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論文目録 上井健史 博士論文題名 ホルミルメチオニンtRNA変換体の合成と活性 1) Comparison of substrate base sequences for RNAligase reactions in the synthesis of a tetradecanucleotide corresponding to bases 21-34 of E. coli tRNA\_ E. Ohtsuka, T. Doi, H. Uemura, Y. Taniyama & M. Ikehara (1980) Nucleic Acids Res. 8, 3909 (E. coli tRNAf<sup>ter</sup>の 21番目から 34番目の塩基に対応的 14merの合成における) (RNA Tigase 反応の基質塩基配列の比較 2) Modification of the anticodon triplet of E. coli tRNA<sub>f</sub> by replacement with trimers complementary tRNA f to non-sense codons UAG and UAA. E. Ohtsuka, T. Doi, R. Fukumoto, J. Matsugi & M. Ikehara (1983) Nucleic Acids Res. 11, 3863 (E. wili tRNAtt のアンチコトントリア・レットを終止コトン UAG, UAAに置換する くことによる修飾 3) Replacement and insertion of nucleotides at the anticodon loop of E. coli tRNA<sub>f</sub> by ligation of chemically synthesized ribooligonucleotides. T. Doi, A. Yamane, J. Matsugi, E. Ohtsuka & M. Ikehara (1985) Nucleic Acids Res. <u>13</u>, 3685 (化学合成りボオリコッタクレオチト"の Tigation による E. coli tRNAfetの アンチコトン) ループの置換と塩基の捜入 Modification of the amino acid acceptor stem of E. coli tRNA<sup>f</sup> by ligation of chemically synthe-sized ribooligonucleotides. T. Doi, H. Morioka, J. Matsugi, E. Ohtsuka & M. Ikehara (1985) FEBS Letters 190, 125 (化学合成川ボオリコッタノレオチト"の ligation に F3 E. coli tRNAtton PE)酸复客) しステムの修飾

5) The in vivo stability, maturation and aminoacylation of anticodon-substituted Escherichia coli initiator methionine tRNAs H. Grosjean, S. DeHenau, T. Doi, A. Yamane, E. Ohtsuka, M. Ikehara, N. Beauchemin, K. Nicoghosian & R. Cedergren (1987) Eur. J. Biochem. <u>166</u>, 325 アンテコドンを置換した E. coli tRNAft min vivo における安定性を成熟化) し及びアミノアシル化 参考論文 1) A new method for 3'-labelling of polyribonucleotides by phosphorylation with RNA ligase and its application to the 3'-modification for joining reactions. E. Ohtsuka, H. Uemura, T. Doi, T. Miyake, S. Nishikawa & M. Ikehara (1979) Nucleic Acids Res 6, 443 (RNA ligaseを用いたり、酸化にあるボーリリボマクレオチトの3標識の新法と) しその結合反応に対する 3、修飾の応用 2) Total synthesis of a RNA molecule with sequence identical to that of Escherichia coli formylmethionine tRNA. E. Ohtsuka, S. Tanaka, T. Tanaka, T. Miyake, A. F Markham, E. Nakagawa, T. Wakabayashi, Y. Taniyama, S. Nishikawa, R. Fukumoto, H. Uemura, T. Doi, T. Tokunaga & M. Ikehara (1981) Proc. Natl. Acad. Sci. U.S.A. 78, 5493 (E. coli tRNAft on 墙基配列を持っ RNA分子a 全合成') 3) Chemical synthesis of the 5'-half molecule of E. coli tRNA<sup>GLY</sup>. E. Ohtsuka, A. Yamane, T. Doi & M. Ikehara (1984) Tetrahedron 40, 47 (E. coli tRNAZ a 5'半分子 a 化学合成)

# 論文内容の要旨

博士論文題名

ホルミルメテオニンtRNA変換体の合成と活性

学位申請者

土井健史

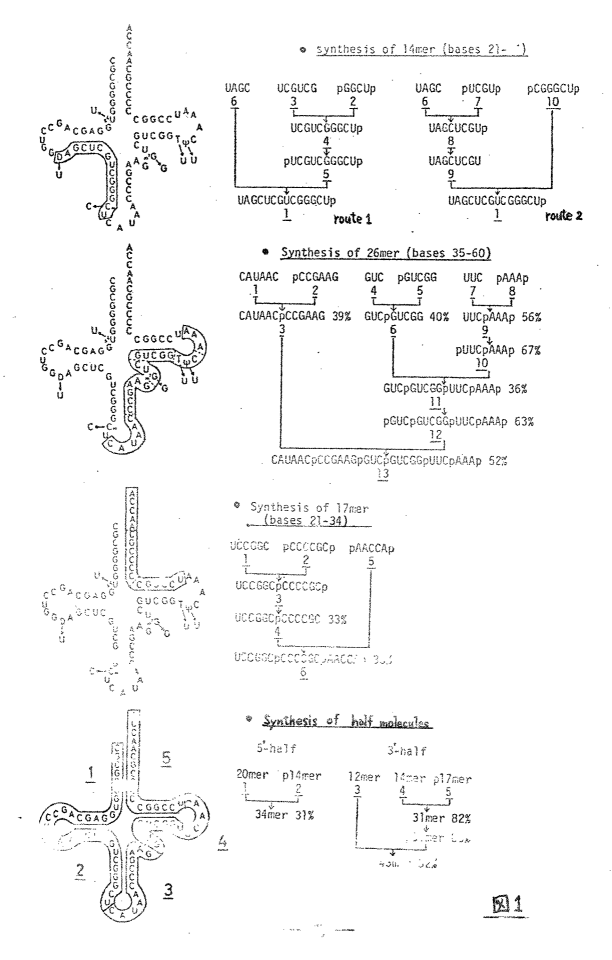
tRNAは鎖長70~80の機能を存するRNAとしては最小の分子である。

著者はE. white tRNAft nascent strand 分子の 3'側半分子, 5'側半分子を化学合成フラグメ ントより RNA ligane"を用いて合成した。 こ の際 これらの RNA ligane 反応における 塩基 綺異性を見い出した。

次にRNA Ligane 反応を用い E. coli tRNA<sup>thet</sup> のアミノアシルステム、TYループ、Dルー プ、アンチコドンループを変換して tRNAを 合成し 主にメテオニル tRNA 合成酵素との 相互作用について調べた。

I RNA ligaseを用いた長鎖オリゴリボ 又クレオチドの合成

図1に示す様に比較的短鎖の化学合成オリ ゴリボヌフレオチドより tRNA 後鎖長分子 を合成した。2)3) さらに これらを用い tRNA 半 分子を合成した。4)



Reaction	conditions	for	joining
----------	------------	-----	---------

acceptor nmol [µM]	donor nnol [µM]	атр µМ	RNA ligase [unit/ml]	time hr	Reaction extent <sup>a</sup> )	Yield %
UCGUCG ( <u>3</u> ) 12[100]	pGGCUp (2) 20[167]	333	250	1	15	4
UAGC(6) 1.2[240]	pUCGUCGGGGCUp( <u>5</u> ) 0.4(80)	200	180	2	27	11 <sup>b)</sup>
UAGC (6) 30 [150]	pUCGUp(7) 20[100]	200	100	3.9	5 88 '	71
UAGCUCGU( <u>9</u> ) 7[100]	pCGGGCUp( <u>10</u> ) 8[114]	200	214	2	52	31 <sup>b)</sup>

Incubation mixtures contained 10% DMSO and the temperature was 25°.

a) Estimated by homochromatography.

汞1

b) The compound was isolated after kination.

RNA ligane 反応において表1に示す様に、 acceptor 分子では 3末端が ビリミジン - ブリ ン配列を有するものが、donor 分子では自己 相補的な構造を有するものが それぞれ反応 し難いことがわかった。30

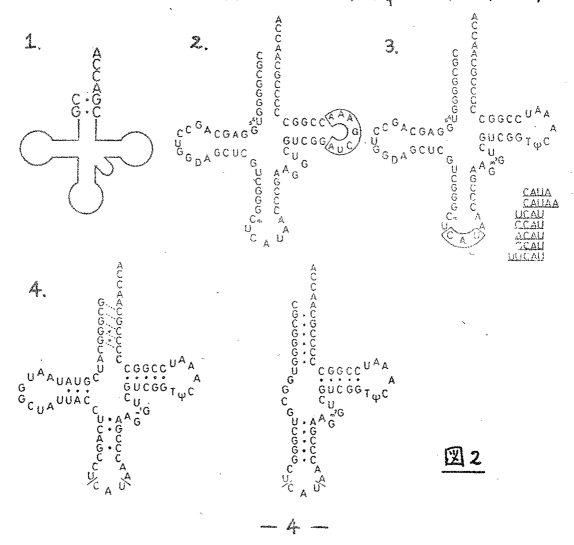
I E. colitRNAT 変換体の合成とその性質

1) E. coli メデオニル +RNA 合成酵素との 相互作用

E coli メチオニル tRNA 合成酵素との相互作用を耐べなために以下のtRNAt 交換体

至合成した。(図2)

- 1、アミノアシルステムの5末端で G-C 塩 基対を形成する tRNAft
- 2. T 4 ループのT 4 C を真核生物のイニシ エーター tRNA に特有の A U C に変換した tRNA<sup>thet 5)</sup>
- 3. アンテコドンの5個隣接位の共通塩基U を他の塩基に変換したtRNAtやアンテコド



ンルーフ<sup>o</sup>を拡大した tRNA<sup>Het</sup>を合成した。<sup>6,0</sup> メテオニル tRNA 合成酵素の アンチュドン 認識については アンチュドン配列 CAU が 重要であることが調べられていた<sup>80</sup>ので、この 配列を残しループを拡大した。

4. Dル-フッの欠除した tRNAft や5個半分子 にE. whi tRNAft の 塩基配列を有する tRNA を合成した。?

合成は天然のtRNAftフラグメントとに学合成オリゴマーとを組みあわせ、これらをRNAliganeで結合し行体、た。

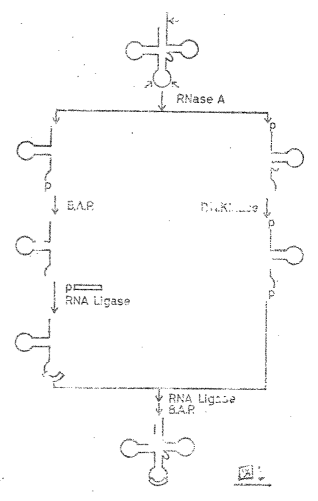
アミノアシルステムでの変換には天然の  $tRNA_{f}^{het}$ の muclease S1 限定分解フラグメント<sup>(0)</sup>を, その他のループの変換には  $tRNA_{f}^{het}$ の RNase A 限定分解フラグメント<sup>30,10</sup> を用い合成した。又,<math>RNA の ジ末端より - 5 展 を除去する方法として週ヨウ素酸酸化を用いた。一例として アンチュドンを変換した時の合成 scheme を図3に示す。

- 5 -

生成物「ホリア クリルアミドゲル 電気泳動で単離し T-。 RNA Ligue 反応 後 IT nearest neighbon analypsis で結合部 位を確認して。

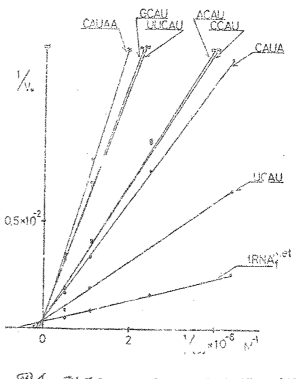
合成した tRNA についてアミノ酸 受容活性を測定し た。

アミノアシルステムを変換した



tRNM はもとのもRNA と下に活性を示したが、 TYループ、Dループを変換したtRNA は ほとくど活性を失った。アンチュドンを実換 したtRNA については、アンチュドンの「側 しを変換したtRNA はしの分子と変わりなか ったが、ループを拡大したtRNA については その付加塩茎と位置により親和仏に差がみら NF。 図4にその 時の Lineweaver-Bunk プロットを示 す。

アンチコドンの 3個と5個で塩基 のこか具合が異な るために5個に しを何加した。RNA がメテオニルセRNA 合成酵素と強い親 礼性を示し、5日



<u>図4</u>テトラマー、ハンタマーはアンテコトン CAU をそれったい温碁配列にき換した もRNAできます。

への、中加や2塩基付かでは弱くなると考えら れる。又、アンテコドンループを拡大した tRNA についてリボソーム上でどのmRNA (トリマー)と結合するかを調べたが、5個にU を付加した時のかもとの、AUG と結合できる ことがわかった。このことよりアンチコドン の5個にCAU の並びをあまり変化させずに Uが入りこむことが可能と考えられる。

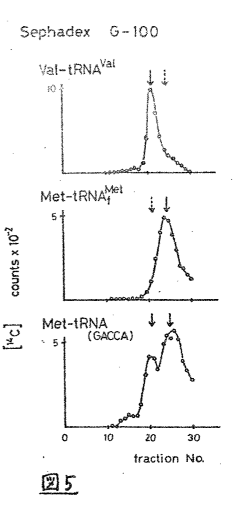
2) EF-Tu との相互 任用

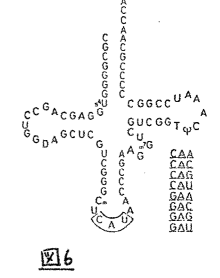
アミノアシルステムで 塩基対を形成して、tRNAft について EF-Tu との結 合実験を行な, Tc。 図5 に示す様に塩基対を形成 しない天然の E. uki tRNAft は EF-Tu に結合しない のに対し, 合成した tRNA は一部複合体を形成する ことがわかった。

> 3) アンチコドンの 3'側 隣接位の塩基修飾

E.uki tRNAftのアンチコ ドンを図6に示す8種類の 塩基に変換したtRNAftを

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合成した。これらの tRNA を Xenopus laevis oryをの細胞質にマイクロインジェクション しアンチコドンの3個が修飾されるかをかを 調べた。 CAU (もとの tRNAft と同じ) と GAU 配列を有する tRNA のみ Aが t<sup>6</sup>A (N-[9-(β-Dnibofuranosyl)purin-6-yl carbamoryl]threonine) に 修飾を受けることがわかった。又, これら2 種の tRNA のみ Xenopus oocyte 中で安定に存在 できることがわかった。

### 結論

- 1. 化学合成フラグメントと RNA ligane を用 い種々の長鎖 RNAフラグメントを合成した。 又, RNA ligane 反応の基質特異性を見出し た。
- 2. E. coli tRNA<sup>nut</sup>のアミノアシルステム,T 4ループ,アンテコドンループ,Dループ を変換したtRNA<sup>nut</sup>を合成しメチオニン受 落活性を調べた。DループとT4ループを

変換した tRNA は ほとんど活性がなくなり, アミノアシルステムの 5'末端で塩基対を形成して tRNA は本来の tRNAt と同様の活性 を示した。アンチュドンループを拡大した tRNA は CAU配列が存在すれば付加塩基の 種類や位置により活性に差はあるがメテオ ニンを受容した。

- 3. E. whi tRNAft のアミノアシルステムで塩 基対を形成した tRNA は塩基対を形成しな いtRNA に比べ E. whi EF-Tu と複合体を形 成しやすいことがわかった。
- 4. アンテコドン配列のちがいにより tRNA の安定性が異なることを E. coli tRNA<sup>hut</sup>変換 体を用い Xenopus laevis ooryte の系で見出 した。又, アンチコドンの 3'側隣接位のA がこの系で修飾されることを見出した。

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# ホルミルメチオニンtRNA 変換体の合成と活性

.

## 土井健史

略語表

Py Pu D T U s <sup>4</sup> U Cm <sup>m</sup> G t <sup>6</sup> A	pyrimidine purine 5,6-dihydrouridine 5-methyluridine pseuduridine 4-thiouridine 2'-O-methylcytidine 7-methylguanosine N-[9-( $\beta$ -p-ribofuranosyl)purin -6-ylcarbamoyl]- threonine 6-( $\Delta^2$ -isopentenyl)adenosine
P.N.kinase	polynucleotide kinase
BAP	bacterial alkaline phosphatase
ARSase	aminoacyl-tRNA synthetase
Met RSase	methionyl-tRNA synthetase
Gly RSase	glycyl-tRNA synthetase
EF-Tu	elongation factor Tu
BSA	bovine serum albumine
Tris HEPES TEAB DTT $\beta$ -ME EDTA NTA TCA DMSO SDS TEMED X.C. B.P.B. *p PEP	tris (hydroxymethyl) aminomethane N-2-hydroxymethylpiperazine-N'-2-ethanesulfonic acid triethylammonium bicarbonate dithiothreitol \$-mercaptoethanol ethylenediaminetetraacetic acid nitrilotriacetic acid trichloroacetic acid dimethyl sulfoxide sodium dodecylsulfate N,N,N',N'-tetramethylethylenediamine xylene cyanol FF bromophenol blue "p paper electrophoresis
PAGE	polyacrylamide slab gel electrophoresis
PAGE(disk)	polyacrylamide disk gel electrophoresis

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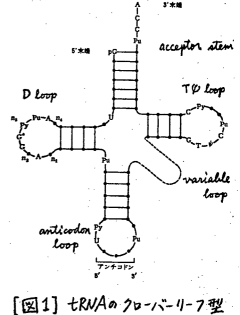
## 緒論

近年のDNA 合成政術の進歩はめまずしく, E.adi tRNAsur の 遺伝子が合成されて以来数多くの遺伝子が合成されている。一 方RNA はその2'水酸基の存在のためDNA に比べ合成が困難 で長鎖オリゴマーの合成には長い時間と労力を要する。Hunwitz 等によりT4 感染大腸菌から発見をれたRNA ligase<sup>2</sup>に一平鎖オ リゴヌクレオチドに作用し、5'末端リン酸基と3'末端水酸基と を結合する酵素で 比較的短鎖のRNA フラグメントを この酵 素を用いてくり返し結合すれば化学合成だけでは得られない長 鎖のRNAの合成が可能となる。

tRNAは鎖長70~80の機能を有するRNAとしては最小のもの で人工合成の標的として非常に興味ある対象である。現在すご 多くの研究者により数目に及ぶ

tRNAの一次構造が明らかにまれ<sup>3)</sup> 一部のtRNAを除き全て共通の フローバーリーフ型二次構造を組 ひと考えられていろ。(図1)又, X線結晶構造解析によりYeastz<sup>~</sup> はtRNA<sup>Phe49,5)</sup>tRNA<sup>Asp 6)</sup>tRNA<sup>fux 7)</sup> E. coli ごにtRNA<sup>fut 8)</sup>の3次構造が 明らかにまれ,それらにいずれも ほぼ天通のL字型構造を頂するこ とがわかった。

生体においてtRNAはmRNA上の情報に従いアミノ酸を運搬しタ



二次構造

- 1 -

ンパク質合成を許すわせるが、特定のセRNA が特定のアミノ酸 を受容し MRNAの情報どわりタンペク噴が合成される機構は興 味深い。セRNA にアミノ酸を受容させるアミノアシルセRNA合 成酵素(ARSase) はこの特異的な認識を行なう酵素で、アミノ酸 の種類により それぞれ独自の酵素が存在する。現在までこの酵 素と セRNA との相互作用について種々の研究ががされているが 今にに認識部位等に関して不明な点が多い。 セRNA の一部を変 腰しに分子を合成し、ARSase との相互作用を調べればセRNA にか けるその部位の役割がより明確になると考えられる。

著者は化学合成オリゴマーと天然のセRNAフラブメントを組みあれて RNA Ligase ご結合し種々の E. coli tRNAt 変換体を合成した。そして それらの分子と主として E. coli メテオニルセRNA 合成酵素(Met RSase)との相互作用を調べた。

以前にSchulmanらによりtRNA分子を化学修飾しMet Risase との相互作用が調べられているい、この方法では目的の箇所以 外でも修飾が生じ特異的な部位の相互作用は説明し難い。

着者はほじめにRNA liganeと化学合成オリゴマーを用い、修飾塩基の欠除した E. whitRNAtt の長鎖フラグメントの合成を試みた。その過程にないて RNA ligane 反応にはかなりの塩基配列 特異性があることがわか、た。

次に化学全滅オリゴマーと天然の E、udi tRNAft より得られ カフラグメントを用い TYル-フ, アンチコドンルーフ, D ルーフ, アクセプターステムなどを変換した tRNA を合成し, Met RSase との相互作用を中心に tRNA 変換体の性質を調べた。 その結果 Met RSase は E. udi tRNAft のアンチコドン部を認識

し、特にそのアンチュドンの CAU という配列が認識に重要であること、又アンチュドン部のみならず tRNA 全体の 3次構造が認識に必要であることがわかった。

さらにアンチコドン部を変換したせRNA についてほ、アフリ カツメガエルの卵母細胞にマイクロインジェクションを行ない この系ごのせRNAの修飾について調べた結果、たむのな系によい ける tRNAの客定性と修飾について今まご知られていなか、た 新しい事実を得ることがでまた。

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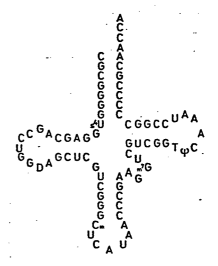
3

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本 論

第一章 化学合成 nascent E. coli tRNAft 737\*×>+の RNA ligase 1= F3 結合反応

+RNAt (図2) は種之のアミノ酸 に対応するtRNAの中でもタンパフ 質合成の開始に関与する特別な分子 ご種之のタンパク質から認識を受け る。このtRNAを人工的に合成する ことは核酸とタンパク質との相互作 用を研究する上で意義があり、又任 意の長鎖RNA が合成可能であるこ との証明となる。

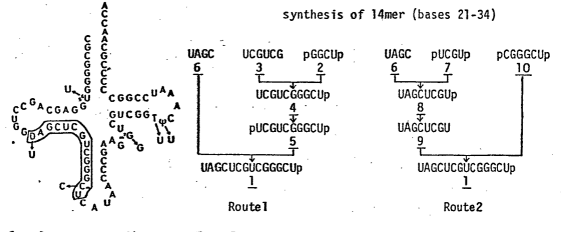


[2] E. whit tRNAH

化学的にRNA鎖を伸展する場合,

鎖長が10を越えると合成がかなり困難となる。 tRNAの様な長 鎖RNAの合成の場合比較的長鎖の化学合成フラグメントを用 いそれらをRNA Ligase ご結合する方法が適当であると考えられ ろ、

