Enzymic and Genetic Control of Polyphosphate Accumulation in *Aerobacter aerogenes*

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SUMMARY

Addition of orthophosphate to *Aerobacter aerogenes* strain A8(0) organisms previously subjected to phosphate starvation induced accumulation of inorganic polyphosphate within the organisms. With resumption of growth the polyphosphate was degraded and served as a source of nucleic acid phosphorus. During phosphate starvation the specific activity of polyphosphate kinase and inorganic polyphosphatase increased five- to tenfold while the amount of alkaline phosphatase increased 50 times. The results suggest that synthesis of polyphosphate kinase and alkaline phosphatase was subject to repression by exogenous orthophosphate. Two mutant strains blocked in polyphosphate accumulation were found to carry defects in the synthesis of these enzymes. Mutants of class Pn-1 contained normal amounts of all three enzymes, but repression of their synthesis was not annulled by phosphate starvation. Mutants of class Pn-2 contained no polyphosphate kinase. It is suggested that synthesis of polyphosphate kinase is controlled by two genes, a structural gene and a regulator gene; the latter gene also appears to control the synthesis of alkaline phosphatase and perhaps polyphosphatase. The patterns of polyphosphate accumulation under various nutritional conditions are discussed in relation to the amounts and activities of the enzymes of polyphosphate synthesis and degradation.

INTRODUCTION

Jeener & Brachet (1944) discovered that addition of orthophosphate to phosphate-starved yeast resulted in rapid uptake of phosphate and accumulation of a basophilic substance within the organisms. Originally thought to be ribonucleic acid (RNA), the basophilic material was later identified as inorganic polyphosphate (Wiam, 1947; Schmidt, 1951). Its accumulation corresponded to the appearance of structures known as 'volutin granules'; both disappeared together when growth of the organisms was resumed. This basic pattern has since been described in various fungi and bacteria (Kuhl, 1960; Harold, 1962; Liss & Langen, 1962; Smith, Wilkinson & Duguid, 1954; Zaitseva, Belozerskii & Novozhilova, 1960; Kaltwasser, 1962). The term 'polyphosphate overplus' is proposed to designate the accumulation of polyphosphate upon addition of orthophosphate to phosphate-starved organisms, in order to distinguish this from polyphosphate accumulation under other conditions of nutrient imbalance (Smith *et al.* 1954). The term corresponds to 'Polyphosphat-Überkompensation' as used by Liss & Langen (1962).

The physiological basis of the polyphosphate overplus has remained obscure despite repeated investigation. Liss & Langen (1962) found an increased rate of
polyphosphate synthesis in yeast previously subjected to phosphate starvation and attributed this to an imbalance of phosphate uptake and utilization. In the present paper evidence will be offered to relate the polyphosphate overplus to the control of the enzymes of polyphosphate metabolism in wild-type *Aerobacter aerogenes* and in mutants of it which carry genetic defects in polyphosphate synthesis. Preliminary accounts of these results have been published (Harold, 1963a, b).

**METHODS**

**Organisms.** *Aerobacter aerogenes* strain A3 (0) was obtained from Professor J. F. Wilkinson. The isolation of auxotrophic mutants requiring uracil (Slu) and methionine (0met) was described by Harold (1963c). Mutants defective in polyphosphate accumulation were isolated by a 32P-suicide technique (Harold & Harold, 1963). These mutants, were of two types: (i) Pn-1 mutants which did not show the overplus effect but accumulated polyphosphate upon sulphur starvation; (ii) Pn-2 mutants which did not accumulate polyphosphate under any conditions. Most of the present experiments were conducted with the strains 1025(Pn-1) and 1023c (Pn-2) previously used.

**Growth experiments.** The bacteria were grown in Tsubertg media as described earlier (Harold, 1963c; Harold & Harold, 1963). This is a mineral medium containing sulphate as sole source of sulphur, glucose as carbon+energy source and buffered with tris (pH 7-6, 10 g./l.). The subscript indicates the phosphorus (P) content as µg. P/ml. All cultures were grown on a rotary shaker at 37°. The population density was normally about 10⁹ cells/ml.; multiplication was followed turbidimetrically at 600 mp. Overnight cultures were grown in T 230 medium; the organisms were then harvested, washed, and resuspended in fresh pre-warmed T medium containing orthophosphate and growth factors as described for the individual experiments. Samples were withdrawn periodically and the organisms centrifuged down. Samples intended for enzyme assay were washed with chilled tris buffer. Phosphorus analyses were performed on unwashed organisms. The fractionation and estimation of phosphorus compounds was described earlier (Harold, 1963c). The bulk of the polyphosphate was acid-insoluble polyphosphate; acid-soluble phosphate, when present, was measured separately and the two fractions added to give the amount of total inorganic polyphosphate.

