Independent Expression of the A Gene of the Tryptophan Operon of 
Escherichia coli during Tryptophan Starvation

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SUMMARY
Two complementary frameshift mutations in the trpA gene of Escherichia coli have been isolated following recombination of the pseudowild double mutant with the wild-type strain. One of these mutations recombines with a third, distal frameshift mutation to give a slow-growing Trp+ strain. Consideration of the relevant RNA codon sequences suggests that the slow-growing recombinant should contain a unique tryptophan codon in the altered reading phase between the frameshift mutations. This prediction has been verified by analysis of the purified tryptophan synthetase A protein from the double mutant. The preferential synthesis of A protein normally associated with prolonged tryptophan starvation is eliminated in the strain which has a tryptophan residue in its A protein.

INTRODUCTION
The biosynthetic tryptophan (trp) operon of Escherichia coli contains five structural genes, trpE, D, C, B and A, which code for the five polypeptide chains involved in the conversion of the aromatic precursor chorismic acid to tryptophan (Yanofsky & Lennox, 1959; Creighton & Yanofsky, 1969). Expression of the operon is controlled by the protein product of an unlinked regulatory gene, trpR (Cohen & Jacob, 1959; Morse & Yanofsky, 1969; Zubay, Morse, Schrenk & Miller, 1972). An operator locus has been identified by the isolation of cis-acting constitutive mutants mapping at the E gene end of the operon (Hiraga, 1969). The five structural genes are transcribed into a single messenger RNA molecule, which is sequentially synthesized and translated from the operator-proximal E gene towards the distal A gene (Imamoto, Morikawa & Sato, 1965; Imamoto & Yanofsky, 1967).

When the trp operon is derepressed during normal growth the enzymes are produced in equimolar quantities (Ito, Cox & Yanofsky, 1969). This situation does not hold, however, when a tryptophan auxotroph is subjected to prolonged tryptophan starvation (Somerville & Yanofsky, 1964). The tryptophan synthetase α subunit, coded by the trpA gene, continues to be synthesized for some 30 h of starvation, whereas the other products of the operon are no longer produced after a few hours. Somerville & Yanofsky (1964) have suggested that this preferential synthesis of the α subunit is due to independent translation of the messenger RNA corresponding to the trpA gene, made possible by the absence of tryptophan from the α subunit (Henning, Helinski, Chao & Yanofsky, 1962). This hypothesis could be directly tested if there were available an α subunit variant which contained a tryptophan codon. This paper describes the isolation of such a variant, and shows that the altered A gene is no longer preferentially expressed during tryptophan starvation.
METHODS

Bacteria and phage. The various trp mutants used in this study were crossed into the W3110 strain of Escherichia coli by transduction with P1kc, using a W3110 (tonB, trpE-A) deletion as recipient, and selecting for growth on plates supplemented with indole (10 \( \mu \)g/ml). Strains trpA466oPR3 and trpA466oPR3FR7 were generously provided by Dr H. Berger: other bacterial strains and the generalized transducing phage P1kc were from the collection of Dr C. Yanofsky.

Media. The minimal medium of Spizizen (1958) was used with glucose (0.2%, w/v) as carbon source. Where possible this was supplemented with 0.05% (w/v) Difco Bacto acid hydrolysed casein. Amino acids were added as required at 20 \( \mu \)g/ml. The rich medium was L-broth (Lennox, 1955) containing (g/l): Difco Bacto Tryptone, 10; Difco Bacto Yeast Extract, 5; NaCl, 5; glucose, 1, adjusted to pH 7.2. Media for plates were solidified with 1.5% (w/v) agar.

Transduction. P1kc lysates were prepared by the confluent lysis technique (Swanstrom & Adams, 1951) and were used for transduction as described by Yanofsky & Lennox (1959).

