ABSTRACT: PHAs are a group of intracellular biodegradable polymer produced by (most) bacteria under unbalanced growth conditions. A series of enzymes are involved in different PHAs synthesis, however PhaC synthases are responsible for the polymerization step. PHAs are accumulated in bacterial cells from soluble to insoluble form as storage materials inside the inclusion bodies during unbalanced nutrition or to save organisms from reduces equivalents. PHAs are converted again to soluble components by another pathways and enzymes for the degradation process. PHAs depolymerases are the responsible enzymes. This review is designed to give the non-specialists a condense background about PHAs especially for researcher and students in medicinal and pharmaceutical field.

KEYWORDS: polyhydroxyalkanoate; PhaC synthase; PHA depolymerase
1. INTRODUCTION

Perhaps, Beijerinck wrote the first report about lucent granules of PHAs in bacterial cells in 1888 (Reported in Chowdhury, 1963) [1, 2]. Poly-\(\beta\)-hydroxybutyrate (PHB) was firstly described as an important bacterial product by Lemoigne (1923). Later, Lemoigne (1927) characterized PHB chemically and observed that it was involved in the sporulation of \textit{Bacillus spp} [3, 4]. Lemoigne and co-workers published 27 articles from 1923 to 1951 [5]. He was the first to observe that these storage materials were not ether soluble, as in lipids. Later, he reported that PHB is the major constituent of these granules. Using microscopic investigation, saponification numbers, and autolysis he proposed a polyester structure with the formula \((C_4H_6O_2)_n\)- and reported that PHB could cast into film like cellulose nitrate material [3-7].

The worldwide demand for degradable plastic is estimated to be 1.4 million metric tons per year by the year 2000, with much of this demand driven by legislation and environmental group [8]. A number of polymers are being developed to meet this demand, with the majority of these being petrochemical based and projected cost at $6.6 kg\(^{-1}\) [9, 10]. Natural products being developed as biodegradable plastics include starch blends, polylactic acid, and bacterial polyhydroxyalkanoates (PHAs). PHAs are described as natural polyesters [11]. Copolymers of poly(\(\beta\)-hydroxybutyrate-\(\text{-co}\)-\(\beta\)-hydroxyvalerate) were produced on a large scale by the Imperial Chemical Industries in Billingham, United Kingdom [12, 13]. This copolymer is a thermoplastic which resembles polypropylene [14] and has been successfully test marketed as shampoo bottles in Europe [15]. Although this natural product has great promise, but its high cost is a limiting factor [10]. The large-scale production of bacterial PHA has a number of interesting problems. Major expenses in the production of PHA are determined by the cost of the fermentation substrate, and the extraction of the polymer from inside the cell [12]. Dunlop and Robards (1973) developed the first models for the structure of the inclusion with intriguing observation that this polymer was stretchable even at very low freeze-fracture temperature [16]. PHB was in fact a thermoplastic with properties much like polypropylene [14]. It was shown that PHB copolymers that contained an amount of \(\beta\)-hydroxyvalerlate and designated P(HB\(\text{-co}\)-HV) could be formed by co-substrate feeding [17]. 3-Hydroxyvalerate synthesis is dependent on the presence of both glucose and propionate [18]. For PHAs nomenclature and abbreviation refer to Amara [2]. Many years have been lost after the first bulk interest with this polymer shown by Lemoigne and his group. However, minor researches have been made. Weibull had isolated the granules of \textit{Bacillus megaterium} by dissolution of the cell wall with a lysozyme [21]. Doudoroff and Stanier and Stainer et al. found that PHB can be produced by photosynthetic assimilation of organic compounds by phototrophic bacteria [21, 22]. They described that the reaction involved in the metabolic pathway is responsible for the biosynthesis of PHB from acetic acid [22, 23]. Lusty and Doudoroff have shown a study on depolymerases, which were able to hydrolyze PHB [24]. Schlegal observed that \textit{Cupriavidus necator} (formally \textit{Wautersia eutropha}, \textit{Ralstonia eutropha} and \textit{Alcaligenes eutrophus}), could accumulate very large amount of PHB in a media with low nitrogen content [25]. The most challenging work on PHAs started when Schlegal working with \textit{C. necator} and Dawes working with \textit{Azotobacter berjerinckii} succeeded to isolate and characterize the enzymes catalyse the PHB monomer synthesis [26, 27]. Isolating
the PhaC synthases was the real step to facilitate the cloning of their responsible genes. The PHA synthase operon of \textit{C. necator} were cloned in three labs nearly in a short span of time [28-30]. Wallen and Rohwedder reported heteropolymers of HB and 3-hydroxyvalerate in chloroform extracts of activated sewage sludge as major constituents with C6 and possibly C7 3-hydroxyacids as minor components [31]. De Smet et al. characterized PHA\textsubscript{MCL} in \textit{Pseudomonas oleovorans} during growth on octane [32]. Huisman et al. proved that PHA\textsubscript{MCL} accumulation was the common criteria of fluorescent pseudomonads [33]. PHAs are also produced by gram positive, negative and phototrophic bacteria as well as archaea [34-39].

