Global dynamics of the chemostat with overflow metabolism

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Abstract
Fast growing E. coli cells, in glucose-aerobic conditions, excrete fermentation by-products such as acetate. This phenomenon is known as overflow metabolism and has been observed in a diverse range of microorganisms. In this paper, we study a chemostat model subject to overflow metabolism: if the substrate uptake rate (or the specific growth rate) is above a threshold rate (different from zero), then secretion of a by-product happens. We assume that the presence of the by-product has an inhibitory effect on the growth of the microorganism. The model is described by a non-smooth differential system of dimension three. We prove the existence of at most one equilibrium (or steady-state) with presence of microorganism, which is globally stable. We use these results to discuss the performance of chemostat-type systems to produce biomass or recombinant proteins.

Keywords Global dynamics · Inhibitor · Overflow metabolism · Chemostat · Recombinant protein · Cooperative system · Global stability

1 Introduction
Escherichia coli (E. coli) is a bacterium that is naturally found in the intestine of humans and other mammals. This bacterium has been a preferred choice for large-scale production of recombinant proteins1 such as insulin, GFP (green fluorescent protein), or the human growth hormone (Baeshen et al. 2015; Huang et al. 2012). For high density cultivation of E. coli, glucose is generally the preferred and most

1 Recombinant proteins are proteins that are artificially made through the recombinant DNA technology.
common carbon and energy source (Bren et al. 2016), since this is inexpensive and readily utilisable. To harvest energy from glucose, *E. coli* combines two different metabolic strategies, aerobic respiration, which needs oxygen, and fermentation, which does not need oxygen Gerosa et al. (2015). Respiration is more energy-efficient than fermentation, nevertheless, in fast growing cells, some energy is also obtained by fermentation. This seemingly wasteful strategy in which cells use fermentation instead of respiration, even in the presence of oxygen, is known as overflow metabolism (Basan et al. 2015). This phenomenon is not only limited to *E. coli*, but to a diverse range of microorganisms (Vazquez 2017). For example, in yeasts, overflow metabolism is known as Crabtree effect (Dashko et al. 2014), and in cancer cells it is known as Warburg effect (Kim and Dang 2006).

Overflow metabolism results in the secretion of fermentation by-products, such as acetate in *E. coli* cultures or ethanol in yeast cultures, which accumulation can have an inhibitory effect on cells growth. For example, glucose uptake is inhibited in *E. coli* and yeast cultures in presence of acetate (Luli and Strohl 1990) and ethanol (Liu and Wu 2008) respectively. Moreover, the formation of these by-products constitutes a diversion of carbon that might have contributed to biomass or protein synthesis. Thus, overflow metabolism can pose a major problem in large-scale production of biomass or recombinant proteins (Eiteman and Altman 2006; Hensing et al. 1995).

Cultivation of *E. coli*, yeasts, and other microorganisms can be done in a chemostat. The chemostat, introduced in the 1950s independently by Monod (1950) and Novick and Szilard (1950), is a perfectly mixed reactor, permanently fed with a nutrient rich medium and simultaneously emptied so that the culture volume is kept constant. Using the chemostat is a way to maintain indefinitely a non-zero growth rate, and therefore to study the organisms under various constant growth rates. The classical chemostat model describes the dynamics of a single population with growth limited by a single nutrient. We refer the reader to Smith and Waltman (1995) and Ajbar and Alhumaizi (2011) for the theory of the chemostat and for different variations of the classical chemostat model. Yano and Koga (1973), Xiu et al. (1998), Heßeler et al. (2006), and Harvey et al. (2014) studied the dynamics of chemostat models with the production of a toxic by-product. Xiu et al. (1998) described the production of the byproduct as a consequence of overflow metabolism. However, in all these works the authors assume that the secretion of the by-product occurs at any growth rate while experimental evidence shows that by-product secretion does not take place at low growth rates Basan et al. (2015).

In this paper, we study the long-term behavior of a chemostat model accounting for the following features of overflow metabolism:

- secretion of a by-product when the substrate uptake rate is above a threshold;
- biomass loss due to secretion of the by-product;
- inhibition of substrate uptake in presence of the by-product.

The model is mainly inspired by the recently proposed model by Mauri et al. (2020) that describes the growth of an *E. coli* culture producing a recombinant protein. In contrast to our model, Mauri et al. (2020) consider growth on the by-product (acetate) and an additional variable describing the dynamics of a recombinant protein concentration. *E. coli* consume acetate only after the glucose (substrate) is totally consumed,
phenomenon known as carbon catabolite repression (Wolfe 2005). Thus, given the continuous supply of substrate in chemostats, we neglect the consumption of the by-product in our model. Note that carbon catabolite repression is also observed in yeasts (Gancedo 1998). With respect to the recombinant protein concentration, in the discussion section we show that our results can be easily extended when considering the dynamics of a recombinant protein.

The chemostat model with overflow metabolism is described by an autonomous system of ordinary differential equations. Using a conservation principle, the model can be reduced to a planar system. Thus, we study the dynamics of the planar system by finding appropriate invariant sets and using results on cooperative systems (Smith 2008). To extend our results to the original model, we use the well known Theorem of Butler-McGehee Smith and Waltman (1995). This technique requires the stability of equilibria, which may be difficult to obtain due to the non-smoothness of the by-product excretion rate function (overflow metabolism). This situation is treated with classical results of the theory of differential equations such as the comparison method (Coppel 1965).

This paper is organized as follows. In Sect. 2, we describe the chemostat model and the main hypotheses. Sections 3 and 4 are devoted to the mathematical analysis of the model. In Sect. 3, we characterize the existence of equilibria and their local stability. In Sect. 4, we present the results on the global behavior of the model. The main result is given in this section (Theorem 1). In the last section, Sect. 5, we begin presenting a brief summary of our mathematical results. Then, we finish the paper with a discussion on the steady state production of biomass and recombinant proteins in chemostat-type systems.

2 Chemostat model

We consider a chemostat (see Fig. 1) with a single population of microorganisms whose concentration is denoted by \( x \). This population grows at a specific growth rate \( \mu(\cdot) \). The specific growth rate considers the carbon gain by substrate uptake and the carbon loss due to metabolic overflow \( i.e. \)

\[
\mu(\cdot) = Y_S r_S(\cdot) - Y_R r_{of}(\cdot),
\]

where \( r_S \) is the substrate uptake rate, \( r_{of} \) is the metabolic overflow rate (or by-product formation rate), and \( Y_S, Y_R \) are yield coefficients. Following Basan et al. (2015), when
By-product excretion rate ($r_{of}$) as a function of the substrate uptake rate ($r_S$)

Fig. 2

$r_S$ is higher than a threshold rate $r_{S0}$, then the excretion of by-product occurs at a rate proportional to the difference between $r_S$ and $r_{S0}$ i.e. $r_{of} = f(r_S)$ with $f$ defined as (see Fig. 2):

\[
f(r_S) = \begin{cases} 
k(r_S - r_{S0}) & \text{if } r_S > r_{S0}, \
0 & \text{if } r_S \leq r_{S0}, \end{cases}
\]

with $k > 0$. The substrate uptake rate $r_S$ is a function of the substrate bulk concentration ($S$) and the overflow metabolism by-product ($R$) i.e. $r_S = r_S(S, R)$. We assume that $r_S$ is continuously differentiable for all $S, R \geq 0$ and that:

\[
r_S(0, R) = 0, \quad \frac{\partial r_S}{\partial S} > 0, \quad \text{and} \quad \frac{\partial r_S}{\partial R} < 0.
\]