Recombination distances between trpA mutants were obtained as described by Yanofsky et al. (1964). When the recipient was cysB, trp, the ratio of cys+trp+ to cys+ transductants was multiplied by 2 to correct for the 50% cysB-trp linkage (Yanofsky & Lennox, 1959). When a hiscysBtrp recipient was used the ratio of cysB+trp+ to his+ transduction was scored. When cys+ and trp+ are selected together, the 50% cys–trp linkage exactly compensates for the relatively inefficient transduction of the his region compared with that of the cys–trp region (Yanofsky & Lennox, 1959). In all crosses the cys+trp+ transductants were selected on minimal plates supplemented with 0.1 \( \mu \)g/ml 5-methyl-DL-tryptophan, which suppresses the growth of leaky mutants without affecting the growth of wild-type recombinants or the frequency of recombination (Yanofsky et al. 1964).

Penicillin enrichment for trp recombinants. The procedure was based on that described by Allen & Yanofsky (1963). After exposure of the cysBtrp+ recipient to transducing phage, organisms were harvested, washed in 0.1 M-sodium citrate buffer, pH 7.0, resuspended in minimal medium containing 0.2% (w/v) glucose and 20 \( \mu \)g/ml L-tryptophan, and grown overnight at 37 °C. The cys+ transductants so selected were diluted 50-fold into fresh minimal medium containing glucose (0.2%, w/v) and tryptophan (20 \( \mu \)g/ml) and grown to about 2 \times 10^8 organisms/ml. Organisms were harvested and washed twice with minimal medium, resuspended at 10^6 organisms/ml in minimal medium + glucose (0.2%, w/v) and grown for a further 3 h at 37 °C. Benzylpenicillin (1000 u/ml) was then added and incubation was continued for a further 2 h before the organisms were washed twice in minimal medium and resuspended in the original volume of L-broth to grow to saturation. Approximately 200 organisms/plate were spread on to glucose-minimal agar plates supplemented with L-tryptophan (20 \( \mu \)g/ml). Plates were incubated at 37 °C for 48 h before replication to unsupplemented glucose-minimal agar plates to screen for trp colonies.

Preparation of extracts. Bacteria were harvested by centrifugation at 0 °C, washed with one-half the original volume of cold 0.1 M-tris-HCl, pH 7.8, and resuspended in 1/50 volume of the same buffer. Organisms were disrupted with a M.S.E. sonifier, and the debris was removed by centrifugation at 38000 g for 15 min.

Enzyme assays. The A and B subunits of tryptophan synthetase were assayed by indole-utilization as described by Smith & Yanofsky (1962). A crude extract prepared from the A gene nonsense mutant trpA96 was used as the source of the B protein. Specific activities...
are expressed as units of enzyme activity/mg of protein. Protein was determined by the procedure of Lowry, Rosebrough, Farr & Randall (1951).

Purification of tryptophan synthetase A proteins. Wild-type and mutant tryptophan synthetase A proteins were purified from 50 l batches of bacteria as described by Henning et al. (1962) as far as the chromatography on G 100 Sephadex. All buffers contained $10^{-3} \text{M} \cdot \beta$-mercaptoethanol and $10^{-3} \text{M}$-EDTA. The precipitated fractions from the G 100 Sephadex column were resuspended in the minimal volume of $0.1 \text{M}$-potassium phosphate buffer, pH 7.2. After centrifuging at 38,000 g for 10 min to remove solids, the supernatant was applied to a 100 $\times$ 2 cm diam. column of DEAE-Sephadex, equilibrated with $0.03 \text{M}$-potassium phosphate buffer, pH 7.2. A linear gradient of $0.03 \text{M}$ to $0.3 \text{M}$-potassium phosphate (700 ml in each vessel) was applied at a flow rate of 20 ml/h, and 12 ml fractions were collected. The peak fractions from this column usually have a specific activity of 2500 to 3500 u./mg protein. Protein from the pooled peak fractions was precipitated with $0.43 \text{g} \ \text{NH}_4\text{SO}_4/\text{ml}$. The precipitate was collected by centrifugation at 38,000 g for 20 min, resuspended in $0.1 \text{M}$-potassium phosphate, pH 7.0, and dialysed for 2 h against $0.05 \text{M}$-potassium phosphate, pH 7.0. After removal of insoluble material by centrifugation at 38,000 g for 10 min, the protein was rechromatographed on a Sephadex G 100 column, 80 $\times$ 2 cm, at a flow rate of 20 ml/h; 5 ml fractions were collected. The peak fractions from this column usually had a specific activity of 3500 to 4500 u./mg protein. Peak fractions from the Sephadex column were precipitated with $0.43 \text{g} \ \text{NH}_4\text{SO}_4/\text{ml}$. The protein pelleted after centrifugation at 38,000 g for 20 min, was resuspended in $0.1 \text{M}$-potassium phosphate, pH 7.0, and again chromatographed on a column of DEAE-Sephadex, 100 $\times$ 2 cm, at a flow rate of 15 ml/h; 5 ml fractions were collected. Specific activities in the peak fractions were usually about 4800 u./mg at this stage.

