Polyhydroxyalkanoate Production by *Pseudomonas putida* KT217 on a Condensed Corn Solubles Based Medium Fed with Glycerol Water or Sunflower Soapstock

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ABSTRACT

*Pseudomonas putida* KT217 was grown on a complex medium comprised of co-products of the ethanol and biodiesel industries to assess the organism’s capability to produce medium-chain-length polyhydroxyalkanoate (mcl-PHA). The growth phase was carried out in a medium containing 400 g/L condensed corn solubles (CCS), supplemented with ammonium hydroxide as a nitrogen source. Following the exponential phase, co-products of the biodiesel industry (soapstock and glycerin) were fed into the reactor to trigger PHA production. When glycerin was added to the bioreactor (75 g/L total addition), the final cell dry weight (CDW) and PHA content were 30 g/L and 31%, respectively. The monomeric composition in the PHA formed was relatively uniform throughout incubation with 3-hydroxydecanoate dominating. When a total of 153 g/L of sunflower soapstock was added to the bioreactor in a fed-batch manner, the final CDW and PHA content were 17 g/L and 17%, respectively. Following addition of soapstock the monomeric composition of the polymer changed dramatically, with the 3-hydroxyoctanoate monomer becoming dominant and greater unsaturation present in the PHA.

Keywords: Polyhydroxyalkanoate; *Pseudomonas putida*; Condensed Corn Solubles; Glycerol; Soapstock

1. Introduction

Polyhydroxyalkanoates (PHAs) are a class of biodegradable polymers that have a wide range of physical properties depending upon the monomeric composition in the polymer [1-4]. For the past 30 years extensive PHA research has been conducted. With recent breakthroughs in PHA production technology and soaring oil prices, PHA and other biopolymers may be at the front of true commercial integration [5]. Biopolymers such as 1 - 3 propanediol, polyactic acid, starch based polymers, and PHA could capture as much as 1.5% - 4.8% of the total plastics market (~260 million tonnes/year). Production of PHA from renewable biomass is providing the “green” alternative to the pollution resulting from use on non-degradable plastics.

The costs of producing and recovering biodegradable polymers have been the key barrier to the marketplace. Numerous methods have been attempted to reduce the cost of extracting and purifying the polymer [6-13]. Similarly, several low cost substrates have been evaluated for PHA production, but low productivities typically result [14-30]. Solaiman *et al.* [31] reviewed the use of vegetable oils, animal fats, dairy whey, molasses, and meat and bone meal as feedstocks for PHA production. The ethanol and biodiesel production industries also generate large quantities of under-utilized byproducts which could be used for PHA production.

According to Renewable Fuels Association [32], the US ethanol production was 13507.9 million gallons from 204 plants and expected to increase by 522 million gallons on completion of 10 more plants under construction. Each gallon of ethanol accompanies with 5% - 7% of condensed solubles. The ethanol production co-product condensed corn solubles (CCS) is normally used in livestock feeds. CCS contains carbon and energy sources such as monosaccharides, oligosaccharides, organic acids, and glycerol, as well as a range of micronutrients and macronutrients such as zinc, iron, manganese, magnesium, sulfur, phosphate and nitrogen. Typically CCS is deficient in nitrogen, but due to its otherwise rich nutritional composition, CCS has been successfully used to grow a range of bacteria and fungi [33-35].

The co-products of biodiesel production include fatty-acids, soapstock, and glycerol water. Soapstock is ob-
tained from alkali refining of oilseeds and is composed of phosphorus lipids, hydratable and non-saponifiable compounds, soaps of free fatty acids (FFA), vitamins A and E, and carotenoids pigments [36-39]. Soapstock is normally added back to oilseed meal for livestock feeding or is disposed of with little or no economic compensation [36].

