Inhibitory Action of TetrathionatEnrichment Broth

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Tetrathionate enrichment broth is a complex mixture of salts including iodides and other polythionates, but only thiosulfate (0.0736 M) and tetrathionate (0.0236 M) in combination were toxic for *Escherichia coli*. Individually, these two salts were not lethal. The lethal action of this thiosulfate-tetrathionate mixture affected only growing cells. A possible relationship between the lethality of the thiosulfate-tetrathionate mixture for a culture and its ability to reduce tetrathionate is suggested.

When isolating salmonellae from sparsely infected material, it is necessary to inoculate the material into enrichment media which allow the growth of salmonellae while restricting the undesirable coliform and other gram-negative bacteria. The enrichment media most often used are the selenite broth of Leifson (7) and the tetrathionate broth of Muller (11). Although these broths have been widely used and often modified, their mode of action is still unknown. Further improvements involving these media could be more effectively developed if the mechanism of inhibition was more fully understood.

Pollock and co-workers (4-6, 16, 17), in a series of studies on the inhibition by tetrathionate broth, concluded that tetrathionate was the most important constituent of the medium. They found that most salmonellae and *Proteus* reduced tetrathionate to thiosulfate, whereas *Escherichia coli* and the shigellae did not. Tetrathionate reductase was demonstrated in washed cells and shown to be adaptive. It was suggested that tetrathionate acted as an alternative hydrogen acceptor to oxygen and therefore extended the log phase of growth. LeMinor (8) and Papavassiliou et al. (13) examined numerous *Enterobacteriaceae* cultures for tetrathionate reductase and found that most salmonellae possessed it whereas *E. coli* did not. They suggested that possession of this enzyme could prove useful in differentiating members of the enteric group.

Several exceptions to the relationship between the ability to reduce tetrathionate and the ability to grow in tetrathionate broth have, however, been observed (4, 22). Although numerous combinations of thiosulfate and tetrathionate were examined for inhibition of growth after 18 to 24 hr, no early growth measurements were made.

This paper is a report on the relationship of tetrathionate reductase production by both growing and resting cells of several cultures of *Enterobacteriaceae* to their inhibition by tetrathionate broth. Further, the basic components of the medium and the polythionates formed by their interaction have been examined to determine their role in this inhibition.

MATERIALS AND METHODS

Cultures. The salmonellae were obtained from Alice Moran, Consumer and Marketing Service, USDA (AM), and from H. Ng, Western Utilization Research and Development Division (WU). Other cultures were obtained from Z. J. Ordal, University of Illinois (UI), and from our own culture collection (EU). Appropriate dilutions of 24-hr cultures grown in Trypticase Soy Broth (BBL) were used as inocula for all studies.

Growth media. The basal medium for all studies was 0.5% proteose-peptone (Difco) plus buffer. When the medium contained thiosulfate and tetrathionate in the proportions found in tetrathionate broth, a white colloidal-sulfur precipitate, indistinguishable from growth, developed. In these cases, 1% CaCO₃ was used as buffer. If growth could be measured turbidimetrically, phosphate buffer (pH 7.6) was used at a final concentration of 0.05 M. The basal medium including any salts except the thionates was sterilized by autoclaving.

Growth was measured either by turbidity in a Beckman model B spectrometer at 540 nm or by surface plate counts on Tryptic Soy Agar (TSA; Difco). Dilutions were made in 0.1% peptone (Difco) water. The plates were incubated at 37 C for 24 hr and counted. Negative plates were incubated longer to check for slow growing colonies.

In studies in which salmonellae and *E. coli* were
mixed, the viable Salmonella count was followed on Brilliant Green Agar (Difco) and the viable E. coli, by the most probable number method in Lactose Purple Broth (Difco).

Preparation of thionates. Sodium tetrathionate (Na₂S₄O₆·2H₂O) was prepared from sodium thiosulfate by iodine oxidation as suggested by Mellor (10). Potassium tetrathionate (K & K Laboratories, Plainview, N.Y.) was used in some studies.

