Utilization of Soluble Starch by Oleaginous Red Yeast

*Rhodotorula glutinis*

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ABSTRACT

Starch containing wastewaters from the food and feed industry have been identified as potential cheap carbon sources for the production of microbial lipids. Due to its high potential lipid content the oleaginous yeast *Rhodotorula glutinis* is often used for fermentations in this field. Moreover it is investigated in the context of microbial carotenoid production, which also requires a cheap source of carbon. Thus, the ability of *R. glutinis* (ATCC 15125™) to degrade and utilize soluble starch for the production of lipids has been assessed in this study. While glucose and fructose were readily consumed from the medium, starch was only slightly reduced in one treatment. The yield of fatty acid methyl esters (FAME) was graduated corresponding to the initial sugar contents, with the highest FAME yield (1.5 g·L⁻¹) at the highest initial sugar content. In the treatment that contained starch as single carbon source, no FAME production was realized. Accordingly, if starchy wastewaters should be used for microbial cultivation with *R. glutinis*, an enzymatic or chemical pretreatment for starch hydrolysis should be applied, to increase the availability of this carbon source.

Keywords: *Rhodotorula glutinis*; Microbial Lipid Production; Starch

1. Introduction

Worldwide, renewable biofuels such as bioethanol and biodiesel contribute a significant share to the overall fuel consumption [1]. Against the background of rising crude oil prices and environmental concerns related with the exploitation of fossil oil resources, politics set strong incentives to further increase the production of biofuels—in Europe mainly biodiesel. So-called 1st generation biodiesel is produced by the transesterification of plant oils like rapeseed and soybean oil with an alcohol—mostly methanol—into fatty acid alkyl (methyl) esters (FAMEs) [2], whereas the terms “FAME” and “biodiesel” are often used synonymously. However, the 1st generation biodiesel derived from agricultural oil crops has been facing increasing criticism in terms of its impact on food prices, climate and ecosystems, so that the question was raised if 1st generation biofuels should actually be further promoted as sustainable energy alternative in the future. Accordingly, research and industry strived to find alternative oil sources for the transesterification into biodiesel. In this context the utilization of oils produced by oleaginous microorganisms has been identified and investigated as one possible approach. These organisms are able to accumulate significant amounts of lipids inside their cells, some as high as 80% of their respective cell dry weight, with similar characteristics as plant oil, when cultivated under nutrient limited and carbon excess conditions [3,4]. In terms of microbial lipid production, the red yeast *Rhodotorula glutinis* has been studied due to its high potential oil content of up to 72% [4]. It has also been investigated for its ability to produce high value carotenoids, namely β-carotene, torulene and torularhodin, which can be utilized as natural colorants or as ingredients of pharmaceutical products due to their antioxidant and pro-vitamin A properties. For both production pathways the cost of the carbon source has been identified as one of the main cost factors. Thus, the utilization of a cheap carbon source could significantly enhance the economic features of both approaches. Hence, a broad range of potential substrates have been tested for 1) lipid production, ranging from industrial by-products like glycerol [5] and molasses [6] over hydrolyzates of agricultural residues [7] to original waste products like municipal wastewater [8] and 2) carotenoid production, including flour extracts, grape must [9], radish brine [10] and whey [11]. Also wastewaters from the food industry with supposedly high organic contents have gained attention [12,13]. A lot of these substrates contain starch as a
principal carbon source. These starch containing waste-
waters bear a large potential as cheap carbon sources for
the microbial lipid production, if the respective microor-
ganisms are able to utilize starch as rather complex po-
lysaccharide in comparison to easily available monosac-
charides. Some molds have shown good results when
cultivated on starch as main carbon source [14,15], where-
as it has to our knowledge not yet clearly been stated if R.
glutinis is actually capable of degrading and utilizing
starch as carbon source. Rubio et al. [16] found that
strains of R. glutinis did not grow on starch and also its
poor performance on starchy wastewaters from potato
processing [17] suggest, that starch should be hydrolyzed
prior to cultivation [18]. In contrast to these findings,
Bhosale and Gadre [19] observed cell growth and caro-
tenoid production by R. glutinis on starch as carbon
source. Elsewhere [20,21] it is stated that starch assimila-
ton of R. glutinis is strain specific. Thus, for the present
study R. glutinis (ATCC 15125TM) was cultivated on
starch and glucose as carbon sources and the production
of biomass and microbial lipids was determined in course
of the cultivation to evaluate the suitability of starch as
carbon source for this specific yeast strain.

