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PERMANENT ADDRESS:

WANDER ALLAND

NL-91 (20-68)



A THESIS



AURINTRICARBOXYLIC ACID INHIBITION OF mRNA: RIBOSOME BINDING AND

PHENYLALANYL-TRNA SYNTHETASE REACTIONS

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THE UNIVERSITY OF ALBERTA

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FACULTY OF GRADUATE STUDIES AND RESEARCH. The undersigned certify that they have read, and recommend to the Faculty of Graduate Studies and Research, for acceptance, a thesis entitled "Aurintricarboxylic acid inhibition of mRNA: Ribosque Binding and Phenylalanyl-tRNA Synthetang Reactions", Submitted by Joan A. Zmean in partial fulfilment of the requirements for the degree of Manter of Science in Biochemistry. Supervisor, Um Paranchych ۶., ٤À DALM ADDALLA AND ADDALLA

ABSTRACT

A study of the inhibitory action of ATA on two different t steps of protein synthesis---the mRNA binding reaction and the phenylalanyl-tRNA synthetase reaction---was undertaken.

Studies of ATA action on the mRNA binding reaction revealed that ATA does not dissociate a preformed mRNA: ribosome complex. However, when added prior to the complex formation, ATA inhibits mRNA binding to 50% at a concentration of 20 µM. It was also confirmed that ATA inhibition is due to its binding to ribosomes. However, it was not possible to deduce the exact site and nature of ATA binding due to the complexity of ribosomal structure and, due to the presence of protein factors closely associated with the ribosomes. An examination of the inhibition from the viewpoint of ATA structure employing various aurin derivatives showed that analogs of ATA are not very effective inhibitors of mena bluding. It appeared that ATA is a unique inhibitor of mena blading due to the presence of hydroxyl and/or carboxyl groups on the Kriphonyimothano akalatal atructure. During the course of the above ARMALON, IR WAA NORAA LINGE ARE ORDER FRANKLONA ADDALLING RO ATA booking the arma blading reaction while the overall proceed of puly-U Alreaged phenylalaning independences in inhibited as a lover concentration of ATA than the wind bluding stap. The search for a reaction sen-ALLIVE, SO ATA LOA UN LO ALBOOVER LAR THE PHENYLALANYL-LENA BYNCHERABE reaction is inhibited by ATA. The ATA inhibition of any aminoacyletRNA synthetase had not proviously been reported. Thus, this novel system yan analyzad. Studian with aurin forlyatives showed that the analogs of ATA wore all potent inhibitors of this reaction, indiasting a different mode of ATA Aution than in the manA binding reaction. In apice of this

the inhibitory activity as was the case in ATA action on the mRNA: ribosome interaction. Studies into the site of ATA action in the synthetase reaction indicated that it binds to the enzyme. The stolehiometry of ATA binding to the synthetase was calculated to be approximately 6 ATA molecules per enzyme molecule. This binding of ATA to the synthtase was found to result in non-competence inhibition of the binding of tRNA to the enzyme. The author wishes to express her appreciation to Dr.S.Igarashi for his guidance and encouragement throughout the course of this thesis work. She would also like to express her sincere gratitude to Miss B. Garbutt for her technical direction in reticulocyte cell preparation.

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ATP	adenosine 5'-triphosphate
€TP	cytidine 5'-triphosphate
GTP	guanosine 5'-triphosphate
poly-U	polyur id yl at e
RNA	ribonucleate
mRNA	messenger ribonucleate
L K NA	transfer ribonucleates
0. V. LRNA	transfer ribonucleate treated with snake venom phosphodiesterase as described in (2-7)
Phe-tRNA	phenylalanyl-transfer ribonucleate
Phy	phonylalanine
۸۳ ۸	A gurthericarboxylle actd
тса	Trichloroaestic actd
Trin	trip (hydroxymethyd) anthomathaue
Drak	dlochylandnoothyl
R. cold	Rachartenta coll
c: k 2m	READANAAR 19 BOOK IN YAAAAR KI
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m l	milliliter 🦻
ا در	microliter
រោយ	millimeter
nın .	nanometer
m£n	minuto
• C	degrees Centigrade
MW	molecular weight
150	concentration of inhibitor multiclent to cause ' inhibition of a reaction).
G	Bravitational force
5	Svedberg unit ,
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The protein synthesizing machinery of living systems is composed of a number of protein and RNA components which interact with each other in a specific manner. These components are: more than 40 kinds of tRNA specific to individual aminoacids, and noacyl-tRNA synthetases which specifically recognize one of 20 amino acids, messenger RNAs whose numbers could be equal to the entire gene contents of an organism, protein factors which control each step of the protein synthesizing machinery (the three initiation factors, three translocation factors, and one or two termination factors), and ribosomes. The ribosome plays a central role in the protein synthesis by providing the binding sites for these reaction components (Haselkorn and Kothman-Denes 1973), The ribosome liself, however, is composed of two subunits, and in fact each aubunit is made of RNA (285 and 55 for the larger subunit and 185 for the small subunit) and more than two dozen proteins. The complexity of this ribosomal structure and the specific recognition by protoin subunity of a appealite alto of elboaomal RNA attll compain a myacory even in the Sta bactorial mystom (non Noviow by Kurland 1972). In fast, the mechaniam of the apaultic procolm: NNA incorrections involved in postein synthesis LA MAL ALMOLDARON LA DALA.

In this regard, the Adaptor Hypotheats by Orick (1958) is perhaps the clearest likestration of the questions that must be answer. It ad to understand the process of protein synthesis, The original problem of how the nucleotide sequence is franklated into an smino acid sequence is now bains replaced by the new question of how to couple the smino acids to specific sequence (tRNAs). After all, the engymes, sminoscyle

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tRNA synthetases, are a sequence of amino acids, and they must recognize the specific combination or sequence of nucleotides of the tRNAs. Therefore, the elucidation of the mechanism controlling these protein:RNA interactions is extremely valuable in understanding protein synthesis. This study deals with an inhibitor of such protein:RNA interactions involved in an early stage of protein synthesis. In particular, it was our intention to study the action of such protein and on the failudacylits derivatived on the mRNA:ribosome interaction and on the failudacyliRNA synthetase reaction (see Fig. 1 for structure of ATA derivatives).

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It was in 1954 that the aurin dyes first drew some attention in blowedlest studies when white at al demonstrated that ATA can be used in the treatment of beryllium polaoning in mice. Their studies into the mechanism of ATA action indicated that ATA likely removes beryllium from the streatment of beryllium polaoning in mice. Their studies into the mechanism of ATA action indicated that ATA likely removes beryllium from the streated of action by chelating the ion (Schubert at al 1952, Lindenbaum et al 1954, Lindenbaum and White 1954). This mechanism appears to analogous to the teshnique of mordanting cotten in dye technology. The test using triphenylmethane compounds other than ATA failed in the polaonid reatment (White and Schubert 1954, Lindenbaum and White 1954). Later studies on the distribution of ATA in the tissue of berylliumpolaoned mice hed to the complusion That the ATA is somplex bound to protects fibers of the fissue examined (Lindenbaum as al 1954, Schutbert 1 and Aindenbaum 1954, Lindenbaum and Lisso (2956).

At about the same time, physical and chemical, shudles on the dystprotectin interaction were progressing with emphasize on developing probes into conformational appears of the proteins (vares 1951, brown 1951, Shimao 1952, Kusunoki 1952). Brown studied the developation of dys solutions by proteins, and appearland that the developation of dys



BOLULIONE may be due to complex formation between the dye and the added protein molecules. It was also found that the oxidation offsh-groups remults in reduced decoloration of dye by protein, indicating the involvement of SH-groups in the dye:protein interaction. Brown further postulated that the site of interaction on the dye molecule was the central carbon atom of the triphenylmethane dye.

It was almost two decades after the work of Brown's group that the scientific value of ATA'was rediscovered. Groliman and Stewart (1968) reported that ATA inhibits the translation of bacteriophage f2 RNA in the <u>E</u>, <u>coll</u> call-free system. They demonstrated that ATA action is exerted at the initiation step of protein synthesis. This was a novel finding since there had been no inhibitor specific to the initiation step at that time. Thus the role of ATA in arresting the reaction at the mRNA; ribosome interaction is of considerable interest in the field of protein synthesis.

Immediately following the report by Grollman and Stewart (1968), the effect of ATA on a variety of muchostidyl transferance reactions and on protein biosynthesis was examined by the members of Dr. Igarashi's laboratory—including myself. The ensymes tested were: DNA-dependent RNA polymeriase of <u>E. coli</u>, R17 RNA-dependent RNA polymerase, tRNA 3'-terminal nucleotidyl transferase from <u>E. coli</u> and L. colis, to aminoacyl-tRNA synthetases and synthetic mRNA binding to ribosomes (all unpublished dats). All of the ensymes susceptible, to ATA suggested to us that ATA is not, a specific inhibitor, contrary to the initial proposal by Grollman and Stewart (1968), Howeyer, all of the reactions tested appear to be sensitive rather specifically to ATA. Other triphenylmethanes tested require concentrations 10 to 100 times that of ATA to exert the same level of inhibition. We thus directed out study toward the elucidation of the mechanism by which ATA uniquely inhibits the enzyme reaction in hope of providing some insight into the specific recognition of RNA by protein. The present study was therefore aimed specifically at the determination of which components of the mRNA binding reaction and of the aminoacyl-tRNA synthetase reaction, both of which pecur is the initial stages of protein synthesis, could complex with ATA. An attempt was also made to determine the effect of ATA on the binding of other ligands to the enzyme.