![Fig. 1: PHAs metabolic different pathways (modified after Amara [2, 19]).](image)

### 2. PHAs PRODUCTION

Examination of the enzymes leading to the formation of PHB has shown that the regulation of PHB synthesis is controlled by acetyl-Co-enzymes A (CoA) acyltransferase [26, 40, 41]. Under balanced growth conditions, acetyl-CoA is fed into the tricarboxylic acid cycle and the resultant CoA inhibits acetyl-CoA acyltransferase and PHB synthesis. Under nutrient limitation (e.g., oxygen limitation) and carbon excess, NADH oxidase activity decrease, NADH increase and inhibits citrate synthase and isocitrate dehydrogenase, and acetyl-CoA
acyltransferase by CoA is overcome. The ensuing condensation reaction forms acetoacetyl-CoA and initiates PHB synthesis [26, 40]. The high NADH/NAD ratio caused oxygen limitation rapidly readjusts as PHB synthesis starts, and PHB assumes the role of an alternative electron acceptor [40]. The polymer is normally accumulated in response to the limitation of an essential nutrient and much of the past work has concerned the control of the synthesis of PHB under in-balanced growth condition [26, 41]. PHB is the most broadly studied PHAs. The PHB biosynthesis pathway consists of three distinct enzymatic reactions, as depicted in Fig. 1. The regulation of PHB synthesis is controlled by acetyl-coenzymes A (CoA) acyltransferase [40, 42]. The first reaction comprises the condensation of two acetyl coenzyme A (acetyl-CoA) molecules into acetoacetyl-CoA by $\beta$-ketoacyl-CoA thiolase ($\beta$-ketothiolase) (encoded by phaA). The second reaction is the reduction of acetoactyl-CoA to (R)-3-hydroxybutyryl-CoA by an NADPH-dependent acetoacetyl-CoA dehydrogenase (encoded by phaB). Finally, the (R)-3-hydroxybutyryl-CoA monomers are polymerized to P(3HB) by P(3HB) synthase (encoded by phaC). Synthesis of PHB is believed to be controlled by the first enzyme, 3-ketothiolase that is inhibited by coenzyme A (CoASH). Synthesis of PHAs other than P(3HB) which are not related to their growth substrates occurs relatively rare in nature [42]. Antonio et al. and Amara et al. produced 3HB-3HHX from cultivation of recombinant E. coli on fatty acid using PhaC class I and acrylic acid [44, 45]. 4-hydroxybutyrate (4HB) was produced in recombinant E. coli using succinyl-CoA from the citric acid cycle, which is converted to 4-hydroxybutyryl-CoA, a precursor for PHA biosynthesis [46].

