.75</td>
</tr>
<tr>
<td>EvO2CO2ac</td>
<td>0.124</td>
<td>pK(_{But}) = 4.82</td>
</tr>
<tr>
<td>EvOH</td>
<td>0 H(^+)/H(_2)</td>
<td>pK(_{Val}) = 4.84</td>
</tr>
<tr>
<td>EvON</td>
<td>-0.113</td>
<td>NAD/NADH</td>
</tr>
<tr>
<td>EvOA</td>
<td>-0.228</td>
<td>Butyric acid</td>
</tr>
<tr>
<td>EvOG</td>
<td>-0.290</td>
<td>Glucose</td>
</tr>
</tbody>
</table>

2. COMPARTMENTS AND FLOWS

Cell wall compartment: CW
\[ \text{dCW/dt} = \text{INCW} - \text{CWGLU} - \text{OUTCW} \]

\[ \text{INCW} = \text{PULSE} (50/180,1,0,1440) \]

\[ \text{CWGLU} = \text{KCW} \times \text{CW} \]

\[ \text{OUTCW} = \text{KOUTCW} \times \text{CW} \]
**Starch compartment: ST**
\[
d\text{ST}/dt = \text{INST} - \text{STGLU} - \text{OUTST}
\]
\[
\text{INST} = \text{PULSE (25/180,1,0,1440)}
\]
\[
\text{STGLU} = K_\text{ST} \times \text{ST}
\]
\[
\text{OUTST} = K_{\text{OUTST}} \times \text{ST}
\]

**Glucose compartment: GLU**
\[
d\text{GLU}/dt = \text{CWGLU} + \text{STGLU} + \text{PYGL} - \text{GLPY} - \text{OUTGLU}
\]
\[
\text{OUTGLU} = 0.0001 \times \text{GLU}
\]

Glucose-pyruvate (1)
\[
\text{GLPY} = 2E10 \times \text{GLU} \times \text{NAD}^2 \times \text{ADP}^2 \times \text{EXP} (C \times \text{DEGLU})
\]
\[
\text{PYGL} = \left(\frac{2E10}{K_{\text{eq1}}} \right) \times \text{PYR}^2 \times \text{NADH}^2 \times \text{ATP}^2 \times \text{EXP} (-C \times \text{DEGLU})
\]

**Pyruvate compartment: PYR**
\[
d\text{PYR}/dt = 2\text{GLPY} - 2\text{PYGL} + \text{ACPY} - \text{PYAC} - \text{LAPY} - \text{PYLA} + 2\text{BUPY} - 2\text{PYBU} + \text{PROPY} - \text{PYPRO} + 2\text{VALPY} - 2\text{PYVAL} - \text{PYMIC} - \text{OUTPY}
\]
\[
\text{OUTPY} = K_{\text{A2}} \times \text{PYR}
\]

Pyruvate-lactate (2)
\[
\text{PYLA} = k \times \text{PYR} \times \text{NADH} \times \text{EXP} (C \times \text{DEPYR})
\]
\[
\text{LAPY} = \left(\frac{k}{K_{\text{eq2}}} \right) \times \text{LAC} \times \text{NAD} \times \text{EXP} (-C \times \text{DEPYR})
\]

Pyruvate-propionate (4) via succinate
\[
\text{PYPRO} = 100 \times k \times \text{PYR} \times \text{NAD} \times \text{ADP} \times \text{EXP} (C \times \text{DEPRO})
\]
\[
\text{PROPY} = \left(100 \times \frac{k}{K_{\text{eq4}}} \right) \times \text{PRO} \times \text{ATP} \times \text{NAD}^2 \times \text{EXP} (-C \times \text{DEPRO})
\]

Pyruvate-acetate (5)
\[
\text{PYAC} = 10000 \times k \times \text{PYR} \times \text{NAD} \times \text{ADP} \times \text{EXP} (C \times \text{DEACE})
\]
\[
\text{ACPY} = \left(10000 \times \frac{k}{K_{\text{eq5}}} \right) \times \text{ACE} \times \text{NAD} \times \text{CO}_2 \times \text{ATP} \times \text{EXP} (-C \times \text{DEACE})
\]