Thus, the by-product $R$ has an inhibitory effect on the substrate uptake rate. An example for $r_S$ is given by (see Mauri et al. 2020):

\[
r_S(S, R) = r_{S,max} \frac{S}{K_S + S} \frac{K_{i,R}}{K_{i,R} + R},
\]

where $r_{S,max}$ is the maximal substrate uptake rate, $K_S$ is a half saturation constant, and $K_{i,R}$ is an inhibition constant. The chemostat is fed at a rate $F > 0$ with a substrate concentration $S_{in}$. The dilution rate is defined as $D = F/V$, with $V$ the volume of the culture. Mass balance equations lead to:

\[
\begin{align*}
\frac{dx}{dt} &= (\mu(S, R) - D)x, \\
\frac{dS}{dt} &= DS(S_{in} - S) - r_S(S, R)x, \\
\frac{dR}{dt} &= -DR + r_{of}(S, R)x.
\end{align*}
\]

Model (4) is that of a standard chemostat with a single species with growth limited by a single substrate, with the added feature that a by-product is produced as a consequence of overflow metabolism.
Throughout the paper we assume:

\[ Y_S - kY_R > 0. \] (4)

This assumption implies that the growth rate function \( \mu \) is strictly increasing in \( S \) and strictly decreasing in \( R \). This follows directly from noting that:

\[
\mu(\cdot) = \begin{cases} 
(Y_S - kY_R)r_S(\cdot) + kY_Rr_{S0} & \text{if } r_S(\cdot) > r_{S0}, \\
Y_S r_S(\cdot) & \text{if } r_S(\cdot) \leq r_{S0}.
\end{cases}
\]

Assumption (4) is satisfied by the parameters given by Mauri et al. (2020). We also assume that

\[ r_S(S_{in}, 0) > r_{S0} \quad (i.e. \ r_{of}(S_{in}, 0) > 0). \] (5)

In the long-term operation with presence of microorganisms, the substrate concentration in the medium cannot be higher than \( S_{in} \). Then, if \( r_S(S_{in}, 0) \leq r_{S0} \), overflow metabolism is not possible in the long-term, and the study of the dynamics of (4) is reduced to that of a classical chemostat model.

Recalling the definition of \( \mu \) and combining (4) and (5), we have the following inequality:

\[ \mu(S_{in}, 0) > Y_S r_{S0}. \]

This inequality allows us to consider dilution rates between \( Y_S r_{S0} \) and \( \mu(S_{in}, 0) \). As we will show in the next sections, in the long-term operation, only when \( Y_S r_{S0} < D < \mu(S_{in}, 0) \) there is presence of the by-product in the culture.

As expected, the domain of biological interest, that is \( \mathbb{R}_+^3 := \{ (x, S, R) \in \mathbb{R}^3 \mid x, S, R \geq 0 \} \), is positively invariant.

The conservation principle for chemostats is satisfied by the variable \( W = x + Y_S S + Y_R R \ i.e. \)

\[
\frac{dW}{dt} = D(Y_S S_{in} - W). \] (6)

We can rapidly verify that

\[ W(t) = W(0)e^{-Dt} + Y_S S_{in}(1 - e^{-Dt}). \] (7)

In view of the definition of \( W \) we have

\[ 0 \leq x(t), Y_S S(t), Y_R R(t) \leq W(t). \]

Since \( W(t) \to Y_S S_{in} \) as \( t \to \infty \), we conclude that (4) is dissipative \( i.e. \) solutions of (4) are attracted by the bounded set \( [0, Y_S S_{in}] \times [0, S_{in}] \times [0, Y_S S_{in}/Y_R] \).
3 Existence of steady states and local stability

Equation (4) admits at most two equilibria. A trivial equilibrium corresponds to the absence of microorganisms. It is given by

\[ E_0 = (0, S_{in}, 0), \]  

and it always exists. The other possible equilibrium is characterized by the presence of microorganisms. The presence of the by-product depends on the dilution rate. The following proposition formally characterizes the existence of this equilibrium.

**Proposition 1 (Existence of the non-trivial equilibrium)**

(a) If \( \mu(S_{in}, 0) > D \), then (4) admits a unique equilibrium \( E^* = (x^*, S^*, R^*) \) with presence of microorganisms (i.e. \( x^* > 0 \)). Moreover,

- If \( Y_{SRS0} > D \), then \( R^* = 0 \) and \( r_S(S^*, 0) < r_{S0} \).
- If \( Y_{SRS0} = D \), then \( R^* = 0 \) and \( r_S(S^*, 0) = r_{S0} \).
- If \( Y_{SRS0} < D \), then \( R^* > 0 \) and \( r_S(S^*, R^*) > r_{S0} \).

(b) If \( \mu(S_{in}, 0) \leq D \), then (4) has no equilibrium with presence of microorganisms.

**Proof** Assume that \( D \leq Y_{SRS0} \). In this case, any positive steady state of (4) has no by-product. Indeed, by contradiction, if \( (x^*, S^*, R^*) \) is a positive steady state of (4) with \( R^* > 0 \), then \( r_{of}(S^*, R^*) = DR^*/x^* > 0 \) (from the third equation in (4)). Thus, from the first equation in (4) we obtain:

\[
\mu(S^*, R^*) - D = (Y_S - kY_R) (r_S(S^*, R^*) - r_{S0}) + Y_{SRS0} - D > 0,
\]

which contradicts the fact that \( (x^*, S^*, R^*) \) is a positive steady state. Hence, any positive steady state of (4) has the form \( (x^*, S^*, 0) \). As in a classical chemostat model (note that \( S \mapsto \mu(S, 0) \) is strictly increasing), (4) admits a unique positive steady state if \( \mu(S_{in}, 0) > D \), and has no positive steady states if \( \mu(S_{in}, 0) \leq D \).

Now assume that \( D > Y_{SRS0} \). In this case, the by-product is present in any positive equilibrium of (4). Indeed, by contradiction, if \( (x^*, S^*, 0) \) is a positive steady state of (4), then \( r_{of}(S^*, 0) = 0 \). But we have \( Y_{SRS}(S^*, 0) = D > Y_{SRS0} \), which implies \( r_S(S^*, 0) > r_{S0} \), and hence \( r_{of}(S^*, 0) > 0 \), which is a contradiction. Then, we look for positive steady states \( (x^*, S^*, R^*) \) with \( R^* > 0 \). If \( R^* > 0 \), then \( r_{of}(S^*, R^*) > 0 \). Thus, we study the following system of equations:

\[
\begin{align*}
0 &= (Y_S - kY_R) r_S(S, R) + kY_R r_{S0} - D, \\
0 &= D(S_{in} - S) - r_S(S, R)x, \quad (9) \\
0 &= -DR + k[r_S(S, R) - r_{S0}]x.
\end{align*}
\]

From the two first equations in (10), we obtain that:

\[
S = S_{in} - \beta_S x, \quad \beta_S := \frac{1}{D} \frac{D - kY_R r_{S0}}{Y_S - kY_R} > 0. \quad (10)
\]
Combining the three equations in (10), we obtain that $x + Y_S S + Y_R R = Y_S S_{in}$ (conservation principle, see (6)). Combining this equation with (10), we obtain:

$$R = \beta_R x, \quad \beta_R := \frac{kY_R D - Y_S r_{SO}}{D Y_S - kY_R} > 0. \quad (11)$$

Combining (10) and (11) with the first equation in (10), we obtain the following equation for $x$:

$$\left( Y_S - kY_R \right) r_S(S_{in} - \beta_Sx, \beta_R x) + kY_R r_{SO} - D = 0. \quad (12)$$

Since $f$ is strictly decreasing, $f(0) = \mu(S_{in}, 0) - D$ and $f(S_{in}/\beta_S) = kY_R r_{SO} - D < Y_S r_{SO} - D < 0$, we conclude that (12) admits a unique solution $x^* \in (0, S_{in}/\beta_S)$ if $\mu(S_{in}, 0) > D$, and has no positive solution if $\mu(S_{in}, 0) \leq D$.