Sixty years after the discovery of PHB, De Smet described the accumulation of the PHAMCL in Pseudomonas oleovorans grown on octane as sole carbon source [32]. PHA from pseudomonads rRNA homology group I were directly related to the structure of the alkane, alkene or fatty acid they used as carbon source [33, 47]. These substrates called ‘related substrates’. When the carbon source consists of fatty acids with 6 to 12 carbon atoms, monomers of the PHA are of the same length as the carbon source or are shortened by 2, 4 or 6 carbon atoms. Pseudomonads are able to produce PHAMCL from sugar. These substrates, which do not resemble the monomers of the accumulated polymer, are named “unrelated substrates”. Pseudomonads of the rRNA homology group I like Psudomonas putida KT2442, accumulate PHA that consists primarily of C10 and C8 monomers, when grown on sugars or gluconate [48, 49]. Experimental evidence suggests that these monomers are derived from intermediates of fatty acid biosynthesis and that the composition of the PHAs is most likely a reflection of the pool of fatty acid biosynthetic intermediates. Reush describes native low molecular mass PHB in yeast and many other eukaryotic cells [50].

Expression of the C. necator PHB biosynthetic pathway was successfully achieved in the yeast Saccharomyces cerevisiae by Leaf et al. [51]. He reported that the expression of PHB synthase in the yeast cytoplasm is sufficient for PHB accumulation and that wild type yeast synthesis D-3HB-CoA could raise from intermediates in fatty acid synthesis or through $\beta$-oxidation [51]. Yeast cells can be used as models to gain information about PHAs synthesis in eukaryotes [52]. It is confirmed that three thiolas exist in S. cerevisiae functioning in mitochondria, cytoplasm and peroxisomes [53, 54]. Safake et al. in their study succeeded to
isolate PHB producing wild type yeast strain from Kombucha tea and reported that the amount of PHB is less than that from prokaryotic and the PHB amount depend upon the type of culture and species [55]. Simple strategy for biosynthesis of polyhydroxyalkanotes in wild type yeasts has been described [56]. Synthesis of PHA<sub>SCL</sub> was established in the peroxisome of a wild-type yeast strain by targeting the <i>C. necator</i> SCL polymerase to the peroxisome [57]. Other strategies for using yeasts other than <i>S. cerevisiae</i> have been reported [58, 59]. Plants are able to produce PHAs directly from CO<sub>2</sub> and solar energy after being transgenic [50]. Steinbüchel and Füchtenbusch reported that the costs of the production of PHAs in transgenic plants could be of the same order of magnitude of their substrate costs (eg. sucrose, lipids and starch), if all the prerequisites are met [60]. PHB has been produced in transgenic <i>Arabidopsis thaliana</i> [61], <i>Brassica napus</i> [62], <i>Gossypium hirsutum</i> [63], <i>Nicotiana tabacum</i> [64] and <i>Zea mays</i> [52]. The copolyester PHB-co-PHV [65] and PHAs consisting of 3HA<sub>MCL</sub> [66] have been produced too. Recently Petrasovits et al. succeeded to produce of polyhydroxybutyrate in sugarcane [67].

3. CLASSES OF PHA SYNTHASES

PHA synthases were classified into four classes based on their substrate specificity and subunit composition. The multiple alignment of the primary structures of these PHA synthases show an overall identity with few conserved amino acids residues (Fig. 2). However, each synthase from the same class is clustered in one group as in Fig. 3. General model for each class can be performed as illustrated in Fig. 4. Only class IV is clustered with class III while class III is so close to it. Class I PHA synthases, with <i>C. necator</i> synthase as prototype, are composed of one single type of polypeptide chain and use mainly (R)-3-hydroxybutyryl-CoA, (R)-3-hydroxyvaleryl-CoA and other short-carbon chain length hydroxyalkanoic acid CoA thioesters including 3-mercaptoalkanoic acid CoA thioesters as substrates [68, 69]. Class III PHA synthases, as represented by the <i>Allochromatium vinosum</i> enzyme, are composed of two different subunits each of about 40 kDa [70]. The substrate specificity of class III is similar to that of class I synthases, although some MCL 3-hydroxyfatty acids are also incorporated [71]. Class II enzymes as well as class I PHA synthases are composed of only one type of subunit. Class I and II were purified, and there <i>in vitro</i> activities have been determined [72-74]. Class II mainly found in pseudomonades such as <i>P. aeruginosa</i>. One major difference between class II and both of class I and III, PHA synthases is the substrate specificity.