Pyruvate-butyrate (6)
\[
\text{PYBU} = 1000 \times k \times \text{PYR}^2 \times \text{ADP}
\]
\[
\text{BUPY} = \left(1000 \times \frac{k}{K_{\text{eq6}}} \right) \times \text{BUT} \times \text{ATP} \times \text{CO}_2^2
\]

Pyruvate-valerate (7)
\[
\text{PYVAL} = k \times \text{PYR}^2 \times \text{NADH}^3 \times \text{EXP} (C \times \text{DEVAL})
\]
\[
\text{VALPY} = \left(k \times \frac{k}{K_{\text{eq7}}} \right) \times \text{VAL} \times \text{CO}_2 \times \text{NAD}^3 \times \text{EXP} (-C \times \text{DEVAL})
\]

**Lactate compartment: LAC**
\[
d\text{LAC}/dt = -\text{LAPY} + \text{PYLA} - \text{LAPRO} + \text{PROLA} - \text{LACA} + \text{ALAC} - \text{OUTLAC}
\]
\[
\text{OUTLAC} = K_{\text{A2}} \times \text{LAC}
\]

Lactate-propionate (3)
\[
\text{LAPRO} = k \times \text{LAC} \times \text{NADH} \times \text{EXP} (C \times \text{DELAC})
\]
\[
\text{PROLA} = \left(k \times \frac{k}{K_{\text{eq3}}} \right) \times \text{PRO} \times \text{NAD} \times \text{EXP} (-C \times \text{DELAC})
\]

Lactic acid formation (12)
\[
\text{LACA} = k_a \times \text{LAC} \times \text{HPLUS}
\]
\[
\text{ALAC} = \left(k_a \times \frac{k_a}{K_{\text{eq12}}} \right) \times \text{ACL}
\]
Lactic acid compartment: ACL
\[ dACL/dt = -ALAC + LACA - OUTACL \]
\[ OUTACL = KA1 \times ACL \]

Propionate compartment: PRO
\[ dPRO/dt = LAPRO - PROLA + PYPRO - PROPY + APRO - PROA - OUTPRO \]
\[ OUTPRO = KA2 \times PRO \]

Propionic acid formation (13)
\[ PROA = ka \times PRO \times HPLUS \]
\[ APRO = (ka / Keq13) \times ACP \]

Propionic acid compartment: ACP
\[ dACP/dt = -APRO + PROA - OUTACP \]
\[ OUTACP = KA1 \times ACP \]

Acetate compartment: ACE
\[ dACE/dt = PYAC - ACPY + COACE - ACECO + AACE - ACEA - OUTACE \]
\[ OUTACE = KA2 \times ACE \]

Acetic acid formation (14)
\[ ACEA = ka \times ACE \times HPLUS \]
\[ AACE = (ka / Keq14) \times ACA \]

Acetic acid compartment: ACA
\[ dACA/dt = -AACE + ACEA - OUTACA \]
\[ OUTACA = KA1 \times ACA \]

Butyrate compartment: BUT
\[ dBUT/dt = PYBU - BUPY + ABUT - BUTA - OUTBUT \]
\[ OUTBUT = KA2 \times BUT \]

Butyric acid formation (15)
\[ BUTA = ka \times BUT \times HPLUS \]
\[ ABUT = (ka / Keq15) \times ACB \]

Butyric acid compartment: ACB
\[ dACB/dt = -ABUT + BUTA - OUTACB \]
\[ OUTACB = KA1 \times ACB \]

Valerate compartment: VAL
\[ dVAL/dt = PYVAL - VALPY + VALA - VALA - OUTVAL \]
\[ OUTVAL = KA2 \times VAL \]

Valeric acid formation (16)
\[ VALA = ka \times VAL \times HPLUS \]
\[ AVAL = (ka / Keq16) \times ACV \]