**Enzyme assay.** Bacterial cells from 30 ml. culture were resuspended in 1 ml. tris buffer (0-1 M, pH 7-0) in a plastic tube. Glass beads (Superbrite, Minnesota Mining and Manufacturing Co., Type 115; 0-50 g.) were added, and the tube exposed for 20 min. to sonic treatment in a Raytheon 10 Kc oscillator cooled with ice water. The tubes were then centrifuged at 4° for 10 min. at 20,000 g; the supernatant fluid was used for the estimation of polyphosphate kinase as described below, and for protein estimation by the biuret method (Layne, 1957).

Assay of polyphosphatase activity in these extracts gave variable results. As a rule, high activity was found only in extracts of bacterial cells which had been allowed to accumulate polyphosphate following phosphate starvation. This apparent production of polyphosphatase in response to polyphosphate accumulation was ultimately found to be an artifact: the enzyme is readily lost, apparently by adsorption to the glass beads, unless inorganic polyphosphate is present. Consequently,
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for the assay of polyphosphatase the bacterial cells were sonically treated in tris buffer containing polyphosphate (synthetic, chain length about 170, 100 µg./ml. P).

**Polyphosphate kinase.** This enzyme was assayed in crude extracts by a modification of the procedure described by Muhammed (1961). It had an obligatory requirement for Mg$^{2+}$ and an optimum at about pH 7. Adenosine diphosphate (ADP) was inhibitory at concentrations above $2 \times 10^{-5}$ M, but polyphosphate itself had little effect at $15 \times 10^{-5}$ M. Each assay tube received 2 µmole acetyl phosphate-$^{32}$P, 2 µmole MgCl$_2$, 1-5-5 units acetokinase (Rose, 1955), 0-2 µmole ADP, 25 µmole glycylglycine or tris buffer (pH 7-0), and 0-05-0-10 ml. enzyme extract in a final volume of 0-38 ml. The tubes were incubated for 10 min. at 37°. At that time, 0-25 ml. N-perchloric acid was added and then 0-50 ml. bovine serum albumin solution (5 mg./ml.) to precipitate the polyphosphate formed. The precipitate was washed twice with 0-5 N-perchloric acid, dissolved in 0-5 N-NaOH and plated for counting. The course of the reaction was linear with time for 20 min. and proportional to enzyme concentration over the range of 0-02 to 4 units/tube. The unit is defined as that amount of enzyme which produces 0-01 µmoles polyphosphate in 10 min. The identity of the product was established by its behaviour in the standard fractionation scheme and by its complete lability to acid (N-HCl, 15 min., 100°).

**Polyphosphatase.** The polyphosphatase activity of crude extracts had a broad pH optimum between 7 and 9. A requirement for Mg$^{2+}$ was observed only in the presence of EDTA, but high concentrations of KCl (about 0-8 M) were necessary for maximal activity. No dialysable cofactors were detected. The assay of polyphosphatase was modified from that described by Muhammed, Rodgers & Hughes (1959). Each assay tube received 100 µmole KCl, 0-25 µmole MgCl$_2$, 1-6 µmole polyphosphate-$^{32}$P, 10 µmole tris buffer (pH 9) and 0-05 ml. enzyme extract in a final volume of 0-30 ml. (the amount of polyphosphate introduced with the extract was neglected). The tubes were incubated at 37° for 30 min., then excess polyphosphate and protein were precipitated by adding 1-0 ml. 0-5 N-perchloric acid and 0-1 ml. bovine serum albumin (10 mg./ml.). The tubes were centrifuged and the supernatant fluids decanted. After addition of ammonium molybdate the $^{32}$P-orthophosphate was extracted with isobutanol (Weil-Malherbe & Green, 1951), plated and counted. The hydrolysis of polyphosphate was linear with time for 60 min. and proportional to the enzyme concentration over a wide range. A unit of polyphosphatase is defined as that amount of enzyme which liberates 0-01 µmole $^{32}$P-orthophosphate in 10 min.

**Alkaline and acid phosphatase.** These enzymes were in general assayed with intact *Aerobacter aerogenes* organisms as described by Torriani (1960). For consistency, the unit is defined as that amount of enzyme which liberates 0-01 µmole orthophosphate from o-nitrophenyl phosphate in 10 min.