2. Materials and Methods

Three treatments were designed which differed in the
type and composition of the carbon source used. Apart
from this, all treatments shared the same basal media,
which contained (per liter) 0.5 g yeast extract; 1.0 g
Na2HPO4 × 12 H2O; 1.0 g KH2PO4; 0.4 g MgSO4 × 7
H2O; 1.0 g (NH4)2SO4; 6 ml FeSO4 solution (4 g·L−1
FeSO4 × 7 H2O), and 10 ml trace mineral solution. The
trace mineral solution consisted of (per liter) 3.6 g CaCl2 ×
2 H2O; 0.75 g ZnSO4 × 7 H2O; 0.13 g CuSO4 × 5 H2O;
0.5 g MnSO4 × H2O; 0.13 g CoCl2 × 6 H2O, and 0.17 g
Na2MoO4 × 2 H2O. Table 1 shows the type and amount
of carbon source used for the respective treatment. “G100”
was set as a control treatment and contained only glucose.
In the other treatments fructose was used as easily avail-
able monosaccharide in order to be able to distinguish
between the initially added carbon source and the glu-
cose resulting out of starch degradation. Generally, the
suitability of glucose and fructose as carbon source for R.
glutinis was assumed to be equal.

Table 1. Type and composition of carbon sources for the experimental treatments.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Glucose [g·L−1]</th>
<th>Fructose [g·L−1]</th>
<th>Starch [g·L−1]</th>
</tr>
</thead>
<tbody>
<tr>
<td>G 100</td>
<td>40.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F + S 50:50</td>
<td>20.0</td>
<td>19.0</td>
<td></td>
</tr>
<tr>
<td>S 100</td>
<td>5.0</td>
<td>33.25</td>
<td></td>
</tr>
</tbody>
</table>

In “S 100” 5 g·L−1 of fructose were added in order to
facilitate the initial adaption of the yeast to the utilization
of starch as carbon source. All media were autoclaved
and afterwards inoculated with 20 mL of R. glutinis seed
culture, which has been cultivated on yeast malt broth (3
g·L−1 yeast extract, 3 g·L−1 malt extract, 10 g·L−1 glucose,
5 g·L−1 vegetable peptone) for 48 h at 25°C on a rotary
shaker at 115 rpm. Rhodotorula glutinis ((Fresenius)
Harrison, anamorph) was obtained from the American
Type Culture Collection (ATCC 15125TM). Fermentation
was carried out in 1000 mL Erlenmeyer flasks, contain-
ing 700 mL of the respective medium, on a rotary shaker
at 115 rpm and 25°C for 240 h. The experiment was car-
ried out in triplicate. 50 mL samples were taken after 0,
48, 120, 168, 216, and 240 h of cultivation and centri-
fuged at 3000 rpm for 10 min. The supernatant was used
for the determination of pH, NH4-N and sugar content.
NH4-N was measured photometrically using reagent
vials from Macherey-Nagel (Düren, Germany). 200 µL
of the sample is added to the test tube. Ammonium reacts
with hypochlorite and salicylate in the presence of so-
dium nitroprussiate as catalyst to form a blue indophenol.
The color strength is then measured with a photometer
(NANOCOLOR 400 D, Macherey-Nagel, Düren, Ger-
many) at a wavelength of 690 nm. The sugar content was
analyzed via HPLC (Merck-Hitachi HPLC system with
DAD detector) using a Phenomenex Rezex™ RPM-
Monosaccharide Pb + 2 (8%) column (300 × 7.8 mm)
with water as mobile phase at a flow rate of 0.6 ml·min−1.
The sugar content was determined by comparison to known standards (Figure 1). The remaining cell pellet was freeze-
dried and weighted to determine the cell dry mass. It was
observed that the pellet also contained a fraction of un-
dissolved starch.