Class II PHA synthases incorporate preferentially 3-hydroxyfatty acids of MCL (C6-C14) into PHAs, and the resulting product is a latex-like polymer [74-76]. These substrates are mainly derived from intermediates of fatty acid β-oxidation or from fatty acid de novo biosynthesis provided fatty acids or simple non-related carbon sources [78, 80-83]. Class II PHA synthases were purified and <i>in vitro</i> activity was determined [83, 84]. A threading model of the <i>A. vinosum</i> PHA synthase was generated [85]. Accordingly, a threading model of the class II PHA synthase from <i>P. aeruginosa</i> was developed with the aid of the SAMT98 algorithm and based on the homology to the structure of the lipase from <i>B. cepacia</i> [86].
Amara and Rhem reported that H479 is not the general base catalyst that activates the nucleophilic C296 for covalent catalysis and that H452 plays a major role [87]. In the multiple alignment of PHA synthases only one cysteine residue (Cys-319) is highly conserved. Therefore, scientists have been searching for several years for the second thiol group. The essential role of Cys-319 of the C. necator PHA synthase (Class I) for the reaction mechanism was obtained from site specific mutagenesis [72]. The weakly conserved Cys-459 was supposed to be involved in the catalytic cycle, providing the second thiol group. However, site specific mutagenesis [72] clearly indicated that this amino acid residue is not essential for catalytic activity. In position 260 of the PHA synthase from C. necator a conserved serine residue was observed. Ser-260 was proposed as target for covalent posttranslational modification by 4-phosphopantetheine, which should provide the second thiol group similar to fatty acid synthase. In order to investigate posttranslational modification by 4-phosphopantetheine, radiolabeling experiments were conducted, expressing PHA synthase gene from C. necator in E. coli SJ16 (panD). Since E. coli SJ16 is β-alanine auxotroph, specific radiolabeling of 4-phosphopantetheinylated proteins occurs, when cells were fed with 2\(^{14}\)C-β-alanine. These experiments indicated that the PHA synthase was labeled by 4-phosphopantetheine [72]. However, detailed analysis revealed that only a small portion of total PHA synthase was labeled [88]. Functional low level expression of PHA synthases from C. necator and P. aeruginosa, respectively, in E. coli SJ16 and also in β-alanine auxotrophic mutants of C. necator with subsequent analysis of 4-phosphopantetheinylated proteins gave no evidence for covalent posttranslational modification by 4-phosphopantetheine [89]. Exchange of amino acid residue Ser-260 against alanine and threonine, respectively, abolished in vivo and in vitro activity of PHA synthase from C. necator [89]. The model of active PHA synthase involves two subunits forming a homodimer considering class I and II PHA synthases, and forming a heterodimer (PhaC and PhaE) in case of class III PHA synthases. Accordingly, class I and II PHA synthases possess two thiol groups provided by conserved Cys-319, and in class III PHA synthases the second thiol group might be provided by conserved Cys-130 of subunit PhaE from A. vinosum [88]. A novel PHA synthase (class V) from Bacillus megaterium required PhaC\(_{Bm}\) and PhaR\(_{Bm}\) for activity in vivo and in vitro was reported [90]. PhaC\(_{Bm}\) showed greatest similarity to the PhaCs of class III in both size and sequence (Figure 3, 4). Unlike those in class III, the 40 kDa PhaE was not required, and furthermore, the 22 kDa PhaR\(_{Bm}\) had no homology to PhaE.