This study employed the incorporation of phenylalanine in the reticulogyte collefree system as directed by poly-U. At the outset, the concentration of ATA required for 50% inhibition of the mRNA binding reaction was established. Further studies showed that the site of ATA action is on the ribosome, but that the preformed mRNA ribosome complex is resistant to ATA section. The determination of the ATA binding site was not possible due to the complexity of the ribosome structure and the presence of associated proteins. An examination of the inhibition from the viewpoint of ATA structure was carried out using ATA analogs The results showed that the presence of carboxy and/or hydroxyl groups is required for inhibitory action.

For the examination of ATA action on aminoacyl-tRNA synthesis, the Phe-tRNA synthetase reaction (as used. The study using ATA analog's in this system suggested a different mode of action from that in the mRNA binding Adaction, even though the carboxyl and/or hydroxyl groups are meassary for the inhibition. The results clearly show that ATA binds to the enzyme and quantitation of the ATA:enzyme interaction indicated that there is a specific number of ATA molecules bound per enzyme molecule, at approximately a ratio of 6:1. An examination of the effect of ATA on the binding of other ligands to the enzyme showed that it interferes with the binding in a non-competative manner.

MATERIALS AND METHODS

(2-1) Radioactively Labelled Chemicals

The following radioactively labelled chemicals, with specific activities as given in perenthesis, were used in this study. ³H-CTP (20.4 C1/mmole) and ³H-phenylalanine (5-16 C1/mmole) were purchased from New England Nuclear. ³H-polyuridylate (52.8 mC1/mmole polynucleotide phosphorus) was from Miles Laboratories ing.

(2-2) Chemicala

ATP, GTP, and CTP were from Schwarz/Mann. Polyuridylate was purchased from Miles Lab. Inc. Phosphoenolpyruyake, Trisma base, heparin, and phenylhydrazine-MCl were from Sigma. Puromyein was from Nutritional (blochemicals. All of the triphenylmethane dyes except ATA (a baker product) were obtained from Marleso. Sepharose, Sephadex, and Agarose were purchased from Pharmaeia Fine Commical Co, DEAE-cellulose was from Carl Schleicher and Schwell Go, PPO (2,5-diphenylosasole) and POPOP (1,4bis(2-(5-phenyloxyzoylbenzene)), which were used in seintillation fluid, were purchased from New England Nuclear, All other ehemicals were reagent grade from Baker.

(2-3) Blological Margriala

New Zealand white rabbits (2-2 kg) were used for the preparation of retiquicayte cells, Yeast tRNA (unfractionated) and phosphodiesterses from <u>Crotalus adamanthus</u> yenom were purchased from Calbiochem. The marker proteins (glucose oxidase, chymotrypsinogen A, and lysozyme) used for molecular weight estimation were obtained from Sigma.

(2-4) Buffered Solutions

The following buffered solutions were used wherever suitable: Buffer A (TT1s 10 mM, KCl 5 mM, MgCl₂ 0s2 mM, sucrose 50 mM, final pH 7.6). Buffer B (Tr1s 50 mM, KCl 5 mM, MgCl₂ 0.2 mM, final pH 7.8). Buffer RM (Tr1s 400 mM, KCl 60 mM, MgCl₂ 10 mM, β -mercaptoethanol 4 mM, final pH 7.8). Buffer RN (Tr1s 25 mM, KCl 40 mM, MgCl₂ 5 mM, β -mercaptoethanol 5 mM, final pH 6.5).

(2-5) Preparation of Rabbit Reticuleyte Polysomes and Monosomes

Rectaulocyce calle were obtained from New Zealand white rabbica (2-3 kg) which had been created with five daily injections of 2.5% noutralized phonylhydrazing HCL at a domage of 0:33 ml per kilogram body weight per injection (Burny and Marbaix 1964). After a two-day reating period, the rabbits were angethetized by injection of 2 ml Nembusal (50 mg/ml) anto the marginal yein of the ear, immediately, here ARIM (2 m) of a 1% solution) was injected. Then blood was collected through the marginal yein of the ear values a vacuum apparatue. The percontage of reticulocytes was examined under a microscope. When the reticulocytes exceeded 85% of the total blood cells, the blood was proceased as follows; the cells were lysed in a hypotonic saling solution , and insoluable materials were removed by centrifugation at 10,000 x G. The resulting supermanant was further clarified by centrifugation at 30,000 x G. This fragiton was then subjected to high speed contribus-ALLON AL 142,000 x G for 40 min in a preparative ultragentrifuse to precipicate polysomes. The resulting supernatant was then incubated at

37° for 40 min in the presence of added puromycin (1 mM) and GTP (0.1 mM) in order to convert residual polysomes to monosomes. The monosomes were precipitated at 142,000 x G for 90 min and suspended in Buffer A. The monosome concentration was adjusted to 100 A_{260} units per ml and stored in a liquid nitrogen refrigerator.

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(2-6) Proparation of Aminoacyl-tRNA Synthetase from Relieulocyte Calla

A mixture of aminoacyl-thNA synthetases was prepared from the supermatant of the high speed centrifugation, post-monosomal supermatant, as described in section (2-5). Two volumes of saturated animonium sulfate in Buffer A were added to one volume of the supermatant, and the resulting solution was allowed to stand in the cold for 60 min. The precipitate was then collected by centrifugation at 15,000 x G for 20 min in a Beckman model J21 refrigerated centrifuge. The precipitals was dissolved in Buffer A to give a protect concentration of 20 mg/ml, and was stored at -20°C until mag. This fraction was designated as the AS66-fraction.

Partial purification of PhantKNA synthetase was performed as follows: a portion of AS66 fraction was dialyzed overmight against Buffer B at 4°C in order to reduce the concentration of the monovalent dation to approximately 20 mM KCl equivalent, The reason for the choice of Buffer B is that the higher Tris concentration is required to maintain constant pH during the subsequent column chromatography. The dialyzed fraction was fractionated on a DEAE cellulose column using a batchwise method, The column was first developed with Buffer B and them with Buffer B containing 0,1 M ammonium sulfate, PhentRNA synthetase activity was found in this 0,1 M ammonium sulfate fraction, while the

majority of hemoglobin was eluted in the Buffer B wash. The fractions containing the synthetase activity were pooled and dialyzed at 4°C for 18 hr against Buffer B containing saturated ammonium sulface. The precipitate formed in the dialysis tubing was collected by centrifugation at 15,000 x G. The precipitate was then dissolved in a minimal volume of Buffer B. This enzyme fraction was then subjected to molecular activing chromatography using Sepharose 68. A 20 mg portion of the enzyme fraction was loaded onto a Sepharose 68 column of bed volume 150 ml and processed with Buffer B containing 150 mM ammonium sulfate. Each fraction was examined for Phe-tRNA synchetase activity , and Active fractions were pooled and dialyzed against saturated annonium aultare in Buffer B overnight. The precipitate was collected by centrifugarion, and diamolyed. In Buffer B to give a final concentration of 10 ms/ml. This fractionstion procedure achieved a 50-fold increase in Q appairie Aurivity of the Aynthetabe from the ASGG fraction.

(2-7) Proparation of Snake Venom Phoaphodiesterase Treated tRNA (A. V. CRNA)

An

In order to provide the substrate for the tritiation of thes, yeast the was subjected to a limited digestion of the 3'sterminal and by snake venom phosphodiesterses. An incubation mixture contained in 2 ml, the 50 mg, phosphodiesterses 0.045 units, lysing 0.1 M (pH 9), MSC12 1.2 mM. After 30 min reaction at 37°C, the reaction mixture was extracted three times with 1 ml of water saturated phenol. From the final squeous layer, RNA was precipitated by the sales methanol method of issrashi and McCalla (1971). The precipitate was washed with ether-methanol, dried, and dissolved in water. The these treated with anake venom phosphodiesterses under the conditions specified

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lacks the 3'-terminal residue -CA and in small proportions -CCA and is designated as s.v.tRNA (Igarashi and McCalla 1971).

(2-8) Radioactivity Measurement

The filters that retained radioactive materials were placed in vials containing 5 ml of the scintillation fluid composed of PPO 6 g, POPOP 0.6 g, in toluene 1 liter. The radioactivity was measured in a Beekman Scintillation System LS-200B. A 10% counting efficiency was obtained for trittium.