The following result shows that the equilibrium with presence of microorganisms is locally stable when $D \neq Y_S r_{SO}$.

**Proposition 2** (Local stability of $E^*$) Assume that $D < \mu(S_{in}, 0)$ and let $E^*$ be the non-trivial equilibrium given by Proposition 1. If $D \neq Y_S r_{SO}$, then $E^*$ is locally stable.

**Proof** If $D > Y_S r_{SO}$, according to Proposition 1, $R^* > 0$ and $r_{of}(S^*, R^*) > 0$. Thus, we can study the local stability of $E^*$ in the following system:

$$\begin{align*}
\frac{dx}{dt} &= [(Y_S - kY_R) r_S(S, R) + kY_R r_{SO} - D]x, \\
\frac{dS}{dt} &= D(S_{in} - S) - r_S(S, R)x, \\
\frac{dR}{dt} &= -DR + k[r_S(S, R) - r_{SO}]x.
\end{align*} \quad (13)$$

Set $\alpha := Y_S - kY_R$. The change of variables $U = x + \alpha S$ and $W = x + Y_S S + Y_R R$ leads (14) to:

$$\begin{align*}
\frac{dx}{dt} &= [\alpha r_S(x, U, W) + kY_R r_{SO} - D]x, \\
\frac{dU}{dt} &= D(\alpha S_{in} - U) + kY_R r_{SO} x, \\
\frac{dW}{dt} &= D(Y_S S_{in} - W).
\end{align*} \quad (14)$$

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with
\[
\hat{r}_S(x, U, W) = r_S\left(\frac{1}{\alpha}(U - x), \frac{1}{Y_R}\left(W - \frac{Y}{\alpha}U\right) + \frac{k}{\alpha}\right). \tag{15}
\]

The Jacobian matrix associated with (15) and evaluated at \(E^*\) is:
\[
J_1 := \begin{bmatrix}
\alpha x^* \frac{\partial \hat{r}_S}{\partial x} & \alpha x^* \frac{\partial \hat{r}_S}{\partial U} & \alpha x^* \frac{\partial \hat{r}_S}{\partial W} \\
kY_Rr_S & -D & 0 \\
0 & 0 & -D
\end{bmatrix}.
\]

It is clear that one eigenvalue of \(J\) is \(-D\). The other two eigenvalues are those of the matrix:
\[
J_1' := \begin{bmatrix}
\alpha x^* \frac{\partial \hat{r}_S}{\partial x} & \alpha x^* \frac{\partial \hat{r}_S}{\partial U} \\
kY_Rr_S & -D
\end{bmatrix}.
\]

We note that:
\[
\frac{\partial \hat{r}_S}{\partial x} = -\frac{1}{\alpha} \frac{\partial r_S}{\partial S} + \frac{k}{\alpha} \frac{\partial r_S}{\partial R} < 0,
\]
\[
\frac{\partial \hat{r}_S}{\partial U} = \frac{1}{\alpha} \frac{\partial r_S}{\partial S} - \frac{Y}{Y_R} \frac{\partial r_S}{\partial R} > 0.
\]

Thus, it is easy to verify that \(Tr(J_1') < 0\) and that \(det(J_1') > \alpha r_{S0} x^* \frac{\partial r_S}{\partial S} > 0\). This implies that both eigenvalues of \(J_1'\) have negative real part. Thus, \(E^*\) is locally stable.

If \(0 < D < YS^{r\prime}_{S0}\), according to Proposition 1, \(R^* = 0\) and \(r_{of}(S^*, 0) = 0\). Thus, we can study the local stability of \(E^*\) in the following system:
\[
\begin{align*}
\frac{dx}{dt} &= [YSr_S(S, R) - D]x, \\
\frac{dS}{dt} &= D(S_{in} - S) - r_S(S, R)x, \\
\frac{dR}{dt} &= -DR.
\end{align*}
\tag{16}
\]

The Jacobian matrix associated with (17) and evaluated at \(E^*\) is:
\[
J_2 := \begin{bmatrix}
0 & x^* YS \frac{\partial r_S}{\partial S} & x^* YS \frac{\partial r_S}{\partial R} \\
r_S & -D - x^* \frac{\partial r_S}{\partial S} & -x^* \frac{\partial r_S}{\partial R} \\
0 & 0 & -D
\end{bmatrix}.
\]
As in the previous case, one eigenvalue of $J_2$ is $-D$. The other two eigenvalues are those of the matrix:

$$J'_2 := \begin{bmatrix} 0 & x^* Y S \frac{\partial r_S}{\partial S} \\ -r_S & -(D + x^* \frac{\partial r_S}{\partial S}) \end{bmatrix}.$$ 

It is clear that $\text{Tr}(J'_2) < 0$ and $\det(J'_2) > 0$. Hence, both eigenvalues of $J'_2$ have negative real part. Thus, $E^*$ is locally stable.  

4 Global behavior and main result

In this section, we aim to prove that if (4) admits an equilibrium with presence of microorganisms, which is unique according to Proposition 1, then any solution to (4) approaches it asymptotically, provided a positive initial population. The first result in this section shows the existence of two positively invariant sets, which will be repeatedly used in this section.

**Lemma 1 (Positively invariant sets)**

(a) The set $\Omega_1 := \{(x, S, R) \in \mathbb{R}^3_+ : x + Y_S S \leq Y_S S_{in}\}$ is positively invariant.

(b) If $D \geq Y_S r_S S_0$, the set $\Omega_2 := \Omega_1 \cap \{(x, S, R) ; r_S(S, R) \geq r_S S_0\}$ is positively invariant.