Results

Recovery of frameshift mutants from a pseudowild revertant strain

Escherichia coli strain trpA46\textsuperscript{asb}PR3FR7, though phenotypically indistinguishable from wild-type, contains two frameshift mutations of opposite phase, between which are two missense mutations in adjacent bases (Berger, Brammar & Yanofsky, 1968). The pertinent amino acid and RNA codon sequences are shown in Fig. 1. An attempt was made to separate the two frameshift mutations by recombination in a genetic cross between A46\textsuperscript{asb}PR3FR7 and a trp\textsuperscript{+} wild-type strain. A P1-lysate grown on A46\textsuperscript{asb}PR3FR7 was used to transduce a cysBtrp\textsuperscript{+} recipient to cys\textsuperscript{+}, and the resulting cys\textsuperscript{+} transductants were screened for the presence of trp recombinants following a penicillin enrichment procedure as described in Methods.

Tryptophan auxotrophs recovered in this way fell into two phenotypic classes (Table 1). Some were able to grow very slowly on minimal agar, while others failed to grow. Mutants of both classes appeared to be frameshifts, since they were stimulated to revert by ICR 191, a powerful frameshift mutagen, but not by NTG, which is known to stimulate largely base replacements (Whitfield, Martin & Ames, 1966). Mutants of both classes fail to grow on plates containing 2 $\mu$g/ml indole $+ 100 \mu$g/ml 5-methyl-DL-tryptophan (Brammar, Berger & Yanofsky, 1967), and show no detectable A protein activity in the conversion of indole to tryptophan (Crawford & Yanofsky, 1958). Their behaviour in both of these tests is characteristic of immunologically non-crossreacting (CRM\textsuperscript{−}) A gene mutants, and is therefore consistent with their classification as frameshift mutants.

To verify that the mutants isolated were derived by recombination from A46\textsuperscript{asb}PR3FR7,
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Strain Wild-type 201 210 213 uu A uu A Pro - Leu - Gln - Gly - Phe - Gly - Ile...

Phenotype Wild

A46 ...CCX UUG CAG GGA UU UGG GC AUC...

Mutant -Glu-

A46asp ...CCX UUG CAG GAU UU UGG UAU C

Mutant, but suppressible -Asp-

A46aspPR3 ...CCX AUU GCA GGA UUU UGC AUC...

Trp+; slow growing

A46aspPR3FR7 ...CCX AUU GCA GGA UUU UGC AUC...

Wild

-Pro - Ile - Ala - Gly - Phe - Cys - Ile-

Fig. 1. Derivation of the multiple mutant strain trpA46aspPR3FR7. The pseudowild strain A46aspPR3FR7 was derived in four successive mutational steps from the wild-type. The first step was a transition of the second G of the gly-210 codon to an A, resulting in the gly → glu replacement of mutant trpA46 (Henning & Yanofsky, 1962). Step (2) was a transversion of the third position A of the mutant glu-210 codon (GAA) of trpA46 to U, converting residue 210 into asp (A46asp). Although the A46asp is itself Trp-, the glu → asp replacement could be selected in the presence of a missense suppressor which misreads asp codons (Berger & Yanofsky, 1967). Step (3) arose as a slow-growing (partial) revertant of A46asp, and was subsequently shown to be a single nucleotide addition frameshift event (Berger et al. 1968). Event (4) was a compensatory deletion of G from the gly 212 codon, restoring the wild-type reading frame and phenotype (Berger et al. 1968).