Valeric acid compartment: ACV
\[ dACV/dt = -AVAL + VALA - OUTACV \]
\[ OUTACV = KA1 \times ACV \]
Carbon dioxide compartment: CO2
\[
d\text{CO2}/dt = \text{PYAC} - \text{ACPY} + \text{CH4CO} - \text{COCH4} + 2\text{ACECO} - 2\text{COACE} + \text{HCO} - \\
\text{COHC} + 2\text{PYBU} - 2\text{BUPY} + \text{PYVAL} - \text{VALPY} - \text{OUTCO2}
\]
\[
\text{OUTCO2} = \text{CLIP (XCO2,0,PRES,1)}
\]
\[
CO2–METHAN (8)
\]
\[
\text{COCH4} = 100 \times k \times \text{CO2} \times \text{NADH}^4 \times \text{ADP} \times \text{EXP} (C \times \text{DECO2CH})
\]
\[
\text{CH4CO} = (100 \times k / \text{Keq8}) \times \text{CH4} \times \text{NAD}^4 \times \text{ATP} \times \text{EXP} (-C \times \text{DECO2AC})
\]
\[
CO2–ACETATE (9)
\]
\[
\text{COACE} = k \times \text{CO}^2 \times \text{NADH}^4 \times \text{ADP}^{0.25} \times \text{EXP} (C \times \text{DECO2AC})
\]
\[
\text{ACECO} = (k / \text{Keq9}) \times \text{ACE} \times \text{NAD}^4 \times \text{ATP}^{0.25} \times \text{EXP} (-C \times \text{DECO2AC})
\]
\[
Dissociation and relation CO2–HCO3 (11)
\]
\[
\text{HCCO} = k \times \text{HCO}^3 \times \text{HPLUS}
\]
\[
\text{COHC} = (k / \text{Keq11}) \times \text{CO2}
\]
Methane compartment: CH4
\[
d\text{CH4}/dt = \text{COCH4} - \text{CH4CO} - \text{OUTCH4}
\]
\[
\text{OUTCH4} = \text{CLIP (XCH4,0,PRES,1)}
\]
Hydrogen compartment: H2
\[
d\text{H2}/dt = \text{HPH2} - \text{H2HP} - \text{OUTH2}
\]
\[
\text{OUTH2} = \text{CLIP (0,0,PRES,1)}
\]
\[
H2–H+ (10)
\]
\[
\text{H2HP} = k \times \text{H2} \times \text{NAD} \times \text{EXP} (C \times \text{DEH})
\]
\[
\text{HPH2} = (k / \text{Keq10}) \times \text{HPLUS} \times \text{NADH} \times \text{EXP} (-C \times \text{DEH})
\]
Bicarbonate ions compartment: HCO3
\[
d\text{HCO3}/dt = \text{SALHCO3} + \text{COHC} - \text{HCO}
\]
\[
\text{SALHCO3} = 0.5 \times 10^{-4} \quad \text{Salivary flux of bicarbonate ions}
\]
H⁺ Compartment: HPLUS
\[
d\text{HPLUS}/dt = \text{H2HP} - \text{HPH2} + \text{AACE} - \text{ACEA} + \text{APRO} - \text{PROA} + \text{ALAC} - \text{LACA} + \\
\text{ABUT} - \text{BUTA} + \text{COHC} - \text{HCO} + \text{VALPY} - \text{VALA}
\]