**Proof** We have that the variable $V := x + Y_S S$ satisfies the following differential equation:

$$\frac{dV}{dt} = D(Y_S S_{in} - V) - Y_R r_{of}(S, R)x.$$  \hspace{1cm} (17)

The proof of (a) follows from the fact that $\frac{dV}{dt}_{V = Y_S S_{in}} \leq 0$ and $\mathbb{R}^3_+$ is positively invariant. For (b), let us consider the variable $y = r_S(S, R)$. Then we have:

$$\frac{dy}{dt} = (D(S_{in} - S) - yx) \frac{\partial r_S(S, R)}{\partial S} + (-D R + r_{of}(S, R)x) \frac{\partial r_S(S, R)}{\partial R}.  \hspace{1cm} (18)$$

Since $\Omega_1$ is positively invariant, it is enough to show that $\frac{dy}{dt}_{y = r_S S_0} \geq 0$ whenever $(x, S, R) \in \Omega_1$. Indeed, we have

$$\frac{dy}{dt}_{y = r_S S_0} = (D(S_{in} - S) - x Y_S r_S S_0) \frac{\partial r_S(S, R)}{\partial S} - D R \frac{\partial r_S(S, R)}{\partial R}.$$ 

Since $\frac{\partial r_S(S, R)}{\partial S} > 0$, $\frac{\partial r_S(S, R)}{\partial R} < 0$, $Y_S S + x \leq Y_S S_{in}$ (inside $\Omega_1$), and $D \geq Y_S r_S S_0$, we obtain

$$\frac{dy}{dt}_{y = r_S S_0} > (Y_S r_S S_0(S_{in} - S) - r_S S_0 x) \frac{\partial r_S(S, R)}{\partial S}.$$
\[ r_{S0}(Y_S S_{in} - x - Y_S S) \frac{\partial r_{S}(S, R)}{\partial S} \geq 0. \]

This completes the proof. \qed

The following result shows that if there is no equilibrium with presence of microorganisms (i.e. \( \mu(S_{in}, 0) \leq D \)), then the population goes to extinction.

**Proposition 3 (Extinction)** Let \( E_0 \) be given by (8). If \( \mu(S_{in}, 0) \leq D \), then any solution of (4) approaches \( E_0 \) asymptotically.

**Proof** Let \((\tilde{x}, \tilde{S}, \tilde{R})\) be a solution of (4), and let \( \tilde{V} = \tilde{x} + Y_S \tilde{S} \). We have that
\[
\frac{d\tilde{V}}{dt} = D(Y_S S_{in} - \tilde{V}) - r_{of}(\tilde{S}, \tilde{R})\tilde{x}.
\]

We can easily verify that:
\[
\frac{d\tilde{x}}{dt} \leq f_1(\tilde{x}, \tilde{V}) := \left[ \mu \left( \frac{\tilde{V} - \tilde{x}}{Y_S}, 0 \right) - D \right] \tilde{x},
\]
\[
\frac{d\tilde{V}}{dt} \leq f_2(\tilde{x}, \tilde{V}) := D(Y_S S_{in} - \tilde{V}).
\]

Now, let \((\hat{x}, \hat{V})\) be the unique solution of
\[
\begin{align*}
\frac{dx}{dt} &= f_1(x, V), \\
\frac{dV}{dt} &= f_2(x, V),
\end{align*}
\]

satisfying \( \hat{x}(0) = \hat{x}(0) \) and \( \hat{V}(0) = \hat{V}(0) \). We note that (19) is cooperative i.e. \( x \mapsto f_2(x, V) \) and \( V \mapsto f_1(x, V) \) are increasing (Smith and Waltman 1995). Then, applying Theorem B.1 from Appendix B in Smith and Waltman (1995), we conclude that
\[
\hat{x}(t) \leq \tilde{x}(t) \text{ and } \hat{V}(t) \leq \tilde{V}(t) \text{ for all } t \geq 0.
\]

Again, due to the cooperativity of (19) we conclude that \((\hat{x}, \hat{V})\) approaches an equilibrium asymptotically. Since the unique equilibrium of (19) is \((0, Y_S S_{in})\), we conclude that \((\hat{x}, \hat{V})\) approaches \((0, Y_S S_{in})\) asymptotically. From (20), \( \tilde{x} \) approaches 0 asymptotically. Now, noting that \( \tilde{S}(t) \leq \tilde{V}(t)/Y_S \) for all \( t \geq 0 \), we have that
\[
\frac{d\tilde{V}}{dt} \geq g(t, \tilde{V}) := D(Y_S S_{in} - \tilde{V}) - Y_R r_{of}(\tilde{V}/Y_S, 0)\tilde{x}(t).
\]
Let $V$ be the unique solution of:

$$\frac{dV}{dt} = g(t, V),$$

(22)
satisfying $V(0) = \tilde{V}(0)$. Thus, by a comparison theorem argument, we conclude that $V(t) \leq \tilde{V}(t)$ for all $t \geq 0$. Let us define $g_0(V) := D(Y_S S_{in} - V)$. Since $|g(t, V) - g_0(V)| = Y_R r_{of}(V, 0)\tilde{x}(t) \to 0$ as $t \to \infty$, we can apply Theorem 1.2 in Thieme (1992) and conclude that $V$ approaches $Y_S S_{in}$ asymptotically. Now, since $V(t) \leq \tilde{V}(t) \leq \tilde{V}$ for all $t \geq 0$, we conclude that $\tilde{V}$ approaches $Y_S S_{in}$ asymptotically. Finally, consider the variable $\tilde{W} = \tilde{x} + Y_S \tilde{S} + Y_R \tilde{R}$. In view of (6), $\tilde{W}$ converges to $Y_S S_{in}$. Consequently, $\tilde{R} = \frac{\tilde{W} - \tilde{V}}{Y_R}$ converges to 0, and the proof is complete.

In view of (6), the solutions of (4) approach the hyperplane:

$$\Omega := \{(x, S, R) \in \mathbb{R}_+^3; x + Y_S S + Y_R R = Y_S S_{in}\}$$

(23)
The set $\Omega$ is positively invariant with respect to (4). This implies that the dynamics of solutions starting in $\Omega$ correspond to that of a two-dimensional system. The following two results describe the dynamics of any solution starting in $\Omega$.

**Lemma 2** If $\mu(S_{in}, 0) > D$ and $D > Y_S r_{S0}$, for any solution $(x, S, R)$ to (4) starting on $\Omega$ with $x(0) > 0$, there exists $t'$ such that $r_{of}(S(t), R(t)) \geq 0$ for all $t \geq t'$.

**Proof** Let $(\tilde{x}, \tilde{S}, \tilde{R})$ be a solution of (4) with $\tilde{x}(0) > 0$ and $(\tilde{x}(0), \tilde{S}(0), \tilde{R}(0)) \in \Omega$, and let $\Omega_1$ and $\Omega_2$ be the positively invariant sets defined in Lemma 1. Since $(\tilde{x}, \tilde{S}, \tilde{R})$ starts in $\Omega$, we have that $\tilde{x}(t) + Y_S \tilde{S}(t) = Y_S S_{in} - Y_R \tilde{R}(t) \leq Y_S S_{in}$ for all $t \geq 0$. Hence, $(\tilde{x}(t), \tilde{S}(t), \tilde{R}(t)) \in \Omega_1$ for all $t \geq 0$. Now, we claim the existence of $t' > 0$ such that $r_S(\tilde{S}(t'), \tilde{R}(t')) \geq r_{S0}$. Indeed, by contradiction, let us assume that $r_S(\tilde{S}(t), \tilde{R}(t)) < r_{S0}$ for all $t \geq 0$. Consider the variable $\tilde{V} = \tilde{x} + Y_S \tilde{S}$. Then $(\tilde{x}, \tilde{V})$ is a solution of the following system:

$$\begin{align*}
\frac{dx}{dt} &= (Y_S r_S \left(\frac{V - x}{Y_S}, \frac{Y_S S_{in} - V}{Y_R}\right) - D)x, \\
\frac{dV}{dt} &= D(Y_S S_{in} - V).
\end{align*}$$