(2-9) Proparation of "H-yeast LKNA

The method used to propare labelled yeast tRNA involved addition of 3M-CMP to the 3'-terminal of A.V. thna catalized by 3'corminal musicocidyl cranefores of Localla (Igereant and Lerrec 1974). The reaction mixture (0.1 ml) contained lyaine (pH 8.5) 100 mMh Macla 1 mM, Mach, 1 mM, KC1 200 mM, A-mercaptoethanol 0.5 mM, in addition, 10 HOL OF 3H-OTTE, I, ME OF M.Y. EKNA AND 100 HE OF ERNA 3'- CARMINAL AND LOGthey cranafaraas were added. This reaction mixture was incubated at 37°C for 60 min. At 60 min, 0,1 moles of CTP per 0.1 ml of reaction MIXENER WAR added, thousand at 37°C for 10 min and finally 0.5 musica AWP POR O.1 MI. REACTION MIXEURA WAR added and incubated addaddigional 20 min at 37"C. To the reaction mixture, an equal volume of water sat unared phanol was added, After 5 min of vigorous mixing using a Vortex mixer, the aqueous layer was separated from the phenol layer by means of contrifugation at 15,000 x C for 5 min at 15°C. In order to ensure clear reparation, the rotor was stopped by coasting. The aqueous layer was reextracted with phenol, Then two volumes of methanol were added to pre-

cipitate tRNA. After one hour of standing in the cold, the precipitate was collected by centrifugation at $10,000 \times G$ for 10 min. The pellet was washed twice with a salt-methanol mixture (1 M KCl : methanol = 3:7, in volume), once with a methanol-ether mixture (1:1) and finally with ether. The final pellet was dissolved in distilled water to a RNA concentration of 10 mg/ml, yeilding 17,000 cpm per microliter. 12

(2-10) Assay Method for Polyphenylalanine Synthesia

Protoin synthesis in the retioulocyte cell-free system was measured by following the poly-U directed incorporation of ³H-phenyl-Alaning into a hot TCA insoluable material. The reaction mixture (0.1 ml) contained: Tris (pH 7.8) 100 mM, MgCl2 10 mM, KCl 20 mM, prmercaptor athanol 5 mM, ATP 1 mM, GTP 5 µM, phoaphoenolpyruvate 0.5 mM. In addicton, 2 µC1 of "H-phanylalaning, 10 µg poly-U, 100 µg of poat-ribosomal proteins and 2 A260 units of ribosomes were added. The reaction rook place at 37°C for 60 min. At intervale, 0.1 ml aliqueta were with-Arown and placed on filter discs. The discs were processed with the hot_{F} TCA wash method, and then subjected to radioactivity measurement as previously described (isarashi and Paranchych 1967). In shore, the filter disca were boaked in gold 10% TCA and allowed to stand in the gold for 40 min. followed by 5% TCA wash swice as room semperature. Diaga were shen heated to 90°C for 45 min in 5% TCA, and washed twice with 5% TCA. They were extracted with ether-ethanol (1:1) mixture for 15 min and finally with other for 10 min. After being completely dried, the diage were placed in a colugne adimitilation fluid. The radioactivity recained on each filter was measured as described in (2-8) "

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(2-11) Assay Method for Phe-tRNA Synthesis

Aminoacyl-tRNA synthetase activity was measured by the incoporation of 3 H-phenylalanine into a cold TCA insoluable fraction. The reaction mixture (0.1 ml) contained: Tris (pH 7.8) 100 mM, MgCl₂ 10 mM, KCl 30 mM, β -mercaptoethanol 5 mM, and ATP 1 mM. In addition, 2 μ Cl of 3 H-phenylalanine, 100 μ g of tRNA and 100 μ g of the synthetase were added to each 0.1 ml of reaction mixture. The reaction took place at 37°C for 20 min. At intervals, 0.1 ml aliquots were removed and processed with the cold TCA wash (lgarashi and Paranchych 1967). In short, the/discs were immersed in cold 10% TCA for 40 min, followed by four suscessive washes with cold 5% TCA for 15 min each time. They were then expacted with ether-ethangl (1:1) for 15 min, and finally with ether for 10 min. After being dried, the discs were placed in a toluene acintification finial and subjected to radioactivity measurement as deseribed in (2-8).

(2-12) AAAAY Mathod for Poly-U Aluding to Ribohoman

³H-poly-U bluding to ribosomes was measured ellipse by the membrane filtration method using Millipse filters DA (pore size 0.65m) (Isaraahi and Paramehyah 1967), or by the sel-filtration method using Asarose A50M. In both methods, ³H-poly-U (20 µmoles polynucleotide phosphorus) man mixed with ribosomes (5 A₂₆₀ units) in 0.5 ml of the reaction mixture containing: Tris (pH 7.8) 50 mM, MgCl₂ 5 mM, KCl 40 mM, and p-meruppeosthanol 1 mM. The reaction mixture was incubated in the cold (4°C) for 5 min, and examined for mRNA binding by one of the two methods mentioned above.

For the Millipore filtration method, the samples were diluted

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with 5 ml of Buffer RM and filtered through the filters using a suction apparatus. The membrane filters were 'further washed with 10 ml of Buffer RM containing bovine serum albumin (1 mg/ml). After being dried, each filter was examined for radioactivity retained as described in (2-8). A control sample, which did not contain ribosomes, was processed similarly. The difference in radioactivities between experimental and control samples represented the amount of ³H-poly-U bound to ribosomes. It should be mentioned that these membrane filters had to be pretreated in order to reduce nonspecific retention of ³H-poly-U by soaking the filters in 0.1 N NaOH, rinsing with distilled water several times; and then soaking in 0.1 N HCL. Finally, they were washed with distilled water unitit mentralized, dried at room temperature, and tested for nonspecific retention of ³H-poly-U retention to a megligible level (less than 200 cpm).

For the gel-exclusion method, an Agarosa ASOM column of bed volume 10 ml was used. The reaction mixture (0.5 ml) was loaded onto the column, and then developed with Buffer A. In order to determine the slution position of ribosomes as well as of poly-U, they were processed separately on the same column under identical slution conditions to those above. Since the slution positions of these two components are quite far apart when processed separately, the radioscrivity which sluted at the same position as the ribosomes was attributed to poly-U , bluding to ribosomes.

(2-13) ABBAY Machod for HatRNA Hinding to PhantRNA Synthetase

²H-ERNA WINDING DO THE AUTOMOTION WAS MEABURED UNING MILLIPORE filters DA. The partially purified synchetase (50 µg) was mixed with ³H-tRNA (10 pmoles) in 0.5 ml of the reaction maxture containing: Tris (pH 6.5) 25 mM, MgCl₂ 5 mM, KCl 40 mM, and (3-mercaptoethanol 5 mM. The reaction mixture was incubated in the cold for 5 min, diluted with 5 /ml of Buffer RN, and filtered through the membrane filters using a suction apparatus. The filters were then washed with 10 ml of Buffer RN containing 1 mg/ml of bovine serum albumin, and examined for radioactivity as described in (2-8).

(2-14) Sepharome on Column Chromatography

Sepharone of column chromatography was used for two types of experiments under identical conditions. The first of these was designed is detect ATA bluding to individual reaction components. The second was to estimate the molecular weight of the enzyme. For both studies, a column of bed volume 5 ml (5 mm x 130 mm) was used and developed with Shuffer RM. For the use of such a small column, the volume of sample was limited to 50 ml.

For the bludles studled, the reaction mixture was chromato-Braphed first, followed immediately by chromatography of individual reaction components. In order to determine the peak positions of these components, the effluent was monitored at suitable wavelengths: 260 nm for these and 508 nm for proteins and 308 nm for ATA.

For the molecular weight determination, the ansyme solution was first processed on the column. Then four proteins of known molecule as weight were chrometographed in order to obtain the correlation beinverse the elution volume vs. molecular weight. These marker proteins wood ware: glucose exidance, 2.86 x 10⁵; hemoglobin, 6.8 x 10⁶; chymokrypainesen A, 2.5 x 10⁶; and typesyme, 1.4 x 10⁶. THE EFFECT OF ATA AND AURIN DERIVATIVES ON WENA; RIBOSOME BINDING

(3-1) Introduction

The initiation of polypoptide synthesis in biological systems involves a merica of reactions leading to the formation of the complex abaignated as the 'initiation' complex (Nomira and Lowry 1907) which the Ludon the ribonous, mena, and anthoneyl-tena an the throg key com-POMPHER, IN Addition, A number of protoin factors control the inter-Autions among these chees reaction components (Reval and Gross 1966). Although the general teatures of the process leading to the formation of this initiation owned are now well understood, the nature of the theoremetion borroom the individual reaction numponements has not you byon cluckdared. One of the reasons for such a delay in understanding the manifember may be attributed to the lask of antiable inhibitors for the Andly Adnal aropa in complex formation. For this reason, the report that ANA waa a apactfic inhibitor of the initiation of polypaptide AYNCHOMEN (OROLLMAN AND SLOWARE 1968) WAS MALKNEELANE WEADAVORY, THE finding was made wains the basisfial self free everem and econ was sonfirmed by other groups nating mammallan and plant coll-tree systems (Loblow as at 1970, Maraun as at 1970). With the hope that a detailed ANALYALA OF ATA INHIBIRION OF INITIALARION COMPLEX FORMARION COULD BANK aoma fisht on the manhantam of interaction between mina and ripodomea. the dealded to undertake a army of the othes, of ATA on the bludtes of POLY-U . W. RADDLE ROLLANDOYEN REDOROMON

The endlose described in this chapter involve the decomman-

tion of the inhibitory concentration of ATA in polyphenylalanine synthesis and in the poly-U:ribosome binding reactions, the determination of the site of ATA action, and the determination of the structural features of ATA responsible for the observed inhibition. These studies led us to conclude that ATA is the most effective inhibitor among the aurin derivatives tested, and that ATA forms a stable complex with ribosomes. ATA thus appears to be a unique inhibitor of mRNA:ribosome interaction. However, as will be described in Chapter 4, it was found that ATA analogs inhibit reactions other than the mRNA:ribosome complex formation, in particular the synthesis of the procursor molecule-aminoacyl-tRNA. Thus, ATA is not a specific inhibitor for mRNA:ribosome interaction (Isarashi and Zmean 1974).