**Chemicals.** Acetyl phosphate-$^{32}$P was synthesized as described by Kornberg (1957), polyphosphate-$^{32}$P by the method of Muhammed *et al.* (1959). Other reagents were purchased from Sigma Chemical Co. and the California Corporation for Biochemical Research.

**RESULTS**

The polyphosphate overplus in wild-type and mutant strains of *Aerobacter aerogenes* A 3(O). *Aerobacter aerogenes* organisms in the logarithmic or the stationary phase of growth contained only traces of polyphosphate. When orthophosphate was added to organisms which had been incubated in phosphate-deficient medium
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for 3–4 hr dramatic accumulation of polyphosphate occurred, together with the appearance of microscopically visible volutin granules (Smith et al. 1954). Induction of this polyphosphate overplus was dependent upon protein synthesis during the starvation phase: no polyphosphate accumulation was observed when orthophosphate was added to strain A3(0) organisms subjected to phosphate starvation in absence of glucose or of sulphur, or to organisms of strain 0\textsubscript{net}-incubated in medium lacking orthophosphate and methionine. Protein synthesis was not, however, required for polyphosphate accumulation itself since chloramphenicol (10 \(\mu\text{g.}/\text{ml.}\)), sulphur deficiency or methionine deprivation had no effect once starvation was complete.

![Graph](image)

Fig. 1. The polyphosphate overplus in Aerobacter aerogenes strain A3(0). The organisms were subjected to phosphorus starvation for 4 hr, collected, washed and resuspended in T\textsubscript{9} medium with and without sulphate. Orthophosphate was added to both flasks at 0 min.

- Solid line, Polyphosphate; Open circle, total nucleic acids.

A typical experiment illustrating the accumulation of polyphosphate and its subsequent fate is shown in Fig. 1. Aerobacter aerogenes A3(0) organisms were subjected to phosphorus starvation in T\textsubscript{9} medium for 4 hr. The bacteria were then harvested, washed, and resuspended in T\textsubscript{100} medium, with and without sulphate. Both cultures rapidly accumulated polyphosphate and smaller amounts of nucleic acids. In the flask supplemented with sulphate, growth of the bacteria resumed and was accompanied by extensive nucleic acid synthesis and degradation of polyphosphate. In the sulphur-deficient culture growth was prevented, and a high polyphosphate concentration was maintained. The degradation of polyphosphate upon resumption of growth was shown to be a consequence of nucleic acid synthesis. In the presence of chloramphenicol protein synthesis and growth were inhibited, but nucleic acids accumulated within the bacteria and polyphosphate was concurrently degraded. In a typical experiment, organisms of the uracil-requiring mutant Sl\textsubscript{u}, were subjected to phosphate starvation; the bacteria were then...
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harvested and allowed to accumulate polyphosphate in radioactive T190 medium without uracil (to prevent nucleic acid synthesis). After 1 hr the 32P-orthophosphate was diluted with excess unlabelled orthophosphate, and uracil + chloramphenicol (10 µg./ml.) were added. Nucleic acid accumulated within the bacteria while the 32P-polyphosphate was degraded and the 32P transferred to the nucleic acid fraction. These results are analogous to those described previously for the degradation of polyphosphate accumulated during nutrient deprivation (Harold, 1963c).

Mutants defective in polyphosphate accumulation showed quite a different pattern of phosphorus metabolism. *Aerobacter aerogenes A3(0), Pn-1 and Pn-2* organisms were subjected to phosphorus starvation for 4 hr. The bacteria were then transferred to sulphur-deficient medium and 32P-orthophosphate (100 µg./ml.) was added. As illustrated in Fig. 2, some nucleic acid was synthesized by all three strains but only strain A3(0) accumulated polyphosphate. Mutant and wild-type bacteria incorporated about equal amounts of 32P into the orthophosphate and acid-soluble organic phosphate fractions.

**Differential synthesis of enzymes of phosphate metabolism in *Aerobacter aerogenes* during phosphate starvation.** From the results presented in the preceding section it is clear that the overplus phenomenon is a consequence of events which occur during the starvation phase and require protein synthesis. Clues to the nature of these events came from studies on the amount of polyphosphate kinase in wild-type and mutant *Aerobacter aerogenes* under various growth conditions. When bacteria of strain A3(0) were incubated in phosphate-deficient medium, the optical density and protein content of the culture increased by about 50 %, and concurrently there was a marked increase in the specific activity of polyphosphate kinase. This increase in enzyme content was abolished by chloramphenicol and, in a methionine auxotroph, was dependent upon the presence of methionine (Fig. 3). The differential increase in the specific activity of polyphosphate kinase was induced specifically by phosphorus starvation: sulphur starvation and amino acid deprivation in auxotrophic mutants had no such effect. Upon addition of orthophosphate to phosphate-starved bacteria the differential rate of enzyme synthesis decreased until the value characteristic of growing bacteria (2-5 units/mg. protein) was attained. These results suggest that the synthesis of polyphosphate kinase is subject to repression by extracellular orthophosphate and thus proceeds at a higher differential rate during phosphate starvation.