Soapstock has been evaluated as a carbon and energy source to produce value added products via fermentation. Hesseltine and Koritala [40] evaluated a 2% soybean source to produce value added products via fermentation. Later on Kaneshiro et al. [41] isolated a bacterium from manure compost (tentatively identified as a *Sphingobacterium* and designated strain NRRL B-14797) that grew on crude soybean soapstock and a soapstock extract, and produced 10(R)-hydroxystearic acid. They found that soapstock extracts were poor nutrients for growth, but were utilized better for hydroxy acid bioconversions than either crude soapstock or pure long chain fatty acids. This is most likely due to removal of some nutrients following purification of the fatty acids. Benincasa et al. [42,43] grew *Pseudomonas aeruginosa* LB1 aerobically on a defined liquid salts medium with 2.5% sunflower soapstock to produce rhamnolipids. Fed batch addition of soapstock gave the most favorable results, with 16 g/L of product after 54 h. Sunflower soapstock contains linoleic acid 50%, oleic acid 25%, palmitic acid 7%, and stearic acid 4%. Bednarski et al. [44] fed soapstock or post-refinery fatty acids to *Candida antarctica* and *C. apicola* to synthesize surfactants (glycolipids). Soapstock resulted in better glycolipid production than fatty acids (7.3 - 13.4 g/L vs 6.6 - 10.5 g/L, respectively), which was similar to the results observed in *Pseudomonas aeruginosa* LBI [41].

According to National Biodiesel Board [45], biodiesel production in the US has reached 1.1 billion gallons. Glycerol is a primary byproduct of biodiesel production. Each gallon of biodiesel accompanies 0.3 kg of glycerol, which is equivalent to 10%, and this has triggered research to develop alternative uses, including microbial processes. For example, Flickinger and Perlman [46] converted glycerol to dihydroxyacetone with a *Gluconobacter* strain. Du-Pont created a process to produce 1 - 3 propanediol that channels glucose through glycerol [47]. It is possible for this product to be created directly from glycerol by microbial conversion via *Clostridium* strains [48]. The fermentation of glycerol by *E. coli* to end products such as ethanol, succinate, acetate, lactate, and hydrogen was found to be pH dependant [49]. Production of PHA from glycerol water has also been tested [21].

The combination of co-products from the ethanol and biodiesel industries could be used to create a medium to support growth and PHA accumulation. Our previous research revealed that *Pseudomonas putida* KT217 grew well on a medium comprised of CCS, supplemented with ammonium as a nitrogen source [50]. In the study reported herein we used this basal medium to test the effects of using a fed-batch feeding strategy with either glycerol water or sunflower derived soapstock to bolster PHA production. Trials were performed in an aerated benchtop reactor at the 2.35 L scale with soybean biodiesel-derived glycerin or sunflower-derived soapstock added fed-batch after 24 h to bolster PHA levels.

2. Materials and Methods

2.1. Bacterial Strain, Maintenance, and Inoculum Preparation

The bacterium used was *P. putida* KT217. Long term storage was via lyophilization, while short term storage was on tryptic soy agar (TSA) slants. The culture was routinely transferred in tryptic soy broth shake flasks incubated for 24 - 48 h at 30°C and 250 rpm. Subcultures were also transferred to shake flasks containing a CCS based medium (described below) at pH 7. A one percent inoculum of a 24 h culture was used to inoculate aerated fed-batch bioreactor trials.

2.2. Experimental Design

The basal CCS medium was prepared by mixing 1.2 kg of CCS from a dry grind ethanol plant with deionized water to a volume of 3 L. The medium pH was adjusted to 7.3 with 30% ammonium hydroxide (after autoclaving the medium pH typically dropped to 6.7). The resulting mixture was centrifuged to remove suspended solids. The liquid portion was then filtered through Whatman 113 filter paper to remove most of the remaining suspended solids and oils. The medium (~2.35 liters each) was then dispensed into a 5 L Bioflo III bioreactor (New Brunswick Scientific, Enfield, CT, USA) and autoclaved.

Trials were conducted aerobically in a fed-batch mode using pH control and dissolved oxygen (DO) was monitored. For pH control a saturated solution of sodium hydroxide and a 20% (v/v) solution of sulfuric acid were used.

