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Stoichiometric model of a fully closed bioregenerative life support system for autonomous long-duration space missions

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Bioregenerative life support systems (BLSS) are vital for long-duration and remote space missions to increase mission sustainability. These systems break down human waste materials into nutrients and CO₂ for plants and other edible organisms, which in turn provide food, fresh water, and oxygen for astronauts. The central idea is to create a materially closed loop, which can significantly reduce mission mass and volume by cutting down or even eliminating disposable waste. In most BLSS studies only a fraction of the resources, such as food, are provided by the system itself, with the rest taken on board at departure or provided through resupply missions. However, for autonomous long-duration space missions without any possibility of resupply, a BLSS that generates all resources with minimal or no material loss, is essential. The goal of this study is to develop a stoichiometric model of a conceptually fully closed BLSS that provides all the metabolic needs of the crew and organisms. The MELiSSA concept of the European Space Agency is used as reference system, consisting of five interconnected compartments, each inhabited by different types of organisms. A detailed review of publicly available MELiSSA literature from 1989 to 2022 revealed that no existing stoichiometric model met the study's requirements. Therefore, a new stoichiometric model was developed to describe the cycling of the elements C, H, O, and N through all five MELiSSA compartments and one auxiliary compartment. A compact set of chemical equations with fixed coefficients was established for this purpose. A spreadsheet model simulates the flow of all relevant compounds for a crew of six. By balancing the dimensions of the different compartments, a high degree of closure is attained at steady state, with 12 out of 14 compounds exhibiting zero loss, and oxygen and CO_2 displaying only minor losses between iterations. This is the first stoichiometric model of a MELiSSA-inspired BLSS that describes a continuous provision of 100% of the food and oxygen needs of the crew. The stoichiometry serves as the foundation of an agent-based model of the MELiSSA loop, as part of the Evolving Asteroid Starships (E|A|S) research project.

KEYWORDS

space exploration, human spaceflight, bioregenerative life support, waste processing, food production, ecosystem modeling, simulation, MELiSSA

1 Introduction

Bioregenerative life support systems (BLSS) will be a key component of future long-duration space exploration, as they can reduce mission mass and volume (Imhof et al, 2017; Audas et al, 2022). A BLSS recycles human waste by feeding it to an artificial ecosystem consisting of a range of different organisms, such as higher plants, microalgae, and bacteria. These organisms gradually break down the waste and in the process provide fresh food, generate oxygen, and clean water for the crew (Nelson et al, 2010; Escobar and Nabity, 2017; Pannico et al, 2022). As a result, there is less reliance on constant resupply from Earth, which greatly reduces the material footprint needed for extended missions in space (Macelroy and Averner, 1978; Häder, 2020; Audas et al, 2022). Because of its constant fresh food provision, a BLSS also solves the problem that essential nutrients in processed and prepackaged food are deficient and degrade over time (Cooper et al, 2017; Carillo et al, 2020). Biological systems are also capable of self-repair, unlike mechanical systems that need separate systems to be repaired. Surviving organisms and tissues in a damaged BLSS can potentially recover to their original size on their own, even after significant damage (Bartsev et al, 1996).

MELiSSA is a BLSS concept developed by ESA with a consortium of 15 international partners and the involvement of approximately 50 organizations (MELiSSA Foundation, 2023). It is an artificial ecosystem consisting of five interconnected compartments inhabited by higher plants, microalgae, microorganisms, and humans (abbreviated as C1 to C5, Figure 1; Table 1). Each of the compartments has its own specific metabolic function within the entire MELiSSA loop. First, all human waste is broken down through a sequence of three bioreactor types: a thermophilic anaerobic compartment (C1), photoheterotrophic compartment (C2) and nitrifying compartment (C3). This results in nutrients and CO₂ for microalgae and plants (C4a and C4b) that, in turn, provide food, oxygen, and fresh water for the crew (C5), thus closing the loop (Hendrickx and Mergeay, 2007; Clauwaert et al, 2017; Vermeulen et al, 2020a). The operational pilot plant at Universitat Autònoma de Barcelona consists of a number of connected MELiSSA compartments and demonstrates a part of the system's entire metabolic loop (Gòdia et al, 2004; Garcia-Gragera et al, 2021). Mathematical models of the MELiSSA loop have been developed and improved since the beginning of the MELiSSA project in 1989 (MELiSSA, 1989; Dussap and Gros, 1991). Some of these models are used for predictive control, to be run alongside the actual physical system. The MELiSSA models help in understanding the dynamics of mass flow fluctuations and achieving long-term reliability of the system (Cornet et al, 2001; Ciurans Molist et al, 2020; Poulet et al, 2020). Such ecosystem models can also be used to simulate the impact of unforeseen perturbations, deliberate interventions, or design changes in the system (Volk and Rummel, 1987; Pilo Teniente, 2015; Vermeulen et al, 2019).

To describe the material flows in an ecosystem model, the stoichiometric relations governing these flows need to be developed (Volk and Rummel, 1987; Begon and Townsend, 2021). Several authors have described mass flow models of biological life support systems (Volk and Rummel, 1987; Garland, 1989; Loader et al, 1997; Finn, 1998), but not all studies contain the underlying

chemical stoichiometric equations. The stoichiometric equations describing mass flows in the MELiSSA BLSS have been published in publicly-available MELiSSA literature, both in peer-reviewed papers and research reports disseminated by the MELiSSA Foundation (Table 1). In less than half of the listed studies, all five MELiSSA compartments were modeled. The other studies describe either a smaller number of compartments, or just one. The loop is never fully closed in any of the listed studies. There are a number of reasons for this. First of all, it may indicate that certain compounds are not part of the regenerative logic of the system and are not generated within the loop, and therefore need to be fully supplied from the outside (Dussap et al, 1993; Poughon et al, 2009). Secondly, it may indicate an output that does not get regenerated, such as a residual indigestible material (Fulget et al, 1999; Poughon et al, 2000). Because of the resulting accumulation of that material, external supply is required to keep the system running. And thirdly, it may be a deliberate system design choice that only a fraction of a particular compound or resource is generated by the BLSS. Several MELiSSA studies, for example, explicitly specify that only a part of the needed food is generated by the loop (Poughon et al, 2000; Guirado and Podhajsky, 2008; Thiron, 2020). The rest is supplied externally. One of the main motivations for such a choice is the limited space that is available in common spacecraft and habitat designs to integrate bioreactors and plant growth chambers (Johnson et al, 2021; Gorce et al, 2022).

Almost three-quarters of the researched MELiSSA studies list all the stoichiometric equations of their models. The other studies limit themselves to a few selected equations as examples. The stoichiometric coefficients in the chemical equations can either be fixed beforehand, or dynamically calculated on the fly, during the simulation run. For example, in simulating the production of *Limnospira indica* in C4a, the elementary composition of its biomass is determined by the variable light irradiation inside the photobioreactor (Guirado and Podhajsky, 2008; Poughon et al, 2009). This then necessitates the recalculation of all subsequent stoichiometric equations that contain this dynamic empirical biomass formula.