The two polyphosphateless mutants exhibited clear defects in the synthesis of polyphosphate kinase. Organisms of strains A3(0), Pn-1 and Pn-2 were grown on T250 medium and then transferred to T0 medium. As shown in Fig. 4, de-repression of polyphosphate kinase occurred only with strain A3(0). Mutant strains of class Pn-1 contained normal amounts of enzyme (2-4 units/mg. protein) in growing organisms, but there was no increase in specific activity during phosphorus starvation. Mutants of class Pn-2 contained no detectable polyphosphate kinase under any nutritional conditions. No evidence was obtained for destruction of polyphosphate kinase in these extracts, nor for the presence of an inhibitor.

The effects of nutritional conditions on the amount of polyphosphatase paralleled those described for polyphosphate kinase. Bacteria from overnight cultures contained very little polyphosphatase (2-4 units enzyme/mg. protein) and incubation of such organisms in medium deficient in sulphur produced no increase in its amount.
Fig. 2. Accumulation of polyphosphate and nucleic acids in *Aerobacter aerogenes* strains $\lambda B (0)$, Pn-1 and Pn-2. The bacteria were subjected to phosphorus starvation for 4 hr, collected, washed and resuspended in $T_6$ medium without sulphate. $^{32}$P-orthophosphate was added to all flasks at 0 min. Closed symbols, polyphosphate; open symbols, nucleic acids. $\bullet$, $\bigcirc$, Strain $\lambda B (0)$; $\Delta$, $\triangle$, mutant Pn-1; $\blacksquare$, $\square$, mutant Pn-2.

Fig. 3. Synthesis of polyphosphate kinase in *Aerobacter aerogenes* strain O$_{net}$ during phosphate starvation. Washed organisms were suspended in $T_6$ medium at 0 hr, and the culture distributed among three flasks. Flask no. 1 (---) was supplemented with methionine (90 $\mu$g./ml.); no. 2 (---) received no methionine; no. 3 (---) received methionine but chloramphenicol (CMP; 10 $\mu$g./ml.) was added after 1 hr.

Fig. 4. Effect of phosphate starvation on the amounts of polyphosphate kinase, polyphosphatase and alkaline phosphatase in *Aerobacter aerogenes* strains $\lambda B (0)$, Pn-1 and Pn-2. Bacteria from an overnight culture were collected, washed and resuspended in $T_6$ medium at 0 hr. $\bullet$---$\bullet$, strain $\lambda B (0)$; $\bigcirc$---$\bigcirc$, mutant Pn-1; $\Delta$---$\Delta$, mutant Pn-2.
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Bacteria harvested during the logarithmic phase of growth contained 6–10 units enzyme/mg protein. During incubation in phosphate-deficient medium, a substantial increase in the specific activity of polyphosphatase occurred in strain A3 (0) organisms and, to a lesser degree, in Pn-2 organisms; the specific activity of polyphosphatase in strains of class Pn-1 was not affected (Fig. 4). The production of polyphosphatase required protein synthesis and thus presumably reflected differential synthesis of the enzyme, but repression upon subsequent addition of orthophosphate was not unequivocally demonstrated.

Differential synthesis of alkaline phosphatase during phosphate starvation in Escherichia coli was demonstrated by Torriani (1960) and by Horiuchi, Horiuchi & Mizuno (1959), and shown to be due to repression of the synthesis of this enzyme by exogenous orthophosphate. It therefore seemed of interest to examine whether the synthesis of alkaline phosphatase and of the enzymes of polyphosphate metabolism might be under joint genetic control. As shown in Fig. 4, phosphate starvation induced extensive synthesis of alkaline phosphatase in organisms of Aerobacter aerogenes strains A3(0) and Pn-2, but not in organisms of strain Pn-1. The amount of acid phosphatase was unaffected.