NAD/NAD
\[
d\text{NAD}/dt = \text{NNH} - \text{NHN}
\]
\[
d\text{NAD}/dt = \text{NHN} - \text{NNH}
\]
\[
\text{NNH} = 2\text{GLPY} + \text{LAPY} + 2\text{PROPY} + \text{PYAC} + \text{PROLA} + \text{H2HP} + 4\text{CH4CO} + \\
4\text{ACECO} + 3\text{VALPY}
\]
\[
\text{NNH} = 2\text{PYGL} + \text{PYLA} + 2\text{PYPRO} + \text{ACPY} + \text{LAPRO} + \text{HPH2} + 4\text{COCH4} + \\
4\text{COACE} + 3\text{PYVAL} + \text{NADHMM} + \text{BESNADHM}
\]
ATP/ADP
\[
d\text{ATP}/dt = \text{ADTP} - \text{ATDP}
\]
\[
\text{ADTP} = 2\text{GLPY} + \text{PYAC} + \text{COCH4} + 0.25\text{COACE} + \text{PYBU} + \text{PYPRO}
\]
\[
\text{ATDP} = 2\text{PYGL} + \text{ACPY} + \text{CH4CO} + 0.25\text{ACECO} + \text{BUPY} + \text{PROPY} + \text{BESATPM} + \\
\text{ATPMM})/(1 + (0.5 \times 10^{-3}/\text{ATP}))
\]
\[
d\text{ADP}/dt = \text{ATDP} - \text{ADTP}
Microbial compartment

dMDM/dt = CRMDM – LYSMDM – OUTMDM
OUTMDM = KM × MDM
CRMDM = MUBAS × MDM × EXP (KMU × (RATDP – 1))
LYSMDM = MUBAS × MDM × EXP (–KMU × (RATDP – 1))
BCRMDM = CRMDM – LYSMDM Apparent Growth
SCRMDM = RAMP(BCRMDM,0)
MUAMDM = 60 × BCRMMD/MDM Apparent growth rate in h⁻¹
MURMDM = 60 × CRMDM/MDM Real growth rate in h⁻¹
UTCM = CRMDM/25 C mol used for growth

ATP utilizing flow by microorganisms
ATPMM = 0.001 × 1.6 × MDM/60 Maintenance
BESATPM = CRMDM/30 Growth

C outflow for microorganism growth
PYMIC = BCRMMD/25 C pyruvate mol used

NADH utilizing flow by microorganisms
NADHMM = 1 × 10⁻⁵ × MDM Maintenance
BESNADHM = 0.5 × UTCM Growth

3. AUXILIARY VARIABLES

Ratio NADH/NAD, ATP/ADP
RNHN = NADH/NAD
RATDP = ATP/ADP

PH Calculation

PH = –Log(HPLUS) pH calculation in a mechanistic way
PHVFA = 6.96 – 6.94 × SOMVFA pH calculation in an empirical way using [VFA]
PHCO3 = pKhco + Log(HCO3/CO2) Calculation for each acid using pKa
PHACE = pKace + Log(ACE/ACA)
PHPRO = pKpro + Log(PRO/ACP)
PHBUT = pKbut + Log(BUT/ACB)
PHLAC = pKlac + Log(LAC/ACL)
PHV AL = pKval + Log(V AL/ACV)
Thermodynamic modeling of ruminal fermentations

**Gas pressures**

\[ \text{PRES} = \text{CH}_4 + \text{CO}_2 + \text{H}_2 \]  
Total pressure in the rumen

\[ \text{XCH}_4 = \frac{\text{CH}_4}{\text{PRES}} \]  
Partial pressures

\[ \text{XCO}_2 = \frac{\text{CO}_2}{\text{PRES}} \]

\[ \text{XH}_2 = \frac{\text{H}_2}{\text{PRES}} \]

**VFA molar proportions**

\[ \text{SOMVFA} = \text{ACE} + \text{ACA} + \text{PRO} + \text{ACP} + \text{Total sum of VFA} \]

\[ \text{BUT} + \text{ACB} + \text{VAL} + \text{ACV} \]

\[ \text{PACE} = 100 \times (\text{ACA} + \text{ACE})/\text{SOMVFA} \]  
Respective proportions of each VFA

\[ \text{PPRO} = 100 \times (\text{ACP} + \text{PRO})/\text{SOMVFA} \]

\[ \text{PBUT} = 100 \times (\text{ACB} + \text{BUT})/\text{SOMVFA} \]

\[ \text{PVAL} = 100 \times (\text{ACV} + \text{VAL})/\text{SOMVFA} \]

\[ \text{ACEsPRO} = (\text{ACE} + \text{ACA})/(\text{PRO} + \text{ACP}) \]  
Acetate to propionate ratio