Whenever creating a stoichiometry to describe mass flows in an ecosystem, choices need to be made about which compounds will be considered, and which level of detail will be used to describe the ecosystem's biochemical processes (Dubitzky et al, 2011; Begon and Townsend, 2021). This choice is influenced by the level of knowledge about the individual metabolism of the different species in the ecosystem, but ultimately depends on the objective and scope of the study. In light of this, the reviewed MELiSSA studies contain varying assumptions concerning the role of different compounds such as lipids, VFAs, and CO₂. This is partially due to the fact that over time the MELiSSA models became more detailed, incorporating more processes. Early MELiSSA papers assume that all lipids consumed by the astronauts are oxidized and no traces of lipids are found in the feces (Dussap and Gros, 1991; Dussap et al, 1993). This is later adjusted, and lipids become an explicit part of the feces input in C1 (Poughon et al, 2000). The mix of VFAs that is generated as output of C1 varies quite significantly between the different studies. In the earliest studies only acetic acid and butyric acid are considered (Dussap and Gros, 1991; Dussap et al, 1993), but in later studies, a broader range of VFAs is included in the stoichiometry (Poughon, 2007b).

chemical equations that are factually listed in the publication. "Stoichiometrically balanced" indicates that the publication includes the stoichiometric coefficients for all fixed equations. Stoichiometric coefficients are described as dynamic when the publication fluctuate throughout a simulation run. The macromolecule composition is described as dynamic when the proportion of CHON TABLE 1 Systematic review of the stoichiometric descriptions of mass flows in the various compartments of the MELISSA loop, as documented in publicly available literature. Only modeling studies containing at least one chemical equation are listed. "Full closure" indicates that all material outputs are utilized as inputs without the need for any external supplies. "Chemical equations" indicates the number of involved

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elements in at least one che	emical formula varies ov	er time. C, compartment; N/A, not a	pplicable.			
MELiSSA compartments	Full closure	Chemical equations	Stoichiometrically balanced	Stoichiometric coefficients	Macromolecule composition	References
C1-C5 ^a	No	Partly	Yes	Fixed	Fixed	MELiSSA (1989)
C1-C5 ^a	No	Complete	Yes	Fixed	Fixed	Dussap and Gros (1991)
C1-C5 ^a	No	Complete	Yes	Fixed	Fixed	Dussap et al (1993)
C1-C5 ^a	No	Complete	Yes	Fixed + dynamic	Fixed + dynamic	Poughon et al (1994)
C1-C5 ^a	No	Complete	Yes	Fixed	Fixed	Fulget (1996)
C1-C5 ^a	No	Examples	Yes	Fixed + dynamic	Fixed + dynamic	Fulget et al (1999)
C1-C5	No	Complete	Incomplete	Fixed + dynamic	Fixed + dynamic	Poughon et al (2000)
C1-C5	No	Complete	Yes	Fixed + dynamic	Fixed + dynamic	Poughon (2007b)
C1-C5	No	Complete	Yes	Fixed + dynamic	Fixed + dynamic	Guirado and Podhajsky (2008)
C1-C5	No	Examples	Yes	Fixed + dynamic	Fixed + dynamic	Poughon et al (2009)
C1-C5	No	Complete	Yes	Fixed + dynamic	Fixed + dynamic	Thiron (2020)
C1-C4a	No info	Complete	No	None	Fixed	Hendrickx et al (2006)
C2+C5	No	Complete	Yes	Fixed	Fixed	Poughon (1994b)
C4a-C5	No	Partly	Yes	Fixed + dynamic	Fixed + dynamic	Poughon (2007a)
C4a-C5	No	Partly	Yes	Fixed	Fixed	Pilo Teniente (2015)
C4a-C5	No	Partly	Yes	Fixed	Fixed	Alemany et al (2019)
C2	N/A	Complete	Yes	Fixed	Fixed	Cornet and Dussap (2000)
C3	N/A	Complete	Yes	Fixed	Fixed	Cruvellier et al (2016)
C4a	N/A	Complete	Yes	Fixed	Fixed	Poughon (1994a)
C4a	N/A	Complete	Yes	Fixed	Fixed	Cornet et al (1998)
C4a	N/A	Complete	Yes	Fixed	Fixed	Cornet et al (2003)
C4a	N/A	Complete	Yes	Dynamic	Dynamic	Poughon et al (2020)
C4b	N/A	Complete	No	None	No info	Maclean et al (2010)
C4b	N/A	Complete	Yes	Fixed	Fixed	Poulet et al (2020)
^a Only C4a (<i>Linnoshira c</i> omnarth	nent), no C4h (higher plant	compartment)				

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The stoichiometric model presented in this paper provides the basis for an agent-based model (ABM) of the MELiSSA loop to study the interactions and emergent behavior of the MELiSSA system from a theoretical perspective. MELiSSA is fundamentally a complex system, and properties such as nonlinearity, stochasticity and chaos may play an important role in the overall behavior of the system, just like in any ecosystem (Bjørnstad, 2015). Because all functions in the MELiSSA loop are tightly coupled and since there are no large buffers in space (e.g., no large atmosphere or water bodies), slight variations in any of the loop's pathways can have dramatic consequences for the entire loop (Macelroy and Averner, 1978; Poulet et al, 2018). It is precisely such tightly coupled systems that are potentially susceptible to irregular fluctuations (Bjørnstad, 2015). ABMs offer a modeling approach that helps to gain insight in such dynamics and helps to explore different design strategies for improved system robustness. ABMs are used to understand those aspects of the behavior of a system, that are not easily represented in population-level differential equations, such as variation among individuals or variation of individuals during their life cycle (Deangelis and Grimm, 2014). Such exploratory models of complex systems lead to insights about essential mechanisms and

principles, a use of modeling that is different from modeling aimed at making accurate predictions (Forrest and Mitchell, 2016). MELiSSA modeling studies to date focus on predictive modeling because this is a critical step to develop a durable and reliable BLSS. The MELiSSA modelling approach is mechanistic, allowing for better understanding and improved predictions. With its exploratory approach, this study can be considered as being complementary to the existing MELiSSA modeling efforts. In order to build an ABM of MELiSSA, a stoichiometric model is needed that describes essential mass flows between all involved agents (bacteria, microalgae, higher plants and humans). The key metabolic processes in all MELiSSA compartments need to be described with enough detail to enable studying the interactions. At the same time, they need to remain abstracted enough to tlead to an efficient model and retain relevance for the objectives of the ABM which are centered around investigating the impact of heterogeneity in the system and capturing nonlinear dynamics.

The research objective of this paper is to develop a stoichiometric model of a hypothetical BLSS that provides a crew with fresh food and oxygen during long-duration space missions, without the need for any resupply. The envisioned stoichiometric model should conform to the following requirements:

- The system provides 100% of all the food and oxygen needed by the crew, and 100% of the resources needed by every other organism in the loop.
- The stoichiometric model is fully closed. Every single output is used as input, and no additional external supplies are needed during steady state.
- No kinetic limitations of the stoichiometric reactions are used. Macromolecules do not have a dynamic composition (e.g., biomass composition does not depend on light), and all stoichiometric coefficients are fixed. Such a static approach is sufficient for the current goals of the ABM of MELiSSA that is developed on top of this stoichiometry.

The current literature (listed in Table 1) has, to the authors' knowledge, not included such a stoichiometric model of the MELiSSA loop. As described above, the loop is always considered to be partially open, primarily for practical reasons. The research reported in this paper is more theoretical in nature, focusing on the internal dynamics of hypothetically fully closed systems.

2 Materials and methods

This section describes the empirical formulas, stoichiometric equations, and the setup of a static spreadsheet model.