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REFARE OF ATA ON POLYPHANYLALANLING SYNCHAALA

The inhibitory concentration of ATA in the poly-U directed polyphenylalanine synthesis was determined using rabbit retionlocyte mononomes supplemented by the ASDO fraction as a source of Phe-tRNA synthetage and other factors, The complete reaction mixture as deseribed in the Methods was used with the presence of varying concentrations of ATA. The reaction was carried out at 37°C for 40 min and the samples were processed as described in the Methods, The level of 3 Hphenylalanine incorporation at 40 min of reaction was expressed as perent of the control experiment which did not contain ATA, Fig. 2 demometrates that ATA inhibits the overall protein synthesizing activity' to 50% at approximately 5 μ M (150 \sim 5 μ M). Thus the inhibition of resign ulocyte protein aynthesize was established. 17



<u>Fig. 2</u> Inhibition of polyphonylalaning synthesis by ATA. Poly-U directed ³H-Pho incorporation was assayed as described in the Methoda (2-10). The incorporation of ³H-phonylalanine in the absence of ATA was taken as 100%. Activity demonstrated in the presence of varying amounts of ATA was expressed as a percent of the control value.

Effect of ATA on the mRNA Binding to Monomeric Ribosomes

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At the time this investigation was undertaken, it was generally believed that ATA was a specific inhibitor of mRNA: Lbosome complex formation. Therefore, an experiment was designed to determine whether the foregoing inhibition of polyphenylalanine synthesis by ATA is attributable solely to an inhibition of the poly-U;ribosome interaction. In order to do so, the interaction between monomeric rabbit reticulocyte rthomomen and "H-poly-U was examined in two ways. The first of these methods involved the use of the membrane filtration method employing Millipore filters DA (pore size 0.65µ). The reaction mixture containing ribosomas (5 A260 mairs) and ³Hopolyou (20 mmalas polymuclauride phoaphorna) was incubated in cold Buffer RM for 5 min, then subjected to membrane filtration. The effect of various concentrations of ATA on the bludlug reaction was examined and expressed by persent inhibition. An Ahova in Fig. 3 the level of ATA regulred for 50% inhibition (150) of the reaction to 20 µM. It to noteworshy that that 150 value to alleners prepare then the 120 for stardy both both and an energy a This difference led us to the investigation of other ATA sensitive reactions in protein synthesis as desorthed in Chapter 4.

The second method used to study the effect of ATA on the binding of mENA to ribosomes was the gel exclusion method using Agarose A50M. The reaction conditions had to be slightly modified from those used above due to the limited sensitivity of this method. The ratio of ATA to ribosomes was set at l_0 (no ATA, 5 A₂₆₀ units of ribosomes), I_{50} (20 µM ATA, 5 A₂₆₀ units of ribosomes), and I_{80} (40 µM ATA, 5 A₂₆₀ units of ribosomes). An excess of poly-U was used to ensure maximum binding of poly-U to ribosomes in the system. Although the column

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Fig. 2 ATA inhibition of ³H-poly-II binding to monomeric ribodomea. Assays were performed as given in the Methods using Millipore filtration. The reaction mixture contained 5 A_{260} units of ribosomes, 20 pmoles poly-U phosphorus, and various concentrations of ATA. The level of ³Hpoly-U bound to monomeric ribosomes in the absence of ATA was taken as 100% setivity. ³H-poly-U binding in the presence of varying smounts of ATA was examined, and the results were expressed as a percent of the control level. 100% corresponds to 18,000 apm.

chromatography method gives rise to qualitive rather than quantitative data, Fig. 4 shows the expected inhibition values, and can be considered as a reconfirmation of the data obtained by the membrane filtration method.

Site of Action in the mRNA: Ribosome Binding Reaction

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The next step in this study was to scortain which reaction component interacts with ATA. If ATA inhibition is mediated by complexthe with one of the reaction components, then increasing the concentration of that particular component in the presence of a fixed concentration of the dye should relieve its inhibitory action. The first component to be examined in this regard, therefore, was poly-U. It was added in increasing amounts to a reaction mixture containing CONALANC amounce of ATA and ribosomes, Fig, 5 shows that for all poly-U concent tractions teaned, the inhibition due to ATA was maintained at a level of about 69%. The offect of varying the ribosome concentration in this ayarom waa axaminad naxe, and the reamire are shown in Fig. 6. It may be seen theorem the theorem concentration was increased, there was a dea grades in the inhibition of poly-U binding. The foregoing two experimanta thus suggest that the ribosome most probably is the component which interacts with ATA. However, the results in Fig. 6 would also be interpreted in terms of ANA binding to impurities in the ribosome solution.

To exclude the latter possibility, a study was carried out whereby the isolation of a ribosome ATA complex was achieved by gel exclusion chromatography, First, poly-U and ribosomes were individually processed through a Sephadex G25 column of 10 ml bed volume, using



<u>Fis. 4</u> ATA inhibition of ³H-poly-U binding to monomeric ribosomes as detested by the sel exclusion method. Assays were performed as given in the Methods using Asarose A50M column chromatography. Ribosomes $1(20 \ A_{260} \ units)$ were mixed with excess ³H-poly-U (200 nmoles phosphorus) in the presence of no ATA (O), 80 µM ATA (\square) and 160 µM ATA (Λ). The mixtures were incubated in cold Buffer KM for 5 min, diluted with the same buffer and chromatographed, 260 nm absorption (—) and radioactivity for each fraction were measured.

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MOLES NUCLEOTIDE PHOSPHORUS

Fig. 5 Effect of ATA on poly-U: ribosome binding at various concentrations of poly-U. The reaction mixture (see Materials and Methods) contained 20 μ M ATA, 5 A₂₆₀ units of ribosomes, and ³H-poly-U as indicated on the abscissa. The mRNA: ribosome complex retained on Millipore filters was estimated by measuring the radioactivity derived from ³H-poly-U bound. Percent poly-U bound (ordinate) indicates the calculation based on the radioactivity in the experimental set over the value in the control set (no ATA present) at a certain concentration of poly-U.

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<u>Fis. 6</u> Effect of ANA on poly-U: Fibosome binding at various concentrant tions of Fibosomes. The reaction mixture contained 20 μ M ATA, 20 nmoles ³H-poly-U phosphorus, and Fibosomes as indicated on the abelses. Reach tion conditions and assay procedures are the same as those in Fis. 5. Percent poly-U bound (ordinate) indicates the calculation based on the radioactivity in the experimental set over the value in the control set (no ATA present) at a given concentration of Fibosomes.
Buffer RM As expected, these two components were excluded from the column due to their large molecular weight. ATA, on the other hand, was included in the column and thus was eluted with the included volume. Secondly, poly-U and ATA were mixed and subjected to column chromatography. The elution profile corresponded to that obtained when these components were chromatographed individually; that is, poly-U appeared in the void volume and ATA appeared in the included volume. However, when the mixture of ribosomes and ATA was chromatographed, there was only one ATA peak in the void volume and none in the included volume. This was taken to mean that the ATA had formed a complex with the ribosome.

An attempt was made to dissociate rabbit reticulocyte polysomes by ATA treatment into monomeric ribosomes and free mRNA. Adthough bluding of ATA to the polysomes did becur as detected by a spectrophotometric measurement, there was essentially no dissociation of the polysomes even at ATA concentrations as high as 200 µM. This result indicated that a preformed mRNA: ribosome complex is protected from ATA inhibitory action.

Although the above results indicated that the ribosome is the site of interaction with ATA, further studies with this system were complicated by the fact that the ribosome fraction used in these studies contained messenger RNA binding factors strongly associated with ribosomes (Igarashi and Dufresse 1974). It was therefore possible that ATA binds to the protein which mediates mRNA binding to the ribosome, or that it binds non-specifically to one or more of the structural components of the ribosome itself. The exact site of ATA action could thus only be determined by purifying the factors and

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ribosomal proteins, and examining the interaction of ATA with each of them individually. However, at the time this study was undertaken, the identification of these factors and the individual ribosomal proteins had not yet been completed. Thus no further study on the site of ATA binding within the ribosomal structure or with factors was possible.