Potassium trithionate (K₂S₅O₆·3H₂O) and potassium pentathionate (K₂S₅O₆·3H₂O) were prepared by the method of Palmer (12). All thionates were stored at −10°C in a desiccator. When used in growth or inhibition studies, they were prepared double-strength, sterilized by membrane filtration, and added to double-strength basal broth.

The purity of the thionates was checked by the method of Jay (3) and by paper chromatography (20). The paper chromatography method also was employed to quantitate the thionate formed by the thiosulfate-catalyzed breakdown of tetrathionate. The intensity of the spots was measured by procedures of Pollard et al. (15) by using a TLC Densitometer System (Photovolt Corp., New York). Because of the instability of the thionates, their purity was checked frequently by paper chromatography.

Tetrathionate reduction. Tetrathionate reductase was measured by two methods. (i) The method of Pollock and Knox (16) is an iodometric titration of the thiosulfate released from tetrathionate by washed cells grown on Tryptic Soy Agar (Difco). In this method, suspensions were mixed with tetrathionate and mannitol, and samples were titrated periodically. Induction time was calculated in the same manner as lag time (2). (ii) In the cultural method of LeMinor (8), a basal medium containing sodium tetrathionate (0.722%) is inoculated with the test organisms. At 24 hr, the broth is titrated with standard iodine solution. The results are compared to the control and expressed as either plus or minus. Some of the cultures also were tested for trithionate and pentathionate reduction by this method.

Resistance to tetrathionate. The gradient plate method (23) was employed to measure resistance to tetrathionate. The bottom layer was phosphate basal broth plus 1.5% agar (Difco) plus 2X tetrathionate (1.422%). The upper layer of phosphate basal broth plus agar was tempered, inoculated with the test organism, and poured over the slanted bottom layer. The length of growth along the concentration axis is a measure of the inhibitory concentration.

Temperature. The tetrathionate reduction and gradient plate studies were performed at both 37 and 43°C. All other studies and incubations were at 37°C.

Salts study. Tetrathionate broth base contains sodium thiosulfate which is converted to tetrathionate upon the addition of iodine dissolved in potassium iodide according to the reaction: 2Na₂S₂O₃·5H₂O + I₂ + [KI] → Na₂S₄O₆·2H₂O + 2NaI + [KI]. In practice, 0.1208 moles of sodium thiosulfate is reacted with 0.0236 moles of iodine dissolved in 0.0301 moles of potassium iodide. This yields 0.0236 moles of sodium tetrathionate plus 0.0472 moles of sodium iodide and 0.0301 moles of potassium iodide, with 0.0736 moles of sodium thiosulfate remaining unreacted. The molarity of sodium thiosulfate were calculated on the assumption that the salt used in preparing the medium is the pentahydrate. The four salts, sodium tetrathionate, sodium thiosulfate, potassium iodide, and sodium iodide, were tested for their effect on various bacteria.

Paper chromatography of the mixture of thiosulfate and tetrathionate showed that the thiosulfate catalyzed the breakdown of tetrathionate to other thionates. These thionates were quantitated, synthesized (12), and added to the basal broth to check their lethality.

Since sulfite is the first product formed when thiosulfate and tetrathionate react (20), the tetrathionate-thiosulfate mixture also was examined for the presence of sulfide and sulfate. Although no free sulfite was found, its lethal effect also was examined in phosphate basal broth.

RESULTS

Tetrathionate reduction. Figure 1 is representative of the responses obtained with different cultures when washed cells were added to the tetrathionate-mannitol suspension. The induction times obtained by this method for 63 strains of bacteria, primarily Enterobacteriaceae, are given in Tables 1 and 2. Most of the cultures were able to reduce tetrathionate, although 12 or more hours was required for some. Salmonellae,
positive appreciably thionatereductase. short the
An either times.
S. Serratia S. Salmonella
S. S. Salmontella S. thompson var. berlin AM40.
S. thompson AM39
S. senfsenberg WU-S8
S. blockley WU2004
S. typhimurium AM9