2.1 Empirical formulas

Table 2 provides an overview of the empirical formulas of the large biomolecules. More detailed descriptions of the origin of each formula are then discussed. The empirical formulas of biomass and feces are provided for comparative purposes only, since both biomass and feces are represented as proportions of carbohydrates, proteins and lipids.

2.1.1 Carbohydrates, proteins, and lipids

The formula for carbohydrates is the general formula for polysaccharides, used in many of the MELiSSA modeling studies

(e.g., in (Dussap et al, 1993) and (Thiron, 2020)). There are, however, a range of different general formulas used for proteins in the MELiSSA literature (Fulget, 1996; Duatis et al, 2008; Thiron, 2020). Our approach was to first calculate the average protein composition of the different organisms generating biomass in our model, and then compare this result to general protein formulas published in multiple BLSS studies. In our stoichiometric model, the higher plants are represented by a generalized "ideal plant", in line with previous MELiSSA studies (Poughon, 2007b; Guirado and Podhajsky, 2008). It is conceived as an average of the nutritional properties of bread wheat, durum wheat, potato and soybean. The CHON composition of the protein of this ideal plant is then averaged with that of Limnospira and Rhodobacter, all using values listed in MELiSSA literature (Dussap et al, 1993; Duatis et al, 2008). This results in $CH_{1.5799}O_{0.3134}N_{0.2558}$. This composition comes very close to $\rm CH_{1.5900}O_{0.3100}N_{0.2500}$ used as general protein formula in studies of the Lunar Palace BLSS (Hu et al, 2010; Fu et al, 2016), and hence this latter formula was selected. Palmitic acid is often used as lipid in stoichiometries of ecosystems (Volk and Rummel, 1987; Guirado and Podhajsky, 2008). However, vegetable oils mainly consist of more complex triacylglycerols (Zambelli et al, 2015). Therefore in this stoichiometric model, lipids are represented by tripalmitin, consisting of one glycerol molecule bonded to three palmitic acid molecules. Just like with palmitic acid, the fermentation of glycerol also leads to VFAs, further contributing to the overall VFA output of C1.

2.1.2 Biomass

Throughout the MELiSSA literature the empirical formulas of the biomass of specific organisms differ to varying degrees. The empirical formula that describes the biomass of the purple sulphur bacteria *Rhodobacter* varies, for example, between (Dussap et al, 1993), (Hendrickx et al, 2006), and (Thiron, 2020). This is to be expected since empirical formulas are always approximations. Moreover, in several MELiSSA modeling studies, biomass composition is dependent on environmental parameters such as light input, and consequently the biomass composition varies throughout the course of a simulation (Table 1). In the current stoichiometric model, biomass is consistently represented as a combination of carbohydrates, proteins, and lipids, and not as one single molecule. This makes it possible to trace the levels of each individual compound throughout the loop. To establish the

TABLE 2 Empirical formulas of all large biomolecules used in the stoichiometric model.

Compound	Empirical formula	References and notes
Carbohydrates	CH _{1.6667} O _{0.8333}	General polysaccharides
Proteins	$\rm CH_{1.5900}O_{0.3100}N_{0.2500}$	Hu et al, 2010, Fu et al, 2016
Lipids	CH _{1,9216} O _{0,1177}	Tripalmitin
Biomass Rhodospirillum rubrum	CH _{1.6472} O _{0.3598} N _{0.1788}	18% carbohydrates, 72% proteins, 10% lipids, literature
Biomass Limnospira sp.	$\mathrm{CH}_{1.6472}\mathrm{O}_{0.3598}\mathrm{N}_{0.1788}$	18% carbohydrates, 72% proteins, 10% lipids, literature
Biomass higher plants (edible)	CH _{1.6889} O _{0.6090} N _{0.05487}	70% carbohydrates, 20% proteins, 10% lipids, literature
Biomass higher plants (non-edible)	CH _{1.6667} O _{0.8333}	100% carbohydrates, 0% proteins, 0% lipids, literature
Feces	CH _{1.6676} O _{0.6028} N _{0.07715}	66% carbohydrates, 28% proteins, 6% lipids, calculated

composition of the different types of biomass, in this study an estimate was made based on averaged values from literature. As validation, the corresponding empirical formula was then compared with formulas published in MELiSSA literature.

In correspondence with the MELiSSA study of (Hendrickx et al, 2006), the same biomass composition was chosen for both *Rhodospirillum rubrum* and *Limnospira sp.* Based on literature (e.g., Dussap et al, 1993; Kobayashi and Kobayashi, 1995; Teuling et al, 2017), the following biomass composition was established (dry weight, or DW): 18% carbohydrates, 72% proteins, and 10% lipids. Both organisms contain high amounts of protein, hence their interesting potential as food sources (Vrati, 1984; Clauwaert et al, 2017; Muys et al, 2019). Using the previously established CHON compositions of carbohydrates, proteins and lipids, the overall biomass composition corresponds to the empirical formula $CH_{1.6472}O_{0.3598}N_{0.1788}$. This is in the same range as the biomass formulas used in the MELiSSA modeling studies of (Hendrickx et al, 2006) and (Thiron, 2020).

Carbohydrates are typically the most abundant compound in plants, encompassing both structural (e.g., cellulose) and nonstructural (e.g., starch and glucose) carbohydrates (Moore and Hatfield, 1994). As explained before, an 'ideal plant' is used in the current model. Four energy- and/or protein-rich crops were selected and combined based on (Paradiso et al, 2013): bread and durum wheat, potato and soybean. Averaging the carbohydrate, protein and lipid fractions of these four crops (Molders et al, 2012; Stasiak et al, 2012; Page and Feller, 2013; Paradiso et al, 2013), leads to the following edible plant biomass composition (DW): 70% carbohydrates, 20% proteins, and 10% lipids. Taking into account the above CHON compositions of carbohydrates, proteins and lipids, this corresponds to the empirical macroformula $CH_{1.6889}O_{0.6090}N_{0.05487}$. This is in line with the "food" descriptions in other MELiSSA studies (Fulget, 1996; Thiron, 2020). The inedible part of the plant is considered to be 100% carbohydrates, assuming that it is only composed of cellulose.

2.1.3 Feces

Just like with biomass, feces are expressed in the model as a combination of carbohydrates, proteins and lipids. The composition of feces was calculated by stoichiometrically balancing the equation for human metabolism (see 5.5). The proportions of carbohydrates, proteins and lipids as reported in literature [review in (Rose et al, 2015)] were used as a guideline to solve the stoichiometry. The resulting empirical formula is $CH_{1.6676}O_{0.6028}N_{0.07715}$.

2.2 Establishing the stoichiometric equations

As a general method to balance the equations that represent the reactions occurring in each compartment, each one is stated as a linear programming problem–or non-linear when the constraints of the particular reaction so require–where the goal is to minimize the total amount of matter used. The variables are defined as the number of moles of each compound on either side of the equation. The objective function is subject to the linear constraint of keeping the number of a given element equal on both sides of

the equation. Additionally for each problem *ad hoc* constraints are described e.g., to keep proportions of VFA's produced similar to experimental data on C2. The solving engine for these problems is MS Excel's built-in optimization tool. For linear problems Simplex is used and for non-linear problems GRN non-linear, both on their default configurations settings. In reporting throughout this paper 4 significant decimals are used, but during calculations in Excel 14 decimals were used.