(24)
The planar system (24) is cooperative, then $(\tilde{x}, \tilde{V})$ approaches a steady state asymptotically. Let $E^* = (x^*, S^*, R^*)$ be given by Proposition 1. System (24) admits two equilibria, $F_0 = (0, Y_S S_{in})$ and $F^* = (x^*, Y_S S_{in})$. Thus, $r_S(\tilde{S}(t), \tilde{R}(t))$ approaches either $r_S(S_{in}, 0) (> r_{S0})$ or $D/Y_S (> r_{S0})$. This implies that $(\tilde{x}, \tilde{S}, \tilde{R})$ enters $\Omega_2$ which contradicts our hypothesis. Then, there is $t' > 0$ such that $r_S(\tilde{S}(t'), \tilde{R}(t')) > r_{S0}$ i.e. $(\tilde{x}(t'), \tilde{S}(t'), \tilde{R}(t')) \in \Omega_2$. Since $\Omega_2$ is positively invariant, the proof is complete. □

The following result describes the global behavior of solutions of (4) starting on $\Omega$.
**Proposition 4** Assume that $\mu(S_{in}, 0) > D$, and let $E^*$ and $E_0$ be the equilibria given by Proposition 1 and (8) respectively. Then, for any solution $(x, S, R)$ to (4) starting on $\Omega$ we have

(a) if $x(0) > 0$, then $(x(t), S(t), R(t)) \to E^*$ as $t \to \infty$,
(b) if $x(0) = 0$, then $(x(t), S(t), R(t)) \to E_0$ as $t \to \infty$.

**Proof** Let $(\tilde{x}, \tilde{S}, \tilde{R})$ be a solution of (4) with $(\tilde{x}(0), \tilde{S}(0), \tilde{R}(0)) \in \Omega$. To prove (a), let us assume that $\tilde{x}(0) > 0$. It is not difficult to see that $(\tilde{x}, \tilde{R})$ is a solution of the following system:

\[
\begin{align*}
\frac{dx}{dt} &= (\mu(\varphi_S(x, R), R) - D)x, \\
\frac{dR}{dt} &= -DR + r_{of}(\varphi_S(x, R), R)x,
\end{align*}
\]

with $\varphi_S(x, R) = S_{in} - (Y_R R + x)/Y_S$. If $D \leq Y_{SR_{S0}}$, then (25) admits only two equilibria, $F_0 = (0, 0)$ and $F^* = (x^*, 0)$. Hence, (25) has no interior steady states, and consequently (in a planar system) no limit cycles. The Jacobian matrix of (25) evaluated at $F_0$ is:

\[
\begin{bmatrix}
\mu(S_{in}, 0) - D & 0 \\
0 & -D
\end{bmatrix}.
\]

Since $\mu(S_{in}, 0) - D > 0$, $F_0$ is a saddle point which can only be reached if $\tilde{x}(0) = 0$. Consequently, by the Poincaré-Bendixson theorem (Theorem 2 in Chapter 3.7 in Perko 2013), $(\tilde{x}, \tilde{R})$ approaches $F^*$ asymptotically, and hence $(\tilde{x}, \tilde{S}, \tilde{R})$ approaches $E^*$ asymptotically. Now, let us assume that $D > Y_{SR_{S0}}$. From Lemma 2, we can assume that $r_S(\tilde{S}(t), \tilde{R}(t)) \geq r_{S0}$ for all $t \geq 0$. Thus, $(\tilde{x}, \tilde{S}, \tilde{R})$ is a solution of:

\[
\begin{align*}
\frac{dx}{dt} &= ((Y_S - kY_R)r_S(S, R) + kY_R r_{S0} - D)x, \\
\frac{dS}{dt} &= D(S_{in} - S) - r_S(S, R)x, \\
0 &= Y_S(S_{in} - S) - Y_R R - x.
\end{align*}
\]

Set $\alpha := Y_S - kY_R$ and consider the variable $\tilde{U} = \tilde{x} + \alpha \tilde{S}$. Then $(\tilde{x}, \tilde{U})$ is a solution of the following system:

\[
\begin{align*}
\frac{dx}{dt} &= (\alpha r_S(\varphi_S(x, U), \varphi_R(x, U)) + kY_R r_{S0} - D)x, \\
\frac{dU}{dt} &= D(\alpha S_{in} - U) + kY_R r_{S0}x,
\end{align*}
\]

with

\[
\varphi_S(x, U) = \frac{U - x}{\alpha} \quad \text{and} \quad \varphi_R(x, U) = \frac{1}{Y_R} \left( Y_S S_{in} - \frac{Y_S U}{\alpha} \right) + \frac{kx}{\alpha}.
\]
Since (27) is a planar cooperative system, \((\tilde{x}(t), \tilde{U}(t))\) approaches either \(F^* = (x^*, x^* + \alpha S^*)\) or \(F_0 = (0, \alpha S_{in})\). The Jacobian matrix of (27) evaluated at \(F_0\) is:

\[
\begin{bmatrix}
\mu(S_{in}, 0) - D & 0 \\
kYrS & -D
\end{bmatrix}
\]

It is clear that \(F_0\) is a saddle point which can only be reached if \(\tilde{x}(0) = 0\). Consequently, as in the previous case, \((\tilde{x}, \tilde{S}, \tilde{R})\) approaches \(E^*\) asymptotically. This completes the proof of part (a). For (b), let us assume that \(\tilde{x}(0) = 0\). Then, \(\tilde{x}(t) = 0\) for all \(t \geq 0\). This implies that \(d\tilde{S}/dt = D(\tilde{S} - S_{in})\) and \(d\tilde{R}/dt = -D \tilde{R}\). Consequently \((\tilde{x}, \tilde{S}, \tilde{R})\) approaches \(E_0\) asymptotically. \(\square\)

According to Proposition 1, if \(D = YSR_{s0}\), then the positive equilibrium \(E^* = (x^*, S^*, 0)\) satisfies \(r_S(S^*, 0) = r_{s0}\) and the function \(r_{of}\) is not differentiable at \((S^*, 0)\). This poses a problem for the study of the local stability of \(E^*\), and consequently for the application of classical arguments (e.g. Butler-McGehe Theorem) to extend Proposition 4 to any initial condition. The following result considers this particular case.

**Proposition 5** Assume that \(\mu(S_{in}, 0) > D\) and let \(E^*\) be given by Proposition 1. If \(D = YSR_{s0}\), then \(E^*\) is stable.

**Proof** Let \(\xi(t) = (x(t), S(t), R(t))\) be a solution of (4) with \(x(0) > 0\) and \(S(0), R(0) \geq 0\), and let \(\Omega_1\) and \(\Omega_2\) be the sets defined in Lemma 1. Consider the following sets:

- \(\Omega_1' := \{(x, S, R) \in \mathbb{R}_+^3 : x + YSS > YSS_{in}\}\),
- \(\Omega_2' := \{(x, S, R) \in \mathbb{R}_+^3 : r_S(S, R) < r_{s0}\} \cap \Omega_1\).