In the glycerol trials, following inoculation the culture was incubated for 96 h at 30°C, with aeration at 1 V/V/min and agitation at 500 rpm. Clerol FBA 3107 (Cognis) was used as an antifoam agent in these trials as needed. In the soapstock trials the agitation and aeration had to be monitored to control foaming along with the antifoam.

After the initial 24 h growth phase, fed-batch additions of sterile glycerol water (obtain from West Central Soy in Ralston, IA) or sunflower soapstock (obtained from

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Cargill) were delivered through the septum port on the fermentor with a sterile disposable 60 mL syringe. Glycerol water could be added as is, however the viscosity of soapstock was too high to deliver directly via syringe. Therefore, 40 g of Sunflower soapstock was diluted with deionized water to 150 mL total volume and autoclaved prior to feeding. Glycerol water was fed based upon HPLC data, to maintain glycerol levels between 5 and 35 g/L during incubation. In total 240 mL of glycerol water was added between 24 and 70 h. Soapstock was fed in response to dissolved oxygen levels rising above 10% following 24 h of cultivation, with 1350 mL (360 g) of soapstock solution added between 24 and 90 h.

2.3. Cell Dry Weight and Colony Forming Unit Measurement

Cell dry weights (CDW) were obtained by harvesting 30 - 50 mL samples and centrifuging at 4000 rpm for 30 min. Cell pellets were washed with 30 mL deionized water, re-centrifuged, and dried to a constant weight in a 60°C oven. Colony forming units (CFU) were determined by serial dilution and plating in triplicate on TSA.

2.4. HPLC Analysis

A Waters HPLC system (Waters Scientific, Milford, MA, USA) with refractive index detector was used to quantify sugars, organic acids, and glycerol in the culture samples. Prior to analysis samples were filtered through 0.2 µm filters, then 50 µl injections were made. The mobile phase was helium-degassed 4 mM sulfuric acid at 0.6 mL/min, through a Biorad HPX-87H (Biorad, Hercules, CA, USA) organic acid analysis column operated at 65°C. Standard solutions of maltose, glucose, lactic acid, acetic acid, propionic acid, succinic acid and glycerol were used for calibration.

2.5. Nitrogen and Phosphorus Analysis

Samples were assayed for nitrogen and phosphorus using Hach test kits and a Hach DR/2010 colorimeter (Hach, Loveland, CO, USA). Samples were filtered through 0.2 µm filters, then diluted 1:100 with double distilled water. Nitrogen was measured as ammonia using the Hach HCT 102 Unicel kit. Phosphorus was measured as free phosphates using of the Hach HCT 122 Unicel kit.

2.6. PHA Analysis

PHA concentrations were determined by first removing 30 - 50 mL samples, centrifuging at 4500 rpm at 10°C for 30 min, then washing and re-centrifuging the cells. Cell pellets were then lyophilized and homogenized prior to PHA extraction.

2.7. PHA Extraction

Standards are not readily available for each of the monomers present in the mcl-PHA produced by P. putida KT217. To overcome this, PHA was utilized from culture samples to create quantitative standards, following the methods describe by Foster et al. [51] and Kim [52]. Lyophilized cell samples from the final culture sample of the soapstock fed trial were extracted with a two-step supercritical fluid extraction [53]. In the first step, neat supercritical carbon dioxide was used to extract non-PHA lipids, followed by an ethanol modified step to extract PHA. Following centrifugation at 4500 rpm to remove excess ethanol, PHA was dissolved in hot chloroform and precipitated by addition of ten volumes of cold methanol. This procedure of dissolving in chloroform and precipitation in cold methanol was repeated three times to purify the PHA. A portion of the purified PHA was dissolved in chloroform and used as a stock solution to prepare quantitative standards. These standards were prepared by hydrolyzing and derivatizing the methyl-esters of the respective monomers included in the PHA.