This stoichiometry is a full revision of a version of that was concisely published in two IAC conference papers (Vermeulen et al, 2018b; 2019) and presented at two MELiSSA conferences (Vermeulen et al, 2018a; 2020b). The revisions were based on a detailed validation session and on further literature study. The main changes were the following:

- All biomass compositions were re-evaluated (*Limnospira*, *Rhodospirillum* and higher plants) to more accurately represent their carbohydrate, protein and lipid content.
- No more distinction between fecal and bacterial protein, in line with the singular use of carbohydrates and lipids in the stoichiometry.
- The VFA spectrum was extended beyond acetic acid and butyric acid.
- Instead of representing urine as NH₃, it is now represented as urea, in order to bring carbon into the urine cycling. Hence, a new subsystem with its own stoichiometry representing the breakdown of urea was added.

2.3 Static spreadsheet model to test the closure level

To calculate the steady state flow values for each stream, each compartment's equation is linked, thus building an equation based model that represents the whole cycle. To solve it, the one *a priori* known value is used: the consumption rate of the crew compartment, which has been set at 3,000 kcal/day/person. Knowing these requirements for C5, the required food outputs of C4a and C4b are calculated. By continuing to work backwards through the entire stoichiometry, the flow rates through the entire loop could be established over a period of 1 day for a crew of six.

Since there is a subcycle between C1 and C2 the exchange flows between them need to be calculated (Figure 2). Having the required steady state flows into C1, and out of C2 (except the subcycle flows into each other), the process of exchange between C1 and C2 can be examined. This requires an iterative approach in which, given an initial output of VFAs from C1 to C2, a subsequent C2 to C1 flow of biomass is calculated. This, in turn, updates the flow from C1 to C2 again. This eventually converges into a stable exchange between C1 and C2, with complete processing of all VFAs. This then results in cycle-wide steady state flow rates.

3 Stoichiometric model assumptions

This section describes the general assumptions behind the model and those per compartment.



3.1 General assumptions

This stoichiometry is based on the four CHON elements; sulfur and phosphor are not yet included. Throughout the entire loop, food and feces are described as a proportion of carbohydrates, proteins and lipids. The water in this stoichiometric model is what is used in the chemical reactions. The actual water needs of a human crew (drinking and hygiene water) have not been included. Net biomass production only occurs in C2, C4a, and C4b because this biomass is a material input for other compartments. For the other compartments it is assumed there is no growth, and hence no net biomass production occurs. This has an impact on the stoichiometric approach for each compartment, which differs when describing a growth metabolism vs. a maintenance metabolism without growth (Cruvellier et al, 2016). It is further assumed that all waste materials are fully digested in C1, and no undigestible residues are left. And there is no limitation in terms of available space and hence the number of bioreactors and growth chambers that can be included.

3.2 MELiSSA loop adjustments

Several adjustments have been made to the official concept of the MELiSSA loop, as illustrated in Figure 1. They can be summarized as follows (more details further down in this section):

- The *Rhodospirillum* biomass that is produced in C2 is not used for human consumption, but is sent to C1 where it is anaerobically digested, together with the other organic waste.
- C2 does not produce any net CO_2 .
- Urea is included as an individual waste product and is processed by C3.
- C3 consists of two subcompartments, C3a and C3b, to respectively process ammonia and urea.

• An auxiliary compartment has been added to burn surplus hydrogen and generate water.

3.3 Assumptions for Compartment 1: anaerobic fermentation

In C1, the thermophilic anaerobic bacteria can convert up to 70% of the waste under experimental conditions (Lasseur et al, 2010). This is because cellulose and hemicellulose can only be hydrolyzed slowly, further hampered by the fact that both are often intermeshed with lignin in lignocellulose, which is even more difficult to break down (Liebetrau et al, 2017). In this model it is assumed that all organic waste is fully converted in C1. In the MELiSSA system up to 71% of the organic nitrogen in the waste is converted into ammonium (Clauwaert et al, 2017). In our stoichiometric model it is assumed that this is 100%. The VFA spectrum produced by C1 consists of four VFAs: acetic acid, propionic acid, butyric acid and valeric acid. These VFAs are characteristically produced during acidogenesis (Jiang et al, 2013; Khan et al, 2016).

3.4 Assumptions for Compartment 2: *Rhodospirillum* culture

In the original conceptualization of MELiSSA, it was assumed that in addition to the microalgae and crops from C4, the edible purple sulfur bacteria from C2 would also serve as a food supply for the crew (Lasseur et al, 1996; Albiol et al, 2000; Figure 1). However, because of close proximity, there is always a chance that pathogens associated with fecal matter in C1 pass through membrane filtration and end up with the edible bacteria in C2 (Clauwaert et al, 2017). Therefore, the produced biomass from this compartment is sent back

Parameter	Value	References and notes	
Daily caloric intake	3,000 kcal	Poughon et al, 1994; Ewert et al, 2022	
Daily caloric intake from higher plants	2,700 kcal	90%	
Daily caloric intake from Limnospira	300 kcal	10%	
Oxygen consumption	0.84 kg/day or 52.50 mol/day	Jones, 2003	
CO ₂ production	1 kg/day or 22.72 mol/day	Jones, 2003	
Urea production	0.1239–0.7758 mol/day	Based on Putnam, 1971; MedlinePlus, 2023.	
Feces production	Stoichiometrically determined	_	

TABLE 3 Assumptions about the human metabolism. Listed values are for one crew member.

to C1 for anaerobic degradation. This is in correspondence with other MELiSSA modeling studies (Poughon et al, 2000; Waters et al, 2004; Thiron, 2020). The overall diet of the crew should only contain a limited amount of microbial single-cell protein, because of increased nucleic acid contents and lower palatability (Tusé and Miller, 1984; Tranquille and Emeis, 1996). The crew is already using another microbial food source (*Limnospira* from C4a, Figure 1), and this might be an additional argument to avoid consuming the bacteria from C2. Theoretically, the assimilation of the VFAs by purple sulfur bacteria in C2 could generate a net CO_2 output (Hendrickx and Mergeay, 2007). However, in practice, this has not been observed yet in MELiSSA bioreactor experiments with C1 effluent as VFA source (Mastroleo et al, 2020).

3.5 Assumptions for Compartment 3: processing ammonia and urea

Analogous to C4, this compartment also consists of two subcompartments, C3a and C3b. C3a processes ammonia and produces nitrate. C3b is a urine treatment unit based on C3a, and also produces nitrate. It is assumed that all incoming ammonia and urea are converted to nitrate.

3.6 Assumptions for Compartment 5: human metabolism

The assumptions about the human metabolism are summarized in Table 3. The reference caloric value of the higher plants is estimated to be 4,000 kcal/kg dry weight (DW). This is based on the reported caloric values of the four crops that constitute the ideal plant (adjusted for their moisture content): bread wheat and durum wheat (Kaleta and Górnicki, 2013; Knowledge4 Policy, 2021; WebMD, 2021), potato (Decker and Ferruzzi, 2013; U.S. Food & Drug Administration, 2017), and soybean (Food Data Central, 2019b). The reference caloric value of *Limnospira* (commercially available as Spirulina) is estimated to be 3,000 kcal/kg DW (Food Data Central, 2019a). Solid output of the human metabolism is described as feces and urea. All other body wastes such as dead skin cells, sweat solids, hair, nails, etc. (Hu et al, 2010) are assumed to be part of the feces. H_2O is only integrated in the stoichiometry as a metabolite, and as such the amounts do not represent actual water consumption. It is assumed that the anabolism and catabolism of the crew are in dynamic equilibrium, and that the crew neither gains or loses weight.