着有は図3の様な反応を触媒する RNA ligane を用い比較的短 鎖の化学合成フラグメント を出発原料にし結合を繰り acceptor donor  $ATP \longrightarrow OH + OP \longrightarrow DH + OP \square DH + OP \square DH + OP \longrightarrow DH + OP \square DH + OP \longrightarrow DH + OP \square DH + O$  第一節 Dループからアンチコドン部位に至ら14men (bases 21-34)の合成 11)



[図4] 14menの合成

[图5] 14 men a 合成経路

図チド示すDループからアンチコドン部位に至る14menを図 5のような2種類のルートを用いて合成した。

はじめにルート1によう結合反応を試みたが ユと 3の結合 収率が非常に悪く数%を越えなかった。 そくごルート 2の様に 結合部位を変えて反応を行なった結果、各段階の Ligation (RNA Ligaseによる結合反応)の収率に大きく向上した。 1) ルート1によう方法

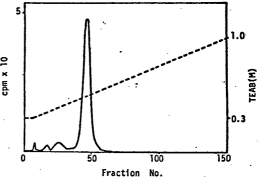
2はGGCUp を kination (P.N. kinase にようち、末端リン酸化 反応: 特に記述しない限り [よ-3p]-ATP を用いリン酸化した。) し得られた分子で、次の ligation における副反応を防ぐためる 末端をリン酸基で保護している。この こを3と ligation したが 反応の進行が悪く最良の条件下でも 15%程度しか進行しなかっ に、20% polyacrylamide disk gel electrophoresio (PAGE(disk))

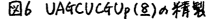
- 5 ----

で単離し、 きからの収率が4%であった。生ほkination 後 Sephadex 4-50 ご精製し 76%の収率で5を得た後点と ligation した。この ligation 反応彼はそのまま次の kination に使用し、 20% PAGE (disk)で pUAGCUCGUCGGGCUp を単離した。 収率は ligation, kination を通して 11%であった。得られた14merの ligation step における結合位置が正しいことは、 nearest neighbon analysis (後述)を行ない確認した。 2) ルートンによる方法

2,10はそれぞれ UCAUp, CAAACUp を trination1, エはイオン 交換カラムクロマトグラフィー, 10はゲル沪週ご猜製1たフラ グメントで, ルート1の場合とは異なる結合位置を有するよう に設計した。

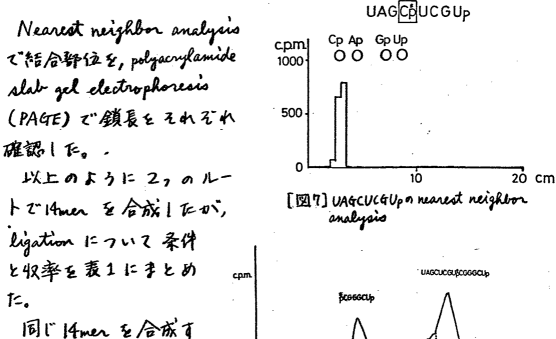
まずらとJをligation 1, DEAE cellulose イオン交換カ ラムクロマトグラフィーご単 離し(図6), 2を好収率ご得 Teo 215 nearest neighbor analysis (ligation product をRNase T2ご完全分解1 下後





pH35の条件下 PEPを行かい3Pが donon 3PUCAUp からaccepton UAAC のCに移り C3pとして検出され 結合部位が正しいこと が確認できる方法)により結合を確かめた。(図7)

215 BAP 処理により3位のリン酸基を除去し、ユルら通算 71%の収率で生を得た。生はしと ligation(20% PAGE(disk) により14men(1)を生から引谷の収率ご単離した。(図8)



3場合においても フラ グメントの組みあわせ で収率がたろく異なる。 この原因IF RNA ligase

fraction number

14merの精製 PAGE(disk) [図8]

acceptor nmol [µM]	donor nnol [µM]	атр µм	RNA ligase [unit/ml]	time hr	Reaction extent <sup>a</sup> )	Yielđ %
UCGUCG ( <u>3</u> ) 12[100]	pGGCUp (2) 20[167]	333	250	1	15	4
UAGC(6) 1.2[240]	pUCGUCGGGCUp( <u>5</u> ) 0.4[80]	200	180	2	27	11 <sup>b)</sup>
UAGC (6) 30 [150]	PUCGUP (7) 20 [100]	200	100	3.5	5 88	71
UAGCUCGU( <u>9</u> ) 7[100]	pCGGGCUp( <u>10</u> ) 8[114]	200	214	2	52	31 <sup>b)</sup>

tion mixtures contained 10% DMSO and the temperature Incuba 25\*.

a) Estimated by homochromatography.b) The compound was isolated after kination.

[表1] 14mer合成 = 5173 ligation の条件と単離収率

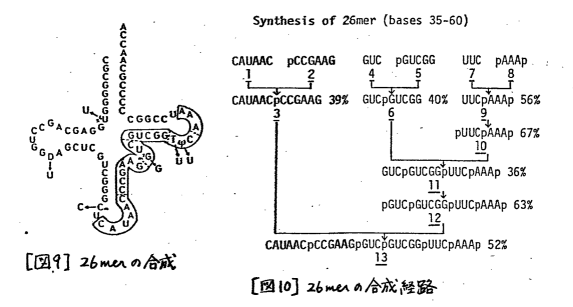
の acception 分子や donon 分子に対する認識の差によるものと考 えられる。 Acception 分子については、 3'末端がち倒からビリミ ジンヌクレオシド(Py)-フリンヌクレオシド(Pu) (ex, C-G) の順に並ぶと ligation の 収率が悪い。これは RNA 分子が右巻 き構造をとるとる'末端の Py-Puの base stacking が弱く末端のPu の conformation が国定これ難くなり RNA ligase との親和性が 低下1 ににめと考えられる。これに対しる'末端が Pu-By (ex.G-C) の場合 base stacking が By-Pu よりも強い構造を有しる'末端分 そが国定これ RNA ligase に対する親和性が増したと思われる。 以上のことから accepton 分子のる'末端スクレオシドの conformation は ligation に大まく影響を及ぼすと推論できる。

Donon 分子については sequence による辞嬰性はあまりから れないが、donon 分子が にとえば自己相補的な sequence で ある場合 分子同工が凝集し ligation が進行しなくなる。

UCGUCG+p64CUpという ligationの場合, accepton 分子の構造のみならず donor 分子についても自己相補的なsequence であるため反応が進行し難く4%という低収率であった。

第二節 アンチコドン部位からT4ループに至る26men (bases 35-60)の合成12)

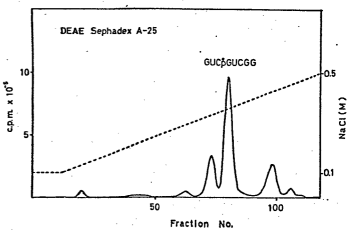
図9に示す様にアンチコドンループからT4ループに至る 修飾審基を含まない26menを図10に示す順序で合成した。これ らの結合反応で 」+2, 生+5 については donon 分子の3' 末端がりン酸基で保護されていなく, 副反応の donon/分子の重



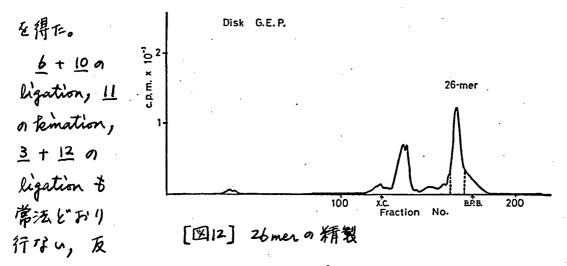
合化を防ぐためにそれぞれ accepton を donon の 10倍, 20倍と過 割に用いた。 1+2ごは反応後20% PAGE (diak)によりまを, 生 +5ごは7M urea存在下45℃でのイオン交換カラムクロマトブ ラフィーによりらをそれぞれ単離した。図11はらを得た時のカ ラムクロマトグラフィーの溶出パターンで目的物のピークの後 にまらに donon 分子が結合した副生成物のピークが見られる。

1+2に収率良く反応が進み,イオン交換カラムクロマトグラフィーにより9を 単離した。

9は Runation 後
 再びイオン交換カラ
 ムクロマトグラブイー
 を行ない定量的に



[四11] GUCGUCG4(5)の精製



応後 それぞれ urea 存在F イオン交換カラムクロマトグラフィー, Sephadex (F-50, 20% PAGE (disk) (図12) で単離した。

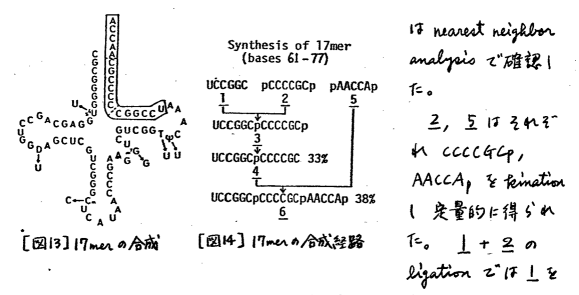
Ligation後はすべて nearest neighbon analysisを行ない, 結合部位が正しいことを確認した。

今回3年端を保護していないdonon分子を用いて ligationを 試みたが acceptor分子を大過利用い反応すれば副反応をほとん どろええ ligationを行なえることがわかった。

第三節 T4ル-70から37末端に至る17men(bases 61-77) の合成

図13に示す3末端部17menを図14の順序ご合成した。以前に この17menは当教室でUCCGG+CCCCG+CAA+CCAというフ ラグメントの組みみわせにより合成されているが、3、末端でB-Pu の配列を持っCCCCCGがacceptonとして用いられた時ligationがほ とんど進行しなかった。そこで今回著者はUCCGGC+CCCCGC+AACCA という組みあわせで行な、た。Ligation後の結合部位が正しい事

--- 10 ----

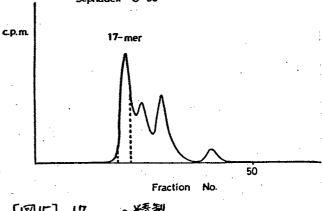


この20~30倍過剰に用いるとほぼ定量的に反応が進行した。しかし、この条件ご大量スケールで反応するにはしの量が不足していたため」をこの24倍しか過剰に用いられずホモクロマトクラフィーによる定量によると約40%しか反応が進行しなかったことがわかった。

Sephadex 年-50によろ分離にかいこも生成物うと原料ユのビー クベト、きり分離せず、1回目の分離ご施粋なえの部分のみを 集め残りのビーク部分を再度 Sephadex 年-50ご分離する操作を行 なった。シロ BAP 処理により 3 末端りン酸基を除えし、こより 海貨コマックルのあままり、

通算33%の収率ご生を 得た。生は互とligation cr により結合しSephadex G-50で単離し 38% の収率ごらを得た。

図1515らを単離する ために行なった Sephadex G-50の溶出パターン

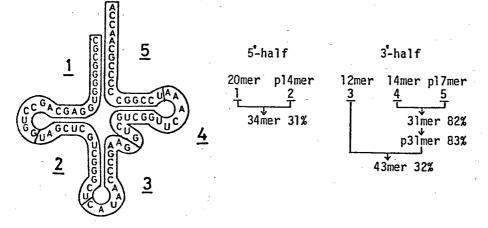


[図15] 17menの猜製

で 生成物 17menを原料の12men, 5men より単離ごきた。

以上のように17menの合成ごはすべて Sephadax 4-50によろ 分離を行なったが、原料と生成物の鎖長(ただし末端のりン酸 残基は1鎖長の効果に相当する。)が相当要なれば分離は良い が 鎖長の接近している分子同工では一度では分離でまないか あるいは分離が不可能であることがわかった。しかし回収率に 関してはこのゲルジ週法が最良と考えられる。 なお、してこの Ngation において 以後の実験より 基質濃度, 酵素濃度を共に 上げれば Ligation は良く進行し、7M urea 存在下イアン交換 カラムクロマトグラフィー により 65%の収率ご えを得ることが でまた。(実験の部参照)

第四節 5°半分子(34men: bases 1-34)及乙" 3°半分子(43men: bases 35-77)の合成



[図16] 5年分子及び3年分子の合成

图16 1=示耳様に5年分子は20men+14men, 3年分子は12men

+ (14men + 17men)の ligation により それそれ 合成 1 に。 1) 5'羊分子の合成

Acceptor 20mer (1)ほ以前に当教室で合成されたフラブメン  $H^{4}$ を用い, donor pl4mer (2)ほ第一節ご合成した14merを抱いれてい しくれを用いた。 Donor 分子に対し acceptor 分子が過剰の方が 反応条件としては好きしいが acceptor 20merの量が限られてい たため 遂に donor 分子を1.5倍週割に用い反応した。反応後 この反応液に直接 BAPを加え末端のリン酸基を除去した後(全 分子合成の acceptor とするため) nitridotriacetic acid (NTA) を加え(最終濃度を5mHに調整) 酵素を失活ませ 20% PAGE 2 分離した。生成物ほゲルより抽出し 31%の収率2、5半分子を得 た。

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2) 3'年后子の合成

3の12men と 生の 14men は第二節で 合成した フラクザメント 2°14men は 図10 の 旦を BAP 20理(調整した。 5の p17men は第三節で合成した 17menを たination し調整した。

まず生+ 5の ligationを行ない, Sephadex 4-50で単離し82%の好収率で 31 menを得た。次にこの31 menをたination し Sephadex 4-50でATPを分離した後 3と ligationした。反応の進行を図17 に示す様に20% PAGE で調べた後 Bio-gel A (0.5m)による43 men と 20% PA4E

1. marker: 44mer

- 2. ligation: 12mer + 31 mer
- 3. marker : 14mer

[図17] 12mer+31mer

Bio-gel A (0.5m) 原料 p31mer の分離を試 みた。(図18) しのし 43mer 20150 220 5°-7 ×5 明確に分離せず p31 men が 少し混在する43men を図 100 に示す範囲で集めた。 Fraction No. 20% PAGE 2 p31mer on

混在の程度を調べ(~30%)

[図18] 43 mer の精製

純粋な34分子と | こは p31men から32%の収率2、反応(集め られていることがわかった。次に混合物の手を feinationを行な い Sephadex G-200 ご精製1 この操作でほとんど混在12いる p31merを除くことがごでた。しかし43merを分析したところ 若干の3年端Aの欠除した42menが混入していることがわかっ た。これな RNA ligaseの逆反応が原因と考えられた。RNA ligase の逆反応は基質に対し酸素を過剰に用いると生じやすいが、今 すご用いてきた条件下ではほとんど生じていず 今回初めて検 出まれた。この43menについては 次の全分子合成後にATP-(CTP): tRNA nucleotidyl transferase により 3 末端が修復これ ろため このまま次の反応に使用できた。

第二章 TUループを変換したE. coli tRNAf の合成

tRNAのTUループは一部のtRNAを除いて共通の塩基配列 TUCが存在する。(図1)

原核生物の initiaton tRNAはこの領域が共通配列と同じT4C であろのに対し、真核生物では 例外的にAUCGである。また このT4ループは 5S riboromal RNAと相補的な塩基配列を有 することからtRNAの riborome への結合に関与しているという 報告間もあり與味ある領域といえる。

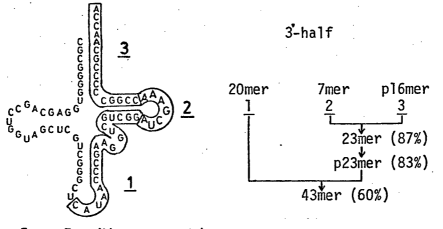
著者にE. wei tRNAft の T4ル- 2° を共通塩基配列 T4C から 真核生物の initiation tRNAに特異的に存在する AUCG へ変換す ることを試みた。

第一節 TUN-701= enkaryotic sequence を有する E. wi tRNAft の合成 18)

E. with tRNAft のTUル-7の塩基配列 TUCAAAU を真核 生物のinitiaton tRNA に存在する AUCGAAA (正確には AUCG miAAA で あろび 化学合成の都合上 AUCGAAAとした) に変換した。

1) 3'羊分子 (43men)の合成

図19に示す様に化学合成フラグメント 1, 2, 3 ×1 3 ×分 子を合成した。 3は16menを fination によりリン酸化し p16men とした分子で ほじめに TY1 ル-7° seguence 2 と ligation した。 反応後 Sephadex 9-50 で分離し 87%の好収率で 23men



[図19] 3洋分子の合成

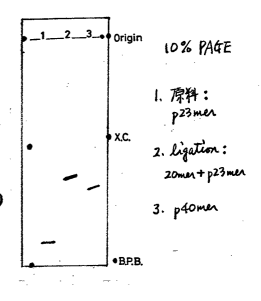
を得た。次に23menをfemation(p23menと1た後 20men (1) とligation(た。図20に示す様に Sephadex 9-100 ご分離し, 60%の収率ご目的の43menを得た。図21にこの時の Ligationの 進行を調べた20% PAGEのパターンを示すが 35年分子が生成 していることが確認ごまた。各 Ligation後は nearest neighbon analysis ご話合を確かめ、20% PAGE ご鎖長を確認した。

第一章 nascent strand tRNAft 合成の Sephadex G-100 場合,短鎖フラグメント A<sub>254</sub> 43mer から出発し何度もfeination, ligation 圣 <り返し下 15、今回の様に長鎖化学 合成フラグメントを用い 20 40 ると短期間に しかも比 Fraction No. 較的容易に目的物を得る [図20] 3年分子の精製 くとができた。(ただし,

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化学合成の立場からは この様な 長鎖オリゴマーの合成には多大へ 時間と労力を要する。) 2)全分子の合成

図22に示す様に全分子の合成 を行なった。5'半分子は E. coti tRNA<sup>thet</sup> を RNase A ご限定分解 することにより 調整した。得ら れた5'半分子は塩基配列を Z次 元ホモクロマト2'ラフィー ご確 認後 3'末端のリン酸基を BAP

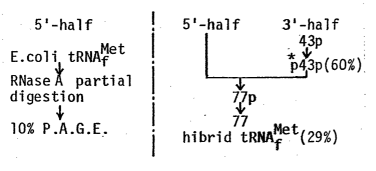


[図21] 20mer + p23mer

処理で除去し ligationの accepton とした。(BAP 処理の前に3' 末端環状リン酸を完全に開裂するために O.IN HCl で4°C 14 時間反応ませた。)

Donon 分子については1) ご述べた3年分子をたかation し Sephadex G-50 を用いATPから分離することにより調整した。 年分子同工の Ligation は accepton を donon の約2倍過剰に

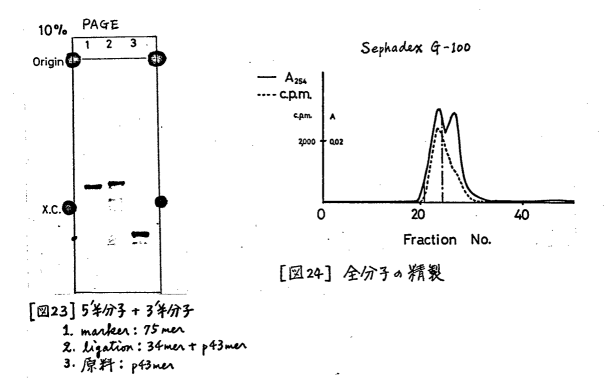
Synthesis of 77mer



[图22] 全分了の合成

用いて反応し,10% PA4E ご反応の進行 を確認した。(図23) 反応混合物をフェ ノール処理で除タン パクし、Sephadex G-1ので脱塩した後 BAP処理を行なった。

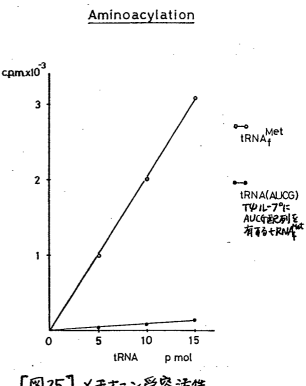
17-



BAP処理後 再びフェノール処理ご除シンハクし, 全長約し mの Sephadex 4-100 ご生成物と原料の分離を試みた。因みに示 す様に2ビークに分かれたが集めたフラクションのチェックを 10% PA4Eご行なったところ若干の短鎖生成物が混入していた。 その比率の測定からタ3%の収率ご目的の全分子を得たことが れかった。Nearest neighbon analysis による結合部位の確認 においても CmUp が 2次元TLC<sup>21</sup>により検出ていそ分子同工が 結合していろことがわかった。

次にこの分子のメテオニン段容活性を E. udiの粗酵素系(核酸成分を除去した S-100 フラクション)を用いて調べた。アミノアシル化後反応激をグラスフィルターにスポットし冷5% Trichloroacetic acid (TCA)で洗浄すると系に加えられた [1℃]-メテオニンのうち tRNAと結合したメテオニンのみがフィ

ルター上に残る。この放 射能をアミノ酸受容活性 として測定した。 合成し たtRNA は図なに示す様 にほとんどメテオニンを 受容できず天然の氏のは tRNAt に比べ4%の活 性しか示さなかった。と れは そのtRNA が安定 な三次構造をとれないた めにこのようび結果が生 じたと思われる。 アルー フィン塩基対を形成すると 考えられるTUループ



[図25] X开オニン受容活性

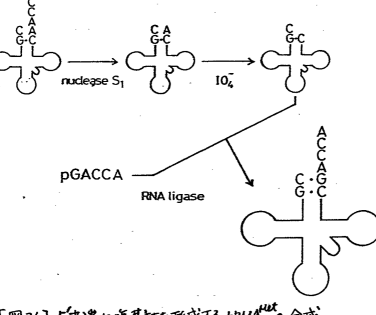
(TUCAAAU)の UC 配列はこの合成 tRNA z"は UC に変換さ れていろが この配列でもカループと塩基対を組ひことができ ろ。しかし TYループ内での T-A 塩基対(TYCAAAU)は合 成+RNAの場合(AUCGAAA)では組めない。この様なる次構 造上の相異いtRNAを不安定化し Met RSase との相互作用を弱 めにと考えられる。又 この 34分子の可変ループの mrf が合 成の都合上非修飾の日でありこのことも影響しているかもしれ ない。