2.8. PHA Hydrolysis and Derivitization

Samples of lyophilized cells or purified PHA (10 - 60 mg) were digested in chloroform:methanol:sulfuric acid (50: 42.5:7.5 % v/v) at 100°C for 4 h, using benzoic acid as an internal standard [54-56]. Following digestion the mixture was washed with distilled water to remove excess sulfuric acid and methanol from the chloroform phase. This procedure creates methyl esters of the 3-hydroxy acids present in the PHA.

2.9. PHA Analysis

The created standards were analyzed on Thermo-Finnigan Trace GC/MS 2000. The column utilized was an RTx-5Sil MS (Restek, 30 meter, 0.25 mmID, 0.5 µm df). The MS detector was utilized first for qualitative analysis of the monomers present in the samples. The FID detector was subsequently utilized for quantitative analysis of the PHA present in the samples. The total areas of the monomers present were correlated to the amount of PHA contained in three different levels of standard and normalized to the methyl ester of benzoic acid as an internal standard to create a calibration curve (R^2 = 0.97 - 0.99). The digests of cell samples were then analyzed and the percentage of PHA contained within determined. For determination of the fractional analysis of PHA monomers, the area of the individual monomers was divided by the total area of the monomers.

2.10. PHA Analysis by NMR Spectroscopy

Samples obtained from SFE were dissolved in deuterated.
chloroform and $^{13}$C-NMR spectra were recorded on a Bruker 400 MHz GRX NMR spectrometer (Bruker AXS, Inc, Madison, WI, USA) at a probe temperature of 27°C.

3. Results and Discussion

3.1. Growth on a CCS Medium with Fed-Batch Addition of Glycerol Water

*P. putida* KT217 was grown in an aerated bioreactor on a 400 g/L CCS-based medium supplemented with 2.2 g/L ammonium hydroxide. The 400 g/L CCS formulation had previously been identified as being optimal for rapid production of cell mass [50]. The average growth curve from two replications, in which glycerol water was added at 29, 34, 48, and 62 h, is shown in Figure 1.

Of the nutrients initially present in the CCS medium, *P. putida* KT217 first utilized the glucose (2.1 g/L), succinic (2.9 g/L), lactic (4.6 g/L), acetic (1.0 g/L), and propionic acids (1.0 g/L) within 20 h (data not shown). Only after these carbon sources were depleted (at approximately 12 h) did *P. putida* begin metabolizing glycerol, which fell from 39.3 g/L initially to 8.4 g/L at 28 h. (Figure 1). Ammonia was exhausted by approximately 28 h, which corresponded to the plateau in cell population ($6.4 \times 10^{10}$ CFU/mL) and CDW (~23 g/L). The CDW included both cell mass and PHA, the latter of which had accumulated to ~2%.

The CDW was produced from ~42 g of carbon, for a CDW yield of ~0.55 g CDW/g carbon utilized. This CDW yield was slightly higher than theoretical and may be attributed to utilization of other medium components in CCS that were not detected by HPLC (e.g., oligosaccharides, amino acids). For example, in a phenol red broth medium with glucose as the carbon source *P. putida* KT217 was found to catalyze deamination of media proteins. It is possible that *P. putida* was able to use proteins and oils in CCS as carbon sources. Viable cell counts increase more rapidly than CDW due to the lower mass of individual cells during exponential growth. However, after the stationary phase was reached, CDW increased as cells filled with PHA.

At 28 h, the first of four glycerol additions was made (Figure 1). These additions maintained glycerol levels between 8 - 35 g/L, and Figure 2 shows the glycerol utilization rate. At 28 h, the glycerol utilization rate peaked at 2.87 g/L/h. During the subsequent additions of glycerol, the utilization rate was maintained between 1 - 2 g/L/h. This lower rate was likely due to several factors including limitations of nitrogen and oxygen, and perhaps the increased salt concentration from the glycerol water. The glycerol utilization rate continued to decline, approaching 0.5 g/L/h at the end of fermentation, as cells filled with PHA.