**4. CALCULATION OF THERMODYNAMICAL VARIABLES**

**Standard potentials of each redox couple at rumen pH**

\[ \text{Ev0pHN} = \text{EvON} - (\text{RT}/(2F)) \times \text{Ln}(10) \times \text{PH} \]

\[ \text{Ev0pHG} = \text{EvOG} - (6\text{RT}/(8F)) \times \text{Ln}(10) \times \text{PH} \]

\[ \text{Ev0pHP} = \text{EvOP} - (\text{RT}/F) \times \text{Ln}(10) \times \text{PH} \]

\[ \text{Ev0pHA} = \text{EvOA} - (\text{RT}/F) \times \text{Ln}(10) \times \text{PH} \]

\[ \text{Ev0pHL} = \text{EvOL} - (\text{RT}/F) \times \text{Ln}(10) \times \text{PH} \]

\[ \text{Ev0pHCac} = \text{EvOCO}_2\text{ac} - (7\text{RT}/(8F)) \times \text{Ln}(10) \times \text{PH} \]

\[ \text{Ev0pHCch} = \text{EvOCO}_2\text{ch} - (\text{RT}/F) \times \text{Ln}(10) \times \text{PH} \]

\[ \text{Ev0pHV} = \text{EvOV} - (7\text{RT}/(6F)) \times \text{Ln}(10) \times \text{PH} \]

**Standard potentials of the reactions at rumen pH**

\[ \text{Ev0pHr1} = \text{Ev0pHN} - \text{Ev0pHG} \]

\[ \text{Ev0pHr2} = \text{Ev0pHP} - \text{Ev0pHN} \]

\[ \text{Ev0pHr3} = \text{Ev0pHL} - \text{Ev0pHN} \]

\[ \text{Ev0pHr4} = \text{Ev0pHCac} - \text{Ev0pHN} \]

\[ \text{Ev0pHr5} = \text{Ev0pHCch} - \text{Ev0pHN} \]

\[ \text{Ev0pHr6} = -177.7 + \text{RT} \times \text{Ln}(10) \times \text{PH} \]

\[ \text{Ev0pHr7} = -6\text{F} \times \text{Ev0pHr7} \]

\[ \text{Ev0pHr8} = -8\text{F} \times \text{Ev0pHr8} \]

\[ \text{Ev0pHr9} = -8\text{F} \times \text{Ev0pHr9} \]

\[ \text{Ev0pHr10} = -2\text{F} \times \text{Ev0pHr10} \]

**Gibbs energies of the reactions (without ATP couplings) in standard conditions**

\[ \Delta G_{0\text{ATPpH}} = \Delta G_{0\text{ATP}} + \text{RT} \times \text{Ln}(10) \times \text{PH} \]
Gibbs energies of the reactions in rumen conditions

Gr1 = G0pHr1 + 2∆G0ATPpH + RT × Ln(((PYR2) × (NADH2) × (ATP2)) / (GLU × (NAD2) × (ADP2)))
Gr2 = G0pHr2 + RT × Ln((LAC × NAD) / (PYR × NADH))
Gr3 = G0pHr3 + RT × Ln((PRO × NAD) / (LAC × NADH))
Gr4 = G0pHr4 + ∆G0ATPpH + RT × Ln((PRO × ATP × (NAD2)) / (PYR × ADP × (NADH2))
Gr5 = G0pHr5 + ∆G0ATPpH + RT × Ln((ACE × NADH × CO2 × ATP) / (PYR × NAD × ADP))
Gr6 = G0pHr6 + ∆G0ATPpH + RT × Ln((BUT × ATP × (CO22)) / (PYR × ADP))
Gr7 = G0pHr7 + RT × Ln((VAL × CO2 × (NAD2)) / (PYR2) × (NADH2))
Gr8 = G0pHr8 + ∆G0ATPpH + RT × Ln((CH4 × (NAD4) × ATP) / (CO2 × (NADH4) × ADP))
Gr9 = G0pHr9 + 0.25∆G0ATPpH + RT × Ln((ACE × (NAD4) × (ATP0.25)) / (CO22 × (NADH4) × (ADP0.25)))
Gr10 = G0pHr10 + RT × Ln((H2 × NAD) / NADH)
Gr11 = RT × Ln((CO2/HCO3)/Keq11)
Gr12 = RT × Ln((ACL/LAC)/Keq12)
Gr13 = RT × Ln((ACP/PRO)/Keq13)
Gr14 = RT × Ln((ACA/ACE)/Keq14)
Gr15 = RT × Ln((ACB/BUT)/Keq15)
Gr16 = RT × Ln((ACV/VAL)/Keq16)