4. PHAs BIODEGRADATION

Lusty and Doudoroff reported the presence of depolymerase activities in Pseudomonas lemoignei ans was able to degrade PHB [24]. While depolymerases have been studied for about 45 years, the first structural gene of a PHA depolymerase (Alcaliginus faecalis, phaZ\(_{Af}\)) was cloned and sequenced only in 1989 by Saito et al [91]. Depolymerases are highly specific for the polymers consisting of monomers in the (R) configuration [92]. PHAs depolymerase are carboxyesterases (EC 3.1.1) and hydrolyze the water insoluble polymer to water-soluble monomers and/or oligomers and lastly to water and carbon dioxide or methane. Early PHB degrading bacteria were isolated by selection for microorganisms able to utilize PHB as the
sole source of carbon and energy [1, 2]. PHAs degrading bacteria differ from each other depending on the type of PHAs they degrade, however some bacteria revealed a rather broad polyester specificity and are able to utilize a wide range of PHAs [93, 94]. Three intracellular PHA depolymerases (PhaZ1, PhaZ2 and PhaZ3) and a 3HB-oligomer hydrolase (previously designated PhaZ2), which hydrolyse PHAs and the cleavage products produced by PhaZs, were cloned and characterized in *C. necator* [94-97]. Molds also could degrade PHAs and their depolymerases have been characterized [98, 99]. PHAs depolymerases show a similar characteristic, they are stable at a range of pH, temperature, relatively small Mr (< 70 KDa), they are inhibited by reducing agent, e.g. dithioerythritol (DTT), which indicates the presence of essential disulfide bonds, and by serine hydrolase disulfide bonds, and serine hydrolase inhibitors such as diisopropyl-fluorophosphate (DFP) or acylsulfonyl derivatives [99].
Fig. 2: Part of the multiple amino acids alignment of different PhaC synthases, shaded lines show different conserved regions.
Depolymerases have three strictly conserved amino acids: serine, aspartate and histidine. The serine is part of the lipase-box pentapeptide Gly-Xaa1-Ser-Xaa2-Gly [100]. The oxygen atom of the serine side chain is the nucleophile that attacks the ester bond [93]. Many strategies were describing using depolymerases in the biotechnological products using efficient microbial strains [19, 101, 102]. Ihssen et al. show a study for using extracellular PHA_MCL depolymerase for targeted binding of proteins to artificial Poly[(3-hydroxyoctanoate)-co-(3-hydroxyhexanoate)] granules [103].

5. PROTEIN ENGINEERING OF PHAC SYNTHASE

Tha PhaC synthases show a great homology (as in Fig. 2) to each other and the conserved amino acids have been highlighted (Fig. 2). Aiming to change the PhaC substrate specificity, improve their activities’, determine their active sites’ and discover new futures, different tools of genetic and protein engineering tools have been implemented to the genes encoded different PhaC synthases [2]. Aiming to enhance the PhaC synthase activity Amara et al. establish a protocol for in vivo random mutagenesis of
the *phaC*<sub>Ap</sub> (*Aeromonac punctata*) [44, 45]. Four mutants exhibited enhanced *in vivo* and *in vitro* PHA synthase activity. The *in vitro* activities of the overproducing mutants ranged from 1.1 to 5-fold of the wild type activity, whereas the amounts of accumulated PHA ranged from 107%-126% to that of the wild type [44, 45]. A list of mutants and their site in the *phaC*<sub>Cn</sub> have been described in details by Rehm et al. and their related effect on PhaC<sub>Cn</sub> structure/activity have been discussed [86]. An attractive observation (in the work done by Amara et al. was the increase in weight average molecular mass of the PHA synthesized by the modified PHA synthase, which suggest a higher processivity of the enzymes with less frequently occurring chain transfer reactions [44, 45]. Slight preference of C4 over C6 was observed in the mutants as compared to the wild-type which keep the door open for the possibility of changing the substrate specificity [44, 45].