Table 1. Characteristics of trpA mutants derived from A46aspPR3FR7

<table>
<thead>
<tr>
<th>Strain</th>
<th>Relative colony size on minimal plates</th>
<th>Growth on indole + 5MT*</th>
<th>Reversion response† to</th>
<th>A/B ratio‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>A46aspPR3FR7</td>
<td>100</td>
<td>+</td>
<td>.</td>
<td>1:0</td>
</tr>
<tr>
<td>A46aspPR3</td>
<td>23</td>
<td>-</td>
<td>.</td>
<td>0</td>
</tr>
<tr>
<td>WB11</td>
<td>Tiny</td>
<td>-</td>
<td>+</td>
<td>±</td>
</tr>
<tr>
<td>WB1</td>
<td>0</td>
<td>-</td>
<td>+</td>
<td>0</td>
</tr>
<tr>
<td>WB13</td>
<td>20</td>
<td>+</td>
<td>.</td>
<td>1:1</td>
</tr>
</tbody>
</table>

* 5MT = 5-methyl-DL-tryptophan.
† ICR = ICR191A [3-chloro-7-methoxy-9-(3-chloroethyl)amino propylamino]-acridine dihydrochloride, generously provided by Dr H. J. Creech. NTG = N-methyl-N'-nitro-N-nitrosoguanidine. Reversion responses were determined in plate-tests as described by Brammar et al. (1967).
‡ The ratio of A-protein activity, measured in the presence of excess B protein, to B-protein activity, measured in the presence of excess A protein, in the conversion of indole to tryptophan.

and were not merely spontaneous tryptophan auxotrophs selected by penicillin enrichment, their map locations were investigated by P1 transduction. Mutants of both classes showed very close linkage to trpA23, a missense mutant affecting the same codon as A46asp (Helinski & Yanofsky, 1962; Berger & Yanofsky, 1967). WB11, the 'leaky' mutant, showed 0.06% recombination with A23, suggesting a separation of three to four codons (Yanofsky, Drapeau, Guest & Carlton, 1967), while the non-leaky mutant Wb1 showed no detectable recombination with A23 (Table 2). A cross between the phenotypically different auxotrophs yielded trp+ recombinants at a rate of 0.17%. Allowing for the fact that this figure includes both wild and pseudowild recombinants, such a recombination frequency suggests a separation between the two mutants of about six codons.
Trp A gene expression

Table 2. Recombination of mutants derived from $A_{46}^{as}P3R3F7$ with $A_{23}$.

<table>
<thead>
<tr>
<th>Donor (his+cys$^+$)</th>
<th>Recipient</th>
<th>His$^+$ or Cys$^+$ transductants</th>
<th>Trp$^+$Cys$^+$ transductants</th>
<th>Recombination (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WB1</td>
<td>his cys trp $A_{23}$</td>
<td>6300*</td>
<td>0</td>
<td>&lt; 0.016*</td>
</tr>
<tr>
<td>WB11</td>
<td>his cys trp $A_{23}$</td>
<td>13950*</td>
<td>9</td>
<td>0.065*</td>
</tr>
<tr>
<td>WB11</td>
<td>cys WB1</td>
<td>41 500†</td>
<td>35</td>
<td>0.17†</td>
</tr>
<tr>
<td>$A_{46}^{as}PR3$</td>
<td>cys WB1</td>
<td>26 450†</td>
<td>16</td>
<td>0.12†</td>
</tr>
<tr>
<td>$A_{46}^{as}PR3$</td>
<td>cys WB11</td>
<td>12 150†</td>
<td>0</td>
<td>&lt; 0.017†</td>
</tr>
</tbody>
</table>

* In crosses with his cys $A_{23}$ as recipient, his is used as the reference marker: Cys$^+$, Trp$^+$ transductants are selected to avoid a high background due to reversion of the trp marker.