第三章 アクセプターステムを変換したE.walt RNAttoの合成

tRNAのアクセプターステムはそのジ末端にアミノ酸が直接 結合する部位でタンペク質合成に関与する種々の因子との相互 作用が考えられる。

一般にtRNA はその5年端で塩基対を形成しているか、原核 生物のinitiaton tRNAだけは形成していない。ポリペアチド鎖 延長因子 EF-Tu は initiator tRNA よりも elongator tRNA に より強く結合することいわれ、これはtRNAの5年端の塩基対に 依存すると思われる。 Schulman らほ このことを確かめるため に5末端塩基をしからしに化学修飾し塩基対が組めるように変 換した。そしてEF-Tu との結合実験を行ない結合能力が上昇 したと報告している。24)しかし この化学修飾の方法では 5本端の みならず他の部位にも修飾が生じている可能性がありちま端の 特異的な影響はわからない。 そこで著者はこのち、末端で塩基対 が形成できるような特異的tRNAを合成し、EF-Tu との結合を 調べた。我々の研究室ではすでにち末端しをひに変換したち末 端から鎖長20のRNAを合成し、天然のその部分が欠除した E. w tRNAft との再構成を行ない EF-Tu との結合が増加し たという報告を行なった。著者は再構成によるのではなく RNA ligaceによろ結合反応を用いて tRNAft の 5年瑞塩基Cに対応 する3側フラグメントAを午に変換したtRNAを合成し、同 様のEF-Tu との結合実験を行なった。

第一節 5末端に温基打至形成す为E. wittRMAft の合成

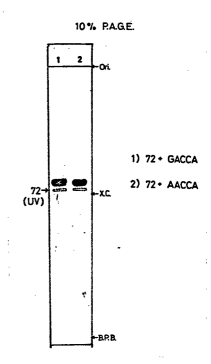


E. whit tRNA<sup>tut</sup> を立体構造を保 みをすてまま れuclease S1ご限 定分解を行なう とつかてたい アフロでの切断 マクー本鎖領域を行 男的に切断し、

[图26] 5末端に這基打を形成了3 tRVAt の合成

3'末端 ACCAの欠除 | たtRNA ボ得られる<sup>26)</sup> このtRNA をまら に修飾塩基ボ障害を受けない条件で 3'末端シスジオールを過ヨ ウ素酸で酸化し β脱離をおこすと 3'末端の一塩基ボ除去でき る<sup>27)</sup> この分子と化学合成オリゴマーとを結合ませると 5'末端に 塩基対を有する tRNA ボ得られる。(図26) 著者にこのルートに 従って tRNA 変換体を合成して。 Nuclease S1 限定分解により 得られる四塩基欠除した tRNA に我々の研究室で調整された試 料を用いた<sup>30)</sup> この tRNA を pH 5.2 の 条件下 NaIO4 で遮光下 4°C, 2時間反応ませた。次に過剰の ラムノースを加え まらに 遮光下 4°C, 0.5時間反応し 未反応の NaIO4 を不活化した。 エタノール 肥殿 により脱塩した後 HCC で pH 8.9.0 に調整し たリジン溶液を用い遮光下 窒温(約20°C), 3.5 時間反応し β脱離をおこした。エタノール沈殿 後 BAP 処理を行ない除タンパクした 後 10% PAGEで精製した。この分子 (72men)の3、末端分析を行なったがC でありAが除去されていることが確認 でまた。

化学合成オリゴマー GACCA と AACCA は Rimation 後 それぞれ Sephadex G-25, Sephadex G-50 ご単離し次の Ligation の donon とした。この 5 men と 72 men を Ligation し 10% PAGE で単離して 目的 の全分子を得た。(図27)



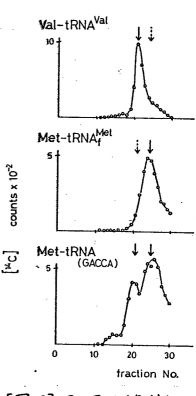
Nearest neighbor analysis ご結合部 [図27] 72men + 5men 位を確認した後 アミノアシル化反応を行なったが、塩基対を 組むようにGACCAに変換したtRNAももとの配列を有するAACCA と同じメテオニン受容活性を示した。このことからtRNAftと Met RSase との相互作用にはこの位置の塩基対は影響しない事 がわかった。

第二節 塩基対を形成1 に変換体と EF-Tu・GTPの相互作用

塩基対を形成した tRNA 変換体(tRNAf ((-4))は kination によりち、末端をリン酸化( Sephadex G-50 z\* 学離した後[化]-メテオニンを用いてアミノアシル化を行なった。 [忆]-Met-tRNAftet(c-4) を低温下 Sephadex G-50 z\* メチオニン より分離し 結合実験に用いた。

コントロールとして 5'末端に塩基 対を組んでいる E、 work tRNA<sup>Val</sup>を [HC]-Val-tRNA<sup>Val</sup> として、塩基対を組ん ていない E. work tRNA<sup>fult</sup> を[C]-Met-tRNA<sup>fult</sup> として それそれ 用いた。 E、 work から 精製された EF-Tu を用い[<sup>3</sup>H]GTP binding により EF-Tu の活性を測定した。

結合反応は EF-Tu・GDPをEF-Tu・GTP に変換した後、アミノアシル tRNA を加え O°C、5分間のインキュベー トご行下った。反応終了後 低温 (4°C)下 Sephadex G-100 によるゲ ル沪過を行ない三重複合体を形成し ているか否かを調べた。 もし三重複 合体を形成していれば EF-Tuの分



Sephadex

G-100

[図28] Tミ/アシレセRNAと EF-Tu・GTPの相互作用

子量が 43,000 dalton であることから、複合体のSephadex G-100 における溶出位置が アミノアシルセRNAの それよりも 先行する。 Sephadex G-100の溶出フラクションをグラスフィルター にスポ ットし 吃のカウントを測定することにより それらの溶出位置 を検出した。

図28に示す様にコントロールのVal-tRNA<sup>Val</sup>は三重複合体を 形成し早く溶出されるが、Met-tRNA<sup>Het</sup>は形成できず遅れて溶 出する。合成により塩基対を形成したMet-tRNA<sup>fut</sup>(c-f)は Val-tRNA<sup>Val</sup>のようにすべて三重複合体を形成することはでき

なかったが、塩基対を生じたことにより確かにEF-Tuと相至 作用することがわかった。

以前に我々の研究室で行なわれたA-U塩基対を有するtRNAf 再構成分子の実験においても同様の結果であった。

以上のことから A-U 塩基対ごあれ G-C 塩基対ごあれ ジ 末端での塩基対は tRNAの EF-Tu・GTP との 三重複合体形成能 に影響していることがわかった。しかし その寄与の程度に関し て tRNA<sup>tet</sup>の EF-Tu に対する解離定数は elongaton tRNAの場 合に比べ数倍程度しかちがわない<sup>28)27)</sup>ということから この塩基対 形成は EF-Tu との相互作用に対し本質的ではないかもしれない。

EF-Tuはち、未端以外にもRNA全体の構造も認識していると考えられるのでち、未端の塩基対の有点による数倍程度の親和性の差をどのように評価するかは問題である。

# 第四章 アンテコドンループを変換した E. coli tRNAft の合成

MRNAの情報を読みとろせRNAのアンチコドン部はせRNAが 機能するにめの不可欠な領域でありせRNA研究上の興味ある標 的といえる。MRNAと邁基対を組ひアンチコドントリフ・レット はせRNAの種類により異なるが、アンチコドンの5個隣接位は ほとんどすべてひであり3個隣接位はPu Rut Pu 誘導体がその 位置を占める。 Ead tRNAttの場合 アンチコドンの5個隣 接位は一般的なひでみるが 3個隣接位に関しては修飾まれてい ないAであり、これはアンチコドンの3字目がひのせRNAでは この位置が修飾まれているのに比べ例外的である。

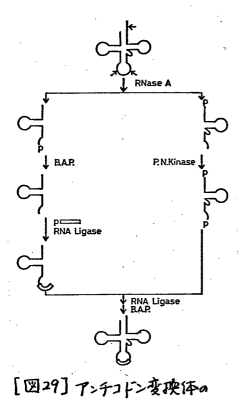
アンチュドントリアレットに関して天然に存在するサアレッ サーセRNAは多くはこの部分に変化が生じたとえば終止コドンを読めるようになり活性を示す。これらのtRNAはアンテュ ドンが変化してもアミノ酸を受容できる。

E. coli の initiaton tRNA について アンチコドン CAU を amber, opal に対応するアンチコドンに変換した tRNA を合成し、E. coli ARS ase により アミノ酸受容活性を調べたが いずれも受容しな かった。<sup>20)</sup> tRNA<sup>Mut</sup>を オプレッサー tRNA に変換する試みは この系にかいては成功しなかった。他のelongaton tRNAを用 い人エサプレッサー tRNAを 合成する試みがなまれたが この tRNA では活性を示した。<sup>30)</sup> これは tRNAの アンチコドン部を 変化ませても ARS ase に認識まれたためで、tRNAや ARS ase が 異なれば その認識のまれ方が異なることを示す。

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著者は E coli tRNAftのアンチュドン部を種々変換(たtRNA を合成し Met RSase との相互作用を調べ tRNAft のアンチュド ン部の働きについて詳しい検討を行なった。

第一節 アンチコドンの隣接位に付加塩基を有する E. whit tRNAft の合成31)



合成経路

E. coli tRNAft a 場合 そのアン デコドントリアレット が他の配列 に変化するとメテオニンを受答し なくなる。<sup>20)</sup> そこご著者は CAUと いうトリアレットは含んごいるが 拡大されたアンテコドンループを 有する tRNAft を合成し, この tRNA が E. coli Het RSase に いか に認識されるかを調べた。

図29に全体の合成経路を示した。 すず 天然の E. whi tRNA<sup>Het</sup> を RNaseA により限定分解<sup>20</sup>(5個) 半分子(1-34)と 3個半分子(38 - 25)を 10% PAGE 2"単離した。

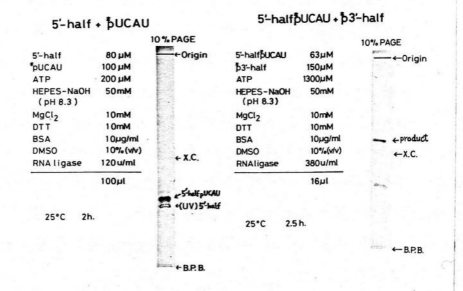
5例半分子についてはBAP処理を行ない 除タンパク,脱塩後,化学合成により得られたアンチコドン部オリゴマー(UCAU, CAUA, CAUAA)と ligation 1 10% PAGE ご订加塩基を有する 5' 側半分子を単離した。 3個半分子については Kination 後 Sephadex

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G-50でATPから分離し、先ほど得られたち倒年分子とLigation した。10% PAGE ご 単離し、酸処理(aIN HQ ご 4°C, 12時間及だ)、 BAP処理を行ない 3'末端 CCA 配列のうち CAの欠除した目的 のセRNAを得た。このCA 部分は後に E. whi S-100 フラクショ ン中に含まれる ATP(CTP): tRNA nucleotidyl Transferance により CCAに修復される。

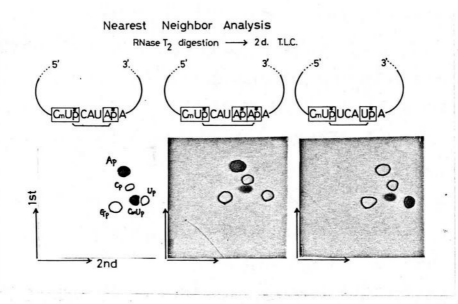
. 国30に一例として CAUをUCAU に変換した tRNAの合成に おける Ligationの条件とPAGEのオートラジオグラフを示した。 他の2種についても全く同様のパターンを示す。

各 ligation後は nearest neighbor analysis を行すい結合 部位を確認した。(図31)



[図30] アンチコドン CAUE UCAU =変換1E tRNAの合成 (ligationの条件とPAGE)

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[図31] アンテンドン に何加塩基を有引tRNAの nearest neighbor analysis

5個羊分子と合成オリゴマーとのligation では収率はあまり 良くないが(単離収率 11~25%)、半分子同工のligation では収 率良く進んだ。(単離収率 39~49%) これは前者の反応にかい て dononの合成オリゴマーの 3末端がリン酸基で保護されてい す そのため わずかではあるが重合体が生じた事と、後者 の反応にかいて半分子同工が 3次構造を組み結合部位が培近し たため ligation が良く進行した事とが考えられる。

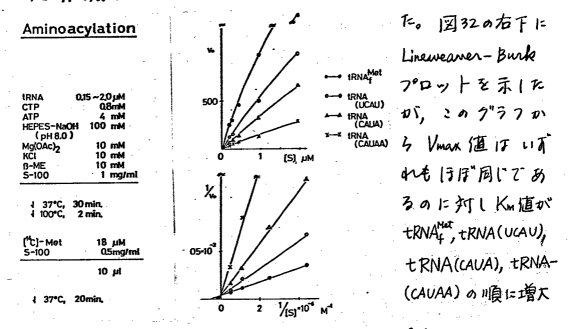
当初このtRNAの合成順序として合成オリゴマーをすず 3個半分子と結合ませることを計画したが、この場合ほとんと Ngationが進行せず図29に示T確な合成経路に変更した。

## 第二節 アンチコドンに何加塩基を有する合成せRNAの 生化学的性質<sup>3)</sup>

第一節ご合成したtRNAについてリアミノアシル化反応, 2)ホルミル化反応, 3)リボソーム上ごのコドンの認識を調べた。 1)アミノアシル化反応

. E. wi 粗酵素系(S-100 フラクション)を用いてメチオニン受容 活性を調べた結果 天然の E. wi tRNAft に比べ UCAU配列を有 する tRNA(tRNA(UCAU)) では 53%, tRNA(CAUA) z は 25%, tRNA(CAUAA) では 13%の活性を示した。

次に図32に示す様にKm, Vmax値を求めた。まず合成 tPNAの 37末端をCCAに修復した後["C]-メテオニンを加え反応を開始し、 一定時間後 グラスフィルターにスポットし酸不溶性放射能を測定し



[図32] アンチコドンに何加這基を有する tRNA a aminoa cylation (Km, Vmax, study)

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Het RSase との親和性がこの順に減少している事がわかった。
 Km 値はそれぞれ ハフルH, 3、3ルH, 6、7ルH, 12、5ルH であった。
 次にこれらのtRNA が他のアミノ酸を受容するかをかを16種のアミノ酸(Ala, Val, Leu, Ile, Phe, Pro, Gly, Sen, Thr, Tyr, Asn, Asp, Glu, Lyo, Hio, Ang)を用い検討したが いずれのアミノ酸
 (こついても アミノアシル tRNA は検出できがかった。
 スルミル化反応

Transformylase TF原核生物のinitiator tRNAと特異的に反応 する酵素で、その認識部位についてはほとんどわかっていない。

合成した 3種の tRNA について Methionyl-tRNA がホルミル 化を受けるかをかを調べた。反応条件は アミノアシル にの条件 と月ぼ等しく、ホルミルドナーとして folimic acid-SFを加え E. coli S-100 フラクションご反応した。反応後 硫酸銅で処理し、ホ ルミル にを受け なかった Met-tRNA を deacyl にした。硫酸銅 で処理した フラクションとしなかった フラクションのカウント の 比を ホルミル にの 比として百分率ご 長わし、 tRNA (UCAU) → 93%, tRNA (CAUA) → 88%, tRNA (CAUAA) → 98% という結 果を得た。 アミノアシル にの 場合 と買なり transformylase は アンチコドン配列を認識していない ことがわかった。 3) リボソーム 上でのコドンの 認識

アンチコドンループを拡大したこれらのtRNAについて, mRNA上のコードを認識するかをかった。

1964年 Nirenberg らが RNA codeword を決定1 に時に用い た方法<sup>33)</sup>に基づき行なった。(四33) mRNA として合成オリゴ マー AUG, UGA, UUA, UAU を用い リボソーム上ごの特異

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#### tRNA-mRNA interaction

tRNA mRNA Tris-HC1(pH7.5) Mg(OAc) <sub>2</sub> KC1 E.coli ribosome 305 or 705	20 mM 50 mM	mRNA	tRNA <sup>Met</sup>	trna Caua	trna Cauaa	trna Ucau
		AUG	100	0	0	33
		UGA	0	0	0	0
	1.8μΜ	UUA	0	0	3	0
	20 µ1	UAU	0	0	0	0
25°C, 30min.		$\mathbf{Z}$ relative to tRNA <sub>f</sub> <sup>Met</sup> -AUG				

[四33] ボソーム上でのコドンの認識

的結合を測定した結果, tRNA(UCAU)のみが AVGと結合し他の合成 tRMAについてはいず れもmRNAと結合しな かった。tRNA(UCAU) のAUGへの話合も天

然のtRNAfのAUGへの結合の約以であった。

tRNAのアンテコドンとmRNAが塩基対を形成するためには、 アンテコドントリプレットボループの外側に出て3塩基が並ば がければならないが、これらのループを拡大したtRNAは行加 塩基の存在によりトリプレット付近のコンホメーションが変化 しmRNAと塩基対を形成することができなくなったと考えられる。

以上の実験結果を基にしてまとめると、Met RSase との相至 作用については tRNA が認識を受けるには アンチコドンの特定 の立体構造と CAU という塩基配列が必要であることがわかっ た。CAU の 5個に Uを何加した tRNAの方が 3個にAを何 加した tRNAより Met RSase に対しより強い親和性を示したが、 これは CAU の 5個に CAU のコンホメーションを ある程度保 持したまま Uが入りこめる余地が存在したためと考えられる。 このことは mRNA との塩基対形成実験においても tRNA (UCAU) のみが AUGと塩基対を形成でまた結果からも推測でする。

tRNAftのアンテコドン部はこの分子にとって機能を果た すために非常に重要ごあることがわかったが、一方ホルミル化

に関しては、この部分は影響しないことも判明した。

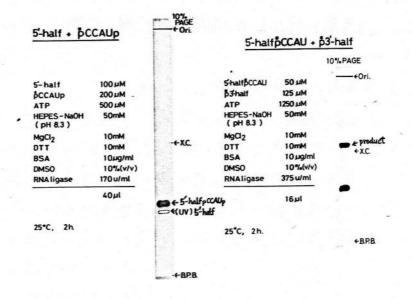
第三節 アンチュドンの 5′側隣接位塩基変換34)

マンテコドンの5個隣接位に何加塩基Uが存在してもアンチ コドン付近のコンホメーションがあまり変化しないことが第二 節ご明らかになった。そこでこの何加塩基Uを他の塩基A,在, C, UUに置きかえたtRMAを合成し, Het RSaseによる認識に Uの場合とどのように異なるかを調べた。さらにアンチコドン のち個隣接位はtRNAに変通のUであるが, この塩基を特異的 に他の塩基に変換したtRNAの合成も試み そのメチオニン受 容活性を調べた。

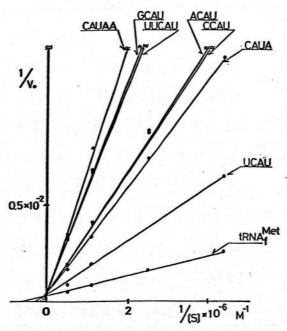
1)5個隣接位にU以外の付加塩基を存するtRNAの合成

全成方法は第二節で用いた方法と同じく RNase A 限定分解よ り得られた天然の E. with tRNAft 5個半分子に公学合成オリコ" マー ACAU, GCAU, CCAU, UUCAU をそれそれ RNA Ligase こ 結合エセ, 最後にこの分子と 3個半分子とを結合エセス全分子 を得た。 結合部位が E { い ことは nearest neighbon analysis により確認 1 た。例と | て 図34に CCAUの 場合における Ligation の条件と 10% PAGE の オート ラジオグラフを示す。今回は合成 オリコッマーの 37末端 ジリン酸基ご保護されていたため、55半分 子 + 合成オリコッマー a ligation において ポリメリゼーション は 防ぐことがでまた。 アンチコドンを有する 5個半分子は 10% PAGE ご単離(, BAP処理により リン酸基を除去(次の ligation の accepton と した。

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[図34] Tンモンドン CAUE CCAUE 変換IE tRNAの合成 (ligationの条件とPAGE)



合成した tRNA について第 二節と同様にメテオニン受容 活性を測定し、今までの拡大 ループを存する tRNA と Km 値の比較を行なった。(図35)

その結果 アンチコドンの 5個への付加塩基はしの時が 最も Met RSase との親和性が 強く, 他のC, A, Gの場合は

[図35] aminoacylation (Km, Vmax study), デトラマー, ペンタマーは アンチンドン CAUを それぞれのオリンマーに変換」に tRNAを示す。

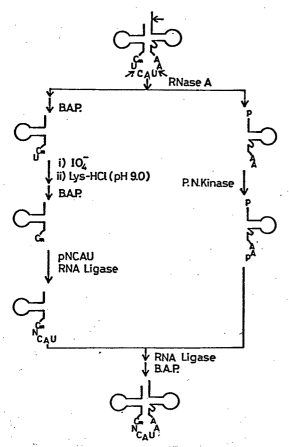
- 33 -

3個にAを付加した時と変わらないか それよりも弱い程度であった。2塩基を付加した場合は 3個,5個いずれも親和性は弱かった。

2) 5個隣接位に存在する共通しの変換

天然の氏いれtRNAftをRNaseAにより限定分解し、得られる 5個半分子をエジに過ヨウ素酸酸化し3末端の共通しの除るを 行なった。過ヨウ素酸酸化は第三章ご用いた方法と国様に行なった。図36に合成経路を示す。当初5個半分子のみを過ヨウ素 酸酸化し、経路に従い合成したが、得られたtRNAはコントロ

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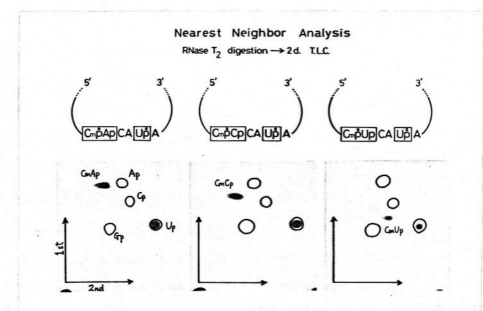