Equilibrium constants of the reactions

Keq1 = EXP (–(G0pHr1 + 2∆G0ATPpH) / RT)
Keq2 = EXP (Ev0pHr2 / (RT/2F))
Keq3 = EXP (Ev0pHr3 / (RT/2F))
Keq4 = EXP (–(G0pHr4 + ∆G0ATPpH) / RT)
Keq5 = EXP (–(G0pHr5 + ∆G0ATPpH) / RT)
Keq6 = EXP (–(G0pHr6 + ∆G0ATPpH) / RT)
Keq7 = EXP (–G0pHr7 / RT)
Keq8 = EXP (–(G0pHr8 + ∆G0ATPpH) / RT)
Keq9 = EXP (–(G0pHr9 + 0.25∆G0ATPpH) / RT)
Keq10 = EXP ((EvOH – EvON) / (RT/2F))
Keq11 = 1/10−pKhco
Keq12 = 1/10−pKlac
Keq13 = 1/10−pKpro
Keq14 = 1/10−pKace
Keq15 = 1/10−pKbut
Keq16 = 1/10−pKval
**Potential values of each couple**

\[
\text{ENAD} = \text{Ev} \text{pHN} + \left(\frac{RT}{2F}\right) \times \ln(\text{NAD} \div \text{NADH}) \\
\text{EGLU} = \text{Ev} \text{pHG} + \left(\frac{RT}{4F}\right) \times \ln(\text{PYR}^2 \div (\text{GLU})) \\
\text{Epyr} = \text{Ev} \text{pHP} + \left(\frac{RT}{2F}\right) \times \ln(\text{PYR} \div \text{LAC}) \\
\text{ELAC} = \text{Ev} \text{pHL} + \left(\frac{RT}{2F}\right) \times \ln(\text{LAC} \div \text{PRO}) \\
\text{Epro} = \text{Ev} \text{pHPP} + \left(\frac{RT}{4F}\right) \times \ln(\text{PYR} \div \text{PRO}) \\
\text{EACE} = \text{Ev} \text{pHA} + \left(\frac{RT}{2F}\right) \times \ln\left(\frac{\text{ACE} \times \text{CO}_2}{\text{PYR}}\right) \\
\text{EH} = \text{Ev} \text{pHH} + \left(\frac{RT}{2F}\right) \times \ln(\text{H}_2) \\
\text{ECO2ch} = \text{Ev} \text{pHCh} + \left(\frac{RT}{8F}\right) \times \ln(\text{CO}_2 \div \text{CH}_4) \\
\text{ECO2ac} = \text{Ev} \text{pHAc} + \left(\frac{RT}{8F}\right) \times \ln(\text{CO}_2^2 \div \text{ACE}) \\
\text{EV AL} = \text{Ev} \text{pHV} + \left(\frac{RT}{6F}\right) \times \ln(\text{PYR}^2 \div (\text{VAL} \times \text{CO}_2)) \\
\]

