DEVELOPMENT AND USE OF ANUCLEATE BACTERIAL CELLS TO ASSAY THE IN VITRO ACTIVITY OF POLLUTANTS

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PROGRESS REPORT

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Title: Development and use of anucleate bacterial cells to assay the \textit{in vivo} activity of pollutants
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Report for research undertaken between April 1, 1981 - August 1, 1982

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I. Report for research undertaken between April 1, 1981 - February 1, 1982

A. Abstract

Experiments were undertaken to determine if misincorporation of $^{35}$S-cysteine into non-cysteine containing T7 proteins could be quantitated by autoradiography. It was found that a minimum of $10^7$ radioactive sulphur atoms were required to sufficiently expose X-ray film so that scanning densitometry could be used for quantitation. Calculations based on this determination indicated that the expected misincorporation of $^{35}$S-cysteine into T7 proteins in infected minicells would be insufficient to permit autoradiography to be used as the assay for misincorporation. It was therefore decided to develop a more sensitive assay based on immune-precipitation of specific T7 proteins synthesized in infected minicells. A modification of a published procedure to isolate T7 0.3 gene product has been developed and we are now in the position to raise antibody to this purified protein.

Experiments have begun to determine if it is feasible to construct a recombinant bacteriophage containing a large (1000-3000 base-pairs) stretch of homopolymer i.e. poly dA-dT or poly dG-dC. The intent is to develop a probe for detection of mistranscription in vivo. These experiments have so far been unsuccessful. It appears that although homopolymers can be constructed in vitro they are not stable in vivo.

B. Detection of $^{35}$S-cysteine incorporation into T7 polypeptides

Anucleate bacterial cells (minicells) were infected with T7 and allowed to incorporate $^{35}$S-methionine or $^{35}$S-cysteine. The radioactively labeled polypeptides, synthesized in infected minicells, were separated by electrophoresis and the resulting gels used to expose X-ray film using the technique of fluorography. As previously observed three T7 proteins (products of genes 0-3, 9 and 16) were not labeled by incorporation of $^{35}$S-cysteine, i.e. they do not contain cysteine residues. Sample volumes were increased to increase the amounts of radioactive proteins on the gels. We were still unable to detect $^{35}$S-cysteine incorporation into 0-3, 9 or 16 proteins at sample sizes so large that gel resolution was impaired. Faint exposures of the X-ray film were obtained using $^{35}$S-cysteine incorporation into 0-3 protein by exposing the gel to the film for 8-12 weeks. This is not, however, a practical laboratory procedure.

Figure 1 shows the results of exposing known amounts of $^{35}$S-cysteine to X-ray film. It can be calculated that a minimum of $10^7$ $^{35}$S-atoms are required to produce an exposure suitable for quantitation by scanning densitometry in two days. Assuming a random amino-acid misincorporation rate of 1 per 10,000 peptide bonds (1) and a polypeptide length of 116 amino-acids (0.3 protein (2)) then one...
Figure 1. Known amounts of $^{35}$S-cysteine were impregnated into a dried polyacrylamide gel prepared for fluorography. The gel was exposed to X-ray film for time periods indicated below each series of spots. The number of atoms of $^{35}$S in each sample are given at the side of the figure.
$^{35}$S-cysteine should be misincorporated into approximately one 0.3 molecule per 2000 molecules synthesized. (This assumes that misincorporation of cysteine occurs at 1/20th the rate of total amino-acid misincorporation.)

If one 0.3 molecule is labeled per 2000 molecules synthesized and $10^7$ $^{35}$S-atoms are required for detection then a minimum of $2 \times 10^{10}$ 0.3 molecules must be loaded onto the gel for there to be a reasonable probability that the misincorporation of $^{35}$S-cysteine will be detected by our autoradiography procedures. Our routine procedures employ $1 \times 10^9$ minicells per sample and therefore each T7 infected minicell would have to synthesize 20 molecules of 0.3 protein to obtain the $2 \times 10^{10}$ molecules per sample needed for minimum detection of $^{35}$S-cysteines misincorporation. Synthesis of 20 molecules of 0.3 protein per infected minicell should occur; however, our experimental results indicate that we do not routinely observe $^{35}$S-cysteine labeling of 0.3 protein. We must, therefore, assume that one or more of the quantitative assumptions used above in making calculations is incorrect. Attempts to stimulate $^{35}$S-cysteine incorporation by addition of streptomycin (known to increase mistranslation by a factor of 2 to 3) to give detectable incorporation were not successful. The conclusion of these studies was that quantitation of $^{35}$S-cysteine incorporation by direct autoradiography of extracts of T7 infected minicells is not sufficiently sensitive to develop as a routine procedure.