[図36] 英通U变换+RNAの合成経路

-ルとして合成した天然と .同等の塩基配列を存する

セRNA でええも そのメチ オニン 愛溶活性をほとんど ママガかった。これはち倒 ギ分子単独の場合, セRNA の様に三次構造をとろこと ができず修飾塩基等が障害 を受けたためと考えられる。 そこで週ヨウ素酸酸化の時 に当量の3個半分子を加え 再構成させて(アンチュドン は欠いていろが) セRNAの 構造を保持させ週ヨウ素酸 酸化を試けた。酸化後, BAP 処理を行体、10% PAGE で3'末端の一塩基を除玉したち'側半分子を単離した。3'末端分析を行ないしが除玉できた事を確認した後 公学合成オリゴマー と hightion した。 化学合成オリゴマー UCAU, CCAU, ACAU は たination しイオン交換クロマトク"ラフィー で単離して hightion の donon とした。 Lightion後 10% PA住 で単離した。 酸処理 (aIN HC& で 4°C, 6時間反応)により末端サイクリック リン酸 を開製した 3'側半分子は reination後 Sephadex (5-50 による精製 を経てアンチコドンを含む 5'側半分子と hightion 1 た。 10% PA住E で単離し BAP処理を行ない目的の tRNA を得た。

図7に全分子合成後a nearest neighbor analysis を示す。 Donor分子の「末端に位置していた3pri acceptor分子に移り, RNase Tz digestion で □ ご囲んだモノマー, ダイマーが 2次元 TLCにより確認できた。最後の ligation でにいずれも CAUの U

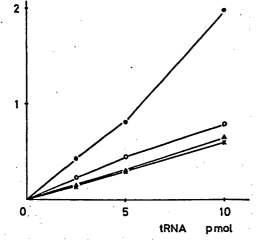


[図37] 英通UE变换IEtRNAs nearest neighbor analysis

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由来のUsip が検出されるが、5個半分子と化学合成オリゴマー の ligation では5個半分子の3末端は2-0-メテルC(Cm)でここ にオリゴマーが結合するとRNase T2 ではCmの3側では切断され ずダイマーCmapNp(N:A,C,U)で検出される。このCmapNpは各々Nの ちがいにより二次元TLCでその位置が異なる。この差異により Nを確認した。合成したtRNAについてメテオニン受容活性を 測定した。その結果を図珍に示す。アンチコドンの隣接位をU 以外の塩基に変換したtRNAにちいてもUのtRNAと同等のメ チオニン受容活性を示し、アンチコドンの5個隣接位の失通U は下ミノアシル化に影響を及ぼしていないことが判明した。

最近UhlenbechらによりYeasttRNAがのアンテコドンとその5例 隣接位Uを変換したサプレッサーセRNAが合成をれてnanslationで はこのUF本質的な役割を果たしていないことが示をれた。今 回の著者の実験も同様の結果を示しておりアミノアシル化の投 Met-IRNA 階で差がないことを示し



cpm x10

> [図38]共通UE要換IF+TRNAのaminoacyletion -- tRNAft, o-o tRNA(CmU→ cmU) control, わらず 完全に構造を回 A-A +RNA(cmU→ cmC), X-X tRNA(CmU→ cmA) 復できずに反応が進み

た。この実験においてア シノアシル化の名が天然 のtRNAteに比べ約半分し い回復していないが, これは過ヨウ素酸酸化の 際にもRNAとしての三次 構造が組めるように3個 半分子を加えてにもかか わらず、完全に構造を回 復できずに反応が進み

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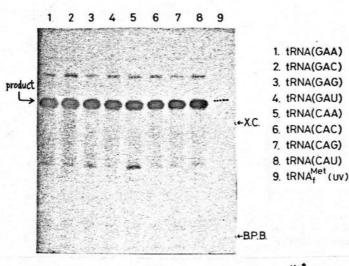
塩基に障害が生じたためと考えられる。

第四節 アンチュドントリフ・レットを種々の配列に 変換した E. coli tRNAft の合成

E. しん tRNAf のアンテュドンの 3個隣接位は例外的に未修飾 のAごある。 E. しん 中ごは修飾をれない tRNAft の このAは 真核生物の系では修飾をれるかをか,又アンチュドンの配列に より修飾のをれ方が変化するかどうか等を調べるために アンチ コドントリプレット CAU を CAA, CAC, CAG, CAU, GAA, GAC, GAG, GAU。各合成オリゴマーに変換した tRNA を合成した。

合成方法は第一節,第三節で記述した拡大ループを有する tRNAftの合成の場合と同様に行なった。各トリマーのをmation は非放射性のATPを用いて行ない、ゴ側半分子のをinationには

10% P.A.G.E.



[図39]アンチコドントリアレットを変換したもRUAはな合成

比活性の高い[8-3p]-ATPを用いた。トリマ -と5個半分子との Digation後10% PAGE で単離し3個半分子 とligation1 た。図39 に半分子同工のLigation における10% PAGEの 泳動パターンを示す。 生成物をゲルより切

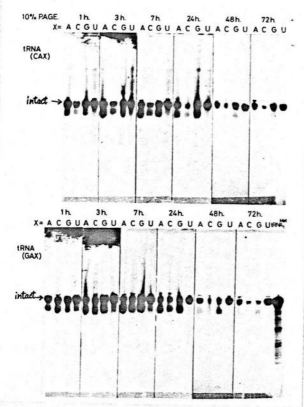
り出し ノサングルあ

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モリジアで、106cpm以上のカウントを有する tRNAを得た。

第五節 アンテコドン変換体の Xenopus laevis oocyte 中における生化学的性質

Xenopus laevis oocyte (アフリカツメガエル卵母細胞)は, mRNA の翻訳が非常に効率良く行なえる系<sup>36)</sup>として多くの研究者 に利用されている。この系を用いると容易に in vivoで、 の実験が行なえ,後処理も比較的簡単であるため著者はこの系 を用い修飾の実験を行



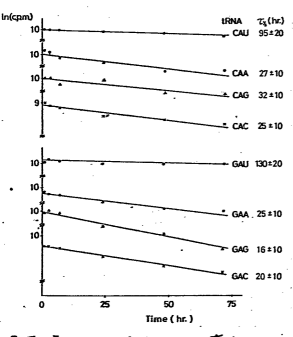
\$7 1=0

第四節で合成15-8 種のtRNAをXenopus laevis oocyteの細胞質 にmicroinjection(下. 一定時間かえに反応 を止め oocyte を破砕 1除タンパク後 10% PAGEにapply ( tRNA の安定性を調べた。

図40にその泳動ペタ -ンを示す。過剰の核 酸成分のため泳動が乱

[图40] or cyte中に injection IE アンチンドン安硬体(インキュベーション時間ごとの PAGE)

Stability of tRNAs in oocytes



[四41] アンデコドン変換体の のひりを中に あいな安実性

れていろか, intact tRNAと 3~4塩基切断の生じたtRNA のバンドが検出てれた。

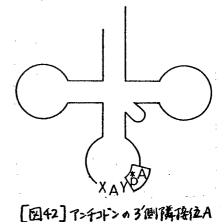
Intact tRNAのバンドを 切り出し そのカウントを測 定してこれらのtRNAsの oocyte中にかける安定性に ついて調べた。(図41)

その結果 アンチュドント リプレットボ CAU, GAU のもRNA は 3~4 塩基切断 されたバンドがほとんど検 出されず非常に安定で, そ れぞれ半減期 てとが 95±

20時間,130エ20時間であった。それに対し他のアンチコドントリプレットを有するもRNAはいずれも切断物が検出されてな

が20~25エ10時間と不安定であっ た。次に切り出したメインバンドの らもRNAを抽出し、エタノール沈殿

(キャリアーと12 Yeast tRNAを約 0,2 A20 unit 加えた。)を2回行ない 脱塩した後, nuclease P1 によろ完全 分解を行なった。反応後サンプルを 2次元TLC上ご展開し、PNの位置 を確認した。図42からわかる様に

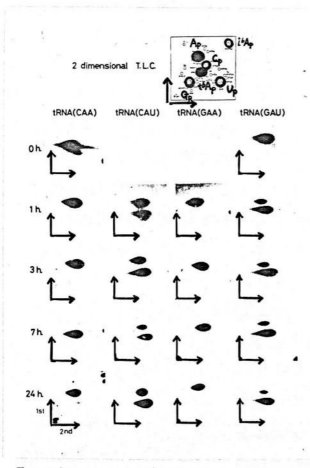


の修飾 X: C, G , Y: A, C, G, U

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<sup>2</sup>Pのラベルは<sup>23</sup>PAも1<は<sup>25</sup>PA<sup>\*</sup>(A<sup>\*</sup>は修飾まれたA)と12 TLC上に検出まれた。図43に 2次元TLCの一部を示すが,Aの 修飾が生じたのは tRMA(CAU)と tRNA(4AU)のみで 他の tRNA に関1 こは何ら修飾を受けなかった。又 tRNA(CAU)と tRNA-(4AU)の場合 いずれもAはt<sup>6</sup>Aに修飾を受けていた。一般に アンチュドンの 3 字目が U ご ある tRNA では 3'倒隣接位のA は ほとんど t<sup>6</sup>Aに修飾を受けることから 予想じ かりの 結果であっ た。又、アンチュドンの 3 字目が A で ある tRNA では 3'倒隣接

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[図43] アンチコドンの3側隣接位Aの同定

位Aは 16A などに修飾 を受けるが、この tRNA-(CAA) と tRMA(GAA)の 場合 全く修飾を受け なかった。 これは 26A に修飾を受ける tRNA に下ンチュドンステム の最も ループ寄り の 漏 すが A-4 であるの に対し tRNAt の場合 G-C ご あるため修飾 酵素に認識まれなかっ たとも 考えられる。

Xenopus laevis oocyte中でのセRNAの 修飾に関して E. wit ではアンチコドンの3 側隣接位Aが修飾されない E.whi tRNAft 2\*\* orcyte 中2\*\*は 他 のアンテコドンのう写目がしてあるtRNAと同様 このAが せんに修飾されることがわかった。Xenopus Laevis orcyte 中 に存在するたん修飾酵素はE.whiの場合と異なり全体のtRNA 構造を認識するのごはなく アンチコドンの3字目の塩基に影 響されることがわかった。E.coliのた人修飾酵素の場合、tRNAft のどみ部分を認識して修飾をしないのかは不明である。

次にXenopus laevis oocyte 中の tRNAの安是性に関ーこに, 今回初めてアンチコドンの配列がtRNAの安定性に影響を及ぼ すことが確認された。アンチコドン配列として CAUと GAUを 頂する tRNAft のみが安定で 他の配列のtRNAft ごは不安定 であった。これは 修飾を受けたために安定になったとは考え られない。なぜなら tRNA(CAU), tRNA(GAU)は修飾を受けた が これら分子 すべてが修飾を受けたわけではなく 未修飾の分 そも存在し、それごもなお安定性を保っていたからであろ。

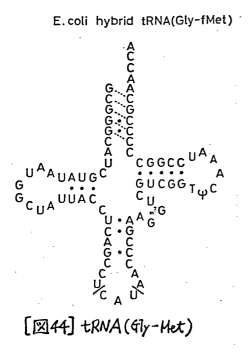
この場合、アンチコドンハ3字目にFリ安定性が規定IAK N、なぜ Xenopus laevis orcyte 中で E. wi tRNAfe a この 位置がUの分子が安定で 他の分子が不安定であ、たかについ ては不明である。 第五章 与侧羊分子を变换した El coli tRNAft の合成37)

5個単分子にE、uditRNA2の塩基配列を有するtRNAft (tRNA(fly-Met))とミトコンドリア中のtRNA<sup>Ser 3010</sup>などにみら れるDル-フッとステムの欠除したtRNAftを合成しそのアミノ 酸受容活性を調べた。

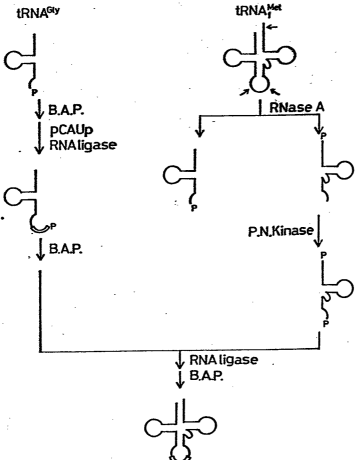
第一節 5個半分子に E. coli tRNA<sup>gly</sup>の 塩基配列を有する E. coli tRNA<sup>flet</sup>の合成

E. coli tRNA<sup>fly</sup> は修飾塩基がかなく 5個半分子には全く含まれていない。このことはこの分子を化学合成する上で非常に有利な点で、最近アンチコドンを含まない E. coli tRNA<sup>fly</sup>の 5個 E. coli hybrid tRNA(Gly-fMet) 半分子(33 men)の化学合成が試み

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られ成功した。39) 著者はこの合成されたせRNA<sup>fly</sup> のち倒単分子の33menを用い,ア ンチコドンと3個半分子にE.cohi tRNA<sup>net</sup>の塩基配列を有するたybrid tRNA(図44)の合成を行なった。



[図45] tRNA(Gly-Met)の合成経路

を得た。結合部位は nearest neighbor analysis ご正(い事 を確認した。 合成 [ F tRNA IZ ついてメチオニン愛 容活性を測定したが 受客活性を示えない った。(第二節図49) 又, グリシン受容活 性についても調べたが活性を示さなかった。合成したtRNAは ゴ末端CCAのCAが欠除しているが「アミノ酸受容活性を示ぎ

を得に。3個半分子

は 天然の E. whi tRNAt

より得られろ分子を

trination (調整(下。

±を RNA ligase z"結

合工七 10% PAGE で

精製 L Rybrid tRNA

これらの年分子国

なかったのは CCA 修復が生じていない事が原因ではないかと 考え E. coli S-100 フラクションご tRNA の 3 末端修復の確認を 行了, TE。修復反标至行了, TE後[5-3p]pCp a single addition を行ない10% PAGEで tRNA%Cp を単離(た。 RNase Tz Z" 完全分解しNB(N=AnCnGnU)を調べた結果、反応前ご

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はCzyが検出されたが修復後Azyに変化し、3本端が修復されていることを確かめた。

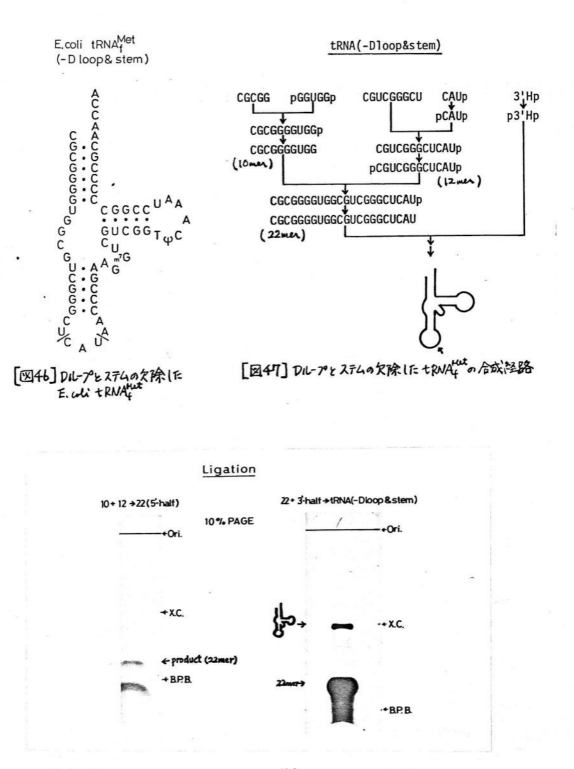
この大yurid tRNA は共通の二次構造に従って塩基対を配引 するとアクセプターステムごニ塩基対しの組むことができない が、図44の破線で示す様に一塩基ずらすと五塩基対組める。 Ryurid tRNA は安定な後者の構造を形成すると予想されるが 全体か形が天然のtRNAft と夏なるため このことが原因ご ARSase に認識されなかったと考えられる。

第二節 Dループとステムの欠除してE Locki tRNAt の合成

ミトコンドリアには一般的なクローベ葉型ニ次構造を有する セRNAと異なり Dループとステムが欠除した特異的なせRNA N存在する。セRNA<sup>Ser</sup> がその例ごこれらの部分が欠除している にもかかわらずセリンを受容しセRNAとしての機能を果たす。 他のクローズ葉型ニ次構造を有するセRNAも Dループやステム がなくても機能するであろうか。着者は E.ooki セRNAft につい て Dループとステムが欠除した セRNAを 全成し、この分子が メテオニンを受容できるか否かについて調べた。

アクセプターステムとアンチコドンステムの開は それぞれ のステムに続く塩基配列を各三塩基,二塩基延長ませ、ミトコ ンドリア tRNA<sup>ser</sup>の場合と同様五塩基の間隔をおいた。(四46) 合成経路を図刊に示す。 5個半分子は顔長が 3から 9 a 化学 合成フラグメントを RNA Lignale ご 結合することにより合成し た。 3個半分子は 天然の たいび tRNA<sup>fert</sup>の RNase A 限定分解

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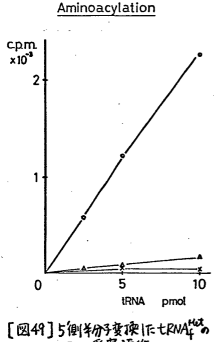
[四48] Dil-7"2 274の欠除 LE tRNAf 合成に Firs ligation

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より得られるフラグメントを用いた。

CGCGG+pGGUGGp, CGUCGGGCU+pCAUpの名ligationより 得られる10men, 12menはいずれもSephadexG-50により猜製 した。10men+12men, 22men+3個単分子の名ligationでは いずれも10%PAGE(図48)で単離し、目的のtRNAを得た。

合成した tRNA についてメテオニン 受容活性を測定した。 図49に結果を示すが Dルーフとステムの欠除した tRNA は 全くティージ しないわけではなく低いながらも わずかにメテオ ニンを受容することがわかった。 Dルーフ・が欠除すると Dルー プとT 4ループの水素結合ができなくなり, tRNA としての 安 定す三次構造が組めないと 予想をれる。 それな Met RSase に認



x开=>受容活住 o--o:tRNA4, a-a:tRNA(-Dloop&stem), x-x:tRNA(Gly-Met) 載これにくくなりほとんどチャージ しなかったと考えられる。Dループが 欠除しているにもかかわらず、天然のtRNAft と同様のL字形三次構造をもつ分子がわ ずかに存在することが推定できる。

最近 Klugらによりミトコンドリ アのtRNA<sup>ser</sup>も安定な構造をとれる 三次構造モデルが提出まれた。<sup>40</sup>この モデルに従いアクセプターステムと アンチコドンステムの間の塩基を変 換しtRNAを合成すれば、安定な三 次構造を有するtRNAが得られメチ オニン受容活性が上昇することが予 測まれる。

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### 結 論

1. 種々の化学合成フラフジントを用いてRNA ligase 反応を行 ないE、colitRNAft の5個半分子、3個半分子を合成した。

又,多くの ligase 反応の結果から RNA ligase に認識され難い 基質の塩基配列を見出した。

- 2. E. odi tRNAft の TUル-プの中で共通塩基配列 TUCをAUC に変換したtRNAft を合成し、そのメテオニン受容活性を調べ たところほとんど活性を示すなかったことより、TUル-プの 塩基配列がE. coli Met RSaseの認識に関与することがわかった。 3. E. coli tRNAft の 5年端位でG-C 塩基対が組める修飾tRNAft を合成し調べたところ、天然のtRNAft と同様のメテオニン受 溶活性を示した。又、E. coli EF-Tu との親和性がこの塩基対 により強まることを示した。
- 4. E. coli tRNAft のアンチコドン部を変換した修飾tRNAft を合成し、Met RSase による認識には CAU配列が必要で、こ の配列が存在すれば他の塩基を3側あるいは5側に付加して も親和性の差はあるにせよ認識されることが明らかとなった。 又、アンチュドンの5個隣接位に共通して存在する塩基U を他の塩基に変換しても Met RSase による認識には影響しな いことがわかった。
- 5、 E. coli tRNAft の アンテコドン部を変換した修飾セRNAft は、 Xenopus laevis oocyte 中でその配列により安定性が異 なろことを見出した。すなわちアンチコドン配列が tRNAの 安定性に影響することを初めて示した。又, E. coli 中では修

飾を受けない塩基がXenopus laevis oocyte中では修飾を受けることを示した。

- 6. E. coli tRNAftの5個半分子をE. coli tRNA21に変換したtRNA を合成し、この分子がMet RSase, Gly RSaseに認識されないこ とがわかった。
- 7. E. witt RNAftのDル-プとステムの欠除したtRNAを合成 し、Met RSaseによりわずかにメテオニンを受容することがわかった。

#### 谢 辞

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### 実験の部

#### Materials

・基質

種々のリボオリゴマー 棄化学教室で合成されたもの E、withNAft アメリカ合衆国 Yale大学 D.Söll博士より 供与されたもの [3P]-正リン酸 日本アイソト- 20協会

[忆]-T:1酸 Amersham Folinic Acid-SF(leucovorin) Lederle製, 武田薬品工業株式 会社

・酵素

RNA ligase 及び"P.N. Kinase (3'-phosphatase free) 藥化学教室で猜想をれたもの

BAP Worthington Biochemical Corp. (BAPF), 及21" 宝酒造株式会社

RNase T2 三共株式会社 RNase A Bockringen - Mannheim 社

ヌクレアーゼ Pi ママガ醤油株式会社

ヌクレアーゼ S1 三共株式会社

E. Loti 粗酵素系 S-100 分画 薬化学教室で調整

E. whi EF-Tu· fipp 東京大学医科学研究所上代淑人 博士より供与されたもの

Xenopus laevis oocyte ベルギー Université Libre de Bruxelles H. Grosjean 博エ 5リ供与 I れにもの

・樹脂

DEAE Sephadex A-25 Pharmacia ZI DEAE Cellulore (DE 23) Whatman ZI Sephadex G-25, G-50, G-100 Pharmacia ZI Bio-Gel P-2 Bio Rad ZI Bio-Gel A (0.5m) (200~400 mesh) Bio Rad ZI DEAE Cellulore plate (polygram cell 300 DEAE/HR-2/15) Macherey Nagel ZI

アビセルSF(セルセース薄層プレート) フナコシ薬品株式会社

#### Method

· RNA ligase 反応

通常50mM HEPES-NaOH (pH813), 10mM DTT, 10mM Mg(lz, 10 μg/ml BSA, 10~15% DH50 (V/V)を含む buffer中 25℃で インキュベートした.