**Determination of the average redox potential in the rumen**

\[
\text{SOMOL} = \text{GLU} + \text{PYR} + \text{LAC} + \text{PRO} + \text{ACE} + \text{VAL} + \text{NAD} + \text{NADH} + \text{H}_2\text{M} + \text{HPLUS} + \text{CO}_2\text{M} + \text{CH}_4\text{M} \\
\text{H}_2\text{M} = 0.0008143 \times \text{H}_2 \text{ Gas concentrations determined thanks to the Henry constant} \\
\text{CO}_2\text{M} = 0.0229 \times \text{CO}_2 \\
\text{CH}_4\text{M} = 2 \times 10^{-5} \times \text{CH}_4 \\
\text{PROPNAD} = (\text{NAD} + \text{NADH}) \div \text{SOMOL} \text{ Calculation of the weighting affected to each redox couple} \\
\text{PROPL} = \frac{(\text{GLU} + (2/7)\text{PYR})}{\text{SOMOL}} \\
\text{PROPL} = \frac{((1/7)\text{PYR} + (1/2)\text{LAC})}{\text{SOMOL}} \\
\text{PROPL} = \frac{(1/2)\text{LAC} + (1/2)\text{PRO}}{\text{SOMOL}} \\
\text{PROPL} = \frac{(1/2)\text{PRO} + (1/7)\text{PYR}}{\text{SOMOL}} \\
\text{PROPL} = \frac{(1/7)\text{PYR} + (1/2)\text{ACE} + (1/5)\text{CO}_2\text{M}}{\text{SOMOL}} \\
\text{PROPL} = \frac{(1/7)\text{PYR} + (1/2)\text{ACE} + (1/5)\text{CO}_2\text{M}}{\text{SOMOL}} \\
\text{PROPL} = \frac{(1/2)\text{CO}_2\text{M} + (1/5)\text{CO}_2\text{M}}{\text{SOMOL}} \\
\text{PROPL} = \frac{(2/5)\text{CO}_2\text{M} + (1/2)\text{ACE}}{\text{SOMOL}} \\
\text{PROPL} = \frac{(1/2)\text{CO}_2\text{M} + (1/2)\text{ACE} + (1/5)\text{CO}_2\text{M}}{\text{SOMOL}} \\
\text{PROPL} = \frac{(2/7)\text{PYR} + \text{VAL} + (1/5)\text{CO}_2\text{M}}{\text{SOMOL}} \\
\text{EPMOY} = (\text{ENAD} \times \text{PROPNAD}) + (\text{EGLU} \times \text{PROPL}) + (\text{Epyr} \times \text{PROPL}) + (\text{ELAC} \times \text{PROPL}) + (\text{Epro} \times \text{PROPL}) + (\text{EACE} \times \text{PROPL}) + (\text{EH} \times \text{PROPL}) + (\text{ECO2ch} \times \text{PROPCOCH4}) + (\text{ECO2Ac} \times \text{PROPCOAC}) + (\text{EV AL} \times \text{PROPL}) \\
\]

**Difference to the average potential**

\[
\text{DEGLU} = \text{EPMOY} - \text{EGLU} \\
\text{DEPYR} = \text{Epyr} - \text{EPMOY} \\
\text{DELAC} = \text{ELAC} - \text{EPMOY} \\
\text{DEPRO} = \text{Epro} - \text{EPMOY} \\
\text{DEACE} = \text{EACE} - \text{EPMOY} \\
\text{DEH} = \text{EH} - \text{EPMOY} \\
\text{DECO2CH} = \text{ECO2CH} - \text{EPMOY} \\
\text{DECO2AC} = \text{ECO2Ac} - \text{EPMOY} \\
\text{DEV AL} = \text{EV AL} - \text{EPMOY} \\
\]

**Potentials of oxydoreduction reactions**

\[
\text{Evr1} = \text{ENAD} - \text{EGLU} \\
\text{Evr2} = \text{EPYR} - \text{ENAD} \\
\text{Evr3} = \text{ELAC} - \text{ENAD} \\
\text{Evr4} = \text{Epro} - \text{ENAD} \\
\text{Evr5} = \text{ENAD} - \text{EACE} \\
\text{Evr7} = \text{EVAL} - \text{ENAD} \\
\text{Evr8} = \text{ECO2ch} - \text{ENAD} \\
\text{Evr9} = \text{ECO2Ac} - \text{ENAD} \\
\text{Evr10} = \text{EH} - \text{ENAD} \\
\]