Given sets \(A, B \in \{\Omega_1', \Omega_2', \Omega_2\}\), we will say that \(\xi\) moves from \(A\) to \(B\), if there are \(t' \geq 0\) and \(\tau > 0\) such that \(\xi(t) \in A\) for all \(t \in (t' - \tau, t')\), \(\xi(t') \in A \cup B\), and \(\xi(t) \in B - A\) for all \(t \in (t', t' + \tau)\). This means that if \(\xi\) moves from \(A\) to \(B\), then there is a time when \(\xi\) is in \(A\) and then later is in \(B\) but not in \(A\). Since \(\Omega_1\) and \(\Omega_2\) are positively invariant (see Lemma 1), \(\xi\) can only move from \(\Omega_1'\) to \(\Omega_2'\), from \(\Omega_1'\) to \(\Omega_2\), or from \(\Omega_2'\) to \(\Omega_2\). Hence, \(\xi\) has one of the following global behaviors:

(a) \(\xi(t) \in A\) for all \(t \geq 0\) with \(A \in \{\Omega_1', \Omega_2', \Omega_2\}\),
(b) \(\xi\) starts on \(\Omega_1'\) and moves either to \(\Omega_2'\) or to \(\Omega_2\),
(c) \(\xi\) starts on \(\Omega_2'\) and moves to \(\Omega_2\),
(d) \(\xi\) starts on \(\Omega_1'\), then moves to \(\Omega_2'\), and then to \(\Omega_2\).

Let \(\epsilon > 0\) be given. We have to prove the existence of a \(\delta > 0\) such that in any situation listed above, if \(||\xi(0) - E^*|| < \delta\) then \(||\xi(t) - E^*|| < \epsilon\) for all \(t \geq 0\). We only give the proof in the situation d) because the proof in the other situations is almost the same. Thus, let us assume the existence of \(t_1, t_2 > 0\) such that \(t_1 < t_2\) and \(\xi(t) \in \Omega_1'\) for all \(t \in [0, t_1]\), \(\xi(t) \in \Omega_2'\) for all \(t \in [t_1, t_2]\), and \(\xi(t) \in \Omega_2\) for all \(t \geq t_2\). For all \(t \geq t_2\), \(\xi(t)\) can be seen as a solution of (14). In such a case, we can study the Jacobian matrix of (14) evaluated at \(E^*\) (as done in the proof of Proposition 2) to conclude the existence of \(\delta_2 > 0\) such that \(||\xi(t) - E^*|| < \epsilon\) for all \(t \geq t_2\) provided \(||\xi(t_2) - E^*|| < \delta_2\). Now
for all $t \in [t_1, t_2)$, $\xi(t)$ can be seen as a solution of (17). In such a case, we can study the Jacobian matrix of (17) evaluated at $E^*$ to conclude the existence of $\delta_1 > 0$ such that \( |\xi(t) - E^*| < \delta_2/2 \) for all $t \in [t_1, t_2)$ provided \( |\xi(t_1) - E^*| < \delta_1 \). Finally, for all $t \in [0, t_1)$, consider the variables $V = x + Y_S S$ and $W = x + Y_S S + Y_R R$. It is clear that:

\[
\frac{dV}{dt} = D(Y_S S_{in} - V) - Y_R r_{of}(S, R)x \leq D(Y_S S_{in} - V). \tag{28}
\]

Using the definition of $\Omega'$ and (28) we obtain:

\[
0 \leq V(t) - Y_S S_{in} \leq V(0) - Y_S S_{in}. \tag{29}
\]

Again, using the definition of $\Omega'$ and (7), we obtain that:

\[
0 \leq W(t) - Y_S S_{in} \leq W(0) - Y_S S_{in}. \tag{30}
\]

Since $S = (V - x)/Y_S$ and $R = (W - V)/Y_R$, we have that:

\[
xh(V(0) - Y_S S_{in}, x) \leq \frac{dx}{dt} \leq xg(V(0) - Y_S S_{in}, x),
\]

with $g(v, x) = \mu(S_{in} + v/Y_S - x/Y_S, 0) - D$ and $h(v, x) = \mu(S_{in} - x/Y_S, v/Y_R) - D$. We note that $g$ is strictly increasing in $v$ and strictly decreasing in $x$ and that $g(0, 0) > 0$. Moreover, for $M(v) := v + Y_S S_{in}$ we have $g(v, M(v)) = -D < 0$. Similarly, $h$ is strictly decreasing in both, $v$ and $x$. It is also clear that $h(0, 0) > 0$ and $h(v, Y_S S_{in}) = -D < 0$. Applying Lemmas 3 and 4 in the Appendix, for any $\epsilon' > 0$ there is $v_{e'}$ such that $|x(t) - x^*| < \epsilon'$ provided $|x(0) - x^*| < \epsilon'/2$, $V(0) - Y_S S_{in} < v_{e'}$, and $W(0) - Y_S S_{in} < v_{e'}$. Thus, from (29) and (30), we conclude that $|x(t) - x^*| < \epsilon'$, $|V(t) - Y_S S_{in}| < \epsilon'$ and $|W(t) - Y_S S_{in}| < \epsilon'$ for all $t \in [0, t_1]$, provided

\[
|x(0) - x^*| < \epsilon'/2, \text{ and } |V(0) - Y_S S_{in}|, |W(0) - Y_S S_{in}| < \min\{v_{e'}, \epsilon'\}.
\]

Choosing an appropriate $\epsilon'$, and writing $S$ and $R$ in terms of $x$, $V$, and $W$, we can find $\delta(\epsilon')$ such that $|\xi(t) - E^*| < \delta_1/2$ for all $t \in [0, t_1)$ provided $|\xi(t) - E^*| < \delta(\epsilon')$. Since $\xi$ is continuous, we conclude that $|\xi(t) - E^*| < \epsilon$ for all $t \geq 0$ provided $|\xi(t) - E^*| < \delta(\epsilon')$. \hfill \Box

**Theorem 1 (Main result)** Let $E^*$ and $E_0$ be the equilibria given by Proposition 1 and (8) respectively. We have:

(a) If $\mu(S_{in}, 0) > D$, then $E^*$ is globally asymptotically stable on $(0, \infty) \times \mathbb{R}_+^2$.

(b) If $\mu(S_{in}, 0) \leq D$, then any solution to (4) approaches $E_0$ asymptotically.

**Proof** Part (b) follows directly from Proposition 3. For (a), let $(x, S, R)$ be a solution of (4) with $x(0) > 0$, $S(0)$, $R(0) \geq 0$. Let us write $P = (x(0), S(0), R(0))$. In view of (7), we have that $\omega(P) \subset \Omega$, where $\omega(P)$ denotes the $\omega$-limit set of $P$ and $\Omega$ is \[ Springer \]
defined in (23). From Proposition 4, the \( \omega \)-limit set of any trajectory passing through \( \Omega \) is either \( E_0 \) or \( E^* \). Consequently,

\[
\omega(P) \cap \{E_0, E^*\} \neq \emptyset.
\]  

(31)

The Jacobian matrix associated with (4) and evaluated at \( E_0 \) is:

\[
J := \begin{bmatrix}
\mu(S_{in}, 0) - D & 0 & 0 \\
-r_S(S_{in}, 0) & -D & 0 \\
r_{of}(S_{in}, 0) & 0 & -D
\end{bmatrix}.
\]