基質量,酵素量については各章ごとに後述する。

· P.N. Kinase Zetos

通常 50mm Tris HCl (pH9.6), 10mm MgCl2, 2.0mm スペルミン, 10mm DTT, 0.1MKCl を含む buffer中 37°C ごインキュ ベートして、

基質量,酸素量については各章ごとに後述する。

• BAP 処理

50mM Tris HQ (pH 8.0)中, 碁質 | nmad に 対し 酵素 0.005 unit 加之, 基質 100~200 µMの奈件で 50°C 1.5 時間 インキュ ベート して 反応を行なった。 反応後は 反応液と同量の 50mM Tris HQ (pH 9.5) 飽和の フェ ノール, クロロ ホルム混液(1:1, %) によって 2度除タンパクし, フェ ノール相を さらに 50mM Tris HQ (pH 9.5)で 3回逆抽出してから水相を約2倍量の エーテルで 5回洗った。溶液中のエーテルを蒸発させた後 Sephadex 6-50 で脱塩した。

· RNase T2 によろ完全分解

40mm 酢酸アンモニウム buffer (pH4.5)中, 基質約2A260 unit (キャリア RNA を含ひ) に対して酵素的 0.05 unit かえ, loyul で 37°C 3時間インキュベートした。反応後 混合物は 沪紙にスポットして pH3.5 の余件下 PEP を行なった。

- ・RNase Aによろ限定分解 第二章の実験参照
- · Nuclease P1 1=F3完全分解

40mM 酢酸アンモニウム buffer (pH5,0)中, 基質約2A260 unit (キャリア RNA を含ひ)に対して酵素約 0,1μg カロン, 10 μl ご 37°c 2時間インキュベートした。 。除タンパク

通常50mm Tris HU (pH25) ご 飽和したフェノール,クロロホルム混液(111,%)を試料に対し国量加之2度抽出操作を行なった。フェノール相は50mm Tris HU (pH25) ご 逆抽出を3度行ないほじめの水相とあわせた。この水相はエーテルご5度洗いフェノール,クロロホルムを除るした。エーテルを留を後エタノール沈殿,又はSephadex (T-50により脱塩した。

。エタノール沈殿

3倍容のエタノールを加え -80℃ご 5分以上冷やし 13,000 rpm, 10分間遠心し上清を除去した。

·沪紙電気泳動(pH 3.5)

東洋決紙No51Aを用い, 0.2 M 酢酸モルホリン buffer (pH 3.5)中, 35 Vcm の電圧で泳動した。泳動後 沪紙を風乾し, 5mmの巾に切って液体シンテレーションカウンターでカウント を測定した。

· 二次元TLC

西村の方法<sup>21)</sup>に従う。 試料をセルロース薄層プレート(ア ビセルSF, 10 m× 10 m)にスポットし、一次元目をイソ酪酸 ーの5Mアンモニア(5:3, 1/2)ご展開し、二次元目をイソプ ロパノールー濃塩酸-水(70:15:15、1/2)で展開した。 •ホモクロマトグラフィー

Sanger等<sup>41</sup>及びWu等<sup>42)</sup>の方法に従ってhomomixture を調 製した。サンプルをDEAE-Cellulose plate にスポットした後, plate をMeOHで洗い サンプルの両脚に色素のマーカー混液 (1%キシレンシアノールFF(青), 2%オレンジ f(黄), 1 %酸性フクシン(桃))をスポットし,上端にはWhatman 3MM 沙紙を吸取紙としてクリップで止めて展開を行なった。(60~70 ℃) 青色マーカー び plate の 先端に達した時点で展開を終え, 風乾後 原点及び を素マーカーの位置に切のマーカーをスポッ トして ラップで包み, X nay 7ィ ルムを接触(感光した。

ホモクロマトグラムから各スポットの量比を算出する場合は、 スポット部分の薄層をかきとりこれをプラステック製ミクロチ ユーブの底に穴をあけ脱脂綿をフめたものにアスロレーターご 吹いこみ Cerenkov法ので測定し算出した。

・二次元ホモクロマトク"ラフィー Wu等, 及び Silverklang 等49 の方法に従う。

・アミノアシル化反応

合成したtRNAのアミノ酸受容活性に Km, Vmax を求める 時以外後に示す条件ご行なった。

反応後 全量をWhatman グラスフィルターにスポットし、冷5% TCA 溶液にひたし 氷冷下 10分間放置した。この洗浄を後2 回計3回行なった後、EttoH: Etz0=1:1 (1/2)溶液を加え 氷冷下 5分間放置した。これを再度行なった後フィルターを赤外

線ランプ下ご乾燥をセ,

トルエンネシンチレータ ーご やのカウントを測 定した。

Km, Vmax を求める時の条件は第四章の実験で 記述する。 アミノアシル化及応の条件

HEPES-Nadh (948.0)	OIM
Mg(CH3 600)2	lomM
KU	IOMM
mercaptoethanol	lomM
ATP	4mM
CTP	ImM
L-[U-14C]-amino acid	9 mM
trna	0,3~2µM
E. whi S-100	Img/ml
	15 pl
in whate at 27° Im	A0

incubate at 37°C for 40 min.

• ポリアクリルアミドゲル電気泳動

10~20%のアクリルアミド溶液(アクリルアミド:ビスアク リルアミド=30:1(W/w),7M urea,45mM Trisborate buffer (pH8:4),2mM EDTA)に週硫酸アンモンとTEMEDを 加えてゲル化して。

Slabgelの場合は厚さがの5~2mm,中20~30cm,長540cm の平板ゲルになるようにガラス板ごわくを作りえこにアクリル アミドを流しこんご国めた。

Diskgelの場合はMiles Laboratories Inc. 製の PREP-DISC を用い 70 mm² × 8 cm のゲルを調製してこ。

サンフッルのapplyの前に2時間程 prerunning をしてゲルの 状態を確かめてから、凍結乾燥したサンフッルに loading solution (9M urea, 10mM EDTA, 40mM Tris borate buffer (pH8:4), ~0.1 % X.C. - B.P.B.)を 2~5 ル 加之 100°C, 2分間 加熱し apply を行すった。

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泳動後 Alabgel ごはガラス 板をはずして ラッファごおおい,Xnay 7イルムを接触させ フリーザー中ご惑光した。Dusk gelの 場合はポンファごくみ出されたフラクションをそのままバイアル に入れて Cerenkov法により測定した。

• ゲルドからの核酸の溶出

. Maxam - Gilbertの方法<sup>67</sup>をかく変えて行なった。溶出用 buffer にはSDSを加えずに調整した。溶出液を濃縮し, Sephadex G-50による脱漏を行ない複酸を単離した。

• DEAE-Cellulore (DE23)カラムクロマトク"ラフィー, unea 非存在下 DEAE-Sephadex A-25カラムクロマトグ"ラフィー

短鎖長(3~8鎖長)のオリゴマーを分離精製するのに用いた。 TEAB buffer の直線濃度勾配により溶出した、流速は約12 ml/氏で7ラクションとして約1.2ml ずっ集めた。

• Unea 标在FDEAE-Sephadex A-25 カラムクロマトグラフ イー

鎖長が8~14のオリゴマーを分離精製するのに用いた。7M UNEA, 20mM Tris HCl (pH2,5)存在下(約50°C), NaClの 直線濃度勾配により溶出した。流速は約5ml/f. で フラクショ ンとして約0.8ml ずっ 集めた。

• Sephadex G-25, G-50 カラムクロマトク"ラフィー 脱塩の目的で使用する時に,長まが20~30cm程度のカラ

いを用い, funation後 ATPとオリゴマーとを分離する時に 15 40 m程度, Ligation後のオリゴマー周エの分離には 長き 90 m程度のカラムを用いた。

カラムにサンプルをapplyする時はできる限り容量を少なくした。溶出は50~100mM TEAB bufferを用いた。流速は約15ml/f、でフラクションとして約1mlずつ集めた。

• Sephadex G-100, Bio-Gel A(0,5m)カラムクロマトク"ラフィー 長さ 90 cm のカラムを用い, の1M TEAB buffer ご溶出した。 流速は約4ml/t. で フラクションとして約1ml ずっ集めた。

。[b-p]-ATPの調整

Glynn-Chappell 活<sup>40</sup>により,杉浦らの方法 を少し変え調 整した。反応前に[<sup>34</sup>7]-正リン酸を沸騰水浴上1時間如熱し, 副生しているポリリン酸を分解した。

反応後は Dowex カラムにapply ( 溶出 [た [>-\*p]-ATP は次に DEAE-Sephadesa A-25のカラムに吸着ませた。50mM TEAB buffer ご脱塩後 0,5M TEAB buffer こ 溶出(た。これ を滅圧乾固( TEAB を完全に除いた後, 10mM Tris HQ (pH 2,8)に溶かして凍結保存した。

・UV吸収の測定

サンプルは水溶液として日立ダブルビーム分光光度計124型, 又は200-10型で測定した。

カラムから溶出されたサンプルの測定にはALTEX Biochemical

UV-VIS Monitor を使用1 E.

。放射能測定

通常 Aloka Liquid Scintillation spectrometer LSC-671 にて 0.4% diphenyloxagole 0.01% triphenyloxagole の トルエン溶液中で測定1 た。 ただしカラムクロマトク"ラフィー のフラクションは試験管ごと Cerenkov 法により測定1た。

. .

 $\frac{1}{2} \left( \frac{1}{2} + \frac{1}{2} \right) = \frac{1}{2} \left( \frac{1}{2} + \frac{1}{2} + \frac{1}{2} \right) \left( \frac{1}{2} + \frac{1}{2} + \frac{1}{2} + \frac{1}{2} \right) \left( \frac{1}{2} + \frac{1}{2} + \frac{1}{2} + \frac{1}{2} \right) \left( \frac{1}{2} + \frac{1}{2} + \frac{1}{2} + \frac{1}{2} + \frac{1}{2} \right) \left( \frac{1}{2} + \frac{1$ 

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# 第一章の実験

· pination

反応条件, 単離収率を表Zにまとめた。Buffer は method に示した。

Substrate			[ <b>r -<sup>32</sup>p]</b> ATP	P.N. Kinase	Volume	Time	Yield
(µM)			(μΜ)	(units/ml)	(µl)	(min.)	(%)
1.	CGGGCUp	100	152	50	500	60	46
2.		100	150	100	1000	30 <b>*)</b>	19
3.		50	100	69	720	45	85
4.		23	75	47	20	60	76
6. 7. 8.	CCGAAG GUCGG AAAp UUCAAAp GUCGUCGGUUCAAAp	400 1000 101 100 80	800 1077 187 143 160	94 250 64 179 64	73 200 692 140 25	60 60 30 45 30 <b>^</b> )	60 88 100 53 88
	CCCCGCp	200	267	100	150	90	99
	AACCAp	200	240	100	50	60	93
12. 13.	UAGCUCGUCGGGCUp UCCGGCCCCCGC- AACCAp	55 68	300 - 150	300 100	40 40	50 <b>*)</b> 90 <b>*)</b>	95 93
14.	GUCGUCGGUUCAAA- UCCGGCCCCCGC- AACCAp	74	148	111	27	90 <b>*)</b>	83

[表2] 第一章 n kination

a) 酵素を加える前に 60°C,5分間 a predeat 処理を行なた。

単離の方法及び条件を次に示す。

1. DEAE-Sephadex A-25 (por7x 150m)

TEAB buffer 0.3M から 1Mまで total 200 ml os gradient で溶出した。

2. DEAE-Sephadex A-25 (\$0,9 × 18 cm) TEAB buffer 0,3 M 105 0,9 M FT total 200 ml os gradient

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で溶出した。

3. Sephadex G-25 (P1.1 × 40 cm)

4, 8, 9, 10, 11, 12, 13, 14.

Sephadex G-50 (\$0.9 × 30 cm)

5. DEAE-Sephadex A-25 (\$0.7 x 20 cm)

TEAB buffer 0.3Mから 0.9M 72" total 150ml os gradient 2" 溶出した。

6. DEAE - Sephader A-25 (\$0.8 × 16 cm)

TEAB buffer 0,3Mから 0.8H F 2" total 200ml n gradient 2" 溶出 1 T=.

7. DEAE - Cellulose (DE 23) (\$0.4×16cm)

TEAB buffer 0.1 M から 0.5 M FT" total 150ml on gradient 2" 溶出 1 TE.

· ligation

友応条件, 単離収率を表3にまとめた。Bufferはmethodに示した。RNAligaseを加える前に60°C,5分間preheatした。 単離の方法及び条件を示す。

- 1,2. 20% PAGE(disk), 定電圧400 V, ポンプ流速30ml/A で フラクションを集めた。
- 3. DEAE- Cellulose (DE 23) (\$ 0.6 × 18 cm)
  - TEAB buffer 0.3Mから1Mまで Total 200ml a gradient 2" 溶出した。
- 4. 20% PAGE (diak), 定電圧 400 V, ポンアの流速 60ml/たで フラクションを集めに。

Acceptor (µM)	Donor (µM)		ATP (µM)	RNA Ligase (units/ml)	Volume (µl)	Time (hr)	Yield (%)
1. UCGUCG 100 2. UAGC 240 3. UAGC 150 4. UAGCUCGU 100	pGGCUp pUCGUCGGGGCUp pUCGUp pCGGGCUp	167 80 100 114	333 200 200 200	250 180 100 214	120 5 200 70	1.0 2.0 3.5 2.0	4 ]] <b>a)</b> 7] 3] <b>a)</b>
5. CAUAAC 1000 6. GUC 2000 7. UUC 400 8. GUCGUCGG 150 9. CAUAAC- CCGAAG 150	pCCGAAG pGUCGG pAAAp pUUCAAAp pGUCGUCGG- UUCAAAp	100 100 200 100 100	200 200 400 200 200	100 50 100 140 115	80 1730 165 320 12	1.0 0.7 1.0 1.0 1.0	39 40 73 36 52
10. UCCGGC 24 11. UCCGGC 132 12. UCCGGC- 101	pCCCCGCp pCCCCGCp pAACCAp	10 100 116	40 200 229	50 100 114	2500 280 70	2.0 2.0 3.0	33 <b>ы</b> 65 38
CGCGGGG- 13. UGGAGC- 74 AGCCUGG	pUAGCUCGUCG- GGCUp	111	227.	535	12	2.0	37 <b>b)</b>
14. GUCGUCG- GUUCAAA 200	pUCCGGCCCCCG- CAACCAp	125	200	150	20	. 2.0	<sup>•</sup> 82
CAUAACC- 15. CGAAG 159	PGUCGUCGGUUC- AAAUCCGGCCC- CCGCAACCAp	100	200	153	17	2.0	32

[表3] 第-章 n ligation

a) kination後の単離収率

b) BAP处理後の単離收率

- 5、20% PAGE (disk), 定電圧 300 V, ホンフ 流速 40 ml/f. で 73 7 ションを集めた。
- 6. Wrea 存在下 DEAE-Sephadex A-25 (中の3×38cm) Nace 0.1Mからの5Mまで total 80mlの gradient (50mM Tris HUE (pH7.5)存在下)で"溶出してい。
- 7. DEAE-Sephadex A-25 (40,6 × 18 cm)

TEAB buffer a3Mから1Mまで total 150ml on gradient 2" 溶出1 T=0

- 8. 1) Urla 存在下 DEAE-Sephadex A-25 (中0.3 × 40cm) Nall 0.25 M から 0.65 M まで total 120 ml の gradient (50 mM Tris HUl (pH 7.5) 存在下) で溶出して。
  2) 20% PAGE (diak), 定電圧 400 V, ボンア流速 56 ml/k. で フラクションを集めた。この時の単雑収率は24%であった。
- 9. 20% PAGE(diak), 定電圧 300V, ポンプ流速 44ml/h. で 7ラクションを集めた。
- 10, 12. Sephadex G-50 (\$0.9 x 82 cm)
- 11. Urla 存在下 DEAE-Sephadex A-25 (90.3×43cm) Nall 0.2Mから 0.6M まで total 100ml a gradient (50 mM Tris Hll (pH 7.5) 存在下) で 溶出 (T=。

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- 13. 20% PAGE
- 14. Sephadex G-50 (\$ 0.9×82cm, \$ 1.0×90cm)
- 15. Bio-gel A (0.5m) (200~400 mesh) (\$0.7x91am)

# 第二章の 実験

· kination

反応条件, 単離収率を表4に示した。 P.N. kinase を加える前 に60℃ 5分間 preheat した。

Substrate · (µM)		[ <b>४-³</b> р]АТР (µМ)	P.N. Kinase (units/ml)	Volume (µl)	Time (min.)	Yield (%)
1. CCGGCCCCCGC- AACCApp	94	150	150	40	80	99
2. AUCGAAACCGGC- CCCCGCAACCAp	100	200	250	30	75	83
3. (43mer)p	100	300	100	12	45	55

[表4] 第二章の kination

単離はすべて Sephadex G-50 (中0,9×30cm)により行なった。

· ligation

反応条件, 単離収率を表5に示した。 RNA ligase を加える前に60°c 5分間 preheat 1 た。

Acceptor (µM)	Donor (µM)			RNA Ligase (units/m])		Time (hr.)	Yield (%)
	pCCGGCCCCCGC- AACCAp	100	200	100	32	2.0	87
2. CAUAACCC- GAAGGUC-148 GUCGG	PAUCGAAACCG- GCCCCCGCAA- CCAp	100	200	100	21	2.0	60
	p(43mer)p	39	94	122	16	2.0	9 <b>a</b> )

[表5] 第二章 n ligation

a) BAP处理後,単離収率

単離の方法及び条件を示す。 1、Sephadex G-50 (中の9×74cm) 2,3. Sephadex G-100 (中の7×90cm)

· E. whi tRNAf o RNase A 1: F3 限定分解20)

E, coli tRNAf Tris HCL (pH 7.5) MgCl 2 RNase A	1,000 A260 50m M 10m M 25,49/ml
•	loope
includance and o c B	or some.
	Tris HCL (pH7.5)

タンパクした(2回)、冷50mM Tris HCe (pH7.5) ご逆抽出を2回行 ない、水相をエーテルで2回洗浄した後エタノール沈殿を 行なった。沈殿をさらに EttoH: Ett20=1:1 混液で洗った後, 乾燥し 10% PAGE で 5′側半分子と 3′側半分子を単離した。

# 第三章▲実験

## · kination

反応条件,単離収率を表るに示した。

Substrate	TP[ <b>ئ-³</b> p]ATP](µM)	P.N. Kinase	Volume	Time	Yield	
(μM)		(units/ml)	(µl)	(min.)	(%)	
t to the second se	57	133	173	30	60	50
	00	157	167	30	60	70

[表 6] 第三章の timation

単離の方法を示す。

- 1. Sephadex G-50 (\$0,8 x 23cm)
- 2. Sephadex G-25 (\$ 1.0 x 26 cm)
- · ligation

反応条件, 単離収率を表7に示した。 RNA ligase を加える前に 60°C 5分間の preheat を行なった。

	Accept (µM)	Donor (µM)		RNA Ligase (units/ml)		Time (hr.)	Yield (%)
	72mer 72mer	pAACCAp pGACCA	1250 1250	212 212	16 16	2.0 2.0	16 30

[表了] 第三章 n ligation

単離はすべて 10% PAGE により行なった。

●過ヨウ素酸酸化による 3'末端一塩基除去 Springlらの方法でに基づき行なった。

nucleane S1 処理により得られた E.wittRNAft の 3'末端より 4 塩基除去された分子 (3 A260 unit)を 50mH 酢酸ナトリウム (pH 5.2), 0.8 mM NaIO4 中 (total volume 60, ul) 遮光下, 4°C 2 時間反応し 3'末端塩基の 2', 3' cio-diol を開裂(た。次に 160 客の 0.5 M ラムノースを加え 遮光下 4°C 0.5 時間反応し 週割の NaIO4 を 不活化した。 EtoH 沈殿 後, さらに沈殿 を EtoH ご 洗, た後 乾燥し, ここに 0.25 M Lys-HCl (pH9.0) を加え遮光下, 窒温(~20°C) 3.5 時間反応した。 反応後 当客の 1 M AcONa (pH 4.6) を加え EtoH 沈殿 を行びい脱塩した。沈 殿 を EtoH ご洗い乾燥した後 BAP 处理をし末端のリン酸を 餘去した。 除タンパフ後 10% PA 4 E で単離した。

生成物は [5-32p]-pCp と RNA Ligase によろ single addition (こより 3'末端にラベルを導入し、この分子を10% PAGEご単 離後、RNase T2で完全分解した。沪紙電気泳動(pH3.5) により ラベルが Azp として検出され末端のCが除去できたことを確 認した。

• アミノアシル tRNA - EF-Tu·GTP 三重複合体の Sephadex G-100 を用いた assay

EF-Tuの活性を測定した後、後に示す条件で三重複合体形成反応を行なった。#

Sephadex G-100 IT buffer (50mM Tris HQ (pH7.8), 150mM NH4(l, 10mM Mg (OAc)z, 10mM mercaptoethanol, 10mM GTP) で平衡化12かす, サンプルをapply 後 同じ buffer で溶 出した。 各溶出フラクションを 500 pl/9分で集めた後,

三重複合体形成の条件 Tris HCl (pH7.8) 50mM Mg(OAc)2 10 mM 150mM NHall mercaptoethanol 10 mM GTP. 30µM phospho (enol) piruvate 13mM pirnvate feinase 3 μg 6.3 pg (30 pmol) EF-Tu·GDP 30 pl

+ incubate at 30°C for 30 min.

+ add ["t]-Met-tRNA" on ["t]-Vel-tRNAVel on ["c]-Met-tRNA (GACCA), 5~ 10 pmol

1 incubate at 0°C for 5 min.