4 Results

This section describes the results of the stoichiometry development, and the simulation results.

4.1 Stoichiometric equations

For each compartment a set of chemical equations with fixed stoichiometric coefficients has been developed, making sure that all types of output are also used as input in the loop, thus enabling full closure. The equations are partially based on existing equations from the MELiSSA literature, but were tailored to serve the needs of this study based on additional literature.

4.1.1 Compartment 1: fermentation with thermophilic anaerobic bacteria

The first compartment of the MELiSSA loop is an anaerobic digester with a consortium of thermophilic bacteria that receives fecal material and other wastes, such as inedible plant material (Michel et al, 2005; Mastroleo, 2009; Poughon et al, 2013). Anaerobic digestion is a natural process that typically consists of four subsequent stages: hydrolysis, acidogenesis, acetogenesis and methanogenesis. Volatile fatty acids are intermediate products in this process, while methane is the main end product (Hendrickx and Mergeay, 2007; Khan et al, 2016). In the MELiSSA system a partial anaerobic transformation is used with a maximal yield of volatile fatty acids (VFAs), all while avoiding the production of methane. The VFAs are useful biomolecules that can be utilized as nutrients for other microorganisms downstream in the MELiSSA loop. Such partial anaerobic digestion is achieved by operating the bioreactor under slightly acidified conditions (pH 5.3) which inhibits methanogenic activity (Mastroleo, 2009; Clauwaert et al, 2017).

Organic waste typically consists of polymers (carbohydrates and proteins), as well as fats (Liebetrau et al, 2017). These cannot be taken up by the microbial cells and have to be broken down into smaller compounds first. During hydrolysis, bacteria excrete hydrolytic enzymes which split these substances into sugars, amino acids, fatty acids, and glycerol. These water-soluble hydrolysis products are then converted into volatile fatty acids, CO_2 and hydrogen during anaerobic fermentation (or acidogenesis) (Lauwers et al, 2013; Liebetrau et al, 2017). C1 receives input from three different sources: feces from the crew, inedible plant material, and biomass from C2. The next paragraphs describe the stoichiometries of the fermentation of, respectively, carbohydrates, proteins, and lipids.

4.1.1.1 Carbohydrate fermentation

Carbohydrate fermentation in the MELiSSA loop has been described in multiple studies, and results in the production of VFAs, CO₂ and hydrogen. In Hendrickx et al (2006) only one generic formula representing all VFAs is used, while in all the other consulted MELiSSA studies just two VFAs are produced (e.g., Dussap et al, 1993; Poughon et al, 2009). In some studies this is complemented with the production of bacterial biomass (Guirado & Podhajsky, 2008; Thiron, 2020). As stated above, in the current stoichiometry, four VFAs are used and no biomass production is needed in the equation of this compartment. As a consequence, a new stoichiometry had to be calculated. Proportions of acetic acid, propionic acid, butyric acid and valeric acid were used from (Yin et al, 2016), describing the anaerobic fermentation of glucose as model carbohydrate at pH 6. In below equation, the proportions of these four VFAs were used since the MELiSSA C1 bioreactor also operates at a low pH of 5.3 (Clauwaert et al, 2017). CO_2 and hydrogen are produced in equal proportions, in line with other published balanced carbohydrate fermentation reactions (Dussap et al, 1993; Yin et al, 2016).

$$\begin{array}{l} CH_{1.6667}O_{0.8333} + 0.1666\,H_2O \rightarrow 0.04651\,C_2H_4O_2 + 0.0711\,C_3H_6O_2 \\ acetic acid & propionic acid \\ + 0.09535\,C_4H_8O_2 + 0.006313\,C_5H_{10}O_2 \\ & butyric acid & valeric acid \\ + 0.2807\,CO_2 + 0.2807\,H_2 \end{array} \tag{1}$$

4.1.1.2 Protein fermentation

Just like with carbohydrates, there are several stoichiometric descriptions of the fermentation of proteins in the MELiSSA literature. Older studies use acetic acid and butyric acid as VFA output (e.g., Dussap & Gros, 1991), subsequent studies use up to 5 VFAs (e.g., Guirado & Podhajsky, 2008; Thiron, 2020). In these latter studies, microbial biomass is also included as an output. As biomass output is not necessary in this part of our stoichiometric model, and as a slightly differing empirical formula for protein is used, a new stoichiometric balance needs to be calculated. Proteins are polymers consisting of amino acids. During anaerobic fermentation, proteins are first broken down into their individual amino acids. The subsequent fermentation of these amino acids can be described using a single reaction step, and results in the production of VFAs, CO₂, hydrogen and ammonia (Ramsay and Pullammanappallil, 2001; Poughon et al, 2009). In stoichiometric equations, CO₂, hydrogen and ammonia are generally being produced in a similar order of magnitude (Poughon, 2007b; Thiron, 2020). Tepari (2019) investigated protein fermentation during anaerobic wastewater treatment, and used bovine serum albumin (BSA) as model protein. At pH 5, only acetic acid (37.4%), propionic acid (13.7%) and butyric acid (48.9%) were produced, and no valeric acid. These proportions were used to calculate the following stoichiometric equation:

$$\begin{array}{c} \mathrm{CH}_{1.59}\mathrm{O}_{0.31}\mathrm{N}_{0.25} + 0.4962\,\mathrm{H_2O} \rightarrow 0.1056\,\mathrm{C_2H_4O_2} + 0.03867\,\mathrm{C_3H_6O_2}_{acetic\,acid} \\ + 0.138\,\mathrm{C_4H_8O_2} + 0.1209\,\mathrm{CO_2}_{butyric\,acid} \\ + 0.037\,\mathrm{H_2} + 0.25\,\mathrm{NH_3} \end{array} \tag{2}$$

4.1.1.3 Lipid fermentation

As described above, tripalmitin is used as model lipid in this model. Since in previous MELiSSA modeling studies, palmitic acid is used, a new stoichiometric equation has to be calculated. When lipids such as triacylglycerols are anaerobically digested, they are first hydrolyzed to glycerol and long-chain fatty acids (LCFAs) (Mackie et al, 1991; Li et al, 2002; Cirne et al, 2007). Subsequently, the glycerol is degraded into VFAs, while the LCFAs are broken down into acetic acid and hydrogen through beta-oxidation (Weng and Jeris, 1976; Ceron-Chafla et al, 2021; Holohan et al, 2022). Firstly, the glycerol and LCFA fermentations will be individually described, after which they will be integrated into one equation.