† When a cys trp recipient is used, cys is used as the reference marker. Trp$^+$Cys$^+$ transductants were selected on plates containing 0.1 µg/ml 5-methyl-DL-tryptophan to prevent the growth of leaky mutants WB11 and $A_{46}^{as}PR3$.

The genetic data are thus in excellent agreement with the expected behaviour of mutants derived by recombination of $A_{46}^{as}P3R3F7$ with wild-type. They would also suggest that the non-leaky mutant WB1 contains both the $A_{46}^{as}$ missense mutation and a frameshift since it fails to recombine with $A_{23}$. The leaky mutant WB11 is probably the base-addition frameshift mutation, since it fails to recombine with $A_{46}^{as}PR3$, but does recombine with $A_{23}$ (Table 2). The ancestral strain $A_{46}^{as}PR3$, which contains both the $A_{46}^{as}$ mutation and the base addition, is distinguishable from the leaky isolate WB11 by its slightly better growth on minimal agar (Table 1).

If the two classes of mutant recovered from $A_{46}^{as}PR3F7$ are complementary frameshifts, then one or other must be of opposite phase to any other frameshift mutant. Thus it should be possible to produce pseudowild recombinants with a third closely linked frameshift mutant, provided the amino acid sequence generated by the altered reading frame between the two mutations is functionally acceptable. Mutant $trpA_{21}$ contains a frameshift mutation of undefined phase located closely distal to the region defined by the two frameshift mutations in $trpA_{46}^{as}P3R3F7$ (see Fig. 2) (Berger et al. 1968). WB11 and WB1 were therefore crossed with $trpA_{21}$, and $trp^+$ recombinants were selected. On subsequent purification and characterization, the WB11/A21 recombinants proved to be of two easily distinguished phenotypes; a fast-growing wild-type strain and a slower-growing pseudowild-type (WB138). In contrast, WB1/A21 recombinants were all phenotypically indistinguishable from wild-type. The pseudowild recombinants produced a CRM$^+$ A protein with normal activity in the indole → tryptophan reaction (Table 1), and were presumed to contain frameshifts of opposite phase, derived from WB11 and $trpA_{21}$. If this were the case, the codon sequence of the pseudowild recombinant would contain an in-phase tryptophan codon (UGG) generated from the third nucleotide of the phe$_{211}$ codon (UUU) and the first two nucleotides of the gly$_{215}$ codon (GGG) (see Fig. 2). Thus, the partially active A protein from the pseudowild recombinant WB138 should contain a single tryptophan residue.

To test this prediction, the tryptophan synthetase A proteins from the WB138 recombinant and the wild-type parent strain were extensively purified as described in Methods and their tryptophan contents were assayed in two different ways. The first method employed the formula of Goodwin & Morton (1946) to calculate the tryptophan and tyrosine contents directly from the u.v. absorption spectra in 0.1 M-sodium hydroxide. This method (Table 3) gave close to 1 mole of tryptophan/mole of the mutant A protein, while giving a suitably low value for the tryptophan content of the wild-type protein. The tyrosine content of the wild-type protein determined this way agreed well with the known value of 7 residues/mole (Yanofsky et al. 1967; Guest, Drapeau, Carlton & Yanofsky, 1967).
Strain

\[ \ldots -\text{Pro} - \text{Leu} - \text{Gln} - \text{Gly} - \text{Phe} - \text{Gly} - \text{Ile} - \text{Ser} - \text{Ala} - \text{Pro} - \text{Asp} - \ldots \]