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Thus, the actual biomass production at the respective sampling times was calculated by subtracting the initially measured weight after 0 h, assuming that the fraction of undissolved starch will remain constant over the cultivation time. The lipid content of the dried cell pellet was determined using a modification [22] of the classical Bligh and Dyer [23] extraction with methanol and chloroform as solvents. The final lipid extract was weighted after the solvents were evaporated under nitrogen blow at 50°C. This extract from the Bligh and Dyer procedure is later referred to as total extractable lipids. Afterward the extracts from sampling times 48 h and 240 h were trans-esterified with methanol with 2% H₂SO₄ as catalyst in a waterbath at 60°C for 2 h. The reaction was quenched with NaHCO₃ solution and the resulting FAMEs were extracted with toluene and analyzed using an Agilent 6890 gas chromatograph with flame ionization detector (GC-FID) equipped with an Agilent J&W GC capillary column (30 m × 0.252 mm ID and 0.25 µm film thickness). The different fatty acids were identified by comparison to responses from a known FAME standard (Supelco® 37 component FAME Mix, Sigma Aldrich, München, Germany) (Figure 2). For quantitative determination of FAME production, the peak areas of fatty acids in the samples were compared with the peak areas of fatty acids in a known amount of FAME standard.

For starch analysis samples of the cultivation media were heated in a waterbath to completely dissolve all starch, then diluted with distilled water, mixed with amylglucosidase (AMG) solution and incubated at 60°C for 1.5 h in order to degrade starch into glucose. The amylglucosidase solution was prepared daily by (per 12 samples) dissolving 10.6 mg of AMG (Amyloglucosidase from Aspergillus niger, ~70 U·mg⁻¹, Sigma Aldrich, München, Germany) in 12 ml buffer solution (pH 4.6). The samples were then filtered and analyzed for their glucose content using HPLC according to the method previously described for the sugar analysis. From the measured glucose contents, the starch content was calculated based on a 90% conversion rate from starch to glucose.

All data was statistically analyzed using Sigma Stat...
3. Results and Discussion

3.1. Sugar and Starch Consumption

As Figure 3 shows, the monosaccharides glucose and fructose were steadily consumed in course of cultivation. Thus, based on the different initial sugar contents, the easily available carbon sources were exhausted after 48 h in S 100 and after 168 h in F + S 50:50. In treatment G 100 glucose was available throughout the cultivation with 75% of total glucose consumed after 240 h.

The initially measured starch content of 18.1 g L\(^{-1}\) in F + S 50:50 did not vary significantly over the duration of the fermentation. In treatment S 100 the starch content decreased significantly from 33.2 g L\(^{-1}\) at 120 h to 27.5 g L\(^{-1}\) at 168 h. This indicates that some type of mechanism for starch degradation must be available in *Rhodotorula glutinis*. Verstraete et al. [24] reported the presence of an amylolytic *Rhodotorula* strain in activated sludge. However, at the same time it is rather incomprehensible that after this point no further decrease in starch was detected. If different carbon sources are offered for microbial growth, the easily available sugars are preferentially utilized. If this source is exhausted, cells switch to the second source, whereas at that time often a certain lag period can be observed, since it needs some time until the required enzymes for the degradation of the respective carbon source are produced. In the present study this lag phase was quite long and could be observed in S 100 between 48 h (fructose exhaustion) and 168 h (starch utilization). In F + S 50:50 fructose was available until 168 h. Thus, it can be assumed that in this treatment after a lag phase a certain degree of starch utilization might occur. To verify this assumption, the cultivation time should be increased in further studies.