| Organism                     | Induction time (hr) |
|------------------------------|---------------------|
| Salmonella senfsenberg 775W  | 1                   |
| S. derby AM20                 | 1                   |
| S. derby AM21                 | 1                   |
| Pseudomonas aeruginosa EU314  | 1                   |
| Escherichia coli D EU306      | 1                   |
| Arizona sp. EU313             | 1                   |
| S. tennessee EU317           | 1                   |
| Salmonella sp. EU315         | 1.5                 |
| Salmonella sp. EU316         | 1.5                 |
| S. typhimurium EU318         | 1.5                 |
| S. bredeney AM112            | 1.5                 |
| S. newport EU320              | 1.5                 |
| S. typhimurium EU319         | 1.5                 |
| S. thompson var. berlin AM40  | 1.5                 |
| S. thompson AM39              | 2                   |
| S. senfsenberg WU-S8          | 2                   |
| S. blockley WU2004           | 2                   |
| S. typhimurium AM9            | 2                   |
| S. chester AM17               | 2                   |
| S. anatum AM78                | 2                   |
| S. cubana AM207               | 2                   |
| S. heidelberg AM16            | 2                   |
| S. typhimurium AM10           | 3                   |
| S. typhimurium AM11           | 3                   |
| S. meleagridis AM209          | 3                   |
| S. saintpaul AM108            | 3                   |
| S. paratyphi B AM3            | 3                   |
| S. oranienburg AM42           | 4                   |
| S. montevideo AM46           | 4                   |
| Klebsiella sp. EU309          | 4                   |
| Proteus sp. I EU310           | 6.5                 |
| Proteus sp. II EU311          | 7                   |
| Proteus sp. III EU312         | 7                   |
| S. typhi AM60                 | 11.5                |
| S. typhimurium AM12           | 12                  |
| S. infantis AM165             | 12                  |

| Organism                     | Induction time (hr) |
|------------------------------|---------------------|
| Pseudomonas sp. EU92         | 0.5                 |
| P. fragi EU43                | 0.5                 |
| P. ovalis EU36               | 0.5                 |
| Serratia marcescens EU279    | 5                   |
| Salmonella enteritidis       | 5.25                |
| Aerobacter aerogenes EU109   | 5.5                 |
| A. aerogenes EU110           | 5.5                 |
| Escherichia coli EU106       | 6                   |
| Gram-negative rod no. 1      | 6.5                 |
| S. choleraesuis AM34         | 7                   |
| S. choleraesuis var. kunzendorf | 7               |
| AM36                         | 7                   |
| E. coli B EU304              | 7                   |
| E. coli E EU307              | 7                   |
| S. abortusovis AM27          | 11.5                |
| Gram-negative rod no. 2      | 11.5                |
| S. pullorum AM75             | 12                  |
| S. paratyphi A AM1           | 12                  |
| Shigella sp. EU321           | 12                  |
| Bacillus subtilis UI-AI      | 12                  |
| B. cereus UI-T               | 12                  |
| A. aerogenes EU300           | 23                  |
| A. aerogenes EU301           | 23                  |
| Enterobacter sp. EU302       | 23                  |
| E. coli CC EU308             | 23                  |
| E. coli C EU305              | >30                 |
| E. coli A EU303              | >30                 |

Table 1. Induction times for tetrathionate reduction by various bacteria found to be positive by the cultural method

Table 2. Induction times for tetrathionate reduction by various bacteria found to be negative by the cultural method

| Organism                     | Induction time (hr) |
|------------------------------|---------------------|
| Pseudomonas sp. EU92         | 0.5                 |
| P. fragi EU43                | 0.5                 |
| P. ovalis EU36               | 0.5                 |
| Serratia marcescens EU279    | 5                   |
| Salmonella enteritidis       | 5.25                |
| Aerobacter aerogenes EU109   | 5.5                 |
| A. aerogenes EU110           | 5.5                 |
| Escherichia coli EU106       | 6                   |
| Gram-negative rod no. 1      | 6.5                 |
| S. choleraesuis AM34         | 7                   |
| S. choleraesuis var. kunzendorf | 7               |
| AM36                         | 7                   |
| E. coli B EU304              | 7                   |
| E. coli E EU307              | 7                   |
| S. abortusovis AM27          | 11.5                |
| Gram-negative rod no. 2      | 11.5                |
| S. pullorum AM75             | 12                  |
| S. paratyphi A AM1           | 12                  |
| Shigella sp. EU321           | 12                  |
| Bacillus subtilis UI-AI      | 12                  |
| B. cereus UI-T               | 12                  |
| A. aerogenes EU300           | 23                  |
| A. aerogenes EU301           | 23                  |
| Enterobacter sp. EU302       | 23                  |
| E. coli CC EU308             | 23                  |
| E. coli C EU305              | >30                 |
| E. coli A EU303              | >30                 |