Wild-type

\[ \ldots \text{CCX UUG CAG GGA UUU GGC AUC } \underbrace{\text{U } \text{AG } \text{UCX } \text{GCX } \text{CCX } \text{GAC}}_{(\ldots - G)} \ldots \]

\[ \ldots \text{CAG GAU } \text{UUU GCA UCA } \underbrace{\text{UCX } \text{CXC } \text{CXC}}_{(\ldots-1)} \ldots \]

\[ \ldots \text{CCX AUU GCA GGG AUU UGG CAU } \underbrace{\text{C } \text{XGC } \text{XCC}}_{(\ldots -1)} \ldots \]

\[ \ldots \text{CCX AUU GCA GGG AUU UGG } \underbrace{\text{CUC } \text{XGC } \text{XCC}}_{(\ldots -1)} \text{GAC} \ldots \]

\[ \ldots \text{CAG GAU } \text{UUU GCA UCA } \underbrace{\text{UCX } \text{CXC } \text{CXC}}_{(\ldots -1)} \ldots \]

\[ \ldots \text{CCX AUU GCA GGG AUU UGG CAU } \underbrace{\text{C } \text{XGC } \text{XCC}}_{(\ldots -1)} \ldots \]

Fig. 2. The probable mRNA codon sequences for the trpA frameshift mutants and the \( \text{wb138} \) recombinant. Codons or nucleotides which differ from the wild-type are underlined. The information for \( A21 \) is taken from Berger et al. (1968). Figures in parentheses represent mutational events.

Table 3. Spectrophotometric determination of tryptophan and tyrosine contents of wild-type and mutant \( A \) proteins

Spectra were determined on \( 1.0 \) mg samples of each \( A \) protein in \( 1.5 \) ml \( 0.1 \) M-sodium hydroxide. Values were calculated using the formula of Goodwin & Morton (1946), using an intersection wavelength of 294.4 nm, and were corrected for background absorption.

<table>
<thead>
<tr>
<th>Source of wavelength (nm)</th>
<th>Tryptophan</th>
<th>Tyrosine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type</td>
<td></td>
<td></td>
</tr>
<tr>
<td>270</td>
<td>0.05</td>
<td>7.4</td>
</tr>
<tr>
<td>280</td>
<td>0.30</td>
<td>7.1</td>
</tr>
<tr>
<td>290</td>
<td>0.03</td>
<td>7.4</td>
</tr>
<tr>
<td>( \text{wb138} )</td>
<td></td>
<td></td>
</tr>
<tr>
<td>270</td>
<td>0.80</td>
<td>8.0</td>
</tr>
<tr>
<td>280</td>
<td>0.90</td>
<td>7.9</td>
</tr>
<tr>
<td>290</td>
<td>0.66</td>
<td>8.2</td>
</tr>
</tbody>
</table>

The tryptophan content of the mutant enzyme was also determined by following the oxidative cleavage of carboxyl-tryptophanyl peptide bonds by \( N \)-bromosuccinimide spectrophotometrically at 280 nm (Patchornik, Lawson, Gross & Witkop, 1960). This method gave a value of 0.86 residues/mole for the mutant \( A \) protein, while showing no detectable tryptophan in the wild-type enzyme. The method was checked using lysozyme, bovine serine albumin and pancreatic ribonuclease, and in each case values obtained agreed very well with those from the literature (Table 4). Thus these determinations of the tryptophan content of the mutant \( A \) protein confirm the prediction that the protein contains a single residue of tryptophan.

\( A \) gene expression in the mutant \( \text{trpA138} \). The recombinant strain \( \text{wb138} \) is itself leaky for tryptophan biosynthesis. Thus, in order to study the expression of the mutant \( A \) gene during tryptophan starvation it was necessary to introduce into the strain a complete genetic block on tryptophan synthesis. This was achieved by constructing a double mutant containing \( \text{trpET3} \), a non-leaky missense mutant in the operator proximal \( E \) gene, together with the \( \text{wb138} \) mutation. The expression of the \( A \) gene of this double mutant was compared with that of a \( \text{trpETpA}^+ \) transductant isolated from the same genetic cross.