Comparative Study on the Inhibitory Action of Aurin Derivatives

An important aspect of ATA action in this system, yet to be explored, was the structural feature of ATA responsible for the observed inhibition. A series of surth derivatives were chosen on the basis of salient differences in their side chain constitution, and tested for their capacity to inhibit the mRNA; ribosome binding reaction using the Millipore filtration method (see Fig. 1 for structures). It is obvious from Fig. 7 that these compounds inhibit mRNA; ribosome binding to some extent, but the 150 values are at least 30 times higher "than that of ATA. However, it is noteworthy that the order of inhibitory power is: ATA ($150 - 20 \mu$ M), Aurin ($150 = 500 \mu$ M), Asure blue b ($1_{50} - 1500 \mu$ M). The foregoing results suggest that both carboxyl and hydroxyl groups are required for effective inhibition.

F18. 7 Effect of various aurin derivatives on poly-U:ribosome binding. ³H-poly-U (20 nmoles phosphorus equivalent) was mixed with ribosomes (5 A₂₆₀ units) in the presence of various aurin derivatives, and the resulting complex was measured by the Millipore filtration method as described in the Methods. For each concentration of the individual dyes, a calculation based on radioactivity in the experimental set over the value in the control set (no dye present) was done. This value is shown in the figure poly-U bound (ordinate) where 100% represents the control set (\bullet) Aurin, (\bullet) Asure blue b, (\blacksquare) Fuchsin Acidic, (\bullet

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The results reported in this chapter indicate that ATA inhibits both polyphenylalanine synthesis and poly-U binding to monosomes in the reticulocyte cell-free system. These results are in good agreement with the report of Grollman and Stewart (1968). However, they concentration required for 50% inhibition of polyphenylalanine synthesis was 5 μ M, while that for 50% inhibition of poly-U binding to ribosomes was 20 μ M. Thus ATA is four times as effective in the incorporation reaction than in the mRNA: ribosome binding reaction. This indicates that there may be steps in the overall protein synthesizing system senallive to ATA in addition to the mRNA: ribosome binding reaction. This prodiction can be supported by the recent reports of Stegelman (Stegelman and Aptrion 1971, and Sloselman 1970) that various stops of protoja synthesis can be inhibited by ATA. Further analysis of our system, as deperibed in Chapter 4, revealed that among the individual stops of prototh Aynthonia, anthonylathia aynthonia is strongly inhibited by ATA. This the very first reaction in protoin synthesis, i.e. the formation of productor is inhibited by ATA. This fact has not been reported to the literature to date and may be the values for the bigher ADMALLIVLEY OF the Indurporation realiton toward ATA. SALLING ANIDO this now finding for discussion in the next chapter, the present dis-AMARION WILL ADMARATARA ON the dys affaat on MNAITIDOROMS bluding ILAALT.

In order to probe the mechanism of ATA Action, three experimental approaches were taken. These included the effect of ATA on preformed polynomes, the relief of ATA inhibition by increasing the concentration of reaction components, and detection of stable complex

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formation between ATA and a reaction component. One experiment, in which the order of adding reaction componeats was manipulated to see whether or not a preformed complex between ATA and a given reaction component changes the degree of ATA inhibition, showed that when ribonomen and poly-U were mixed prior to adding ATA, the inhibition was mosligible, This fact indicates that ATA interfores only with a process leading to the stable complex formation between mRNA and ribosomes. In fast, when invisited polynomes from reticulocytes were incubated with ATA, there was virtually no dissociation of polynomes into monosomes. This realerance of polynomes towards ATA action aproon with the above observation that ATA is less attactive on preformed poly-U; ribosome complexes. Recent studies (Rhoads of al 1973, Henshav of al 1973, and RINNAR CLIMMAR AL AL 1973) WATCH ANDPORT THIA COMMINATION, domination of that the addition of low levels of ATA does not affect the structure of the bollynomen at feast for the time reduited to complete nearent bolls. pepkide alialine.

The second set of experiments was designed to determine the site of ATA sector, and consisted of changing the concentration of individual reaction components in the preschoe of a fixed concentration of ATA. Early studies of Brown (1951) and Lindenbaum (1956) indicated that ATA would bind to proteins. However, no studies were reported that ATA would bind to proteins. However, no studies were reported examining the binding of ATA to RNA. The results reported here showed that varying the concentration of poly-11 at a fixed concentration of ATA does not alter the effectiveness of the dye action, but that inereasing the concentration of ribosomes, showed a definite decrease in the ATA inhibition of poly-11 binding to ribosomes, it is worth mentioning that the protein synthes'sing reaction requires Ms⁺⁺ as a cofactor.

Therefore, the possibility that the ATA inhibition is the result of chelating $Mg^{\pm\pm}$ should be considered carefully. The $Mg^{\pm\pm}$ concentration in the reaction mixture was 1 mM or higher depending on the reaction, whereas the effective concentration of ATA is between 5-20 μ M. Thus, to significantly reduce the $Mg^{\pm\pm}$ concentration to produce the observed inhibition of the reaction, each ATA molecule would have to chelate at least 20 $Mg^{\pm\pm}$ ions, such an occurance is chemically improbable and it was therefore concluded that the ribosomes are the alte of ATA inter-

The chird apt of experimence provided further support for the above conclusion. Get exclusion chromekosraphy of mixtures of ATA and poly-U or ATA and elboarman alward that ATA binds to the elboarman to form a atable complex which witherande the Bel exclusion proceed. A Almillar atable complex between ATA and poly-U was not detected. Other LABORALORINA DAVE REPORTED ALMILLAR REALIZE (GROLLMAN AND SLOWARE 1970, and Halbars at AL 1971). It to thus aloar that ATA blading to ribosomen namaga an inhibitions of the poly-U binding reaction. This observation streaking the nature of Alla blughtone: (1) what to the mature of Alla blugting to the ribosomes? (2) how many APA molecules will blud per ribosome? At the point howayar, difficultion ware encountared in our scudies due to the complexity of the ribosome Atructure. Further delineation of the binding airs must differentiate between ribosomal KNA, ribosomal protein, and the protein factors associated with the ribosomes. Separation and purification of these escuetural components of the ribosomes would be naveragely of complete of an analgument of the blading alte of ATA which to directly reaponable for inhibition of the mana binding reaction. An ALLAMPT WAN FIRST MADE TO PROPARS REDICULOSYLA MONOBOMOR FRAS OF

poly-U binding factor using a method which was later applied to the rabbit liver monosomes successfully (Dufresse and Igarashi 1974). However, it was not effective in the reticulocyte system, since the method gave only about 50% removal of the poly-U binding factor as compared to 99% removal from liver monosomes. The reason for this difference between liver and reticulocyte ribosomes is as yet unknown. Since the poly-U binding factor of the reticulocyte systemwas very strongly associated with the ribosome, it was not possible to carry out fighter studies on the nature of ATA binding to reticulocyte ribosomes,

The accoupt to clusidate the mature of the ATA binding alto WAR LIMITED TO THE OXCOME JURE REPERTING IN the proceeding eaction. HOWEVER, ONE MORE IMPORTANT APPERT OF THE INHIBITOR BLURY MURE DE COMaldered, I.e. the armetural alement of ATA reapenables for the observad hamberlion. In this reserve the effect of various antia derivatives on the winderthoacome blocking reaction was examined. The results showed that all of the compounds costed are capable of inhibiting the reaction to nome extent. They are, however, much lease attactive than Alla--consoleration of at logat 25 times that of AXA is required to produce a comparable level of inhibition. The analysis of these data allows us to probe the mechanism of ANA action one step further. The aurin deriva-ELVER WICH IS VALUER CLOBER TO THAT OF ALL ARE AUTH AND AQUED BLUE B. A common feature of these two compounds is that they are negatively sharbad at physiological ph. It is known shat ATA forms "lakas" by complexing with metal lone (Lienko es al 1971). Therefore the possibility of alialation of the Mg. ion once asain comes to light, However, as ALALAA Aboya, whe ATA molecule must chelate at least 25 Mg TT long under

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the experimental conditions used. This is physically impossible and therefore, the mechanism of ATA action cannot be chelation.

This conclusion led us to a close examination of the structure of ATA, Aurin and Azure Blue B. It is evident that the important functional groups common to these compounds are hydroxyl and/or carboxyl groups. These groups are not found on the other triphenylmethanes tested—Fuchain Basic, Fuchain Acid and Erichrome Cyanine— which are less effective inhibitors of the reaction, with 150 values of greater than 5 mM. From this observation, we can conclude that the presence of hydroxyl and/or carboxyl groups as functional groups has a role in the inhibitory action. The observation for this apparent requirement is as yet unclear.

These eacly studies on the ATA inhibition led us to a series of studies on the effect of this compound on various encymic reactions. The following chapter describes one of these studies involving the effect of ATA on the phenylalanyl-tRNA synthetase reaction.