with few exceptions, had very short induction times. The induction time was independent of the weight of cells, although the rate was proportional to cell mass once reduction had begun. An excess of thiosulfate equal to that present in the complete enrichment broth did not affect either induction time or rate of reduction by selected cultures. Incubation at 43°C did not appreciably affect induction time.

Also given in Tables 1 and 2 are the results from the cultural method of determining tetrathionate reductase. In general, bacteria with short induction times for washed cells were also positive by this method (Table 1). Several cultures were examined by the cultural method for their ability to reduce tri- and pentathionate; cultures positive for tetrathionate reduction were positive for the other two thionates and vice versa.

The Pseudomonas sp. and Aerobacter sp. were exceptions in that they were negative by the cultural method, but had short induction times (Table 2). When checked at frequent intervals during the 24-hr incubation period, no iodine-reacting compounds were detected even though the cells grew well.

**Gradient plate studies.** All cultures, tested at both 37°C and 43°C, grew at least half way across the gradient plates. This indicated the ability to grow in a concentration of tetrathionate equal to that found in tetrathionate broth.
Effect of enrichment-broth constituents on growth. The gradient plate studies and the ability of most cultures to reduce tetrathionate suggested that the inhibitor in tetrathionate broth was not the tetrathionate alone. Therefore, the ability of various combinations of the four salts in tetrathionate broth to inhibit $S$. derby and $E$. coli EU106 was examined. The combinations were: basal (no salt), NaI (0.0472 m), KI (0.0301 m), thio (Na$_2$S$_2$O$_3$·5H$_2$O, 0.0736 m), tet (Na$_2$S$_4$O$_6$·2H$_2$O, 0.0236 m), NaI-KI, KI-thio, thio-tet, NaI-tet, NaI-thio, KI-thio, NaI-KI-thio, NaI-KI-tet, NaI-thio-tet, KI-thio-tet, and NaI-KI-thio-tet. The only combinations showing inhibition of $E$. coli EU106 were those containing both thiosulfate and tetrathionate. $S$. derby was not inhibited by any combination. The viable cell count for $S$. derby increased from $10^4$ to $2 \times 10^4$ per ml in 20 hr. The viable cell count for $E$. coli EU106 increased in a similar manner, except in those combinations containing both thiosulfate and tetrathionate. In this instance the viable cell count at 20 hr was more than four log cycles below that of the basal. Since the combination of thiosulfate and tetrathionate markedly inhibited $E$. coli EU106, while not affecting $S$. derby, these two salts were examined for their effect on the growth of several other cultures. Representative data are presented in Fig. 2. Data similar to those for $E$. coli A were obtained for seven other strains of $E$. coli, Shigella sp., $S$. paratyphi A, and $S$. choleraesuis. Data similar to those obtained for $S$. typhimurium were obtained for six other strains of Salmonella, $S$. paratyphi B, Pseudomonas aeruginosa, two strains of Proteus, four strains of Aerobacter, Klebsiella sp., and Arizona sp. In most instances, the viable cell count at 12 hr was the same as that at 20 hr. Results were similar with the commercially prepared tetrathionate and the one prepared in our laboratory. It also was observed that tetrathionate, although not inhibiting $E$. coli, stimulated the total growth of salmonellae.

Figure 3 shows the typical response of a mixture of salmonellae and $E$. coli in basal broth with thiosulfate and tetrathionate. Viable $E$. coli declined markedly, whereas the salmonellae showed typical growth.