The specific activities of tryptophan synthetase \( A \) and \( B \) proteins were followed over 40 h.
Fig. 3. The effect of tryptophan starvation on the synthesis of tryptophane synthetase A protein in \textit{trpA}+ and \textit{wB138} strains. Both strains carried the \textit{trp}– missense mutant, \textit{trpET3}. Cultures were grown in minimal medium containing 0.05 \% (w/v) acid-hydrolysed casein, 0.3 \% (w/v) glucose and a growth-limiting concentration of L-tryptophan (4 \mu g/ml). Samples were harvested when growth stopped due to depletion of tryptophan (time 0), and at intervals for a further 36 h. The data are plotted as the ratio of A protein activity to B protein activity to emphasise the differential synthesis of A protein. ○--○, \textit{trpET3}, \textit{wB138}; ●—●, \textit{trpET3}, \textit{A}+.

Table 4. Tryptophan contents determined by titration with \textit{N}-bromosuccinimide

Samples (2 mg) of each protein were dissolved in 1.3 ml 8 M-urea, adjusted to pH 4.0 with glacial acetic acid. Samples were titrated by addition of 10 \mu l samples of 10 mm-\textit{N}-bromosuccinimide, and the decrease in absorption at 280 nm was followed. The calculations of the tryptophan contents were based on the following molecular weights: Lysozyme, 15,000 \((a)\); bovine serum albumin, 65,000 \((b)\); pancreatic ribonuclease, 14,800 \((c)\); tryptophan synthetase A protein, 29,500 \((d)\).

<table>
<thead>
<tr>
<th>Protein</th>
<th>Determined</th>
<th>Reported</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lysozyme</td>
<td>5.3</td>
<td>6*</td>
</tr>
<tr>
<td>Bovine serum albumin</td>
<td>1.5</td>
<td>2†</td>
</tr>
<tr>
<td>Pancreatic ribonuclease</td>
<td>0</td>
<td>0‡</td>
</tr>
<tr>
<td>A protein: wild-type</td>
<td>0</td>
<td>0§</td>
</tr>
<tr>
<td>A protein: \textit{wB138}</td>
<td>0.86</td>
<td></td>
</tr>
</tbody>
</table>

* Canfield \& Liu (1965).  
† Hirs, Stein \& Moore (1960).  
‡ Peters \& Hawn (1967).  

in the absence of tryptophan. The data, expressed as the ratio of A protein to B protein, are shown in Fig. 3. It is evident that the preferential synthesis of A protein observed with the \textit{trpA}+ strain, and with all other \textit{A} gene mutants examined, is eliminated in the strain which has a tryptophan residue in its A protein.
DISCUSSION

The frameshift event which gave rise to \(A46^{os}\)PR3 created a tryptophan codon in the wild-type reading frame. The tryptophan synthetase A protein from \(A46^{os}\)PR3, although feebly active \textit{in vivo}, could not be detected or analysed \textit{in vitro} (Berger et al. 1968). Unfortunately, the mutational event which gave rise to the pseudowild revertant \(A46^{os}\)PR3FR7 eliminated the \textit{trp} codon and created a \textit{cys} codon in restoring the wild-type reading frame. By separating and recovering the component frameshift mutants from \(A46^{os}\)PR3FR7, and then recombining one of them with another, more distal frameshift mutant (A21), it has been possible to create a double mutant which contains a \textit{trp} codon in its \(A\) gene.

The nature of the u.v.-induced mutational event in \textit{trpA21} had not previously been precisely defined: it could have been a deletion of either one or two nucleotides (Berger, Brammar & Yanofsky, 1969). Because A21 is complementary in phase to the single nucleotide addition in \(A46^{os}\)PR3, the mutational event in A21 must be a single nucleotide deletion.

In principle, it should be possible to recombine \(A46^{os}\)PR3 itself with A21 to generate a multiple mutant and restore the wild-type reading frame. In practice, however, this would be very difficult to achieve, because \(A46^{os}\)PR3 itself grows relatively well on minimal medium. It is interesting that \(A46^{os}\)PR3, which contains a frameshift mutation and two base-change mutations, grows very much better than WBII, which has only the frameshift event. This must be because the \(A46^{os}\)PR3 protein retains the wild-type \textit{phe} residue at position 211, whereas in the WBII protein it must be replaced by an \textit{ile} residue.