CHAPTER 4

THE EFFECT OF ATA AND AURIN DERIVATIVES ON PHENYLALANYL-ERNA SYNTHETASE REACTION

(4-1) Introduction

The studies described in Chapter 3 led us to speculate that there must be some ATA sensitive reaction other than the mRNA: ribosome interaction in the overall protein synthesizing process. This prediction is based on the observation that ATA is more effective in inhibiting the overall incorporation of phenylalanine that in the more specific mRNA: ribosome binding reaction. Thus the search for at least one more ATA sensitive reaction was conducted. In the course of the incorporation reaction, we always set aside an aliquot of the reaction mixture for detection of aminoacyl-tRNA synthesis to ensure the formation of auffletent presence of ATA, aminoacyl-tRNA synthesis is indicated that, in the presence of ATA, aminoacyl-tRNA synthesis is markedly reduced. The inhibitory effect of ATA on the aminoacyl-tRNA synthesis. "A an of ATA inhibition of the phenylalanyl-tRNA synthesis."

At the time this study commenced, the nature of aminoacylation reactions was well established for both bacterial and mammalian systems. It is a two-step reaction involving the formation of aminoacyl-AMP and the subsequent transfer of the aminoacyl molety to a specific receptor tRNA. This two-step reaction is catalyzed by a single enzyme specific to each amino acid (Novelli 1967). However, the structure of the enzyme from mammalian origins was not fully elucidated at that time.

The overall molecular weights of aminoacyl-tRNA synthetase vary extensively (Lanks et al 1971, Fasiolo et al 1970, and Weinstein et al 1971). Moreover, the subunit structure of the enzymes varies considerably to give rise to either $\alpha 2$ or $\alpha 2\beta 2$ forms (Le Meur et al 1972, Penneys et al 1974, Fasiolo et al 1970, and Tacherene et al 1973). The present investigation on the action of ATA on the phenylalanyl-tRNA synthetase reaction included: the determination of the inhibitory concentration of ATA, the determination of the structural features responsible for the observed inhibition, and the stoichiometry of the ATA; enzyme interaction. Although the lack of precise information regarding the molecular weight and subunit structure of phenylalanyl-tRNA synthetase introduced some ambiguity into these analyses, the studies nonetheless lad us to complude that ATA is a non-specific inhibitor which interacts not only with ribosomes but also with aminoacyl-tRNA synthetase.

(4-2) RAAULCA

REFAUL OF ALLA OR the AMINGAGYI-TRNA Synthetase Reaction

In order to study the effect of ATA on the synthetase reaction, At WAS necessary to establish the optimal conditions for this reaction. In particular, the effect of salt concentration required special attention because the grude reticulocyte synthetase fraction was precipitated by summonium sulfate at 66% saturation and suspended in an equal volume of Buffer A. Thus this fraction contained a significant smount of monovalent cations necessitating an adjustment in the standard assay, depending on the volume of enzyme solution added. Fig. 8 shows that the optimal concentration of KCl is 30 mM and of MgCl₂ is 12 mM when the ATP concentration is 1 mM. The addition of p-mercaptoethanol at 1 mM is also essential to the reaction, Based on this data, the synthetase



ATP (0.1)umoles), and AS65 enzyme were held constant. H-phenylalanyl-tRNA synthesized at 20 min of Teaction at 37°C was plotted on the ordinate.

reactions in the following experiments were carried out as specified in the Methods $(2-11)^{2}$.

The effect of various concentrations of ATA on the phenylalanyl-tRNA synthetase reaction was then studied using the tabbit reticulocyte synthetase fraction AS66. The complete reaction mixture (see Methods) included various concentrations of ATA and was incubated at 37° C for 20 min. The samples were processed with the cold TCA wash as described in the Methods. The level of H-Phe-tRNA synthesized after 20 min reaction in the presence of ATA was compared with that in the control set; which contained no ATA, and expressed as percent activity. Fig. 9 demonstrates that ATA inhibits the Phe-tRNA synthetase reaction, reducing it by half at a concentration of 50 μ M ($1_{50} - 50 \mu$ M).

Effor of Aurin Derivatives on the Phe-thna Synthetase Reaction

In order to investigate the nature of the ATA inhibition of the phenylalanyl-tRNA synthetase reaction, an inhibitor study was conducted using autim derivatives. The 1_{50} level for each of the triphenylmethaues tested was established using the method previously described for ATA. The structures of these compounds are given in Fig. 1. Table 1 shows the 1_{50} values of the autim derivatives tested for the phenylalanyl-tRNA synthetase reaction. It is clear that all of the compounds tested are potent inhibitors of the aminoacyl-tRNA synthetase reaction unlike that observed for the mRNA:ribosome binding reaction described in Chapter 3. It must be pointed out that the charge properties of the side chains are not critical in this system since both Aurin (negatively charged) and Fuchsin Basic (positively charged) exhibit inhibitory action at approximately the same concentration-250 μ M. It is also noticeable that compounds possessing carboxyl groups are \sim



F18. 9 Inhibition of phenylalanyl-tRNA synthetase by ATA. Assays were performed as given in the Methods. The incorporation of ³H-phenylalanine into ³H-phenylalanyl-tRNA at 20 min incubation was taken as 100% activity. The activity in the presence of varying amounts of ATA is indicated as the percent of activity relative to the control value. (100% = 4.2 pmoles Phe). The protein concentration was 100 μ g per 0.1 ml reaction mixture.

Т	дb	le	1

Aurin Derivative	1 ₅₀ Value
Αμτια	250 µM
Fuchain Acid	250 juM
Puchain Baale	' 250 µM
PARONE BLUO	150 µM
Asuro Blug B	μ. 50 μ.Μ.
Aurimericarboxylic acid	50 jum

Inhibition of PhestRNA Synthesis by Aurin Derivatives

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a ghtly stronger inhibitors than those possessing sulfonyl groups. A reduction in the number of conjugated resonance groups from three (ATA) to two (Azure Blue B) does not alter the inhibitory effect at all. These observations suggested to us that neither the skeletal structure of triphenylmethanes nor the side chains are important for their inhibitory action. This information then suggested to us that the aurin derivatives may inhibit this reaction through a different mechanism from that observed in protein synthesis. It is possible that the phenyl group of the aurin derivatives itself may compete with phenylalanine towards its binding site, However, this possibility is remote since all of the sminoacyl-tRNA synthetase reactions tested were inhibited by ATA. A more detailed analysis of the mode of action will be dealt with in a later section.

Site of ATA Agrion on the Phe-tRNA Synthetase Reagtion

In order to investigate the possibility that ATA exerts its inhibitory agtion by interacting with one of the reaction components, a preincubation study was undertaken. When ATA at a concentration of 20 µM was preincubated 5 min at 4°C with yeast tRNA, the subsequent assay for PhentRNA synthesis at 37°C showed 887 activity as illustrated in Table 2. When ATA was preincubated with both the enzyme and tRNA, 907 activity was shown. When ATA was preincubated with the enzyme alone, 767 activity was detected. These results show that preincubation of ATA with the enzyme fraction produces a slightly higher inhibition. Nowever, they do not clearly distinguish whether the site of inhibition is tRNA or the enzyme.

In order to provide more definitive data, a concentration



Table 2



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effect was examined, where the concentration of one reaction component was altered while the concentration of the other components and ATA (50 μ M) remained constant. If changing the concentration of a substance alters the level of inhibition produced by a given concentration of the inhibitor, then it is an indication that that substance is the site of the inhibitory action. The four reaction components of the synthetase reaction were examined—tRNA, ATP, Mg⁺⁺, and synthetase. Fig. 10 shows " the effect of varying the tRNA concentration at a constant level of ATA. The curve shows essentially the same percent activity whether 30 or 300 µg of tRNA are added to 0.1 ml of reaction mixture. From these reaults, we conclude that the concentration of tRNA does not have a pronounced effect on the inhibition equaed by ATA. Similarly, changes In the ATP and Mg concentrations did not alter the level of inhibition achieved by ATA at a concentration of 50 µM. However, in the case of the experiment in which synthetape concentration was varied, the results yore much different from those described above. Fig. 11 demonstrates that, by ralaing the concentration of synthetase above 100 µg per reaction mixture in the presence of 50 µM ATA, inhibition levels of leas than 50% are observed. In fact, ralaing the concentration of the anayma to 120 he or higher ber reading mixture offeringly reacores the level of ³H-PhentRNA formation to 100% in the presence of 50 µM ATA. Conversely, lowering the concentration of synthetase to levels balow 100 mg produces levels of inhibition greater than 50%. Of the reaction components casted, only synthetake showed such a concentra-١ sion affect. These reaults clearly show that increasing the enzyme concentration serves to overepme the ATA inhibition, Therefore, we Arow a concative gongluaton shat ATA interacts in some way with



F18. 10 Effect of tRNA concentration on inhibition of PhertRNA formation by ATA, Assays were performed as described in the Methods, except that tRNA was added to start the reaction. PhertRNA synthesis at 20 min of reaction in the absence of ATA was taken as 100% solvity. The level of activity for each tRNA concentration in the presence of μ M ATA was measured, and relative activity to the control value (for that tRNA concentration) was expressed as percent activity.