![Figure 1. Growth and PHA production of *P. putida* KT217 on a CCS medium with fed-batch glycerol addition. Average of two fermentations with error bars representing standard deviation.](image-url)
After 28 h, the CDW continued to increase to around 30 g/L at 85 h. Since the viable cell population remained constant during this time period, the increase of 7 g/L in CDW was primarily due to the accumulation of PHA. The increase in CDW of 7 g/L from use of ~65 g/L of glycerol, corresponds to a CDW yield of 0.11 g/g glycerol (CDW is sum of PHA and cell mass). This lower yield indicates that the majority of the glycerol was being used for maintenance energy in the culture. The various processes resulting in the utilization of carbon source for maintenance energy have been reviewed [57]. Additionally, an increase of PHA of 8 g/L was measured by gas chromatography. This represents 31% of the total CDW. This is greater than the total increase in cell mass following limitation and may indicate that there was significant turnover of cell mass in the medium during this time resulting in a cell mass with varying PHA content.

The composition of the PHA produced by *P. putida* KT217 on a CCS medium fed with glycerol water changed with time as evidenced by gas chromatographic analysis (GC-MS and GC-FID). The overall concentration of 3-hydroxydodecanoate and 3-hydroxydodecanoate in the polymer decreased as the incubation proceeded, whereas the amount of 3-hydroxyhexanoate, 3-hydroxyoctanoate, and 3-hydroxydecanoate all increased (Figure 3). Polymer composition was dominated throughout by 3-hydroxydecanoate which consistently represented more than 60% of the polymer. The concentration of 3-hydroxytetradecenoate, 3-hydroxytetradecanoate, and 3-hydroxyhexadecanoate were all detectable, but in very small concentrations (Figure 3). These conclusions are supported by $^{13}$C-NMR data shown in (Figure 4). Chemical shifts are similar to previously reported data for PHA analyzed in this method [15,58,59].

### 3.2. Growth on a CCS Medium with Fed-Batch Addition of Sunflower Soapstock

*P. putida* KT217 was grown on 400 g/L CCS-based medium supplemented with ammonia as a nitrogen source (Figure 5). Carbon sources initially utilized for growth (glucose, lactic, succinic, acetic, and propionic acids) followed the same pattern as observed in Figure 1, and when these were depleted (~20 h), glycerol utilization began. Viable cell counts plateaued at 15 h, but rose from $4.6 \times 10^9$ CFU/mL to $9.44 \times 10^9$ CFU/mL by 40 h. This corresponded to an increase in CDW from ~11 to 24 g/L over the same period. Between 15 and 35 h, the average glycerol utilization rate was 1.75 g/L/h, which was comparable to the average rate observed (1.4 - 2.9 g/L/h) in the previous trials in the same time frame. Glycerol consumption continued until depletion at 39 h, which corresponded to the peak CDW level. Ammonia depletion occurred at 32 h. Following glycerol depletion in the medium the CDW decreased until 80 hours.

When glycerol fell to 10 g/L (~28 h) sunflower soapstock additions were initiated. Because soapstock was viscous, 40 g of soapstock was blended with water to a volume of 150 mL prior to each addition. To compensate for this added volume, 150 mL of culture broth was removed prior to the diluted soapstock addition. It was not possible to monitor the level of soapstock, since it is a complex of several lipids. However, Lee et al. [60] previously reported that dissolved oxygen levels can be used to monitor carbon levels. Therefore when the dissolved oxygen saturation rose above 10%, soapstock additions were made. This caused oxygen saturation to drop to near zero, and as it was consumed, oxygen saturation gradually increased. A total of 360 g of sunflower soapstock was added in a total volume of 1.35 L, at 6 - 10 h intervals through 86 h.

We had previously observed that *P. putida* KT217 could utilize the components in soapstock as a carbon and energy source during aerated shake flask trials in a defined medium (data not shown). The foaming problem in these trials was consistent with previous reports [42,43]. Soapstock also increased foaming in the aerated
Figure 3. Time profile of monomeric composition determined by gas chromatography in PHA derived from fed-batch fermentation of *P. putida* KT217 on a syrup medium fed with glycerol water.