C. Purification of 0.3 protein

Based on our calculations that quantitation of $^{35}$S-cysteine misincorporation into 0.3 protein could not be successfully obtained by scanning densitometry we decided to purify the 0.3 protein so that antibody could be raised against the protein and used to specifically precipitate 0.3 protein from T7 infected minicells. The misincorporation of $^{35}$S-cysteine could then be quantitated by scintillation counting. In addition, antibody against 0.3 protein could be used to precipitate 0.3 protein synthesized in T7 infected cells, as well as in minicells, increasing the potential alternative systems in which 0.3 protein could be used to assay errors in protein synthesis.

A procedure for purification of 0.3 protein has been published (2) and we initially followed the procedure precisely as described. In essence the purification depends on the ability of 0.3 protein to bind to DEAE even in the presence of 0.3M NH$_4$Cl plus the unusual property that 0.3 protein is soluble in ethanol. Our experiments confirmed that 0.3 can be separated from most proteins in an extract of T7 infected cells by binding to DEAE at 0.3M NH$_4$Cl and elution at 0.7M NH$_4$Cl. Our problems arose in our attempts to repeat the published procedures to dissolve 0.3 protein in ethanol following TCA precipitation. A very time consuming series of experiments were undertaken varying salt concentration, pH, sequence of additions, etc. to obtain pure 0.3 protein as determined by SDS-gel electrophoresis using silver nitrate staining. We now have resolved these problems and have 0.3 protein in sufficient amounts to begin vaccination procedures for antibody production. Figure 2 is a flow chart of our procedure for isolating 0.3 protein from T7 infected cells.
Figure 2. Purification of T7 0.3 protein. Modification of the procedure described by Mark and Studier (1981).

Extract of T7 H3; lam193; LG3 infected E. coli cells

- DNase digestion. Bring to 0.3M NH₄Cl
- Slow speed centrifugation
- High speed centrifugation
- Pass supernatant through DEAE column in 0.3M NH₄Cl. Most proteins do not bind.
- Elute with 0.3M-1M NH₄Cl gradient
- Test fractions for 0.3 protein by running 10-20% polyacrylamide gel and AgNO₃ staining
- Add 10% TCA to 0.3 containing fractions
- Collect precipitate
- Dissolve in Tris buffer + 0.3M NH₄Cl
- Add 4 volumes ethanol; pH2
- Discard precipitate
- Add 8 volumes ethanol
- Discard supernatant
- Dissolve pellet (pure 0.3) in 0.1M Tris buffer
D. Construction of a lambda based recombinant

Only a limited number of experiments have been undertaken on this part of the project as Dr. Alice Desmyter, who was hired to carry out these experiments, did not arrive in Columbus until December 1981. The problem of recruiting trained personnel for recombinant DNA studies was the basis for the no-cost extension granted to the project. Our experiments to date have been designed to determine the procedural conditions to synthesize long (1000-3000) base-pair homoduplexes using DNA polymerase I in the absence of template DNA and using terminal transferase. Based on the results of incorporation of radiochemically labeled precursors and gel electrophoresis of products it does appear that molecules of the appropriate length can be synthesized in vitro. We have not, so far, been able to clone these molecules into lambda vectors. The preliminary experiments already completed indicate that homopolymers may not be stable in vivo. A similar conclusion has been reached by Prof. R.D. Wells (University of Wisconsin; Personal communication 1981).
A. Abstract

The T7 0.3 gene product (0.3 protein) was purified by a modification of the published procedure (2), and used to raise antibody to this protein. A radioimmune precipitation (RIP) assay was developed which could be used to estimate the increased misincorporation of cysteine into 0.3 protein. Parameters of the RIP assay were varied to make the RIP-polyacrylamide gel electrophoresis (RIP-PAGE) assay specific for the 0.3 protein. A single protein band was, however, never achieved although increased misincorporation of cysteine into the 0.3 protein can now be estimated by RIP-PAGE combined with scanning densitometry.

Experiments to determine the normal level of mistranslation in vivo are currently in progress. All necessary preliminary experiments for these assays have been completed. We have devised a labeling medium, determined the minimum saturating levels of cysteine and developed satisfactory methods to purify small amounts of labeled 0.3 protein, to demonstrate nonconversion of cysteine to methionine and to reliably determine the specific activity of purified, labeled 0.3 protein.