↓ gel filtration on Sephadex G-100 at 4°C (40.7×46cm)

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フラクションをグラスアルター にスポットし乾燥後、ト ルエン系シンテレーターで カウントを測定して。 この条件ではVal-tRNA<sup>bal</sup> は複合体を形成できない。

# 第四章の実験

## · kination

反応条件、単離収率を表8に示した。

Substrate (µM)	[ <b>४ -³</b> þ]ATP (µM)	P.N. Kinase (units/ml)	Volume (µl)	Time (min.)	Yield (%)
1. 3' Hp       67         2. 3' Hp       78         3. CAUA       150         4. CAUAAp       99         5. JICAU       100         6. ACAUp       106         7. GCAUp       100         8. CCAUp       100         9. UUCAUp       104         10. CAA       11. CAC	100 117 100 150 100 83 78 90 80	67 122 50 99 50 37 87 90 80	120 50 100 80 300 141 150 167 125	120 70 90 180 60 120 120 120	56 85 59 63 56 47 83 93 89
12. CAG 13. CAU 14. GAA 15. GAC 16. GAG 17. GAU	260	180	50	90	70 99

[表8] 第四章 a kination

単離の方法及び条件を示す。

1, 2, 4, 9. Sephadex G-50 (\$ 0.8 x 23 cm)

3. DEAE-Sephadex A-25 (Q0.5×6m)

TEAB buffer 0,2Mから0,7Mまで total 80ml a gradient 2" 溶出した。

5, 10. DEAE-Sephadex A-25 (\$0.5x6cm)

TEAB buffer 0.2Mからの7Mまで Total 60ml a gradient で 溶出した。

- 6, 11, 12, 14, 15, 16, 17. DEAE-Sephadex A-25 (\$0,5×6cm) TEAB buffer 0,2M から 0,8M まで total 60 ml os gradient で 溶出して=。
- 7,8. DEAE- Sephadex A-25 (中0,5×6cm) TEAB buffer 0,3Mから0.9Mまで Total 60ml or gradient z" 溶出して。
- 13. DEAE-Sephadex A-25 (中0.5×6cm) TEAB buffer 0.1Mから 0.6Mまで total 60mla gradient 2" 溶出して.
- · ligation

反応条件, 単離収率を表9に示した。 RNA ligase を加える前に 60°C, 5分間 preheat 1 た。

単離はすべて 10% PAGEにより行なった。

·アミノアシル化反応(Km, Vinax study) 「たに示す条件で行なった。

tRNAの3<sup>\*</sup>末端 CCA 配列を修復し た後,[<sup>14</sup>C]-Metを加えて反応した。 反応後の処理は method に記述し た方法と同一に行なった。

trna	0.15~2.0µM
СТР	0.8mM
ATP	4 mM
HEPES-NOOH (PHB.O)	100 mM
Mg(OAc)2	LOMM
Kil	10 mM
mercaptoethanol	IOMM
E. coli S-100	1 mg/ml

+ incubate at 37°C for 30min. + Reat at 100°C for 2min.

\$ incubate at 37°C for 20 min.

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Accepto (µM)	Donor (µM)		ATP (µM)	RNA Ligase (units/ml)	Volume (µl)	Time (hr.)	Yield (%)	
1. CAUA 2. 5 'H 3. 5 'H 4. 5 'H 5. 5 'HpCAUA 6. 5 'HpCAUA 7. 5 'HpUCAU	50		95 66 125 167 120 125 156	250 133 300 300 480 875 938	143 108 100 120 241 376 375	40 80 40 100 25 16 16	2.0 2.0 2.0 2.0 2.0 2.0 2.0 2.0	4 11 20 16 42 40 49
8.5'H 9.5'H 10.5'H 11.5'H 12.5'HpACAU 13.5'HpGCAU 14.5'HpCCAU	100 100 100 100 63 56	pACAUp pGCAUp pCCAUp pUUCAUp p3'Hp p3'Hp p3'Hp	125 240 200 200	500 500 500 500 1250 1250 1250	130 130 173 173 376 376 375	40 40 40 16 16 16	2.0 2.0 2.0 2.0 2.0 2.0 2.0 2.0 2.0	25 <b>a</b> ) 23 <b>a</b> ) 25 25 25 15 7 21
15. 5' HpUUCAU 16. 5'H-1 17. 5'H-1 18. 5'H-1 19. 5'H-1-pACAU 20. 5'H-1-pCCAU 21. 5'H-1-pUCAU	50 67 67 67 60 50	p3'Hp pACAUp pCCAUp pUCAUp p3'Hp p3'Hp p3'Hp		1250 670 670 670 200 200 200	375 201 201 201 344 344 344	16 30 30 30 10 10 10	2.0 2.0 2.0 2.0 2.0 2.0 2.0 2.0 2.0	30 30 25 5 17 17 21
22. 5'H 23. 5'H 24. 5'H 25. 5'H 26. 5'H 27. 5'H 28. 5'H 28. 5'H		pCAA pCAC pCAG pCAU pGAA pGAC pGAG pGAU	300 433	667	200		2.0	17 17 10 22 30 12 13 28
30. 5 ' HpCAA 31. 5 ' HpCAC 32. 5 ' HpCAG 33. 5 ' HpCAU 34. 5 ' HpGAA 35. 5 ' HpGAC 36. 5 ' HpGAG 37. 5 ' HpGAU	13	p3'Hp p3'Hp p3'Hp p3'Hp p3'Hp p3'Hp p3'Hp p3'Hp p3'Hp	3	600	300	3	2.0	

[表9] 第四章 a ligation

a) BAP処理後の単離収率

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・tRNAff の RNase A による限定分解 第二章の実験で記述したとおり行なった。

• 不儿ミル化反応49)

右に示す条件で行なった。

反応後 反応液を2等分し,一 方はそのキキクラスフィルターにスポットし 他方は等量の 50mm Cus04-0、3M Tris HQ (pH7,5)溶液を加え, 37℃ 10分間反応しグラスフィルター にスポットした。グラスフィルター 後5% TCA 溶液で10分間3回,

trna	0.7 µM
HEPES-NAOH (PH8.0)	DIM
Mg(OAc)2	10 mM
KU	10 m M
mercaptoethanol	10 m M
ATP	4 mH
CTP	1 mM
[ <sup>14</sup> C]-Met	9 µH
calcium leucovorin	0.7 mM
E. whi 5-100	Img/ul
۲.	Jone

incubate at 37°C for 30 min.

冷 FtoH-Etzo(1:1,5)液でち分間2回洗浄(, 乾燥後トルエン系シンテレーターでカウントを測定した。 CuSO4処理にまり fMet-tRNA は deacyl にされないが Met-tRNA は deacyl にされることから CuSO4処理を行なった方のカウントの行なわ なかった方のカウントに対する比をホルミルにの効率として %で表わした。

・リボソーム上でのmRNAとtRNAの相互作用

Nirenbergらの方法<sup>33)</sup>にFリ行 th 73, た。リボソームはE.woli A19 Tri 株より調整した。<sup>50)</sup> ち、末端を[b-37] KC ATPとP.N. kinaseによりラベルした <u>E.</u> tRNAを用いたの条件で反応した。

insubate at 25°C for 3	zoµl
E. whi ribosome 30 Sor 70 S	1.8µM
KÜ	50 mM
Mg(OAc)z	20 mM
Tris H(l (pH7,5)	OIM
mRNA	Z5 MM
tRNA	0.5 JuH

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mRNA としてトリマーを用いた。

反応後 冷 buffer (の1M TrisHle (pH7.5), 20mM Hg(OAL)2, 50mM K(l)を850ml 加え ミリボアフィルター ご沪週した。 同様の buffer 2ml で 3回フルターを洗浄した後 乾燥し、トルエン系シンテ レーター ごカウントを測定した。

非特異的な結合によるカウントを差し引き Endit RNAft と AU午 a 特異的結合を 100 %とし、それぞれの特異的結合を示して。

- •過ヨウ素酸酸化にFろ tRNAftの5個半分子の3'末端-塩基除去
  - 第三章の実験で記述した方法に従った。

5個半分子の他の箇所の損障をおさえるために及応前に当量の3個半分子を加えてRNAの三次構造を組ませた後反応した。 (3個半分子に10%PAGEで5個半分子と反応後分離できる。)

Xenopus laevis oocyte への microinjection<sup>36</sup>(図50) アフリカツメガ"エルより 卵巣をとり出し、Barth solution<sup>51)</sup>
(2mM Tris HCl (pH7.6)、89mM Nacl, 1mM KCl, 0.33mM Ca(NO3)2、 0.41mM Cacle2、0.82mM Hq504、2.4mM NaHCO3、0.01g/L penicilline、 0.01g/L streptomycine) 中でビンセットを用いて卵母細胞に 綺麗し、この solution 中 20°C下 injection を行けうまで保存 した。

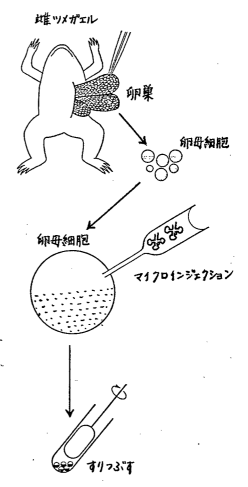
サンプルは injectionを行なう直前に 13,000 vpm, ち分間遠心し dustを除去した。(マイクロビペットがっまろのを防ぐた

ぬ)

印母細胞 E Barth solution on 入, ドシャーレに移し, 1, a サンフ・ルニフラ5個の卵母細胞 a細胞質に 50 ml t'o inject 1下。(1印母細胞みたり0.0025 purel on tRNA ze inject ( TC . ) Inject後 エッペンドルフチ ューブ(1.5ml)にち個の卵母網 胞と Barth solution を移し、20℃ でインキュベートした。

1, 3, 7, 24, 48, 72 時間 それぞ れ1271 ベート後 Barth Nolution E除き, ガラス棒ご卵 田畑胞を破砕した。Buffer (0.2 M NaOAc (pH4.5), IOMM Mg(l2, ImM EDTA, 2% SDS(W)) 200 plを直ちにかえ suspend 1 フェノール:フロロホルム= 1:1 溶液(a2M NaOAc 飽和) 200ml ご除タンパクした。

フェノール相をきほどの SDS E含ひ buffer 100ml ご 逆抽出 し、 あわせた水相にEtoH 700 pl 加え EtoH 沈殿を行なった。 沈殿を EtoH: 0.3M Na OAc (pH5,2)=4:1 溶液ご洗净1 乾燥 17-1後 10% PA4E 1= apply 1 T=.



[图50] 卵母細胞~のmicroinjection

# 第五章の実験

## · kination

# 反応条件、単離収率を表しに示した。

Substrate		۲۹[¶ <sup>ב</sup> ً¥]	P.N. Kinase	Volume	Time	Yield
(µM)		(Mµ)	(units/ml)	(µl)	(min.)	(%)
1. CAUp	250	150	75	200	90	47
2. 3'Hp	81	119	81	62	90	44
3. CGUCGGGCUCAUp	100	334	100	100	90	80

[表10] 第五章 a kination

単離の方法及び条件を示す。

1. DEAE - Sephadex A-25 (Φ 0,5 x 6 cm)

TEAB buffer 0.3Mから 0.8Mまで total 60ml or gradient で溶出した。

Z, 3. Sephadex G-50 (\$0.8 × 23 cm)

· ligation

反応条件, 単離牧率を表11 に示してー。 RNA ligaseを加え ろ前に 60°C 5分間 preheat してー。

単離の方法を示す。

1, 2, 5, 6. 10% PAGE

3, 4. Sephadex 4-50 (\$ 0.7 × 90 cm)

Acceptor (µM)	^	Donor (µM)		ATP (µM)	RNA Ligase (units/ml)		Time (hr)	
1.5'H(G1y)	100	pCAUp	200	400	100	70	2.0	7
2.5'H(G1y)-	31	p3'H(Met)p	138	1000	375	16	2.0	2
3. CGUCGGGCU	67	pCAUp	77	267	113	300	2.0	25
4. CGCGG	300	pGGUGGp	150	400	150	200	2.0	33
5. CGCGGGGU- GG	200	pCGUCGGGCU- CAUp	100	600	260	50	2.0	20
6. CGCGGGGU- GGCGUCG- GGCUCAU	75	рЗ'Нр	150	1000	340	10	2.0	3

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[表11] 第五章の ligation

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	ni Doi, Haruki Uemura, Yoshi	o Taniyama and Morio Ikehara

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#### Received 8 July 1980

#### ABSTRACT

A tetradecanucleotide U-A-G-C(U-C-G)G-G-C-Up corresponding to bases 21-34 of a nascent sequence of formylmethionyl tRNA of <u>E. coli</u> has been synthesized by the joining of two combinations of chemically synthesized oligonucleotides: 1) U-A-G-C + U-C-G-U-C-G + G-G-C-Up and 2) U-A-G-C + U-C-G-U + C-G-G-G-C-Up. In reaction 1) the extent of joining \*pG-G-C-Up to U-C-G-U-C-G was only 15.4% and the last ligation of the decamer to U-A-G-U proceeded to 27%. In reaction 2) joining between U-A-G-C and pU-C-G-Up gave a high yield (88%). The ligation of this octamer and \*pC-G-G-G-C-Up also gave a satisfactory yield (52%). These reactions suggest that sequence preferences in RNA ligase reactions may arise from the structure of the 3'-end of acceptor molecules.

#### INTRODUCTION

We have previously reported the joining of chemically synthesized tRNA fragments with RNA ligase to yield  $3^{2}$ - and  $5^{3}$ quarter molecules of <u>E. coli</u> tRNA<sup>Met</sup>. In the present paper we describe syntheses of a tetradecanucleotide U-A-G-C-U-C-G-U-C-G-G-G-C-Up which correspond to bases 21-34 of a nascent sequence of tRNA<sup>Met</sup> of <u>E. coli</u> (Fig. 1). Two different sets of synthetic oligonucleotides were used as substrates for RNA ligase to yield the tetradecanucleotide. As has been observed in previous joining reactions of synthetic ribooligonucleotides,<sup>4,5</sup>) the extent of joining differed in each case. Some structural preferences of acceptor molecules in ligase reactions were observed in these reactions.

### MATERIALS AND METHODS

### Nucleotides

All oligonucleotides used in this study have been synthe-

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Comparison of substrate base sequences for RNA ligase reactions in the synthesis of a tetradecanucleotide corresponding to bases 21-34 of E. coli tRNA $_{f}^{Met \ 1}$ 

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Received 8 July 1980

#### ABSTRACT

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### INTRODUCTION

We have previously reported the joining of chemically synthesized tRNA fragments with RNA ligase to yield  $3^{,2}$ - and  $5^{,3}$ quarter molecules of <u>E. coli</u> tRNA<sup>Met</sup><sub>f</sub>. In the present paper we describe syntheses of a tetradecanucleotide U-A-G-C-U-C-G-U-C-G-G-G-C-Up which correspond to bases 21-34 of a nascent sequence of tRNA<sup>Met</sup><sub>f</sub> of <u>E. coli</u> (Fig. 1). Two different sets of synthetic oligonucleotides were used as substrates for RNA ligase to yield the tetradecanucleotide. As has been observed in previous joining reactions of synthetic ribooligonucleotides,<sup>4,5)</sup> the extent of joining differed in each case. Some structural preferences of acceptor molecules in ligase reactions were observed in these reactions.

#### MATERIALS AND METHODS

### Nucleotides

All oligonucleotides used in this study have been synthe-

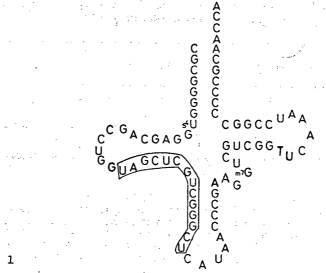


Fig. l

sized chemically." Enzymes

RNA ligase was purified as described previously.<sup>7</sup> Polynucleotide kinase was isolated from <u>E. coli</u> infected with T4 PseTIamN8SP62 which was a generous gift of Dr. A. Sugino. Other enzymes were obtained as described previously.<sup>2,4</sup> <u>Kination and Ligation</u>

Reaction conditions for 5'-phosphorylation using polynucleotide kinase and  $[\gamma^{-32}p]$ ATP were as described previously. A two fold excess of ATP with respect to donor molecules was used in RNA ligase reactions unless otherwise specified, in the presence of 50 mM Hepes-NaOH (pH 8.3), 10 mM DTT, 10 mM MgCl<sub>2</sub> 10 µg/ml BSA.

# Chromatography, Electrophoresis and Other Methods

Paper chromatography of nucleotides in nearest neighbor analysis, gel electrophoresis and paper electrophoresis for purification and identification of nucleotides were described previously.<sup>2,3</sup> Homochromatography<sup>8</sup> was carried out using Homomix I-VI<sup>9</sup> and partial digestion with nuclease Pl was as described previously.<sup>2</sup> The joined products were isolated either by ionexchange chromatography on DEAE-cellulose (DE-23, Whatman) or

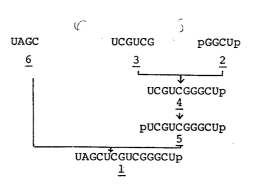
electrophoresis on 20% acrylamide gel using a disk aparatus.<sup>3</sup> Desalting was performed by gel filtration on Sephadex G-50 equilibrated with 0.05 M triethylammonium bicarbonate (pH 7.5).

Enzymatic digestion of oligonucleotides for characterization was as described previously.<sup>2,3</sup>

#### RESULTS

.Synthesis of the tetradecanucleotide U-A-G-C-U-C-G-U-C-G-G-G-G-C-Up (1) using the tetranucleotide pG-G-C-Up (2)

The oligonucleotides 2, 3 and 6 were used for the synthesis of the tetradecanucleotide 1 as shown in Chart 1. The 3',5'bisphosphorylated tetranucleotide (2) was prepared by phosphorylation of G-G-C-Up with  $[\delta^{-32}P]ATP$  and polynucleotide kinase. The 3'-phosphomonoester served as a blocking group to prevent self-polymerization in the next RNA ligase reaction. The tetramer (2) was joined to the hexamer with RNA ligase using the conditions shown in Table I. The extent of the reaction was not higher than 15% in spite of an excess of the enzyme. Alteration of temperature and other conditions did not improve the yield. The decamer 4 was isolated by a disk gel electrophoresis and phosphorylated enzymatically to give 5. The decamer (5) was then joined to the tetramer  $(\underline{6})$ . The result and reaction conditions are shown in Table I. The tetradecamer (1) was isolated by a preparative disk gel electrophoresis after treatment with polynucleotide kinase and  $[\Upsilon^{-32}P]ATP$ . The overall yield was 11%. The elution profile from the gel is shown in Fig. 2.



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Table	I	Reaction	conditions	for	joining
			00110T 0T 0110	~ U -	10

acceptor nmol [µM]	donor nnol [µM]	атр µМ	RNA ligase [unit/ml] %&	hr	Reaction extent <sup>a</sup> )	Yield %
UCGUCG ( <u>3</u> ) 12[100]	pGGCUp ( <u>2</u> ) 20[167]	333	250 (20	<b>1</b> پېژ	15	4
UAGC(6) 1.2[240]	pUCGUCGGGGCUp( <u>5</u> ) 0.4[80]	200	180 专	2	27.	11 <sup>b)</sup>
UAGC ( <u>6</u> ) 30 [150]	pUCGUp( <u>7</u> ) 20[100]	200	100	3.5 o	88	71
UAGCUCGU( <u>9</u> ) 7[100]	pCGGGCUp( <u>10</u> ) 8[114]	200	214 ໃ	2	52	31 <sup>b)</sup>

Incubation mixtures contained 10% DMSO and the temperature was 25°.

a) Estimated by homochromatography.

b) The compound was isolated after kination.

Alternative synthesis of the tetradecanucleotide (1) using the hexanucleotide pC-G-G-G-C-Up (10)

The tetradecanucleotide  $(\underline{1})$  was prepared by using a different set of synthetic oligonucleotides as shown in Chart 2. The yield

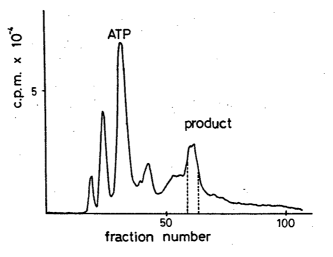


Fig. 2 Elution profile of the products obtained by joining of U-A-G-C and \*pU-C-G-U-C-G\*pG-G-C-Up, followed by phosphorylation with polynucleotide kinase and ATP, from a preparative disk gel (70 mm<sup>2</sup> x 80 mm) at 600 V with a suction rate of 32 ml/hr. The last peak contained \*pU-A-G-C\*pU-C-G-U-C-G\*pG-G-C-Up.

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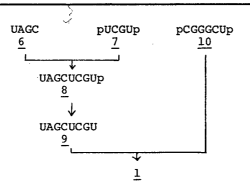


Chart 2

in joining <u>6</u> and <u>7</u> was nearly quantitative and that in joining the octamer (<u>9</u>) and the hexamer (<u>10</u>) was also satisfactory (Table I). The first joining product (<u>8</u>) was isolated by ionexchange chromatography on DEAE-cellulose as shown in Fig. 3 and the 3'-phosphate was removed by phosphatase treatment to give <u>9</u>. The nearest neighbor analysis of <u>1</u> was performed by digestion with RNase T2 followed by identification of C\*p and U\*p in acidic electrophoresis (Fig. 4). The result showed complete transfer of the 5'-phosphate to the tetramer (<u>7</u>) to the 3'-end of <u>6</u> and the 5'-phosphate of <u>10</u> to the 3'-terminal U of

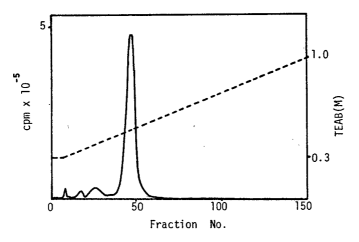


Fig. 3 Chromatography of the joined product, U-A-G-C\*pU-C-G-Up on a column (0.6 x 18 cm) of DEAE-cellulose (bicarbonate) equilibrated with 0.15 M triethylammonium bicarbonate. Elution was performed with a linear gradient of triethylammonium bicarbonate (0.3-1 M, total 200 ml). The main peak contained the product.

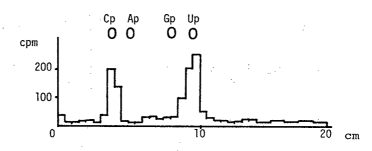
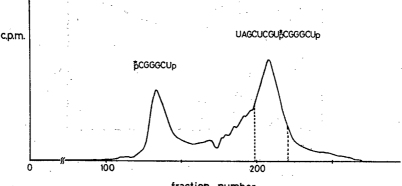


Fig. 4 Nearest neighbor analysis of the joined product U-A-G-C-\*pU-C-U-C-G-G-G-C-Up with RNase T2. The digested mixture was electrophoresed at pH 3.5 with nucleotide UV markers.

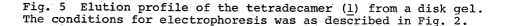
<u>9</u>. The isolation of the joined product (<u>1</u>) in the reaction in the reaction between <u>9</u> and <u>10</u> was performed using preparative disk gel electrophoresis. The elution profile is shown in Fig. 5. The tetradecamer (<u>1</u>) was characterized by slab gel electrophoresis for comparison of chain length.