The hydrolysis of tripalmitin can be written as follows:

$$\begin{array}{cc} C_{51}H_{98}O_6 + 3 H_2O \rightarrow & C_3H_8O_3 + 3 C_{16}H_{32}O_2 \\ \text{tripalmitin} & & \text{glycerol} & \text{palmiticacid} \end{array} \tag{3}$$

In experiments by Yin et al, 2016, glycerol was subjected to (nonstrict) anaerobic fermentation at pH 6. The study presents three individual stoichiometric reactions, each describing the degradation of glycerol into acetic acid, propionic acid, and butyric acid. The experiment resulted in a production of 32% acetic acid, 52% propionic acid, and 16% butyric acid. Integrating the three individual stoichiometric reactions, taking into account the above mentioned proportion, leads to the following single equation:

$$\begin{array}{l} C_{3}H_{8}O_{3}+0.2727\,H_{2}O \rightarrow 0.2727\,C_{2}H_{4}O_{2}+0.4481\,C_{3}H_{6}O_{2}\\ & \text{acetic acid} \end{array}$$

The stoichiometric equation for palmitic beta-oxidation is described by Li et al (2018):

$$\begin{array}{c} C_{16}H_{32}O_2 + 14\,H_2O \rightarrow 8\,C_2H_4O_2 + 14\,H_2 \\ \text{palmitic acid} \end{array} \tag{5}$$

Integrating Eqs 3–5 leads to:

$$CH_{1.9216}O_{0.1177} + 0.8789 H_2O \rightarrow 0.4759 C_2H_4O_2 + 0.008786 C_3H_6O_2$$

acetic acid propionic acid
+ 0.002737 C_4H_8O_2
butyric acid
+ 0.01082 CO_3 + 0.8505 H_3 (6)

4.1.2 Compartment 2: VFA processing by photoheterotrophic bacteria

The purpose of MELiSSA C2 is to recycle the VFAs produced in C1. The purple non-sulphur phototrophic bacteria *Rhodospirillum*

rubrum is utilized for this. Under anaerobic conditions, it uses light as an energy source to convert VFAs into biomass (Mastroleo, 2009; De Meur et al, 2020). Because we use a custom composition of *Rhodospirillum* biomass in this model, a new stoichiometry had to be calculated. Anaerobic digestion of organic matter may lead to fluctuating proportions of different VFAs (Alloul et al, 2018). Because of the variable VFA spectrum coming out of C1, it was decided to develop a individual equation for each VFA. The assimilation process consumes CO_2 and NH_3 , and for acetic acid also hydrogen (Dussap et al, 1993; Fulget, 1996; Thiron, 2020). As explained above, no net CO_2 production occurs in the following equations.

$$\begin{split} & C_{2}H_{4}O_{2} + H_{2} + 0.2774 \text{ CO}_{2} + 0.4073 \text{ NH}_{3} \\ & \rightarrow \left[0.3327 \text{ CH}_{1.6667}O_{0.8333} \\ & + 1.6291 \text{ CH}_{1.5900}O_{0.3100}N_{0.2500} + 0.3155 \text{ CH}_{1.9216}O_{0,1177} \right]_{\text{Rhodospirillum}} \\ & + 1.7353 \text{ H}_{2}O \end{split}$$

$$\begin{array}{l} 0.8556 \quad C_{3}H_{6}O_{2} + 0.1611 \, \text{CO}_{2} + 0.4879 \, \text{NH}_{3} \\ \rightarrow \left[0.3986 \, \text{CH}_{1.6667}O_{0.8333} + 1.9515 \, \text{CH}_{1.5900}O_{0.3100} \, \text{N}_{0.2500} \\ & \text{carbohydrates} \end{array} \right]_{\text{proteins}} + 0.3780 \, \text{CH}_{1,9216}O_{0,1177} \\ \left]_{\text{lipids}} + 1.052 \, \text{H}_{2}O \end{array} \tag{8}$$

$$\begin{array}{l} 0.4084 \ C_{5}H_{10}O_{2} + 0.3762 \ CO_{2} + 0.4325 \ NH_{3} \\ \\ \rightarrow \left[0.3533 \ CH_{1.6667}O_{0.8333} + 1.7299 \ CH_{1.5900}O_{0.3100}N_{0.2500} \\ \\ + 0.335 \ CH_{1,9216}O_{0,1177} \\ \\ \\ I_{lipids} \end{array} \right]_{Rhodospirillum} + 0.6991 \ H_{2}O \tag{10}$$

4.1.3 Compartment 3: nitrification of ammonia and ureolysis

In some cases, ammonia and urea can be taken up directly by plants or microorganisms grown for food production. However, in a BLSS it is desirable to convert ammonia and urea to nitrate because it is a less volatile and less reactive molecule (Clauwaert et al, 2017; De Paepe et al, 2018; Sachdeva et al, 2021). MELiSSA C3 receives ammonia from C2 and urine from C5, and transforms both into nitrate for the plants and microalgae in C4 (Duatis and Moreno, 2009; 2010). Ammonia is processed in a nitrifying bioreactor, while urine is processed separately in a modified version of the nitrifying bioreactor, the so-called Urine Treatment Unit (UTU) (Guirado and Podhajsky, 2008). As both bioreactors process different compounds, two separate compartments are defined in the stoichiometric model: 3a and 3b.

4.1.3.1 Compartment 3a: nitrification of ammonia by nitrifying bacteria

The nitrifying bioreactor as defined in the MELiSSA project is a cylindrical packed-bed bioreactor with polystyrene beads that are colonized by a biofilm of *Nitrosomonas europaea* and *Nitrobacter winogradsky* (Cruvellier et al, 2016; Garcia-Gragera et al, 2021). The nitrification of ammonia happens in two steps, each step carried out by one of the bacterial species (Cruvellier et al, 2016):

$$2 \operatorname{NH}_{3} + 3 \operatorname{O}_{2} \to 2 \operatorname{HNO}_{2} + 2 \operatorname{H}_{2} \operatorname{O}$$
(11)
Nitrosomonas

$$2 \text{HNO}_2 + \text{O}_2 \rightarrow 2 \text{HNO}_3$$
(12)
Nitrobacter

Taken together this results in:

$$\mathrm{NH}_3 + 2\mathrm{O}_2 \to \mathrm{HNO}_3 + \mathrm{H}_2\mathrm{O} \tag{13}$$

4.1.3.2 Compartment 3b: breakdown of urea by ureolytic heterotrophic bacteria

In a recent MELiSSA study of a UTU concept (Christiaens et al, 2019), showed how human urine can be nitrified with the use of a synthetic microbial community of ureolytic heterotrophic bacteria and nitrifying bacteria cultivated in a continuous stirred tank reactor. First, during ammonification, the ureolytic heterotrophic bacteria break down urea into ammonia and CO_2 , a process catalyzed by the enzyme urease (Nicolau et al, 2010; De Paepe et al, 2018). Subsequently, the ammonia is further converted into nitrate by the nitrifying bacteria as described above. The stoichiometric equation describing the ammonification of urea can be found in (Guirado and Podhajsky, 2008):

$$CO(NH_2)_2 + H_2O \rightarrow 2NH_3 + CO_2$$
(14)
urea

Taken together with (13), this results in:

$$CO(NH_2)_2 + 4O_2 \rightarrow 2HNO_3 + H_2O + CO_2$$
(15)
urea

4.1.4 Compartment 4: food production 4.1.4.1 Compartment 4a: photoautotrophic cyanobacteria growth

The edible microalgae *Limnospira platensis* and *indica* (previously *Arthrospira platensis* and *indica*) are cyanobacteria that consume nitrate and CO_2 that are produced in other compartments of the loop, and generate oxygen and food for the crew (Clauwaert et al, 2017; Alemany et al, 2019; Sachdeva et al, 2021). *Limnospira* is grown inside a gas lift photobioreactor with illumination and a gas injection system (Garcia-Gragera et al, 2021; Poughon et al, 2021). Since a custom biomass composition is used in our model, a new stoichiometric balance had to be calculated (generated biomass is dry mass):

$$CO_{2} + 0.1788 \text{ HNO}_{3} + 0.7342 \text{ H}_{2}\text{O}$$

$$\rightarrow \left[0.1461 \text{ CH}_{1.6667} \text{ O}_{0.8333} + 0.7154 \text{ CH}_{1.5900} \text{O}_{0.3100} \text{ N}_{0.2500} \right]_{\text{proteins}} + 0.1385 \text{ CH}_{1,9216} \text{O}_{0,1177} \left]_{\text{Limnospira}} + 1.4554 \text{ O}_{2}$$
(16)