DISCUSSION

Accumulation of inorganic polyphosphate in micro-organisms is generally associated with nutritional conditions unfavourable to growth. Two quite distinct procedures for the induction of polyphosphate accumulation in Aerobacter aerogenes were described by Smith et al. (1954). On the one hand, polyphosphate accumulates when growth of the organisms ceases because of lack of certain essential nutrients, e.g. in sulphur starvation. Deposition of polyphosphate under these conditions has been shown to be due primarily to the cessation of nucleic acid synthesis while assimilation of phosphate from the medium continues (Harold, 1963a). On the other hand, polyphosphate accumulates upon addition of orthophosphate to a phosphate-starved culture. This phenomenon, for which the term 'polyphosphate overplus' is proposed, is the subject of the present paper. The patterns of polyphosphate accumulation under these two nutritional conditions are quite different. Polyphosphate accumulation due to nutrient deprivation is relatively slow, and is prevented by concurrent nucleic acid synthesis as well as by other environmental factors which accelerate polyphosphate degradation (Harold & Sylvan, 1968). The polyphosphate overplus is more rapid and is quite independent of nucleic acid synthesis (Fig. 1) and of changes in the composition of the medium. The physiological basis of these differences appears to reside in the amount of polyphosphate kinase. Wild-type A. aerogenes organisms normally contain a low concentration of polyphosphate kinase, and the rate of polyphosphate synthesis in vivo during nutrient deprivation agrees well with that calculated from the amount of enzyme present. During phosphate starvation the specific activity of polyphosphate kinase increases up to tenfold, and a corresponding increase in the rate of polyphosphate accumulation is observed when orthophosphate is made available to the starved organisms. It thus appears that the rate of polyphosphate synthesis is directly proportional to the amount of polyphosphate kinase within the organisms. Results with mutants blocked in polyphosphate accumulation (Harold & Harold, 1968) support this conclusion. Aerobacter aerogenes mutants of class Pn-2 lack polyphosphate
kinase, and do not accumulate polyphosphate under any conditions. Hence polyphosphate kinase catalyses the main, if not the only, reaction responsible for polyphosphate synthesis. Mutants of *A. aerogenes* class Pn-1 contain normal amounts of polyphosphate kinase and are therefore capable of slow polyphosphate accumulation when subjected to nutrient deprivation. However, repression of the synthesis of this enzyme is not annulled by phosphate starvation and hence no polyphosphate overplus occurs upon addition of orthophosphate.

The intracellular concentration of polyphosphate must be a function of the rates of synthesis and of degradation. From the present results and from those reported earlier (Harold, 1963c), it seems clear that polyphosphate degradation is coupled to nucleic acid synthesis. Except under special nutritional conditions (Harold & Sylvan, 1963), little polyphosphate degradation occurs until growth and nucleic acid synthesis resume. The molecular basis underlying the control of polyphosphate breakdown is unknown. Studies with cell-free extracts have provided evidence for four possible routes for the mobilization of inorganic polyphosphate: transfer of phosphate to ADP by reversal of polyphosphate kinase (Kornberg, 1957); transfer of phosphate to adenosine monophosphate (Winder & Denneny, 1957); direct phosphorylation of hexoses by polyphosphate (Dirheimer & Ebel, 1962; Szymona, Szymona & Kulesza, 1962); hydrolytic degradation to orthophosphate by inorganic polyphosphatase. The limited information available from work with intact organisms (Hughes & Muhammed, 1962; Harold, 1962) seems to favour hydrolysis as the chief route of polyphosphate degradation. Should this prove to be the case in *Aerobacter aerogenes*, an explanation will be required for the accumulation of polyphosphate in organisms subjected to phosphate starvation despite their elevated content of polyphosphatase (Fig. 4).

It remains to consider the genetic control of polyphosphate metabolism. During phosphate starvation differential synthesis of alkaline phosphatase, polyphosphate kinase and polyphosphatase occurs in wild-type *Aerobacter aerogenes*; the synthesis of the first two enzymes is repressed when orthophosphate is restored. Two mutations were found to affect this metabolic region. Mutants of class Pn-1 contain all three enzymes but repression of their synthesis is not annulled by phosphate starvation. Mutants of class Pn-2 lack polyphosphate kinase, but synthesize alkaline phosphatase and a small amount of polyphosphatase in phosphate-deficient medium. It is thus reasonable to postulate two genes, a regulator gene which controls the synthesis of all three enzymes (Pn-1) and a structural gene for polyphosphate kinase (Pn-2). The diminished formation of polyphosphatase in Pn-2 shows that this simple picture is no more than a first approximation. Nevertheless, the finding that these three enzymes form a unit of genetic regulation suggests that they constitute also a physiological unit, functionally concerned with phosphate storage (Harold, 1963b).

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REFERENCES