FIE: 11 Effect of the concentration of PheneRNA synthetage on the inhibition of PheneRNA formation by ATA. Assays were performed as dear eribed in the Methoda, except that tRNA was added last to start the. reaction.¹ The ³H-PheneRNA synthesis at 20 min of reaction in the absence of ATA for each ensyme concentration was taken as 100% activity. The level of activity for each ensyme concentration in the proseduce of 50 µM ATA was measured and the percent activity calculated as in Fig. 10.

phenylalanyl-tRNA synthetase.

Binding of ATA to PheriRNA Synthetase

If the observed inhibitory action of ATA on the Phe-tRNA synthetase reaction is truly caused by the interaction of ATA with the enzyme, a stable complex formation between them may be detected as was done in the mRNA; ribosome binding reaction. Initially, an attempt) was made to separate the enzyme; ATA complex by a molecular serving method as was done in Chapter 3 for the detection of a ribosome; ATA complex. However, a problem was encountered due to the fact that the AS66 enzyme contained a large proportion of hemoglobin which possesses an absorption spectrum overlapping with that of ATA. In addition, the molecular weight of hemoglobin is in a similar range to that of the synchetase—as will be described later. These problems prevented adequate measurement of the ATA; synthetase complex. Therefore, purification of Phe-tRNA synthetase was required. After achieving the removal of hemoglobin as described in the Methods, the following experiments were performed.

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First of All, the 1_{50} value had to be re-natabilished for the partially purified emaynes at a level of 100 µg per reaction mixture and 7 in the presence of various concentrations of ATA. Assays were performed, as given in the Mathods. The level of ³M-Rhe-tRNA synthesis in the presence of various concentrations of ATA was compared. From Fig. 12, the 1_{50} value for the pertially purified enzyme was estimated to be 13.5 µM. This value represents a shift in 1_{50} from that for the erude Phy-tRNA synthetase (AS66) fraution. The should be mentioned that



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Fig. 12 inhibition of partially purified Phe-tRNA synthetase by AlA. Assays were performed as given in the Methods. The incorporation of ³H-Phe into ³H-Phe-tRNA at 20 min of reaction in the abgence of dye was taken as 100% activity (30 pmoles Phe). Varying amounts of ATA were adds ad, the activity measured, and the results were expressed as a percent of the control activity. The protein concentration was 20 μ g per 0.1 ml of reaction mixture.

the ratio of these two I_{50} values (approximately 4) is very close to the ratio of protein concentrations used in these two sets of experiments: 100 μ g/20 μ g = 5. It thus appears that the reduction in the I value of the partially purified enzyme is due to the reduction in protein concentration. This is what one would expect if ATA has affinity toward many proteins to about the same extent.

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In order to detect the formation of a stable complex between ATA and the Phe-tRNA synthetase, a mixture of ATA and the enzyme was chromatographed on a Sepharose 68 column. The elution profile of the column was determined by an assay of each fraction for synthetase acttivity. The peak position of the enzymatic activity coincided with that for a protein fraction as monitored by 280 µm absorption. This indicated that the enzyme was reasonably pure in terms of charge properties and of molecular weight. The position of ATA was then determined by monitoring absorption at 308 pm, which provides a minimum overlap with the absorption apentum of protein and RNA.

To determine whether or not ANA binds to synthetase, a solution of 500 µg emayme in Buffer RM was made to 40 µM in ATA. Fifty microliters of this mixture was loaded onto a Sepharose 68 column of bed volume 5 ml which was pre-equilibrated with Buffer RM. Five drop fractions of volume 0.175 ml were collegted and diluted to 0.5 ml with water, Each fraction was examined for absorption at 280 nm and 308 nm using a spectrophotometer. These results, illustrated in Fig. 13, show that there is significant absorptance at 308 nm at the position which

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Fig. 13 Sepherone 68 analysis of partially purified Phe-tRNA synthetase: ATA mixture. A solution containing 500 μ g enzyme and 40 μ M in ATA in 50 μ l was layered onto a Sepherone 68 column of bed volume 5 ml (5 mm x 130 mm). Materials were eluted from the column using Buffer RM. Five drop fractions (0.175 ml) were collected and diluted with water to a volume of 0.5 ml. Thereafter, absorption at 280 pm for the enzyme and at 308 μ m for ATA was determined for each fraction. Absorbance at 280 μ m (\blacksquare) Absorbance at 308 μ m (\blacktriangle).

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corresponds to the enzyme peak. This indicates that ATA binds to the enzyme to form a stable complex detectable by column chromatography.

A similar experiment was then performed using a mixture of yeast tRNA and ATA. For this study, 50 μ g of tRNA was made to 40 μ M ATA in Buffer RM. The procedure used was as described above except that the diluted samples were examined at 260 nm for tRNA and at 308 nm for ATA. As shown in Fig. 14, no absorbance at 308 nm occurs in the tRNA region. From this observation, we conclude that tRNA does not bind to form a stable complex with ATA.

Semiguantitation of ATA: Synthesaae Interaction

As the first step in determining how many ATA molecules can blud per enzyme molecule, it was necessary to determine the molecular weight of the PheetRNA synthetase. This enzyme apparently migrated as a single peak on Sepharose 6B as detected by assay of the fractions for PheetRNA synthetase activity. Fig. 15 shows the relative elucion pattern of marker proteins of known molecular weight and the enzyme, The molecular weights of the marker proteins are as follows: Glucose oxidase, 186,000; Chymotrypsinogen A, 25,000; Hemoglobin, 68,000; and Lysozyme, 14,000. Fig. 16 shows the replot of the molecular weight and the elucion positions of the individual proteins from the column, From this data, the position of the PheetRNA synthetase activity corresponds to a molecular weight of 70,000 daltons.

The next step was to calculate the number of ATA molecules bound per molecule of Phe-tRNA synthetase. Tube #24 from Fig. 13, containing the enzyme ATA, was used for this purpose. The smount of enzyme

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<u>F18.14</u> Sepherose 6B analysis of tRNA-ATA mixture. Fifty microliters of a tRNA: ATA mixture (50 μ g yeast tRNA, 40 μ M ATA) was layered onto a Sepherose 6B column of bed volume 5 ml (5 mm x 130 mm). Materials were eluted from the column using Buffer RM and five drop fractions were collected (volume 0.175 ml). Samples were diluted to 0.5 ml and examined for absorption at 260 nm for tRNA and at 308 nm for ATA. Absorbance at 260 nm (**m**) Absorbance at 308 nm (**A**).

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F18. 16 Molecular weight eatimation of partially purified Phe-tRNA aynthetase on a Sepharose β B column. The relative positions of marker proteins and the enzyme, as determined in Fig. 15, are plotted against the molecular weight in the logarithmic scale. The molecular Awight of, each marker protein used is inserted in the figure: 1. Glucose oxidase, 186,000; 2. Fhe-tRNA synthetase; 3. Chymotrypsinogen A, 25,000; and 4. Lysozyme,14,000. The peak positions were determined by 280 pm absorbance except in the case of the synthetase where the peak position was determined by assay of the fractions for synthetase activity. This data says rise to a molecular weight of 70,000 daltons for Phe-tRNA synthetase.

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extinction coefficient of $1 A_{280}/mg/ml/10$ mm light path together with the estimated molecular weight of the enzyme as 70,000. The amount of ATA present in the tube was determined from its absorbance at 308 nm using its extinction coefficient of 15.2 $A_{308}/mg/ml/10$ mm light path. The result of these calculations indicated that for each enzyme molecule, there are 6 ATA molecules present. Since the extinction coefficient of Phe-tRNA synthetase had not been established, the above value may be slightly different from the true value. However, extinction coefficients of various natural proteins deviate between 0.4 to 1.2 (Wetlaufer 1962). Therefore, the number of ATA molecules bound per molecule of the enzyme would range from 2.5 to 7 molecules.

The Mode of Inhibition in the Phe-tRNA Synthetame Reaction

It is valuable to determine the effect of ATA: enzyme complex formation of the binding of other ligands involved in the reaction. Based on the finding that Phe-tRNA synthetase can be retained on Millipore filters under appropriate conditions (see Methods), the effect of ATA on tRNA binding was studied using radioactively labelled yeast tRNA. A mixture of the partially purified enzyme and ³M-tRNA was tested for binding to Millipore filters in the presence and absence of ATA (30 μ M). The amount of ³M-tRNA bound to the enzyme in the presence of ATA was reduced to 20% rejetive to that in the absence of ATA. The results shown in Table 3 indicate that ATA does interfere with the binding of tRNA to Phe-tRNA synthetase. However, the result does not distinguish the possible mechanism of inhibition due to either direct

Reaction mixture	ty ,	CPM Retained
³ H-tRNA alone		394
³ H-LRNA † enzyme		12,854
H-LKNA + PRAYME + ATA (30 µM)		3,978

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Binding of ³H-tRNA to Phenylalanyl-tRNA Synthetase

effect. An attempt was made to study the effect of ATA on ATP and phenylalanthe binding to the enzyme under the same conditions as described above. However, there was no detectable complex formation between these ligands and the enzyme on Millipore filters. Their dissociation constants must be much lower than that of tRNA.