It is clear that \( J \) has two negative eigenvalues and one positive eigenvalue. Let \( \Omega_0 \) be the two-dimensional subspace spanned by the eigenvectors corresponding to the negative eigenvalues i.e. \( \Omega_0 := \{0\} \times \mathbb{R}^2 \). It is clear that \( \Omega_0 \) is positively invariant and that any solution starting on \( \Omega_0 \) approaches \( E_0 \) asymptotically. Since \( \Omega_0 \) is a manifold trivially tangent to \( \Omega_0 \) at 0, we conclude that \( \Omega_0 \) is the stable (global) manifold of (4) at \( E_0 \) (see Chapter 2.7 in Perko 2013). Since \( P \notin \Omega_0 \), we have that \( \omega(P) \neq \{E_0\} \). Now, let us assume that \( E_0 \in \omega(P) \). According to the Theorem of Butler-McGehee (see for example page 12 in Smith and Waltman 1995), \( \omega(P) \) intersects \( \Omega_0 \) in a point other than \( E_0 \). The (whole) trajectory of that point, say \((0, S_0, R_0)\), is given by

\[
\gamma(t) = (0, S_0 e^{-Dt} + Y_S S_{in} (1 - e^{-Dt}), R_0 e^{-Dt}), \quad t \in \mathbb{R}.
\]

It is clear that \( \gamma \) is unbounded (as \( t \to -\infty \)). Consequently, \( \omega(P) \) contains an unbounded trajectory. However, \( \omega(P) \) is a bounded set because the solutions to (4) are ultimately bounded (see Lemma 3.1.2 in Hale 2010). This contradiction implies that \( E_0 \) cannot be in \( \omega(P) \). Hence, from (31), we conclude that \( E^* \in \omega(P) \). From Propositions 2 and 5 we have that \( E^* \) is stable, hence \( \omega(P) = \{E^*\} \). This completes the proof.

\[\square\]

5 Discussion

5.1 Summary of our mathematical results: survival, extinction, and stability

The chemostat with overflow metabolism, described by (4), admits at most two equilibria. An extinction equilibrium, denoted by \( E_0 = (0, S_{in}, 0) \), that corresponds to the absence of microorganisms and always exists. The other possible equilibrium, denoted by \( E^* = (x^*, S^*, R^*) \), is characterized by the presence of microorganism i.e. \( x^* > 0 \). Our main result (Theorem 1), states that if \( E^* \) exists, then any solution to (4) with a positive initial population approaches (asymptotically) \( E^* \). That is, given a solution \((x(t), S(t), R(t))\) of (4) with \( x(0) > 0 \), we have that

\[
\lim_{t \to \infty} x(t) = x^*, \quad \lim_{t \to \infty} S(t) = S^*, \quad \text{and} \quad \lim_{t \to \infty} R(t) = R^*.
\]
On the other hand, the non-existence of $E^*$ implies that any solution to (4) approaches the extinction equilibrium asymptotically: meaning that $\lim_{t \to \infty} x(t) = 0$. Proposition 1 in Sect. 3 gives necessary and sufficient conditions for the existence of $E^*$. Indeed, $E^*$ exists if and only if $\mu(S_{in}, 0) > D$. The survival of microorganisms (existence of $E^*$) does not ensure the presence of the overflow metabolism by-product in the medium. According to Proposition 1, $R^* > 0$ if and only if $Y_{SR} < D < \mu(S_{in}, 0)$.

Overflow metabolism, and the consequent presence of a by-product, does not generate multistability. That is, if $E^*$ exists, there are no solutions with positive initial population converging to $E_0$. Xiu et al. (1998) observed the multiplicity of stable steady states. However, apart from taking $r_{S0} = 0$, they assume that excess of substrate inhibits the growth rate. Thus, the existence of multiple steady states is due to substrate inhibition and not to overflow metabolism.

5.2 Acetate formation and productivity in *E. coli* cultures

In *E. coli* cultures, the by-product corresponds to acetate. According to Proposition 1, the presence of acetate in the non-trivial equilibrium $E^*$ depends on the dilution rate. This is illustrated in Fig. 3a. Indeed, in presence of bacteria, $R^* > 0$ if and only if $D > r_{S0} Y_S$. This relation between the acetate steady state concentration and the dilution rate has been observed experimentally by El-Mansi and Holms (1989). This may suggest an optimal operation of the chemostat at dilution rates lower than $Y_{SR} Y_S$ to avoid the presence of acetate in the culture. Indeed, different authors have shown that preventing acetate formation in fed-batch leads to higher density cultures (Korz et al. 1995; Babu et al. 2000).

To evaluate this strategy in chemostat cultures, let us consider the (steady state) productivity defined as $P^* = D x^*$, with $x^*$ the steady state concentration of bacteria at the dilution rate $D$. $P^*$ quantifies the biomass that is produced per unit of time at steady state. To determine $P^*$ numerically, let us assume that $r_S$ is given by (2), and consider the parameters estimated by Mauri et al. (2020) (see Table 1). Figure 3b shows that the steady state productivity is maximal at a value of the dilution rate higher than $Y_{SR} Y_S$ (continuous line). This suggest that preventing acetate formation is not a good strategy in chemostat cultures, in contrast to fed-batch cultures. The veracity of this observation depends on the choice of parameters. For instance, for low values of $K_{i,R}$ (strong inhibition), the maximal productivity is reached at $D = Y_{SR} Y_S$ (see dashed line in Fig. 3).

As shown in Fig. 3b (continuous line), maximal productivity of the system is accompanied by the secretion of acetate. A natural strategy to increase this maximal productivity is removing acetate from the culture during fermentation. This can be done with a dialysis reactor (Nakano et al. 1997), or with macroporous ion-exchange resins (Huang et al. 2012). However, these methods tend to remove nutrients that are necessary for cell growth. A promising alternative consists in introducing an additional *E. coli* strain (a cleaner), which has been metabolically engineered to consume acetate. Thus, two different *E. coli* populations coexist in the culture: one producing biomass, and one reducing the presence of acetate. Experimental results have shown an increase of the productivity with this strategy (Bernstein et al. 2012). A few math-
Acetate concentration (a) and productivity (b) evaluated at steady state for different dilution rates. The function $r_S$ is taken as in (2). The continuous line is obtained with the parameters from Table 1. The dashed line is obtained with the parameters from Table 1 after replacing the value of $K_{i,R}$ by 0.052 g/L.

**Table 1** Kinetic parameters and yield coefficients taken from Mauri et al. (2020). DW stands for dry weight

| Parameter   | Value | Unit     |
|-------------|-------|----------|
| $r_{S,\text{max}}$ | 1.53   | h\(^{-1}\) |
| $K_S$       | 0.09   | g/L      |
| $K_{i,R}$   | 0.52   | g/L      |
| $k$         | 0.17   | –        |
| $r_{S0}$    | 0.7    | h\(^{-1}\) |
| $Y_S$       | 0.44   | gDW/gS   |
| $Y_R$       | 0.3    | gDW/gR   |

Mathematical works have studied the dynamics of such communities (Heßeler et al. 2006; Harvey et al. 2014). However, as mentioned in the introduction, the authors assume that overflow metabolism always occur (i.e. $r_{S0} = 0$). Thus, our results give a basis to understand the dynamics of such microbial communities when $r_{S0} > 0$.