4.1.4.2 Compartment 4b: higher plant growth

Edible crops are grown in C4b of the MELiSSA loop (Pannico et al, 2022). The goal of this compartment is in line with that of C4a: recycling nitrate and CO_2 , and producing oxygen and food (Lasseur et al, 2010; Peiro et al, 2020). In the current model, a harvest index of 50% is used, which means that half of the plant biomass is edible, while the other half is not. The non-edible part is considered as waste and is used as feed for the thermophilic anaerobic bacteria in C1. This can be summarized in the general equation:

$$CO_2 + H_2O + HNO_3 \rightarrow [biomass]_{higher plants edible}$$

+ 0.8965 [biomass]_{higher plants non-edible} + O_2 (17)

This corresponds to the following stoichiometric equation (generated biomass is dry mass):

```
2.9101 \text{ CO}_2 + 0.0842 \text{ HNO}_3 + 2.4 \text{ H}_2\text{O}
```



4.1.5 Compartment 5: human metabolism

The general equation for the human metabolism can be summarized as follows (Guirado and Podhajsky, 2008):

$$[biomass]_{higher plants edible} + [biomass]_{Limnospira} + O_2$$

$$\rightarrow [feces] + CO(NH_2)_2 + H_2O + CO_2$$
(19)

To determine the final stoichiometric equation, a few preparatory calculations are needed. Firstly, the right proportion between edible higher plant biomass and Limnospira biomass needs to be established. In the model, 90% of the caloric content of the food comes from the higher plants, and 10% comes from Limnospira. Higher plants and Limnospira are considered to have a caloric content of respectively 4,000 and 3000 kcal/kg. This amounts to a target daily food intake of 675 g DW (27.86 mol) of higher plant biomass and 100 g DW (4.56 mol) of Limnospira biomass per crew member. Using the biomass compositions, this can then be translated into corresponding values for carbohydrates, proteins, and lipids. For oxygen, CO₂, and urea, reported daily values were transformed into moles (Table 2), and this was then used for calculating a stoichiometry with realistic proportions between these compounds and biomass uptake. The amount of feces is calculated by solving the stoichiometric equation. Below equation describes the metabolism of one crew member:



4.1.6 Auxiliary processes

4.1.6.1 Excess hydrogen burn-off

One auxiliary process has been added to the loop with the function to burn any excess hydrogen. This is in correspondence with other MELiSSA modeling studies such as (Poughon et al, 2000).

$$2 H_2 + O_2 \rightarrow 2 H_2O$$

4.2 Simulation results

The results of the stoichiometric spreadsheet model running in steady state conditions can be found in Table 4. The overall input and output values of all compounds in the loop are listed, for a crew of six over a period of 1 day (the time unit used in the ABM for which this stoichiometry was developed). All different stoichiometric equations were successfully balanced with each other, which resulted in a very high closure level with 12 out of 14 values reaching full closure, with no material loss. Oxygen displays an almost neglectable loss of 0.00021%, and CO₂ 0.00001%. This is assumed to be caused by the numerical limitations of the software tools (e.g., number of decimals and rounding). The values correspond to respectively 2.13 mg O2 and 1.47 mg CO2 material loss per person per day. Over the course of a year, this would amount to 4.67 g O₂ and 3.21 g CO₂ for the entire crew. The VFA spectrum (weight-based) at steady state consists of 57.20% acetic acid, 12.59% propionic acid, 29.98% butyric acid and 0.23% valeric acid.

5 Discussion

This study presents a stoichiometric model of the MELiSSA BLSS, which describes the main metabolic processes in all compartments using a series of static chemical equations. In steady state, 100% of the necessary food and oxygen for the crew, as well as 100% of the compounds required for all other organisms in the loop, are generated. Using the assumption that all waste compounds in the loop can be broken down again, full closure was achieved in the static model by balancing the size of each compartment relative to all the other compartments. This is the first time a fully closed static MELiSSA model is described that can provide all the food and oxygen for astronauts. The model is a useful tool for theoretical

Compound	Input (mol)	Output (mol)	Material loss(mol)	Material loss(%)
Carbohydrates	1832.2836	1832.2836	0.0000	0.00000%
Protein	7587.0675	7587.0675	0.0000	0.00000%
Lipids	1490.3754	1490.3754	0.0000	0.00000%
Acetic acid	1570.2506	1570.2506	0.0000	0.00000%
Propionic acid	426.4730	426.4730	0.0000	0.00000%
Butyric acid	1207.5497	1207.5497	0.0000	0.00000%
Valeric acid	10.8735	10.8735	0.0000	0.00000%
Urea	4.6548	4.6548	0.0000	0.00000%
NH ₃	1882.6600	1882.6601	0.0001	0.00000%
HNO ₃	14.1068	14.1068	0.0000	0.00000%
H ₂ O	5591.1996	5591.1994	-0.0001	0.00000%
H ₂	2004.4967	2004.4967	0.0000	0.00000%
O ₂	391.0724	391.0716	-0.0008	-0.00021%
CO ₂	1543.0613	1543.0611	-0.0002	-0.00001%

TABLE 4 Input and output values of the different compounds, after reaching steady state. Values for a crew of six, over a period of 1 day.

research on autonomous BLSSs for long-duration space exploration, allowing tracing of the elements over the entire cycle.

However, in practice, no system can be 100% closed. There are always potential outfluxes from the system, for example, through a buildup of recalcitrant organic matter (Hendrickx and Mergeay, 2007; Lasseur et al, 2010; Zhang et al, 2018) or the accumulation of different precipitates such as carbonates and phosphates (De Paepe et al, 2018; Christiaens et al, 2019). Recalcitrant organic matter consists mainly of plant fibers that do not rapidly biodegrade. Plant cell walls are composed of cellulose, hemicellulose, pectin, and lignin and are among the least degradable polymers in a BLSS (Hendrickx and Mergeay, 2007; Zhang et al, 2018). As a result, up to about 70% of the organic waste could be digested in MELiSSA lab experiments (Farges et al, 2008; Lasseur et al, 2010). The MELiSSA community has investigated additional methods to break down the remaining recalcitrant matter using hydrothermal and chemical oxidation, and anaerobic and hyperthermophilic cellulose-degrading bacteria (Hendrickx & Mergeay, 2007; Lasseur et al, 2010). The most efficient results were obtained by using supercritical water oxidation (SCWO) in which up to 98% of all organic matter could be broken down (Hendrickx and Mergeay, 2007; Zhang et al, 2018). Hydrogen peroxide was used as an oxidizer in these tests, which is a compound that can be generated within a BLSS loop (Tikhomirov et al, 2011; Vijapur et al, 2017; Nelson et al, 2020). Nevertheless, it is safe to assume that, in reality, there will always be some level of material loss in any BLSS loop. It would therefore be interesting to investigate what accumulates over a long period of time and then calculate the dimensions of the reserve storage required to mitigate the resulting material shortages.