Because of the limited information available from the above experiment, we analyzed some already available data somewhat differently to probe the mode of ATA inhibition. Namely, the data given in Fig. 10 , concerning the effect of substrate concentration on ATA inhibition, was submitted to a kineweaver-Burke plot (plot of the reciprocal of the initial velocity vs. the reciprocal of the concentration of a substrate when the concentrations of the other substrates are fixed). Fig. 1/ clearly indicates that ATA is a non-competentive inhibitor for tRNA. This analysis led us to conclude that ATA binds to the synthetase at a alte other than the tRNA binding site, .

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The results reported in this chapter indicate that ATA inhibited Phe-LENA synthesis at a concentration of 50 μ M. This reaction is not more sensitive than the mENA:ribosome binding respired (150 ~ 20 μ M). Nevertheless, inhibition of Phe-tENA synthesis by ATA has not been previously reported, and the study of ATA setion in this system could provide some information about the nature of its action.

In order to probe the mechaniam of ATA inhibition in the Phen the Annihetane reaction, an experimental approach similar to that used in (Mapter 3 for the menarchoad pluding reaction washemployed. First of All, the structural requirement for the described inhibition by ATA



<u>Fig. 17</u> Linewayar-Burka plot of the phenylalanylatting synthetian readtion from Fig. 10: the realprocal of the initial velocity against the realprocal of tRNA concentrations in the presence and abagnes of a fixed level of ANA. (\uparrow) no ANA, (\blacklozenge) 20 µM ANA.

was sought by comparing the relative inhibition by various aurin derivatives. Then the site of ATA inhibition in the synthetase reaction was studied in detail.

A comparative study with aurin derivatives indicated that all of the compounds tested were potent inhibitors of the reaction. This finding is quite different from what was observed in the mRNA: ribosome binding reaction as reported in Chapter 3 and suggests that the mode of ATA action may be different in the two reaction systems. The fact that Aurin (megatively charged) and Fuchain Baate (positively charged) inhibit the PhartRNA synthetage reaction at the same concentration of 250 µM, clearly indicates that the charge of the molecule is not important for this inhibitory action. However, the type of aubacktuting groups present on the triphenylmethane arelaton appears to attent the level of their inhibition. Those compounds possessing hydroxyl BROUDD ARD DEFOREIVO INDIDIEDEN. IN Addielon, the NARDONYL BROMP ONhamong the inhibitory action. For example, Ametic, which has three hydroxyl and no varboxyl brough had an 150 - 150 pM, while, ATA, which has three hydroxyl and three narboxyl broups has an 150 5 50 µM. Thus the breadnes of both hydroxyl and carboxyl broups blyon-rise to the ALTONBOAK ANDILLORY ANTION ON THIS ANALYNG AVALAM.

The study of the mechanism of ATA sector in the Phe-tRNA synthesis reaction included; the addition of reaction components in different order, relief of ATA inhibition by increasing the concentration of reaction components, and detection of a stable complex formation between ATA and individual reaction components, by changing the order of adding the reaction components, it was hoped to probe the site of ATA interaction with a specific reaction component, The results showed o 60 ഔ

that when ATA and tRNA were mixed prior to addition of the enzyme, the inhibition was lower than when ATA and the enzyme were mixed prior to the addition of tRNA. This indicates that the inhibitory action of ATA is more effective before the formation of the synthetase:tRNA complex. Bluementhal et al (1973) reported a similar finding from their studies uaing E. coll RNA polymerase. In the second set of experiments, the concentration effect of the individual reaction components on the level of inhibition of a fixed concentration of ATA was examined. If the observed inhibition was due to the interaction of ATA with a certain reaction component, then rateling its concentration should serve to overf come the lightblition. Of the four reaction components examined in this experiment (Mg , ATP, tRNA, and the synthetase), only alteration of the anyme concentration had an effect on the lovel of inhibition by ATA. Reducing the enzyme concentration from 100 µs to 50 µs reputted in an Increase in Inhibition from 50% to 90%, while increasing the engyme CONSIGNTEALLON to 150 MB REPAILED IN REPAILEDILY NO INHIBITION. THERES FORD WE COMPLEADE CHAR CHE ALLE OF ANA MARGEAUCION IN CHE PMAYME. The reaults from the third set of experimence confirmed this gonelusion. HARA, MAXENEAN AL PARTIALLY PARTLAN ANAYMO AND AND AND AN AN HOLL AN OC ERNA and ATA ware proveded by a gal exclusion abromatography method to LOAS TON BLADIO DOMPLAN FORMATION DELYOON ATA AND REASTION DOMPONENTS. The reaults showed that stable complex formation could be demonstrated between ATA and the synthetase but not between ATA and SRNA. At the 1. time these studies were conducted, there was no direct evidence for Acable complex formation between energing and ATA to result in the observo ad inhibition, although there were some reports that AIA would bind to proseine (Lindenbaum and Schubers 1956).

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We went further to study two basic questions: (1) what is the nature of ATA binding to the synthetase? and (2) how many ATA molecules can bind per enzyme molecule? These questions are the basis for the next step in the study of the mechanism of ATA action on the phenylalanyltRNA synthetase reaction.

In order to probe further into the nature of ATA binding to aynthetase, an experiment was carried out to determine whether or not the bound ATA would interfere with the binding of various ligands to the ensyme. Binding studies using Millipore filtration and radioactively labelled precursors demonstrated that the presence of ATA at a concentration of 30 µM reduced the binding of tRNA to the enzyme to 30%. Binding of ATA and phenylalaning to the enzyme could not be demonstrated under these conditions and so the effect on these ligands penales for further when y

With the information that ATA binds to phenylalanyl-tRNA synthetase so as to interfere with the binding of other ligands, an attempt was made to obtain the stolchiometry of the ATA-ensyme interaction. For this purpose, a molecular weight determination of the partially purified synthetase had to be done. The ensyme migrated as a single peak on Sepharose of chromatography as detected by synthetase activity. Thus, knowing the relative position of marker proteins of known molecular weights, the molecular weight of the enzyme could be determined. The results show that the molecular weight of the synthetase is 70,000 daltons. The molecular weights reported for purified phenylalanyl-tRNA synthetase by other laboratories range from 150,000 to 280,000 daltons with a four subunit atructure ($\alpha_2\beta_2$), Fasiolo (1973) reported a molecular weight of 220,000

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with subunits of 56,000 and 63,000 for yeast phenylalanyl-tRNA synthetase while Weinstein et al (1973) reported a molecular weight of 280,000 with subunits of 69,000 and 74,000 for the same enzyme in rat liver. Purification of this enzyme in our laboratory subsequent to this thesis work established a molecular weight of 275,000 with subunits of 58,000 and 75,000 (Igaraphi 1974). From these findings, it would appear that the above figure of 70,000 daltons for the synthetase is in the range of aubunits rather than the intact enzyme and yet its position was established by assaying the fractions cluted from the Sepharose 6B column. The possible explanation for this discrepancy is as follows; if the conditions of the partial purification of the enzyme were such that the synthetase was dissociated into its subunits, then a mixture of the two types of aubunits must have been subjected to get exclusion chromatography. The Sepherone 60 column used in these experiments is not capable of clearly apparatilis the mixture of proteins differing by less than 50,000 daltons the molecular weight, although it allows determination of peak positions of PURE PROTECTION PROCESSED ASPARACELY, The REALL of the chrometography of the mixed subunits of molecular veloce 75,000 and 58,000 would result in only a partial apparation of the aubunits with compiderable overlapping at sparagion aquivalant to 70,000 dalcanarcha malqaular waisht datarminad for the synchetase. Thus, the tube containing this overlapped region contains both aubunits, so that reconstitution of an astive form of the anzyme can Transcur, in this particular fraction. Leaving the detailed acudy of anyme acrueture to a further study, the number of ATA molecules bound per syme shecane molegule was eastmaned assuming the molegular weight of the ensyme is

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70,000 daltons. For this purpose, the fraction from the Sepharose 6B column chromatography containing the ATA:enzyme complex was used. The ratio of ATA to the enzyme in this complex was determined by their extinction coefficients at 308 nm and 280 nm respectively. The results show that there are 6 ATA molecules bound per enzyme of 70,000 daltons. This estimated value will be modified somewhat due to lack of critical information regarding the extinction coefficients used here were for a typical protein solution ($1 A_{280}/mg/ml/10$ mm light path) and for a free solution of ATA. Though the value for the number of ATA molecules bound per synthetase is not likely the correct value, it does indicate that there is a limited number of sites on the enzyme where ATA can bind.

The results presented above remain somewhat qualitative in nature. However, the finding that sminoacyl-tRNA synthetase is sensitive to ATA was a movel one, and led to further detailed studies of the kinetic features of ATA action in this laboratory. In this respect, it is noteworthy that the kinetic analysis shows that ATA is a non-competative inhibitor at least for tRNA. This is a new finding and different from our original idea that the phenyl group with a suitable side chain may compete with the phenylalamine binding site, Obviously, the enzyme conformation changes upon binding with ATA and the binding sites for some of the ligands become inscensible. Further studies on the conformational changes of the enzyme is anticipated.

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