3.2. Biomass Production

The results of sugar and starch consumption were in line with the data obtained from biomass and lipid production (Figure 4). Until 168 h the treatments G 100 and F + S 50:50 showed almost an identical increase in cell mass production along with a simultaneous decrease of the ammonium content, with ammonium as nitrogen source being completely exhausted after 120 h. The further increase in biomass was probably facilitated by the utilization of the yeast extract, which was initially added to the medium as secondary nitrogen source. In G 100 the cell mass continued to increase until 216 h, reaching a biomass yield of 12.3 g L\(^{-1}\). Since glucose was still available at this point it can be assumed that nitrogen was the limiting factor for further biomass production. In F + S 50:50 biomass production ceased after 168 h at 9.1 g L\(^{-1}\). This can be explained by the limitation of carbon, since the entire fructose was consumed from the medium and...
starch as secondary carbon source was not utilized. In treatment S 100, fructose was exhausted after 48 h of cultivation, while no starch utilization could be observed until 120 h. Despite this limited availability of carbon the cell mass increased more or less steadily until 168 h.

However, compared to the other treatments biomass production was much lower, yielding less than half (4.0 g·L⁻¹) of the cell mass compared to G 100 and F + S 50:50 after 168 h. Due to the fact, that nitrogen was utilized for the formation of biomass, ammonium was consumed from the media more slowly in S 100, being exhausted after 168 h of cultivation. The increase of biomass production after 216 h in F + S 50:50 and S 100 cannot be coherently explained. However, it is noticeable that the increase was almost identical for both treatments containing starch and it was also confirmed by photometrical cell density determination. Thus, it is unlikely since nitrogen was exhausted at this point. Since the cell pellet after freeze drying was contaminated with residues of undissolved starch, the biomass production was calculated by subtracting the initially measured pellet weight at 0 h. For this, a stable proportion if residual starch was assumed. Thus, a decrease in the fraction of undissolved starch could appear as an increase in biomass production. However, this assumption is not supported by the fact, that no decrease in the starch content was detected.

### 3.3. Lipid and FAME Production

Regarding lipid production, it needs to be differentiated between the gravimetrically determined extractable lipid fraction as obtained by the Bligh and Dyer extraction (Figure 4) and the amount of saponifiable lipids represented by the amount of FAMEs as detected using GC analysis (Figure 5), because next to this saponifiable fraction of lipids the Bligh and Dyer extracts contain many other compounds that will dissolve in chloroform, e.g. certain carotenoids.

Still the gravimetical results for lipid production are valuable indicators regarding the cycle of lipid production during microbial cultivation. Furthermore it needs to be considered, that not all extracted lipids are necessarily derived from the process of De novo lipid synthesis, which is defined as the production of storage lipids from the cell membranes, an increasing lipid yield can be to some extent associated to a rise in biomass production. The gravimetical lipid production (Figure 4) can roughly be divided into three phases, which are similar for all treatments: 1) an increase in lipid production during the first 48 h of growth, 2) a stagnation of lipid production in the middle of cultivation until 168 h, and 3) an increase in lipid production at the end of cultivation after 168 h (G 100 and S 100) and 216 h (F + S 50:50), respectively. The initial increase is probably related to the simultaneous onset of biomass production as described previously and thus cell growth induced. The second phase of lipid production 3) just starts after ammonium is exhausted from the medium (Figure 4). This is characteristically for De novo lipid synthesis, since carbon is only channelled into the production of storage lipids, when a limitation of a growth required nutrient (mostly nitrogen) restricts cell growth processes [4]. With 3.0 g·L⁻¹ after 240 h, G 100 showed the highest lipid production. Other reports where shake flask cultivations of R. glutinis were applied observed similar yields [25-27]. However, treatments F + S 50:50 (2.4 g·L⁻¹) and S 100 (2.5 g·L⁻¹) had only slightly lower values for lipid production. This is rather surprising, since at that time in both treatments fructose was exhausted and starch was not or only slightly degraded, thus a carbon limitation could be assumed, which would usually hamper the production of biomass and lipids. The effect of the differences in the initial amounts of easily available sugars in the different treatments becomes obvious with the chromatographically determined FAME production (Figure 5). Since only glucose and fructose have been consumed as carbon sources, the FAME production follows the pattern of initial sugar content, whereas G 100 with the highest sugar content yields the highest amount of FAMEs, followed by F + S 50:50 with around half the initial sugar content and FAME production. S 100 had the lowest sugar content and only small amounts of starch were utilized. Thus, carbon was limited and no significant FAME production could be observed.