The interaction between compartments C1 and C2 represents a subcycle in this model. The *Rhodospirillum* biomass generated in C2 is fed back into C1 as organic waste. The goal of this subcycle is to produce a surplus of CO_2 , H_2 , and NH_3 that is needed in the other compartments of the MELiSSA loop. Steady state of this subcycle is

reached gradually until enough biomass from C2 is sent back to C1 to ensure that the exact amount of VFAs is produced to sustain a stable Rhodospirillum culture. If too few VFAs are provided, C2 will underperform due to a lack of Rhodospirillum growth. If too many VFAs are provided, the Rhodospirillum culture will not be able to process all of them, leading to accumulation and negative impacts on the mass flows and efficiency of the entire loop. At steady state, this subcycle generates the following surplus (per day, for a crew of six): 213 mol CO₂, 434 mol H₂, and 126 mol NH₃. The subcycle also explains the dominant amount of protein present in the loop, despite the food provided to the crew containing much less protein (Table 4). Due to the need for a significant mass of Rhodospirillum to process the VFAs and their high protein content (72%), a lot of protein is present in the loop in general. Bio-electrochemical oxidation is a potentially more efficient approach to convert the VFAs into CO₂. In a study by Luther et al, 2018, using bio-electrochemical oxidation within a microbial electrolysis cell, 80%-100% of the VFAs in the effluent of MELiSSA's C1 were converted into CO₂ in 7 days. With further process development and optimization, bio-electrochemical oxidation can potentially entirely replace the use of Rhodospirillum in C2.

(Jiang et al, 2013) conducted a series of experiments on anaerobic digestion of food waste. At a pH of 5.0, the resulting VFA spectrum consisted of 60.40% acetic acid, 8.32% propionic acid, 31.13% butyric acid, and 0.15% valeric acid, which falls within the same range as the results achieved in our model at steady state. Another study by Lim et al, 2008 also involved anaerobic digestion of food waste. At a pH of 5.5, the resulting VFA spectrum had a more variable composition, but acetic acid was generally the dominant VFA. These reported results are consistent with the results of our static model, which suggests that the theoretical VFA outcome is a potentially realistic scenario. However, it must be noted that VFA spectra generated during anaerobic fermentation are highly variable and depend on various factors such as substrate composition and inoculum (Poughon et al, 2013; Khan et al, 2016), pH and temperature (Shin et al, 2004; Jiang et al, 2013; Khan et al, 2016), and bioreactor design and operation (Lim et al, 2008; Bharathiraja et al, 2016). For instance, in experiments using food waste by Shin et al (2004), and in MELiSSA experiments using a synthetically composed organic waste (Luther et al, 2018; Zhang et al, 2019), both acetic acid and butyric acid were the main VFAs, with butyric acid being dominant.

Growing enough plants to meet the food requirements of a human crew typically results in the production of more oxygen than the crew needs. This is because the harvest index of the crops is often less than 100%, meaning that only a portion of the grown biomass is used for food (Jones, 2003). However, the unused biomass still generates oxygen. In a closed loop system, this could lead to an accumulation of excess oxygen (as e.g., observed in Poughon et al, 2000; Gros et al, 2003) and, eventually, the collapse of the system. In our model, the excess oxygen is used to hydrolyze solid waste in C1, in the form of H_2O . The hydrogen-burning auxiliary helps to transform any remaining oxygen gas into H_2O . Since oxygen originates from splitting H_2O during photosynthesis, the amount of surplus oxygen matches the amount of surplus hydrogen in the system.

In our model, the human crew consumes 21.51 mol/person/day of CO₂, which is very close to the target value of 22.72 mol/person/day used for calculating the stoichiometry. However, the human crew only uses 24.29 mol/person/day of oxygen, much lower than the target value of 52.50 mol/person/day. The proportion between consumed oxygen and produced CO₂ should be 2.31, but in our model, this proportion is 1.13, precisely because of the lower oxygen consumption. Remarkably, other BLSS modeling studies show the same tendency. In MELiSSA studies, the proportion is 1.20 (Fulget, 1996; Thiron, 2020) and in Lunar Palace studies, it ranges from 1.13 to 1.38 (Hu et al, 2010; Fu et al, 2016). This suggests that the description of the O₂ consumption in all of these studies is not entirely complete. This could be due to the fact that in reality, more compounds are oxidized than represented in the simplified stoichiometric equation for human metabolism. The total mass of consumed food per person per day corresponds to 781 g DW/day. This is slightly higher than the range of baseline values used in MELiSSA studies and reference documents. Adjusted to a daily food intake of 3,000 kcal/person, the values in these studies and documents range from 509 to 748 g DW/day (e.g. (Dussap et al, 1993; Poughon et al, 2000; Thiron, 2020)).

The composition of feces was calculated by solving the stoichiometric equation for human metabolism, using the general characterization of feces reported in the literature as a starting point (Rose et al, 2015). After solving the equation for human metabolism, the resulting feces composition was found to be 66% carbohydrates, 28% proteins, and 6% lipids, corresponding to the empirical formula CH_{1.6676}O_{0.6028}N_{0.07715}. This formula is similar to the one used in BLSS studies by Hu et al, 2010 and Fu et al, 2016, which is CH_{1.70}O_{0.60}N_{0.05}. Because the MELiSSA diet is a vegan diet, it is expected that there is a high fiber intake and a resulting high amount of fiber in the feces (Brodribb et al, 1980; Kay, 1982; Forsum et al, 1990). As a result, the overall carbohydrate fraction could be quite significant. It should be noted that the fraction of protein in the feces could be an underestimation, as bacteria can compose up to half of the fecal solids and may have a significant protein content

(Cummings, 2001; Rose et al, 2015). Intestinal bacterial growth is not included in the human stoichiometry, which is justifiable as it means more digestion is done by the same groups of microbes in C1. The amount of feces produced is 253 g DW/person/day. Jenkins et al, 2001 measured 160 ± 24 g DW/person/day for a vegetarian diet. It should be noted that no other solid body wastes have been included in the stoichiometry, and as such the 'feces' in the equation represent a more general representation of all wastes combined.

Further development of the stoichiometric model could focus on increasing its granularity. This could involve expanding the model to include more elements beyond CHON, incorporating a greater diversity of crops, and using a more detailed description of human metabolism. Nitrogen, phosphorus, and potassium are the three major macro-elements for plants (Usherwood and Segars, 2001; Kanazawa et al, 2008). Therefore, including the latter two elements in the next version of the model seems like a logical step. Fertilizers typically contain these three elements, but it is crucial that they are applied in the correct ratio (Usherwood and Segars, 2001; Chun et al, 2017), which must also be factored into the model. Diverse plant crops could be used instead of the concept of an 'ideal plant', allowing for a more detailed description of the crew's daily diet. Integrating water usage beyond it's role as a metabolite will result in a more accurate description of the entire water budget. For example, water transpiration, water excretion, and free water inside the plant could be factored in. By integrating solid body wastes other than feces, the stoichiometry of human metabolism could also be further refined.

Data availability statement

The raw data supporting the conclusion of this article will be made available by the authors, without undue reservation.

Author contributions

AV and AP conceived and designed the study and performed the MELiSSA literature review. AV did the background research for the stoichiometry, while AP developed the tool in Excel to balance the chemical equations. The stoichiometric spreadsheet model was created in Excel by AP. AV wrote the first draft of the manuscript, and AP contributed sections. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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