Figure 4. $^{13}$C-NMR of the PHA extracted from the final sample of the glycerol water fed batch fermentation.
Figure 5. Growth and PHA production of *P. putida* KT217 on a CCS medium with fed-batch sunflower soapstock addition. Arrows indicate addition of 40 grams of sunflower soapstock. Average of two fermentations with error bars representing standard deviation.

Bioreactor, even with antifoam addition. Therefore we reduced the aeration rate from 1 vvm to 0.5 vvm.

After soapstock additions began, CDW decreased, even though viable cell populations remained steady. Evidently, the broth removed prior to soapstock additions contained cells at least partially filled with PHA, and these were replaced by newly formed cells that didn’t contain as much PHA. This did not occur in the prior glycerol fed-batch process because only 240 mL of the concentrated (85%) glycerol water was added, compared to the more dilute soapstock (50% FFA) [37]. Acetic acid levels began to accumulate after 60 h, eventually rising to 7 g/L. This was likely due to fatty acid degradation into acetyl-CoA, coupled with the decreased aeration rate and hence reduced acetic acid being excreted from the cells. It is likely that the Krebs cycle could not process acetyl-CoA at the same rate it was being produced, thus diverting some acetyl-CoA to acetic acid. This build up in acetyl-CoA may have activated the fatty acid synthesis pathway as well, explaining the increase in PHA.

PHA accumulated at a slow rate throughout incubation, reaching a concentration of 2.8 g/L at the end, representing 17% of the total CDW. The composition of the PHA changed dramatically during incubation (Figure 6). Initially, the composition favored 3-hydroxydecanoate monomers due to the utilization of glycerol in the CCS. Following soapstock addition, polymer composition shifted towards 3-hydroxyoctanoate monomers. The final PHA contained 3-hydroxytetradecanoate and 3-hydroxytridecanoate. 13C-NMR analysis of the final PHA produced during the incubation supported the information derived from GC-MS and GC-FID analysis showing an increase in unsaturation in the polymer (Figure 7). This is evident by examining the peaks in the chemical shift range of 120 - 150 ppm which has been shown previously [15,55].

4. Conclusions
PHA production was observed over 100 h of aerated incubation when *P. putida* KT217 was grown on a basal medium of 400 g/L CCS (wet basis) and 2.2 g/L ammonium hydroxide, supplemented in fed-batch mode with either biodiesel co-products glycerol water (240 g) or soapstock (390 g). Foaming was a problem during soapstock trials, but was of less concern when using glycerol.
Figure 6. Time profile of monomeric composition determined by gas chromatography in PHA derived from fed-batch fermentation of *P. putida* KT217 on a syrup medium fed with sunflower soapstock.

Figure 7. $^{13}$C-NMR of the PHA extracted from the final sample of the sunflower soapstock fed batch fermentation.
feeding strategy. A majority of the carbon source fed to the P. corrugata grown on soybean molasses carbohydrates also resulted in 3-hydroxy acyl-CoA monomers of this chain length. In contrast, sunflower soapstock resulted in 3-hydroxydecanoate generated by the de-oxidation, de novo fatty acid synthesis and fatty acid elongation pathways. This phenomenon was also noticed with P. putida when grown on a biodiesel co-product where the FFA's composition shifting from 3-hydroxydecanoate to 3-hydroxyoctanoate over time due to β-oxidation, de novo fatty acid synthesis and fatty acid elongation pathways.

The maximum concentration of PHA in cell mass that we observed was only 31%, following a lengthy incubation when glycerol water was used as the carbon source. A majority of the carbon source fed to the P. putida was evidently used for maintenance energy (or possibly other products not detected). This low PHA level would likely lead to poor extraction economics, as current industrial processes target a 90% PHA level. However these processes are primarily based on PHB-PHx copolymers, and utilize starch-derived sugars for growth. Use of lower costs carbon sources may lead to more economically PHA production processes, however additional research is needed to improve microbe performance and feeding strategy.

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