The fatty acid composition (Table 2) showed a pre-
that this strain of reduction of starch was measured, it cannot be ruled out, report starch utilization by starch hydrolysis. However, since some literature sources visible to conduct an additional pretreatment step for wastewaters are used as fermentation medium it is ad-
of starch degradation and utilization. Thus, before these experiments the cultivation time should be increased ther experiments the cultivation time should be increased to observe if starch is further utilized. However, in the context of starch containing wastewaters as potential carbon source for microbial lipid and biodiesel production, it can be stated that a suitable enzymatic or thermo-
chemical pretreatment for starch hydrolysis would increase the amount of available carbon and thus potentially increase the lipid yields. Other possible measures for increased starch utilization could be the co-cultivation with amylolytic yeast strains (e.g. [18]) or the use of specific strains of R. glutinis, which are adapted to the utilization of starch.

4. Conclusion

Wastewaters from food industry with high starch contents are a promising cheap carbon source as feedstock for microbial lipid and biodiesel production and also of potential interest for the microbial production of carot-
enoids. When red yeast R. glutinis was cultivated on soluble starch as carbon source, it yet showed a poor ability of starch degradation and utilization. Thus, before these wastewaters are used as fermentation medium it is advisable to conduct an additional pretreatment step for starch hydrolysis. However, since some literature sources report starch utilization by R. glutinis, the present results might be strain specific.

Table 2. Fatty acid composition [%] of microbial lipids from Rhodotorula glutinis after 240 h of cultivation on different carbon sources.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>16:0</th>
<th>18:0</th>
<th>18:1</th>
<th>18:2</th>
<th>18:3</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>G 100</td>
<td>27.9</td>
<td>11.6</td>
<td>43.7</td>
<td>10.7</td>
<td>3.5</td>
<td>2.6</td>
</tr>
<tr>
<td>F + S 50:50</td>
<td>30.2</td>
<td>8.8</td>
<td>44.0</td>
<td>11.9</td>
<td>1.6</td>
<td>3.5</td>
</tr>
<tr>
<td>S 100</td>
<td>12.6</td>
<td>5.8</td>
<td>42.7</td>
<td>33.9</td>
<td>2.8</td>
<td>2.2</td>
</tr>
</tbody>
</table>

dominance of palmitic (C 16:0) and oleic (C 18:1) acid, which is characteristic for R. glutinis [26,28]. Generally in all treatments the long-chain C 16 and C 18 fatty acid account for the vast majority with over 97% of total fatty acids. Thus, the microbial oils can be transesterified into biodiesel just like conventional plant oil. Moreover, the high share of oleic acid has beneficial effects on bio-
diesel fuel quality [29]. The comparably high content of linoleic acid (C 18:2) in S 100 and the fact that linoleic acid is often found in the lipids of cell membranes indicate that in this treatment biomass associated lipids derived from cell membranes prevailed.

The obtained results generally suggest that the mechanisms of starch utilization are poorly developed in R. glutinis. However, since in treatment S 100 a significant reduction of starch was measured, it cannot be ruled out, that this strain of R. glutinis over time develops the required enzymatic systems for starch utilization. For further experiments the cultivation time should be increased to observe if starch is further utilized. However, in the context of starch containing wastewaters as potential carbon source for microbial lipid and biodiesel production, it can be stated that a suitable enzymatic or thermo-
chemical pretreatment for starch hydrolysis would increase the amount of available carbon and thus potentially increase the lipid yields. Other possible measures for increased starch utilization could be the co-cultivation with amylolytic yeast strains (e.g. [18]) or the use of specific strains of R. glutinis, which are adapted to the utilization of starch.

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