### 5.3 Recombinant protein production

Following Mauri et al. (2020), and using the notation of this paper, the dynamics of a recombinant protein, which concentration is denoted by $H$, follows from:

\[
\begin{align*}
\frac{dH}{dt} &= Y_H\mu(S, R)x - DH, \\
\frac{dx}{dt} &= [(1 - Y_H)\mu(S, R) - D]x, \\
\frac{dS}{dt} &= D(S_{in} - S) - r_S(S, R)x, \\
\frac{dR}{dt} &= -DR + r_{of}(S, R)x.
\end{align*}
\]
Here, $Y_H$ is the protein yield coefficient representing the carbon diversion to protein production. Let $(H, x, S, R)$ be a solution of (32) with $x(0) > 0$, $H(0), S(0), R(0) \geq 0$. The dynamics of $(x, S, R)$ is independent of $H$ and can be described by Theorem 1. Indeed, if $(1 - Y_H)\mu(S_{in}, 0) > D$, then there is $x^* > 0$ such that $\lim_{t \to \infty} x(t) = x^*$. Now, it is easy to verify that the variable $y := \frac{Y_H (1 - Y_H) x - H}{1 - Y_H}$ satisfies $\frac{dy}{dt} = -Dy$. Therefore, $\lim_{t \to \infty} y(t) = 0$, which implies that $\lim_{t \to \infty} H(t) = \frac{Y_H x^*}{1 - Y_H}$. Thus, we define the steady state protein productivity as:

$$P_H^* = D \frac{Y_H (1 - Y_H) x^*}{1 - Y_H}.$$  (33)

Note that the value of $x^*$ depends on the values of $Y_H$ and $D$ and that $P_H^*$ only exists if $0 < D < (1 - Y_H)\mu(S_{in}, 0)$. These results allow to illustrate the impact of $Y_H$ on the protein productivity. If $Y_H = 0$, there is no production of $H$, and consequently $P_H^* = 0$. On the other hand, if $Y_H$ approaches 1, it can be shown that $P_H^*$ approaches 0. Indeed, using the restriction over $D$ we obtain $P_H^* < \mu(S_{in}, 0)Y_H x^*$, where it is clear that $\lim_{Y_H \to 1} x^* = 0$. This shows the existence of an intermediate value of $Y_H$ maximizing $P_H^*$. Now, for each value of $Y_H \in [0, 1)$ we compute the maximal productivity with respect to the dilution rate i.e. $\max\{P : 0 < D \leq (1 - Y_H)\mu(S_{in}, 0)\}$. These results are depicted in Fig. 4. We observe that the optimal value of $Y_H$ is 0.505, suggesting that protein productivity is maximal ($0.373 \text{ g L}^{-1} \text{ d}^{-1}$) when 50% of the absorbed substrate, that is not excreted in form of acetate, is diverted into protein production.

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2 Set $\mu'(\cdot) = (1 - Y_H)\mu(\cdot)$, and note that $\mu'(\cdot) = Y'_S S(\cdot) - Y'_R \rho_f(\cdot)$ with $Y'_S = (1 - Y_H)Y_S$ and $Y'_R = (1 - Y_H)Y_R$. Thus, Theorem 1 applies directly to (32) when replacing $(1 - Y_H)\mu(\cdot)$ by $\mu'(\cdot)$.

3 From (32), the intuition says that $x^*$ approaches 0 as $Y_H$ approaches 1. This can be proved determining explicitly $x^*$ when $D \leq Y_S (1 - Y_H)$ and using the upper bound for $x^*$ given in the proof of Proposition 1 when $D > Y_S (1 - Y_H)$ (last paragraph).
Appendix

We present two results on differential inequalities. We only give the proof of the first result since both proofs are very similar. Consider the following differential equation:

$$\frac{dy}{dt} = f(y, u)y,$$  \hfill (34)

with \( f(y, u) : \mathbb{R}_+^2 \rightarrow \mathbb{R} \) a continuous function such that (34) admits a unique solution for all \( t, u \in \mathbb{R}_+ \) and any initial condition \( y(0) = y_0 \geq 0 \).

Lemma 3 Consider (34) and let \( x : \mathbb{R}_+ \rightarrow \mathbb{R}_+ \) be a function satisfying

$$\frac{dx(t)}{dt} \leq f(x(t), u)x(t),$$  \hfill (35)

for all \( t, u \in \mathbb{R}_+ \). Assume that \( f(y, u) \) is strictly decreasing in \( y \) and strictly increasing in \( u \). Moreover, assume that \( f(0, 0) > 0 \) and that for any \( u \in \mathbb{R}_+ \) there is \( M = M(u) > 0 \) such that \( f(M, u) < 0 \). Then there is a unique \( x^* > 0 \) such that \( f(x^*, 0) = 0 \), and for any \( \epsilon > 0 \) there is \( u_\epsilon \) such that \( x(t) - x^* < \epsilon \) for all \( t \geq 0 \) provided \( x(0) - x^* < \epsilon/2 \) and \( u < u_\epsilon \).

**Proof** For any \( u \geq 0 \) we have \( f(0, u) \geq f(0, 0) > 0 \) and \( f(M, u) < 0 \). Then, for any \( u \geq 0 \) there is a positive real number \( a = a(u) \) such that \( f(a(u), u) = 0 \). This proves the existence of \( x^* := a(0) \). Now note that \( \lim_{u \to 0^+} a(u) = x^* \) and that \( a(u) \) increases with respect to \( u \). This implies that for a given \( \epsilon > 0 \), there is \( u_\epsilon > 0 \) such that \( a(u_\epsilon) < x^* + \epsilon/2 \). Let \( \tilde{y}(t) \) be the solution to

$$\frac{dy}{dt} = f(y, u_\epsilon),$$  \hfill (36)

with \( \tilde{y}(0) = a(u_\epsilon) + \epsilon/2 \). Since (36) is an autonomous differential equation, \( a(u_\epsilon) \) is the unique positive equilibrium of (36), and \( f(\tilde{y}(0), u_\epsilon) < 0 \) we conclude that \( \tilde{y}(t) \leq a(u_\epsilon) + \epsilon/2 < x^* + \epsilon \) for all \( t \geq 0 \). Now, if \( x(0) < x^* + \epsilon/2 \), then \( x(0) < a(u_\epsilon) + \epsilon/2 = \tilde{x}(0) \). In view of (37) Applying Theorem B1 in Smith and Waltman (1995), we conclude that \( x(t) < \tilde{x}(t) < x^* + \epsilon \) for all \( t \geq 0 \). \( \square \)

Lemma 4 Consider (34) and let \( x : \mathbb{R}_+ \rightarrow \mathbb{R}_+ \) be a function -satisfying

$$\frac{dx(t)}{dt} \geq f(x(t), u)x(t),$$  \hfill (37)

for all \( t, u \in \mathbb{R}_+ \). Assume that \( f(y, u) \) is strictly decreasing in both, \( y \) and \( u \). Moreover, assume that \( f(0, 0) > 0 \) and that for any \( u \in \mathbb{R}_+ \) there is \( M = M(u) > 0 \) such that \( f(M, u) < 0 \). Then there is a unique \( x^* > 0 \) such that \( f(x^*, 0) = 0 \) and for any \( \epsilon > 0 \) there is \( u_\epsilon \) such that \( x^* - x(t) < \epsilon \) for all \( t \geq 0 \) provided \( x^* - x(0) < \epsilon/2 \) and \( u < u_\epsilon \).
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