One of the two frameshift mutants isolated, WBII, shows a significant reversion response with the alkylating mutagen NTG (see Table 1). NTG has been previously observed to stimulate the reversion of some frameshift mutations (Yourno & Heath, 1969), and it has been postulated that it does so by causing deletions of single base-pairs (Oeschger & Hartman, 1970). The NTG-stimulated reversion of WBII is not necessarily due to frameshift mutagenesis, however, since WBII could give rise to partial revertants like \(A46^{os}\)PR3 by base-change mutations.

The purified tryptophan synthetase A protein from the recombinant strain WB138 was shown to contain 1 residue of tryptophan/mole of protein, both from its spectrum and by titration with N-bromosuccinimide. Control experiments with the wild-type enzyme and with three other proteins demonstrate the validity of the methods. These determinations could be criticized on the grounds that the protein preparations were not pure. The preparation of mutant A protein used for these determinations had a specific activity of 4000 u./mg, suggesting a purity of about 80%. Care was taken to use a wild-type fraction with a similar specific activity. Since the two enzymes were purified by the same method it is probable that the two preparations contained the same impurities, and that the determined difference in tryptophan content is a real one. This conclusion is supported by inspection of peptide maps of tryptic digests of the two A proteins. The relevant wild-type peptide, TP 3, (Guest, Carlton & Yanofsky, 1967) is missing from the peptide map of the mutant protein, but a new peptide is evident which shows the U.V. fluorescence characteristic of tryptophan-containing peptides (unpublished results).

The preferential synthesis of tryptophan synthetase A protein during tryptophan starvation is eliminated in the strain which contains a tryptophan codon in its \textit{trpA} gene. This result directly confirms the suggestion that the preferential synthesis of A protein is due to the latter's lack of tryptophan. Since Morse, Mosteller & Yanofsky (1969) have shown that tryptophan messenger RNA synthesis is normal during acute tryptophan starvation, the synthesis of A protein under these conditions must be due to independent translation of the
Trp A gene expression

A gene region of messenger RNA. During tryptophan starvation ribosomes prior to a tryptophan codon would be arrested at the preceding codon (Capecchi, 1967; Bretscher, 1968a). Thus the ribosomes which translate the A gene messenger RNA must do so by internal attachment to the messenger RNA, at or near the beginning of the A gene region. The conclusion that ribosomes can attach to internal positions in a messenger RNA molecule has already been obtained by several different methods (e.g. Lodish, 1968; Spahr & Gesteland, 1968; Bretscher, 1968b; Morse et al. 1969; Webster & Zinder, 1969). It is not yet clear whether there is a specially coded ribosome-attachment site at the beginning of each gene, or whether a single initiation codon (AUG) will suffice.

It might be expected that acute starvation for a particular amino acid, in blocking ribosome movement, would act like a nonsense mutation and produce strong polarity. Morse & Guertin (1971) have recently shown, however, that amino acid deprivation very effectively relieves polarity in Escherichia coli by a mechanism which involves stabilization of the unprotected messenger RNA.

The absolute rate of A-protein synthesis during prolonged tryptophan starvation is of interest. In these experiments the specific activity of wild-type A protein increased from 25 u/mg at 6 h to 73 u/mg after 24 h of starvation. This represents an increase of 48 u/mg in 18 h (or 2.7 u/mg/h), which corresponds to the synthesis of about 1350 molecules of tryptophan synthetase/organism/h. On derepression with indole propionic acid, a log-phase culture of the w3110 strain synthesizes about 12000 molecules/organism/h (Morse, Baker & Yanofsky, 1968). Thus, under conditions in which the synthesis of other proteins has stopped due to lack of tryptophan, tryptophan synthetase A protein is synthesized at about 10% of its maximum rate.

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REFERENCES