#### DISCUSSION

As shown in Table I there were large differences in yields between the reactions shown in Chart 1 and 2. This may indicate that ribooligonucleotide fragments used as substrates for RNA ligase in the synthesis of larger RNA fragments should be pre-







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pared according to the preferences in recognition between the enzyme and oligonucleotides. It was found that the 3'-hydroxy groups of oligoadenylates were much better acceptor substrates<sup>4</sup> than those of oligouridylates and various trimers showed different extents of joining when they were treated with the same 5'phosphorylated trinucleotide donors in RNA ligase reactions.<sup>5</sup> In those trimer reactions C-C-G was the poorest acceptor when \*pC-C-A was used as a donor. The trinucleotide \*pC-C-A itself was also a poor acceptor molecule, since no polymerization occurred in the conditions used and C-C-A was joined to \*pC-C-A in a yield of 38% which was about one half the efficiency compared to C-A-A or C-C-C. Thus the pyrimidine-purine sequences at the 3'-termini effect the efficiency of reactions with 5'phosphorylated oligonucleotides. This may be due to unstacked structures at the 3'-ends of acceptors when a 3'-linked pyrimidine nucleoside is present next to a 5'-linked purine nucleoside (e.g. C-G). In a right-handed helical strand the pyrimidine ring of the 3'-linked pyrimidine nucleoside is overlapped with the imidazole ring of the 5'-linked purine nucleoside. This overlapping is less favorable the comparable overlapping between pyrimidine rings, which may result from purine-pyrimidine sequences (e.g. G-C). Guanosine is known to adopt the syn conformation to a significant extent<sup>10</sup> and this conformation may be unfavorable for RNA ligase reactions if it exist at the 3'-terminus of acceptors. However oligonucleotides having the sequence G-G at the 3'-terminus have been good acceptor molecules in the RNA ligase reactions so far tested. Thus purine-purine sequence seems to be function well on acceptor molecules. Oligonucleotides bearing C-G at the 3'-end, e.g. C-G-C-G, <sup>3</sup> C-C-C-C-G, <sup>2</sup> have been found to be poor acceptors in the synthesis of  $tRNA_{f}^{Met}$  quarter molecules. It has also been reported previously that in singlestranded oligonucleotides guanine has a tendency for unstacking especially when adjacent to uridine residues probably due to the formation of syn and anti comformers.<sup>11</sup>

### ACKNOWLEDGEMENTS

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Modification of the anticodon triplet of *E.coli* tRNA $_{1}^{Met}$  by replacement with trimers complementary to non-sense codons UAG and UAA

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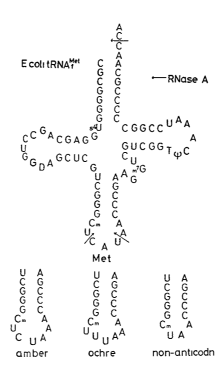
### ABSTRACT

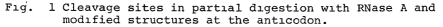
E. coli tRNA<sup>Met</sup> was hydrolyzed with RNase A using a limited amount of the enzyme to give two half molecules lacking the anticodon trimer and 3'-terminal dimer. Chemically synthesized trimers CUAp and UUAp were joined to the 5'-half molecules by phosphorylation with polynucleotide kinase plus ATP followed by treatment with RNA ligase. These modified tRNA species had anticodons complementary to the termination codons UAG and UAA. Two half fragments were joined by a similar procedure to yield a molecule lacking the anticodon trimer and the 3'-dimer. Methionine acceptor activity of these tRNA was tested under conditions in which the CAU inserted control  $tRNA_f^{Met}$  accepted methion It was found that all three modified molecules were accepted methionine. not The recognized by the methionyl-tRNA synthetase from E.coli. other sixteen amino acids were not incorporated with partially purified aminoacyl-tRNA synthetases.

## INTRODUCTION

We have previously reported enzymatic joining of chemically synthesized ribooligonucleotides with T4 RNA ligase to yield the <u>E.coli</u> formylmethionine tRNA.<sup>2</sup> The basic methodology used in the synthesis of RNA of this size can be applied to the synthesis of modified tRNAs which are useful for studies on structure-function relationship of tRNAs. Anticodon triplets in tRNAs play an essential role in decoding messenger RNAs and suppression of nonsense mutations is known to occur by the action of aminoacyl-tRNAs having anticodon triplets complementary to non-sense codons.<sup>3</sup> Synthesis of tRNAs with anticodons complementary to nonsense codons is of interest in testing properties in decoding systems. Replacement of the anticodon of yeast phenylalanine tRNA was reported by Uhlenbeck and his coworkers by removal of Y base and partial RNase A digestion followed by ligation of a new oligonucleotide.<sup>4a</sup> They found a

sequence-specific contact between the anticodon loop of yeast tRNA<sup>Phe</sup> and the phenyl alanyl-tRNA synthetase.<sup>4b</sup> Recognition of tRNA by aminoacyl-tRNA synthetases seems to differ in each amino acid and cognate tRNA. Modification of the anticodon loop does not affect aminoacylation of tRNA in certain cases.<sup>6</sup> The wobble position of E.coli tRNA<sup>Met</sup> has been converted to uridine and this modification seemed to affect the aminoacylation.<sup>5a</sup> Replacement of the anticodon of <u>E.coli</u> formylmethionine tRNA in the present study by joining of RNase A fragments of <u>E.coli</u> tRNA<sup>Met</sup> with CUA or UUA abolishes the recognition with the methionyl-tRNA synthetase of <u>E.coli</u>. (Fig. 1). Removal of the anticodon triplet also caused no-charging. Recently replacement of the same tRNA was reported.<sup>5b</sup>





## MATERIALS AND METHODS

## Materials

<u>Nucleotides</u>. Trinucleotides CAU, CUA and UUA were synthesized either by the phosphodi- or triester methods.<sup>2</sup> <u>E.coli</u> tRNA<sup>Met</sup> was a gift of Dr. D. Söll and had a specific acceptor activity of 1.4 nmol/A<sub>260</sub> unit.

Enzymes. T4 RNA ligase was purified as described.<sup>7</sup> Polynucleotide kinase and <u>E.coli</u> alkaline phosphatase were obtained from Takara Shuzo Co.. 3'-Phosphatase-free kinase was isolated from T4 Pse T1-amN82SP62 as described.<sup>8</sup> RNase T1, RNase T2 and Nuclease S1 were obtained from Sankyo Co. Nuclease P1 was obtained from Yamasa Shoyu Co.. Other enzymes for characterization of products were obtained as described.<sup>8-12</sup>

Amino acids. <sup>14</sup>C L-Amino acids were purchased from Amersham International plc: Specific activities were Ala, 285 Ci/mol; Val, 285; Leu, 150; Ile, 150; Phe, 225; Pro, 125; Gly, 50; Ser, 165; Thr, 100; Tyr, 225; Asn, 100; Lys, 340; His, 150; Arg, 150; Asp, 225; Glu, 255. Methods

Phosphorylation, ligation and dephosphorylation were performed as described previously. <sup>8-12</sup> Complete digestion by nucleases was performed in the presence of carrier RNA (yeast RNA, sigma Co. Type IV) in 10 µl solution. RNAase A (1 µg), carrier RNA (0.3  $A_{260}$  unit) in 50 mM Tris-HCl (pH 7.5) and EDTA 1 mM at 37° for 1 hr; RNase T2 (2U), carrier RNA (0.2  $A_{260}$ ) 50 mM Tris-HCl (pH 7.5) and EDTA 1 mM at 37° for 20 min, nuclease Pl (0.2 µg), carrier RNA  $IA_{260}$  in 50 mM ammonium acetate (pH 5.0) at 37° for 1 hr; RNase T2 (2U) carrier RNA (0.8  $A_{260}$ ) in 50 mM sodium acetate (pH 4.5) at 37° for 30 min. Oligonucleotides recovered after homochromatography contained ca. 200 µg of RNA/ cm<sup>2</sup>.

Partial digestion of tRNA (100  $A_{260}$ ) was performed in 50 mM Tris-HCl(pH 7.5)-20 mM MgCl<sub>2</sub> at 0° for 30 min using RNase A (25 µg). The enzyme was removed by extraction with phenol and nucleotides were precipitated with ethanol. The precipitate was subjected to acrylamide gel electrophoresis. Two bands were eluted from gel and desalted by gel filtration on Sephadex G-50. The yields was ca. 20%.

Aminoacylation of tRNA was performed in 50 mM HEPES-NaOH (pH 8.0), 10mM magnesium acetate, 10 mM potasium chloride 10 mM  $\beta$ -mercaptoethanol, 4mM ATP, 0.5 mM CTP, 9  $\mu$ M L- U-<sup>14</sup>C Met, 0.5  $\mu$ M tRNA and E.coli S-100<sup>13</sup> at 37°C for 40 min in 20  $\mu$ l and counted as described previously.<sup>2</sup>

## RESULT AND DISCUSSION

Removal of the anticodon triplet of tRNA<sub>f</sub><sup>Met</sup> with RNase A

A single strand specific nuclease, nuclease S1, has been used to hydrolyze anticodon loops of tRNAs.<sup>14</sup> Later it was reported by Wrede et al. that  $\underline{\text{E.coli}} \text{tRNA}_{f}^{\text{Met}}$  was hydrolyzed at the unique position presumably because of a different conformation at the anticodon region.<sup>15</sup> We found that digestion of E.coli  $tRNA_{f}^{Met}$  with nuclease Sl gave heterogeneous products. On the other hand partial digestion of the same tRNA with a limited amount of RNase A occurred at specific points. Two large fragments with chain length 34 and 38 were isolated by 20% acrylamide gel electrophoresis in a yield of 20%. The 5'-terminal sequence of these oligonucleotides was determined by the mobility shift analysis as shown in Fig. 2. The 5'-ends of the 5'- and 3'-halves were found to have cytidine and adenosine, respectively. The 3'-end of the 5'-half molecule was analyzed by two dimensional thin layer chromatography of the complete digest after labeling with 5'-32P pCp and RNA ligase. It was found to be CmU\*p. The 3'-terminus of the 3'-half was analyzed after the 3'-labeling as shown in Fig. 3, which indicate a loss of two nucleotides from the terminus.

Construction of  $tRNA_f^{Met}$  with modification at the anticodon

Two ribotrinucleotides pCUAp and pUUAp which were complementary to nonsense codons UAG and UAA were inserted between the 5'and 3'-half molecules obtained above. The methionine anticodon pCAUp was also inserted to obtain a control molecule. The 3', and 5'- halves were also joined to yield a tRNA<sup>Met</sup><sub>f</sub> analog lacking the anticodon triplet.

A typical example of ligation is shown in Fig. 4. The 3',5'-diphosphorylated trimer pCAUp was joined to the dephosphorylated 5'-half molecule and the product (37 nucleotides) was isolated by 20% gel electrophoresis after treatement with phos-

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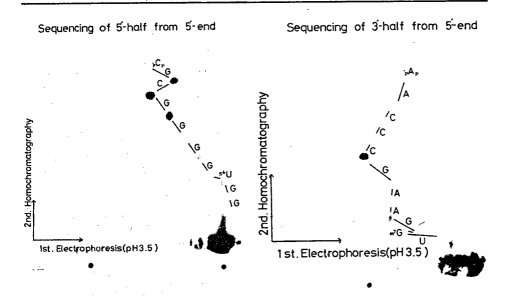


Fig. 2 Mobility shift analyses of the 5'-half (a) and the 3'half (b) by labeling at the 5'-end.

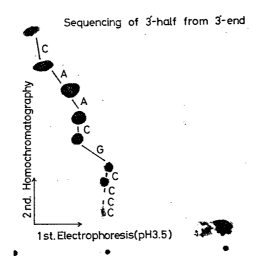
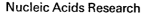


Fig. 3 The sequence of the 3'-end of 3'-half (38-75) developed after labeling at the 3'-end with 5'- <sup>32</sup> p pCp and RNA ligase.



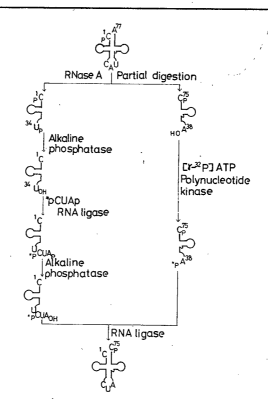
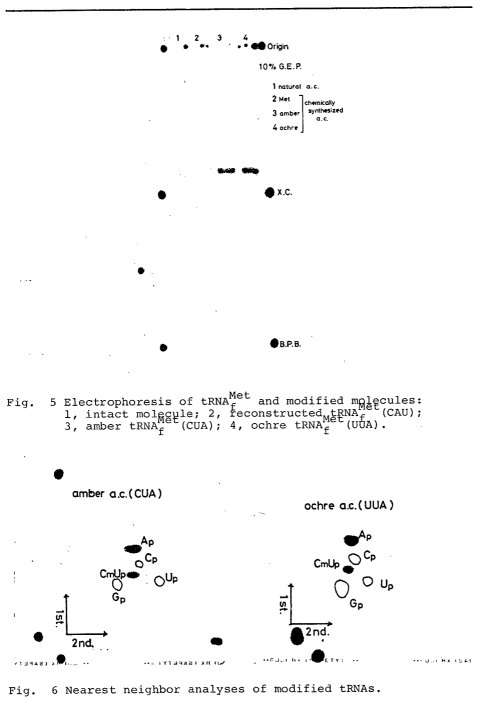


Fig. 4 Scheme for reconstruction of  $tRNA_f^{Met}$ .

phatase. The 3'-half was 5'-phosphorylated with polynucleotide kinase and isolated by 20% gel electrophoresis. These two fragments were joined and treated with phosphatase. The molecule lacking CA at the 3'-position (75 nucleotides long) was purified by 20% gel electrophoresis and identified by nearest neighbor analysis.

The amber suppressor tRNA containing CUA anticodon and ocher suppressor tRNA (UUA anticodon) were constructed by the same procedure. Electrophoretic mobilities on 10% acrylamide gel of these molecules are shown in Fig. 5 and the result of nearest neighbor analys by digestion with RNase T2 is shown in Fig. 6. The tRNA<sup>Met</sup> lacking the anticodon CAU (72 nucleotides) was also isolated by electrophoresis on 20% acrylamide gel. Aminoacylation of Modified tRNA<sup>Met</sup>

Incorporation of methionine to these tRNA was tested using



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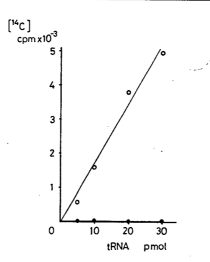


Fig. 7 Charging of methionine to the intact  $tRNA_{f}^{Met}$  (o--o) and the amber  $tRNA_{f}^{Met}$  (CAU) (o--o).

partially purified <u>E.coli</u> aminoacyl tRNA synthetase containing ATp(CTD) tRNA nucleotidyl transferase to repair the 3'-terminal CA. The 3'-terminus was analyzed by labeling with  $5'-{}^{32}P$  pCp and RNA ligase.<sup>15</sup> It was found that more than 95% count was transfered to pA. As summarized in Table I, the reconstructed

Table I <sup>14</sup>C Met-acceptor activity of modified tRNAs

trna	pmol <sup>14</sup> C Met charged	% relative to reconstituted
	per 10 pmol	tRNA <sup>Met</sup>
tRNA <sup>Met</sup> f	4.1	
reconstituted (CAU)	2.3	100
amber (CUA)	0.04	2
ochre (UUA)	0.14	6
non-anticodon	0.06	3
'3'(38-75) and 5'(1-34)-halves	0.08	3
3'-half (38-75)	0.09	4

 $tRNA_{f}^{Met} \{tRNA(CAU)\}$  was charged 55% with respect to the natural The tRNA containing CUA anticodon {amber tRNA(CUA)}, molecule. ocher tRNA (UUA), non-anticodon tRNA a mixture of two halves and the 3'-half fragment were not aminoacylated by the synthetase preparation. An example of charging experiment using increasing amounts of amber tRNA(CUA) is shown in Fig. 7 together with the natural tRNA<sup>Met</sup> control. This result indicated that the E.coli methionyl-tRNA synthetase did not recognize  $tRNA_{f}^{Met}$  modified at the anticodon loop. We have observed previously that replacement of the fourth nucleotide from the 3'-end did not affect the charging by the E.coli methionyl-tRNA synthetase significantly.<sup>17</sup> The enzyme may recognize the shape of the anticodon region more restrictively than that of the amino acid acceptor stem part.

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Replacement and insertion of nucleotides at the anticodon loop of *E. coli* tRNA  $_{f}^{Met}$  by ligation of chemically synthesized ribooligonucleotides

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## ABSTRACT

Insertion of the four major nucleotides at the 5'-side of the anticodon triplet of E. coli tRNA<sup>ME</sup> was performed by joining of the half molecules obtained by limited digestion with RNase A and the chemically synthesized tetranucleotide pN-C-A-U using RNA ligase. Insertion of U-U at the 5'-side or A and A-A at the 3'-side of the anticodon were also performed using U-U-C-A-U, C-A-U-A and C-A-U-A-A. The constant U next to the 5'-side of the anticodon was replaced with A and C by ligation of A-C-A-U and C-C-A-U to the 5'-half molecule which had been treated with periodate plus lysine, followed by joining to the 3'-half. These modified tRNAs were tested for their ability to accept methionine with the methionyl-tRNA synthetase of E. coli. The affinity of these analogs for the synthetase decreased more extensively when the insertion was at the 3'-side of the anticodon triplet. Insertion of mononucleotides at the 5'-side or replacement of the constant U next to the 5'-side of the anticodon did not affect aminoacylation drastically. This may mean that the 3'-side of the anticodon loop of tRNA is one of the major recognition sites for the methionyl-tRNA synthetase.

## INTRODUCTION

Recognition of tRNA by aminoacyl-tRNA synthetases is a particularly good example of the specific interaction of proteins with nucleic acids. By using tRNA analogs with specific modifications at certain regions, information on the recognition of tRNA by the synthetase should be obtainable. We have previously reported that the anticodon triplet of <u>E</u>. <u>coli</u> tRNA<sup>Met</sup><sub>f</sub> could be replaced by ligation of chemically synthesized trimers to partial RNase A digestion products of the tRNA and that the C-A-U sequence of the anticodon was indispensable for recognition by <u>E</u>. <u>coli</u> methionyl-tRNA synthetase.<sup>1)</sup> This methodology using RNA oligomers and T4

RNA ligase to obtain tRNA analogues is very useful for studies on structure-function relationships. Since the discovery of T4 RNA ligase, this methodology has been used in various studies.<sup>2-19</sup>)  $\langle \cdot \rangle$ 

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The anticodon is one of the most important regions in the interaction of tRNA with mRNA during protein synthesis. There are more than 20 species of tRNAs each with their own anticodon which can form hydrogen bonds with complementary codons in mRNA. Furthermore, the 5'-side of the anticodon is almost always occupied by uridine and the 3'-side is a purine or a purine derivative. It is not clear why the residue on the 5'-side of the anticodon should be uridine. In the X-ray crystal structures of tRNAs for yeast tRNA<sup>Phe 20,21</sup> and <u>E</u>. <u>coli</u> tRNA<sup>ff</sup>, the 5'-side of the anticodon consists of the so called "U-turn" structure. Recently Uhlenbeck et al. showed that this uridine in yeast tRNA<sup>Tyr</sup> was not essential for amber suppression.<sup>13</sup>

In this paper, we describe the synthesis of several mutant  $\underline{E}$ .  $\underline{coli}$  tRNA<sup>Met</sup><sub>f</sub>'s having additional bases at the 3'- or 5'- side of the anticodon, and investigate whether these additional bases have an influence on the methionine acceptor activity. In addition we have changed the constant U to the 5'-side of the anticodon in order to ascertain the importance of this structural requirement for aminoacylation of the tRNA.

## MATERIALS AND METHODS

## Materials

Trinucleotides; A-U-G, U-A-U, U-G-A, U-U-A, tetranucleotides; C-A-U-A, A-C-A-U, C-C-A-U, G-C-A-U, U-C-A-U, pentanucleotides; C-A-U-A-A, U-U-C-A-U, were prepared by the triester method.<sup>3)</sup>  $I\gamma^{-32}P$  JATP (specific activity; 2,900 Ci/mmol) and L-  $[U^{-14}C]$  Methionine (specific activity; 282 mCi/mmol) were obtained from NEN and Amersham, respectively. E. <u>coli</u> tRNA<sup>Met</sup><sub>f</sub> was a gift of Dr. D. Söll. (specific activity; 1,400 pmol/A<sub>260</sub> unit) Folinic Acid-SF as a formyl-donor was obtained from Lederle. T4 RNA ligase was purified from <u>E</u>. <u>coli</u> A19 strain infected with T4 phage as described.<sup>23)</sup> Polynucleotide kinase and alkaline phosphatase from <u>E. coli</u> Al9 were obtained from Takara Shuzo Co., RNase A, RNase  $T_2$  and Nuclease  $P_1$  were obtained from Boehringer Co., Sankyo Co. and Yamasa Shoyu Co., respectively. Crude <u>E</u>. <u>coli</u> aminoacyl-tRNA synthetase was prepared as described by Nishimura et al.<sup>35)</sup> <u>E. coli</u> ribosomes were prepared as described by Nishizuka et al.<sup>24)</sup>

## Methods

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Phosphorylation, dephosphorylation and ligation were performed as described previously.<sup>25-29)</sup> Phosphorylated tetra and pentanucleotides corresponding to the anticodon were purified by DEAE Sephadex A-25 ion exchange column chromatography and Shephadex G-50 gel filtration, respectively. 5'- and 3'-half molecules of <u>E. coli</u> tRNA<sup>Met</sup><sub>f</sub> were prepared by partial digestion with RNase A followed by purification by 10 % polyacrylamide gel electrophoresis (PAGE), and sequenced by mobility shift analysis as described previously.<sup>1)</sup> After RNA ligase reactions, the products were purified by 10% PAGE and the binding sites between two RNA molecules were identified by nearest neighbor analysis.

Removal of one base from the 3'-end of the 5'-half using periodate plus lysine was performed as described by Sprinzl et al.<sup>30)</sup> We added an equal amount of 3'-half to the 5'-half in the reaction mixture to form a complex with the secondary structure of tRNA<sup>Met</sup><sub>f</sub>. After the reaction, the 5'-half lacking one base at the 3'-end was purified by 10% PAGE.