Taguchi et al. have been used another strategy based on *in vitro* mutagenesis for *phaC*<sub>Cn</sub> using PCR [105, 106]. Seven site directed mutations have been performed in PhaC<sub>Pa</sub> where five conserved residues were replaced by site directed mutagenesis in order to identify the function of these amino acids in catalysis. The substitution of W398 by alanine abolished PHA<sub>MCL</sub> synthase activity indicating that W398 is essential for enzyme activity [87]. The substitution of H479 by glutamine did not affect the PHA<sub>MCL</sub> activity, which indicates that H479 is not the general base catalyst that activates the nucleophilic C296 for covalent catalysis. The substitution of C296 with serine, which is the general base catalyst in lipases, did not abolish PHA<sub>MCL</sub> synthase activity while still exhibiting 20% activity in comparison to the wild type. Substitution of C296 with alanine abolished PHA<sub>MCL</sub> synthase activity which is a strong evidence that serine may be able to replace cysteine as catalytic nucleophile for changing substrate specificity of PHA synthase. For further investigation, another site directed mutagenesis has been done in another conserved histidine (H452), which was replaced with glutamine. The H452Q mutant was highly impaired in PHA<sub>MCL</sub> activity, which indicates that H452 plays a major role as a general base catalyst instead of H479 [87]. Hu et al. (2007) reported an improve for fractions by using site directed mutagenesis. The recombinant *E. coli* harboring plasmid pETJ1 (L65A), pETJ2 (L65V) or plasmid pETJ3 (V130A) synthesized the enhanced 3HHx fractions of PHBHHx from dodecanoate, indicating that Leu-65 and Val-130 of PhaJ(Ah) play an important role in determining the acyl chain length substrate specificity. The mutated PhaJ (Ah) (L65A, L65V, or V130A) provided higher 3HHx precursors for PHA synthase, resulting in the enhanced 3HHx fractions of PHB-co-HHx [107]. Fusion proteins composed of the N terminal part of the PHA synthase from *P. aeruginosa* and the C terminal part of the PHA synthase from *C. necator* indicated that fusion points located in the α/β-hydrolase fold region are not tolerated. Furthermore, these fusion points were located in predicted and structurally conserved α-helical regions [104].

Matsumoto et al. reported a chimeric enzyme composed of polyhydroxyalkanoate (PHA) synthases from *C. necator* and *Aeromonas caviae* enhances production of PHAs in recombinant *E. coli*. [107]. An *P. putida* mutant was isolated which is able to grow normally on any carbon source tested and which synthesized PHA<sub>MCL</sub> from fatty acids but not from gluconate or glucose [78]. Biochemical studies on the affected enzyme revealed that this enzyme catalyses the transfer of *(R)*-3-hydroxydecanoyl moieties from the acyl carrier protein: coenzyme A transferase. The respective gene, which complemented the mutant, was referred to as *phaG* [78], which has been also identified and cloned from *P. aeruginosa* [81] and *P. oleovorans* [82]. The PhaG protein overproduced in *E. coli* as a C-terminal His<sub>6</sub>-tagged fusion protein. The His<sub>6</sub>-tagged PhaG was purified to homogeneity by refolding of PhaG obtained from inclusion bodies, and a new enzyme assay was established. Gel filtration chromatography analysis in combination with light scattering
analysis indicated substrate-induced dimerization of the transacylase. In the presence of the substrate (R)-3-hydroxyacyl-CoA the transacylase appeared in apparent molecular weights of 45.4, 83.7 and 102 kDa, which indicated substrate-induced conformational changes and oligomerization of the transacylase [108]. A threading model of PhaG was developed based on the homology to an epoxide hydrolase (1cq2). In addition, the alignment with the α/β-hydrolase fold region indicated that PhaG belongs to α/β-hydrolase superfamily [108]. Accordingly, CD analysis suggested a secondary structure composition of 29% α-helix, 22% β-sheet, 18% β-turn and 31% random coil that support the PhaG protein model. Using hanging drop methods Amara, (2003) obtain crystals from the purified PhaG [20]. PhaG is responsible for diverting intermediates of fatty acids to PhaC synthase to produce new polymers of PHA_{SCL/MCL}.

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