Effect of components of thiosulfate-tetrathionate mixture on growth of $E$. coli. When thiosulfate is mixed with tetrathionate in the proportions found in the complete broth, 0.0736 moles and 0.0236 moles, respectively, colloidal sulfur and polythionates are formed (1, 15, 20). Paper chromatography of the solution after mixing the tetrathionate and thiosulfate showed that it did contain, in addition to the starting compounds, tri- and pentathionates and traces of compounds
assumed to be hexa-, hepta-, and octathionates on the basis of their position on the chromatogram. A quantitative analysis of the mixture is shown in Table 3. Paper chromatograms showed that the various thionates were formed within 2 hr after mixing.

Trithionate, pentathionate, and a mixture of tri-, tetra-, and pentathionates were found to have no effect on the growth of E. coli. The higher thionates were not tested due to difficulties in isolation and purification (18).

The mixture was examined for other sulfur compounds by the column technique of Levinthal and Schiff (9), and no detectable amounts of sulfide or sulfate were found. However, since sulfite is the first compound formed when thiosulfate and tetrathionate react (20), its effect on E. coli and salmonellae was investigated. When added to the basal medium in concentrations as high as 0.04 M, the only effects were that the lag times of both cultures increased slightly and their maximum populations were decreased slightly. Removal of colloidal sulfur from the thiosulfate-tetrathionate mixture by filtration did not affect the subsequent lethality of the filtrate when added to basal medium and inoculated with E. coli.

The data in Table 3 indicate that thiosulfate catalyzed the breakdown of tetrathionate. To assess the quantitative relationships, 1.83 grams of thiosulfate per liter (0.00736 M) was mixed with 7.22 grams of tetrathionate per liter (0.0236 M) in basal broth and allowed to react for 40 hr and analyzed by paper chromatography. Quantitative analysis showed that this low thiosulfate mixture contained the same concentrations of thionates as the regular thiosulfate-tetrathionate mixture (Table 3). However, this low thiosulfate mixture showed no lethality towards E. coli. A mixture of thiosulfate and low tetrathionate (1/10x) also had no lethal effect on E. coli.

The effect of adding graded amounts of thiosulfate to tetrathionate in basal broth on the viable count is given in Fig. 4. These data indicate that the thiosulfate concentration must be at least 0.06 M to have any great effect. The maximum lethal effect on E. coli occurs at a 3 to 1 molar ratio of thiosulfate-tetra-thionate (0.0736 M and 0.0236 M, respectively).

**Studies with growing and nongrowing E. coli.** Since the thiosulfate-tetrathionate mixture was lethal to growing E. coli, its effect on nongrowing cells was investigated. E. coli A was grown for

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**Table 3. Quantitative analysis of the thiosulfate-tetrathionate mixture**

| Compound       | Before mixing (moles/liter) | After mixing (moles/liter) |
|----------------|-----------------------------|-----------------------------|
| Thiosulfate    | 0.0736                      | 0.0662                      |
| Trithionate    |                            | 0.0121                      |
| Tetrathionate  | 0.0236                      | 0.0131                      |
| Pentathionate  |                            | 0.00103                     |
| Hexathionate   | Trace                       | Trace                       |
| Heptathionate  | Trace                       | Trace                       |
| Octathionate   |                            | Trace                       |

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**Fig. 4. Influence of different concentrations of thiosulfate (μ thio) in basal broth containing 0.0236 μ tetrathionate on the viable cell count of E. coli A.**

**Fig. 5. Influence of the thiosulfate-tetrathionate mixture on washed cells of growing and nongrowing E. coli A.**
24 hr in phosphate basal broth and washed once with 0.05 M phosphate buffer (pH 7.6). The cells were resuspended in buffer and added to 0.5% glucose (nongrowing cells) or to 0.5% proteose-peptone (growing cells). Thiosulfate and tetrathionate were added, and the suspensions were incubated at 37°C. Periodically, samples were placed on XL agar (Difco). After 24 hr, cells in buffered glucose showed no change, whereas the growing cells showed the typical decline and regrowth (Fig. 5). Cells in buffer alone also showed no change in number.