B. Detection of $^{35}$S-cysteine incorporation into 0.3 protein by radioimmune precipitation

1. Preliminary experiments. Rabbit antibody was raised to purified 0.3 protein. This was allowed to react with $^{35}$S-methionine labeled lysates of T7-infected E. coli cells. In our initial RIP experiments protein A-containing S. aureus was used to precipitate immune complexes. On visualization of the immune complexes by fluorography following PAGE, 0.3 protein was the major band present; however, other polypeptides were also present. Exhaustive attempts were made to remove these extraneous polypeptides by varying conditions of the assay. Improvements were seen, especially when protein A Sepharose was used instead of S. aureus for precipitation of immune complexes (fig. 3). We were, however, unable to completely eliminate the other polypeptides. This may be due to nonspecific adherence of these polypeptides to protein A Sepharose or to the presence in our 0.3 antiserum of rabbit antibody to E. coli or to other T7 proteins.

2. Specificity of the RIP-PAGE assay for the 0.3 protein. Specificity was demonstrated by reacting normal or immune serum with lysates of E. coli strains infected with a T7 amber mutant (CRI0b) in the 0.3 gene. These strains were either capable or incapable of suppressing the nonsense mutation in the 0.3 gene (fig. 4). With normal serum, protein was not seen in the 0.3 protein position in either case, but with immune serum the 0.3 protein was seen, and only when the nonsense mutation-suppressing strain was used.

3. Increase in misincorporation of cysteine into 0.3 protein with drugs. The RIP-PAGE assay was used to detect an increase in $^{35}$S-cysteine incorporation into the 0.3 protein after treatment with streptomycin or gentamicin. An increase in incorporation was detected by both scintillation counting and autoradiography using either drug at all concentrations tested (5, 10, 20 and 40 μg/ml). In the absence of drugs $^{35}$S-cysteine incorporation into 0.3 protein was detectable by autoradiography in only some experiments and in those cases exposure times in excess of 2 weeks were required. When
FIGURE 3

1a = Uninfected E. coli + normal serum + S. aureus
1b = Uninfected E. coli + immune serum 1 + S. aureus
1c = Uninfected E. coli + immune serum 2 + S. aureus
2a = T7 infected E. coli + normal serum + S. aureus
2b = T7 infected E. coli + absorbed immune serum 1 + S. aureus
2c = T7 infected E. coli + immune serum 1 + S. aureus
2d = T7 infected E. coli + absorbed immune serum 2 + S. aureus
2e = T7 infected E. coli + immune serum 2 + S. aureus
3a = T7 infected E. coli + normal serum + protein A Sepharose
3b = T7 infected E. coli + absorbed immune serum 1 + protein A Sepharose
3c = T7 infected E. coli + immune serum 1 + protein A Sepharose
3d = T7 infected E. coli + immune serum 2 + protein A Sepharose
4a = Extract of T7 infected E. coli B cells used in this experiment
4b = Extract of T7 infected E. coli B cells used in this experiment
Specificity of the ribosome A site for S. Protein

**FIGURE 4**

T7 infected *E. coli* labelled with $^{14}C$ leucine was allowed to react with normal or immune serum, and then protein A Sepharose.

1a = T7 infected *E. coli* B + normal serum
1b = T7 infected *E. coli* B + immune serum
2a = T7 (0.3 amber mutant) infected Su$^+$ strain of *E. coli* + normal serum
2b = T7 (0.3 amber mutant) infected Su$^+$ strain of *E. coli* + immune serum
3a = T7 (0.3 amber mutant) infected Su$^-$ strain of *E. coli* + normal serum
3b = T7 (0.3 amber mutant) infected Su$^-$ strain of *E. coli* + immune serum
4 = Extract of T7 infected *E. coli*
streptomycin or gentamycin were added, however, \( \text{S}^{35} \)-cysteine incorporation was always detectable and the darkness of the band (amount of \( \text{S}^{35} \)-cysteine incorporation) was proportional to the concentration of the drug used (fig. 5). To quantitate the increase in misincorporation of \( \text{S}^{35} \)-cysteine into 0.3 protein we will subject this autoradiogram to scanning densitometry. An isolated cysteine-containing protein will be used to correct for the overall decrease in protein synthesis resulting from the use of gentamicin at these high concentrations.

C. Determination of the normal mistranslation level in vivo

To estimate the increased error frequency in translation in vivo when drugs or toxic chemicals are present, it is necessary to know the natural error level. The T7 0.3 protein system provides an advantage over other systems used to estimate mistranslation levels in vivo. The T7 0.3 protein is made only after T7 infection and therefore all the 0.3 protein made is under the conditions of the experiment, i.e., all 0.3 protein synthesized is labeled. It is therefore unnecessary to determine the amount synthesized during the labeling period relative to the amount existing before label was added. In addition, we have simplified the purification procedure for 0.3 protein and made it suitable for the small amounts of protein obtained in labeling experiments.