**DISCUSSION**

When washed cells were incubated with tetrathionate, there was poor correlation between induction time and inhibition in tetrathionate broth except with certain salmonellae and *E. coli*. With the cultural method for detecting reduction, a closer relationship was observed, but there were several exceptions. When these observations are coupled with the lack of a lethal action of tetrathionate alone on any of the cultures, it appears doubtful that the production of tetrathionatereducing enzyme is the primary reason for the inhibitory action of tetrathionate broth.

The stimulatory effect of tetrathionate on the growth of bacteria which reduce it as reported by Pollock and co-workers (4, 5) was also observed in our studies. As they noted, however, this was not an effect on growth rate, but on total growth, and was never more than one log cycle.

When the various combinations of compounds that are either added to or produced in tetrathionate broth were investigated, the lethal effect was observed only when both tetrathionate and thiosulfate were present. Although it was demonstrated that other polythionates were formed when tetrathionate and thiosulfate were mixed, the tri-, tetra-, and pentathionates were shown to have no effect. The higher thionates were not available as pure compounds for testing, but their possible involvement was eliminated when a low level of thiosulfate was mixed with tetrathionate. The quantities of polythionates produced by this combination were essentially the same as found in the regular tetrathionate broth, yet no lethality occurred. The report by Smith (21) that di-, tri-, and pentathionates would relieve selenite-inhibited growth of *E. coli* is added evidence that these compounds do not have a significant role in the inhibitory action of tetrathionate broth.

Knox et al. (5) found that iodide-free tetrathionate was less inhibitory than tetrathionate made from iodine and thiosulfate, whereas Muller (11) got better inhibition of *E. coli* when he used iodide-free tetrathionate. Our data indicated that iodides have no significant role in the lethal action of tetrathionate broth.

Preuss (19) proposed a new tetrathionate broth containing only calcium tetrathionate. However, when mixed cultures containing salmonellae or *Proteus* are inoculated into it, reduction of the tetrathionate will occur and thus it would become like regular tetrathionate broth, i.e., a lethality due to thiosulfate and tetrathionate. This medium might prove better in that reduction would begin later in the enrichment period either because of few cells or the slow growth of injured cells. This late reduction would then yield the thiosulfate necessary to kill susceptible cells.

*Proteus, Enterobacter,* and *P. aeruginosa* are similar to salmonellae in not being inhibited by the thiosulfate-tetrathionate mixture. These organisms also come through the tetrathionate enrichment and present problems during the isolation of salmonellae from meat and meat products (Alice Moran, personal communication).

The data in Fig. 4 and 5 show the typical decline and regrowth of *E. coli* in thiosulfate-tetrathionate broth. The possibility that this regrowth is due to selection of a mutant was tested by picking colonies from the TSA plate at 24 hr and reinoculating them into thiosulfate-tetrathionate broth. These cultures showed the typical decline and regrowth and thus we discounted the idea of a mutant. This regrowth could be caused by a decrease in the toxic substance(s). Although *E. coli* does not reduce tetrathionate, there is a chemically caused decrease in the tetrathionate concentration. This chemical change alters the importance of the tetrathionate-thiosulfate ratio. Thus we have further support for the importance of the tetrathionate-thiosulfate ratio.

The lethality of tetrathionate broth is directly related to the concentrations of thiosulfate and tetrathionate in the medium. Decreasing the concentration of either salt reduces the lethal action of the mixture. Concentrations of 0.0736 M and 0.0236 M of thiosulfate and tetrathionate, respectively (3:1 ratio), appear to be optimal. The ability of a microorganism to reduce tetrathionate may play a role in decreasing the inhibition for certain species. When part of the tetrathionate is reduced, the ratio would be altered.

The lethality is a growth-related phenomenon since no killing occurred with nongrowing cells in the presence of the thiosulfate-tetrathionate mixture (Fig. 5). The actual mechanism of this lethal action is not known. However, tetrathionate is known to react with free sulfhydryl groups of enzymes and to cause their inactivation (14). Thiosulfate can also react with sulfhydryl...
groups (18). Thus, it is suggested that tetrathionate broth interferes with the synthesis, the activity, or both, of sulfur-containing enzymes or cell wall and membrane components.

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