1. Purification of small amounts of labeled 0.3 protein. Two methods were compared: affinity chromatography using rabbit anti-0.3 antibody conjugated to CNBr-activated Sepharose 6B and DEAE cellulose anion exchange chromatography followed by ethanol extraction (scaled down from our procedure to purify milligram amounts of 0.3 protein). We found that the 0.3 protein, eluted from the immunopreaffinity column, was impure and required additional ethanol extraction steps for a complete purification. In addition, the capacity of the immunopreaffinity columns was much less than that of the DEAE columns and not sufficient to retain all the 0.3 protein made in a labeling experiment. We, therefore, decided against the use of immunopreaffinity chromatography and have used anion exchange chromatography for the initial purification steps.

2. Experimental conditions necessary for calculations of mistranslation levels. To calculate the normal mistranslation frequency, i.e., the number of cysteine residues per molecule of 0.3 protein, it is necessary to know the specific activity of cysteine available to the cell for protein synthesis. This means that the amount of radioactively labelled cysteine provided should not be diluted by endogenous cysteine synthesis nor should the precursor \( \text{S}^{35} \)-cysteine be converted to methionine. An excess of methionine must be provided to block cysteine conversion into methionine by feedback inhibition and the endogenous pathway of cysteine biosynthesis must be repressed by providing a saturating level of cysteine in the absence of sulfate. Experiments were performed to determine minimum saturating levels of cysteine and methionine. In Figures 6 and 7 cysteine and methionine uptake can be seen to be at the saturating level at \( 5 \times 10^{-4} \) M. Additional experiments showed that the minimum saturating level for both amino acids was \( 2 \times 10^{-7} \) M.

3. Demonstration that \( \text{S}^{35} \) is not transferred from cysteine to methionine. If \( \text{S}^{35} \)-cysteine were converted to \( \text{S}^{35} \)-methionine the calculated error level in the synthesis of 0.3 protein would be higher
FIGURE 5

<table>
<thead>
<tr>
<th>Lanes</th>
<th>Gentamicin (µg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a,1b</td>
<td>0</td>
</tr>
<tr>
<td>2a,2b</td>
<td>10</td>
</tr>
<tr>
<td>3a,3b</td>
<td>20</td>
</tr>
<tr>
<td>4a,4b</td>
<td>40</td>
</tr>
</tbody>
</table>

Lanes 1a, 2a, 3a + 4a are extracts of T7 infected E. coli B, labelled with 35S-cysteine, and allowed to react with normal serum and then protein A Sepharose. Lanes 1b, 2b, 3b + 4b are extracts of T7 infected E. coli B, labelled with 35S-cysteine and allowed to react with antiserum and then protein A Sepharose.
than the true error level since methionine is present in 0.3 protein. It is, therefore, necessary to show that methionine in the 0.3 protein contains no $^{35}$S when $^{35}$S-cysteine is supplied and that all the radioactivity is present in cysteine. Thin layer chromatography (TLC) was chosen as the best way to determine this. Several solvent systems and stationary phase solid supports were examined in preliminary experiments to determine the best way for separation of methionine and cysteine. We found that two-dimensional TLC provided no advantage over one-dimensional TLC and that N-propanol: ammonium hydroxide was the solvent system best suited to our needs. Silica gel, cellulose, paper can be used as the solid support. The specific activity of $^{35}$S-cysteine-labeled 0.3 protein is naturally very low and therefore long exposure times of the TLC chromatograms to x-ray film are needed for satisfactory autoradiography.

We expect to obtain a precise determination of the normal error level for translation in vivo during the synthesis of 0.3 protein within the next few months.

III. References

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EDUCATION:
University of Birmingham, England, B.Sc., 1968, Bacteriology
University of British Columbia, Canada, Ph.D., 1971, Microbiology

THESIS TITLE: Cell division in a temperature-sensitive mutant of E. coli.

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1981 - Professor, Department of Microbiology, The Ohio State University
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1971-1973 - Postdoctoral fellow with Dr. Neil H. Mendelson, Department of Microbiology, University of Arizona, Tucson. Research in bacterial cell growth. Isolation and characterization of mutants including the minicell mutants of B. subtilis.

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USPHS, NIH, 1-ROI-CA30338-01, FeLV leukemogenesis and pre-neoplastic lesions, co-investigator, 3 yrs., $180,805 total, 11/81-10/84. (I wrote the entire grant proposal. Richard G. Olsen is listed as P.I. because of University regulations.)

The Ohio State Canine Research Funds, "Immunoprevention of Parvovirus-Induced Diarrhea of the Dog", 2 yrs., $20,000/yr., 1981-1982. (I wrote the entire grant proposal. Richard G. Olsen and C. Stephen Klinko are listed as P.I.'s because of University regulations.)
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