Aminoacylation of tRNA was performed as described previously<sup>1)</sup> except for measurement of the Km. Km studies were carried out by two step reactions; restoring the C-C-A end and aminoacylation. The C-C-A restoring reaction on the synthesized tRNA was performed in 100mM HEPES-NaOH (pH 8.0), 10 mM magnesium acetate, 10 mM potasium chloride,10 mM  $\beta$ -mercaptoethanol, 4 mM ATP, 0.8 mM CTP, 0.15-2.0  $\mu$ M tRNA and 1 mg/ ml <u>E. coli</u> S-100 at 37°C for 30 min in 10  $\mu$ l. After the C-C-A restoring reaction the mixture was heated at 100°C for 2 min and slowly cooled to room temperature. Aminoacylation was started by adding 200 pmol of L-[U-<sup>14</sup>C] Met and 2  $\mu$ g of <u>E. coli</u> S-100. This reaction mixture was incubated at 37°C for 20 min and quantified as described previously.<sup>3)</sup>

Formylation was performed in aminoacylation buffer containing 0.7 mM Folinic Acid-SF at 37°C for 30 min in 30  $\mu$ l. After the reaction one half of the reaction mixture was spotted onto a glassfilter disk and to the other half was added the same volume of 50 mM CuSO<sub>4</sub>-0.3 M Tris-HCl (pH 7.5). Incubation was continued at 37°C for 10 min<sup>31)</sup> and this mixture was also spotted on another glassfilter disk. These disks were washed three times with cold 5% trichloroacetic acid for 10 min, twice with cold ethanol-ether (1:1, v/v) for 5 min and dried and counted.

A binding study involving the codon-anticodon interaction on <u>E</u>. <u>coli</u> ribosomes was performed as described by Nirenberg et al.<sup>32)</sup> except that  $5'-^{32}P$  labelled tRNAs were used instead of aminoacyl-tRNAs.<sup>33)</sup>

## RESULTS

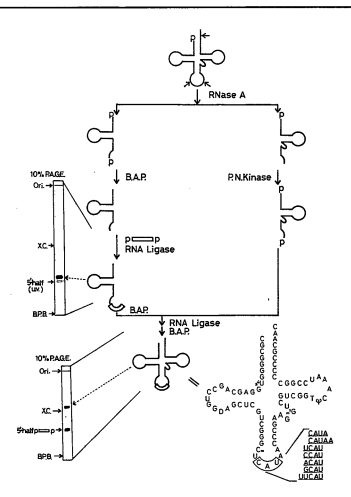
## Synthesis of modified tRNAs with extra bases at either side of the anticodon triplet

We synthesized several <u>E</u>. <u>coli</u>  $tRNA_f^{Met}$  derivatives having additional bases either side of the anticodon as shown in Fig. 1. We removed the anticodon trimer of <u>E</u>. <u>coli</u>  $tRNA_f^{Met}$  by partial digestion with RNase A. In these conditions we could obtain a 5'-half molecule (bases 1-33) and a 3'-half molecule (bases 37-75) as major products. These halves were purified on 10% PAGE. Both halves were sequenced as described previously.<sup>1)</sup> The 3'-end phosphate of the 5'-half was removed by treatment with bacterial alkaline phosphatase after opening the 2',3'-cyclic phosphate.

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Tetra or pentaribooligonucleotides containing the anticodon sequence were synthesized by the triester method. Three kinds of RNAs; the 5'-half, an anticodon sequence and the 3'-half, were joined with T4 RNA ligase to reconstruct the whole structure of a tRNA. Firstly, phosphorylated anticodon-oligomers were ligated to the 5'-half molecules and the products were isolated on 10% PAGE. After



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Fig. 1 Scheme for syntheses of modified <u>E</u>. <u>coli</u> initiator tRNAs and the pattern of 10% PAGE in each ligation step. Open square shows the synthesized tetra- or pentaribooligomers corresponding to the anticodon sequence. These autoradiographs of 10% PAGE show the case in which C-C-A-U is the anticodon sequence. In the cases of other ribooligomers, all the PAGE patterns<sub>3</sub>were the same as that of C-C-A-U. The asterisk denotes <sup>32</sup>P-phosphate.

dephosphorylation, 5'-half plus anticodon-oligomer molecules were ligated to the phosphorylated 3'-half molecules and the final product was isolated on 10% PAGE. After each step of ligation, we carried out nearest neighbor analysis to confirm that the binding site between the donor and the acceptor was

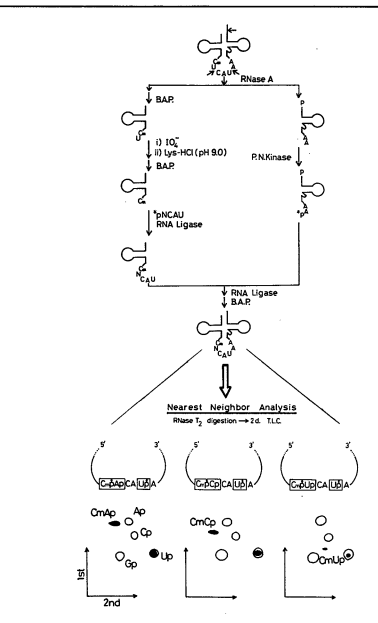
not incorrect. By this strategy we obtained seven kinds of tRNA which have anticodon sequences of A-C-A-U, C-C-A-U, G-C-A-U, U-C-A-U, U-U-C-A-U and C-A-U-A-A instead of C-A-U. The yield of ligation between 5'- and 3'-halves was about 50% after isolation on 10% PAGE. This was relatively satisfactory compared to the yield for ligation of the 5'-half molecule and anticodon sequences (20-25 %). Replacement of constant U

The 5'-half molecule which was generated by digestion of E. coli tRNA<sup>Met</sup> with limited amounts of RNase A was treated with phosphatase to remove the 3'-phosphate and with NaIOA at pH 5.2. The 3'-uridine was removed by  $\beta$ -elimination with lysine. The 3'-terminal analysis showed that the U residue was removed and that the Cm residue to the 5'-side of U was the new 3'-terminal base. Three tetramers; U-C-A-U, C-C-A-U and A-C-A-U, were phosphorylated with polynucleotide kinase and ATP, and joined to the 5'-half lacking the 3'-terminal U. These 5'-half molecules plus tetramer and the phosphorylated 3'-half molecule were joined and the products were purified on 10% PAGE. Fig. 2 shows the scheme for the replacement of constant U and nearest neighbor analysis of these products.  ${\rm Cm}^{32}{\rm pNp}$  from the first step of the ligation and  ${\rm U}^{32}{\rm p}$  from the second step were detected on two dimensional TLC.<sup>34)</sup> These experiments also showed that the 3'-side of Cm was changed to the planned base as judged by the position of Cm<sup>32</sup>pNp on TLC.

## Aminoacylation of synthesized tRNAs

Synthesized tRNAs which have extra bases in the anticodon loop and another base at the 5'-side of the anticodon instead of constant U were investigated for their methionine acceptor activity using partially purified <u>E</u>. <u>coli</u> aminoacyltRNA synthetase. This crude enzyme contained ATP(CTP)tRNA nucleotidyl transferase to repair the 3'-terminal C-A sequence of the synthesized tRNAs and in fact this repair was almost complete as described previously.<sup>1)</sup> All tRNAs having extra bases in the anticodon loop accepted methionine. Kinetic studies showed that the addition of two bases beside the anticodon increased the Km value of these tRNAs for the

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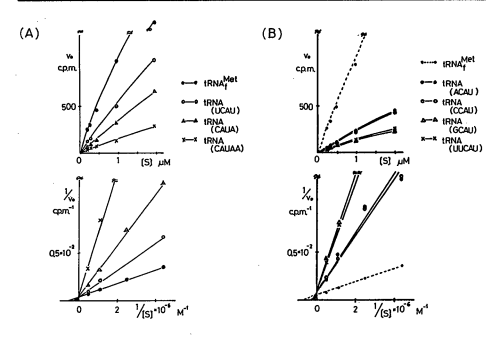
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Fig. 2 Scheme for the replacement of constant U and nearest neighbor analyses of these products. The asterisk denotes  $^{32}P$ -phosphate. N; A, C, U.



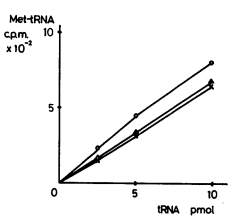
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Fig. 3 Aminoacylation of synthesized tRNAs. tRNA(UCAU), tRNA(CAUA), tRNA(CAUAA), tRNA(ACAU), tRNA(CCAU), tRNA(GCAU) and tRNA(UUCAU) indicate the tRNAs whose anticodon sequences are U-C-A-U, C-A-U-A, C-A-U-A-A, A-C-A-U, C-C-A-U, G-C-A-U and U-U-C-A-U, respectively. Lower graphs show the Lineweaver-Burk plots.

methionyl tRNA synthetase considerably; the Km value was 12.5  $\mu$ M (the Km value of natural tRNA<sup>Met</sup> was 1.7  $\mu$ M). In cases where one base was added, the addition of U at the 5'-side of the anticodon preserved the high affinity for the synthetase; the Km value was 3.3 µM. On the other hand, A at the 3'-side increased the Km value; the Km value was 6.7 µM. (Shown in The addition of A, C or G at the 5'-side of the Fig. 3.A) anticodon also increased the Km value. (Shown in Fig. 3.B) The synthesized tRNAs replacing the constant U with A or C at the 5'-side of the anticodon were also tested for their methionine acceptor activity. (Fig. 4) Both tRNAs had the same level of methionine acceptor activity as the control  $tRNA_{f}^{Met}$ , whose constant U was first removed by periodate treatment then replaced. However this control tRNA<sub>f</sub><sup>Met</sup> did not attain the same level of activity as natural E. coli

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Fig. 4 Methionine acceptor activity of tRNAs with changed constant U. o-o; control tRNA<sup>f</sup><sub>f</sub>,  $\Delta$ - $\Delta$ ; tRNA(C-33) meaning that the constant U is changed to C, x-x; tRNA(A-33).

tRNA<sub>f</sub>Met This may be due to periodate treatment of the 5'-half molecule. Initially we carried out the periodate treatment of the 5'-half molecule without adding the 3'-half reconstructed tRNAs using these periodate treated 5'-halves had little activity as methionine acceptors. Therefore for subsequent periodate treatments we added 3'-half molecules to the reaction mixture in order to protect the 5'-half by forming a secondary structure as in tRNA. Following this procedure, the methionine acceptor activity of reconstructed tRNAs increased but did not reach the natural level. It seems that not all 5'-half molecules were protected by secondary structure formation as in  $tRNA_f^{Met}$  during the periodate treatment and that some bases were damaged to decrease the methionine acceptor activity.

The interaction on ribosomes between triribonucleotides and the anticodons of  $t_{\rm RNA}^{\rm Met}_{\underline{f}}$  sequences having larger anticodon loop

We examined the interaction between ribotrinucleotides and some synthesized tRNAs with larger anticodon loops on <u>E</u>. <u>coli</u> ribosomes to investigate the structural role of the anticodon. Using the chemically synthesized ribooligomers, A-U-G, U-G-A, U-U-A, or U-A-U as mRNA, we measured the

Table I							
	tRNAf pmol(%)	tRNA(CAUA) pmol(%)	tRNA(CAUAA) pmol(%)	tRNA(UCAU) pmol(%)			
-mRNA AUG UGA UAU UUA	0.45 0.95(100) 0.46(~0) _	0.50 0.49( 0) 0.50( 0) 0.45( 0) 0.49( 0)	0.42 0.41(0) 0.38(0) 0.39(0) 0.43(~0)	0.36 0.51(33) 0.37(0) 0.35(0) 0.33(0)			

Codon-anticodon interactions on E. coli ribosomes Each  $^{32}$ P-labeled tRNA (10 pmol) was added to the reaction mixture containing mRNA and E. coli 70S ribosomes. -mRNA indicates the condition of no mRNA in this reaction mixture. % when the specific binding of natural tRNA<sub>f</sub> to A-U-G is 100%.

specific binding of tRNAs whose anticodon sequences were C-A-U-A, C-A-U-A-A, U-C-A-U instead of C-A-U to mRNA on <u>E</u>. <u>coli</u> ribosomes. Table 1 shows that only  $tRNA_f^{Met}$  having U-C-A-U at the anticodon; tRNA(UCAU), binds specifically to A-U-G and that this binding was 33% with respect to the binding of natural  $tRNA_f^{Met}$  to A-U-G. This result showed that the structure of the anticodon trimer in tRNA(UCAU) is similar to that in natural  $tRNA_f^{Met}$  notwithstanding the insertion of U at the 5'-side of the anticodon. It is presumed that three bases of the anticodon in tRNA(UCAU)stand in line outside the phosphate backbone of the loop. Formylation of synthesized  $tRNA_f^{Met}$  species

Formylation is an event specific to the initiator tRNA in prokaryotes. The part of the initiator tRNA recognized by the transformylase is unknown. Three tRNAs having enlarged anticodon loop were tested for formylation by adding the formyl donor to the aminoacylation system. After the reaction, the mixture was divided to two parts. One part was treated with 50 mM  $CuSO_4-0.3$  M Tris HCl (pH 7.5) in order to deacylate non-formylated methionyl-tRNA, and then spotted on a glass filter. The other was spotted on a glass filter without treatment. The ratio of  $CuSO_4$  treated sample radioactivity to non-treated sample radioactivity was defined as the formylation ratio. The percentages for tRNA(CAUA), tRNA(CAUAA) and tRNA(UCAU) were 88%, 98% and 93%,

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respectively. All these values were considered to represent quantitative formylation within the limits of experimental error. These values suggest that the transformylase does not interact with the anticodon area of  $tRNA_{e}^{Met}$ .

## DISCUSSION

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We have synthesized seven kinds of tRNAs which have larger anticodon loops. In the synthetic procedure, we first tried to join the anticodon sequences to 3'-half molecules but the yields in ligation were too low for the expected product to be obtained. This may be due to the bulkyness of the 3'-half as a donor in the RNA ligase reaction. In cases of ligation of the 5'-halves containing the anticodon sequence to the 3'-half, we could obtain relatively high yields. In this reaction, the 5'-half and 3'-half molecules form tRNA secondary structure and therefore the 3'-end of the 5'-half is located close to the 5'-end of the 3'-half. Essentially this ligation between the two halves behaves as an intramolecular joining reaction across the anticodon.

Modified tRNAs having large anticodon loops were previously shown to accept methionine.<sup>16)</sup> We have further modified the structure around the anticodon of <u>E</u>. <u>coli</u>  $tRNA_f^{Met}$ . The additional U at the 5'-side of the anticodon does not change the conformation of the anticodon loop more than the additional A at the 3'-side of the anticodon. Even if U was added to the 5'-side of the anticodon, this tRNA could form a complex with A-U-G on ribosomes. It may be assumed that the additional U does not disturb the so called U-turn structure with the anticodon trimer in line.

The 5'-side of the anticodon in tRNAs is almost always occupied by a U residue. Uhlenbeck et al. showed previously that this constant U and the anticodon sequence in yeast tRNA<sup>Tyr</sup> could be substituted by another base.<sup>13)</sup> We have showed similar results in this report. Thus the tRNA<sup>Met</sup> which had other bases instead of constant U could be charged with methionine to the same level as tRNA<sup>Met</sup> having constant U. We conclude that the constant U at the 5'-side of the anticodon is not indispensable for the amino acid acceptor

activity of tRNA. Presumably even if the 5'-side of the anticodon is another base than U, the anticodon structure is maintained as in natural tRNA Met.

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## ACKNOWLEDGEMENT

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# Modification of the amino acid acceptor stem of *E. coli* $tRNA_{f}^{Met}$ by ligation of chemically synthesized ribooligonucleotides

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The single-stranged region of the amino acid acceptor stem corresponding to the 3'-end of *E. coli* tRNA<sup>Met</sup> was replaced by ligation of chemically synthesized ribooligonucleotides, in order to change the length of the single-stranded CCA terminus. The chemically synthesized ribooligomers, CCA, ACCA, AACCA and CAACCA, were ligated to nuclease-treated *E. coli* tRNA<sup>Met</sup>, which lacked the ACCA sequence at the 3'-end. The methionine acceptor activities of these modified tRNAs were examined using *E. coli* methionyl-tRNA synthesized pentamer (AACCA) to the acceptor terminus restored the methionine acceptor activity, whereas ligation of the hexamer (CAACCA) or trimer (CCA) to the acceptor terminus did not Modification of the acceptor terminus had no effect on the formylation of accepted methionine.

Aminoacylation

Formylation Elongat Sv

Elongated aminoacylation end Synthetic oligonucleotide Truncated aminoacylation end ligation

## 1. INTRODUCTION

Transfer RNA is one of the smallest nucleic acids that has a biological function in living cells. In many laboratories structure requirements for the recognition of tRNAs by aminoacyl-tRNA synthetases have been concerned with the interaction between nucleic acids and proteins [1–14].

We have synthesized various analogues of E. coli tRNA<sup>Met</sup><sub>f</sub> by ligating chemically synthesized ribooligomers to natural tRNA<sup>Met</sup><sub>f</sub> fragments using T4 RNA ligase [12-14].

Replacement of the anticodon trimers in *E. coli*  $tRNA_f^{Met}$  caused a large decrease in the amino acid acceptor activity with *E. coli* methionyl-tRNA synthetase [14]. On the other hand, changes in the discriminator position of the amino acid acceptor stem had little influence on the affinity for the synthetase [13].

\* To whom correspondence should be addressed

*E. coli* tRNA<sup>Met</sup> has a unique structure at the anticodon loop and the amino acid acceptor stem [15,16] where the last 5 bases from the 3'-terminus curl back in contrast to other tRNAs [16]. We assumed that the acceptor end directly binds methionine, and therefore investigated the correlation between the length of the protruding 3'-terminus and the methionine acceptor activities of analogs having a different protruding end. Here, we describe the syntheses of several modified *E. coli* tRNA<sup>Met</sup><sub>f</sub> formed by ligating CCA, ACCA, AACCA or CAACCA to a tRNA<sup>Met</sup><sub>f</sub> lacking the 3'-ACCA. The methionine acceptor activities of these analogues were examined using the *E. coli* methionyl-tRNA synthetase.

## 2. MATERIALS AND METHODS

Ribooligonucleotides were synthesized by the triester method [17].  $[\gamma^{-32}P]ATP$  and L-[U-<sup>14</sup>C]-methionine were obtained from NEN and Amer-

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sham, respectively. *E. coli* tRNA<sup>Met</sup> was a gift from Dr D. Söll. T4 RNA ligase was purified as described [18]. Polynucleotide kinase and *E. coli* alkaline phosphatase were obtained from Takara Shuzo and nuclease S1 from Sankyo. Other enzymes used for the analyses of products were obtained as in [14,19,20]. Crude *E. coli* aminoacyltRNA synthetase was purified as described [21]. Folinic acid-SF as a formyl donor was obtained from Lederle.

Partial digestion of *E. coli* tRNA<sub>f</sub><sup>Met</sup> with nuclease S1 and aminoacylation reaction were performed as in [13]. Phosphorylation with polynucleotide kinase, dephosphorylation with alkaline phosphatase and ligation with T4 RNA ligase were performed according to [12,14]. Formylation was performed under the conditions for aminoacylation containing 0.7 mM folinic acid-SF at 37°C for 30 min in 30  $\mu$ l formylation reaction mixture. Half of the mixture was treated with the same volume of 50 mM CuSo<sub>4</sub>, 0.3 M Tris-HCl (pH 7.5) at 37°C for 10 min.

## 3. RESULTS AND DISCUSSION

A scheme for the synthesis of modified E. coli  $tRNA_{f}^{Met}$  is shown in fig.1. E. coli  $tRNA_{f}^{Met}$  was treated with a limited amount of nuclease S1 to remove the 3'-terminal ACCA. The remaining fragment with 73 nucleotides [tRNA(73)] was isolated on 10% polyacrylamide gel electrophoresis (PAGE). Four ribooligonucleotides, CCA, ACCA, AACCA, CAACCA, were synthesized by the triester method as in [17]. These chemically synthesized ribooligonucleotides were phosphorylated with  $[\gamma^{-32}P]ATP$  and polynucleotide kinase and then joined to the tRNA(73) with T4 RNA ligase. The ligated products were isolated on 10% PAGE and subjected to a nearest neighbor analysis to confirm the junction point. The yields of these ligase reactions were below 10%.

The ligated molecules, tRNA(76), tRNA(77), tRNA(78) and tRNA(79), were dephosphorylated with alkaline phosphatase and tested for their methionine acceptor activities (fig.2). The tRNA(77)

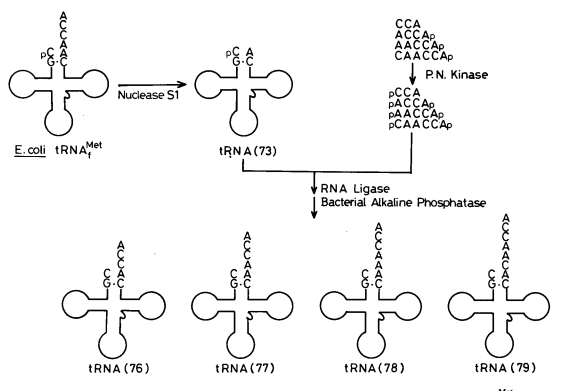


Fig.1. Scheme for the modification of the amino acid acceptor stem of E. coli tRNA<sub>f</sub><sup>Met</sup>.

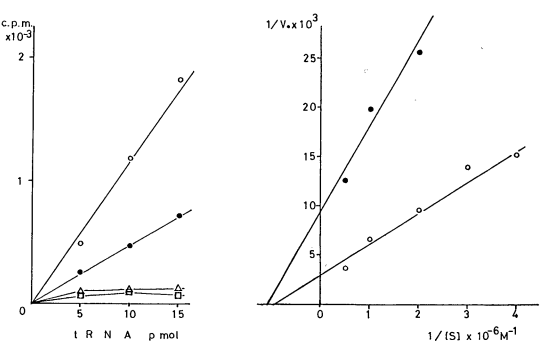


Fig.2.Aminoacylation of modified tRNAs. Methionine acceptor activities of tRNAs (left) and kinetic studies for aminoacylation (right). (○—─○) Control tRNA(77), (△—△) tRNA(76), (●—●) tRNA(78), (□—□) tRNA(79).

molecule was synthesized as a control since it had the same sequence as the natural tRNA<sup>Met</sup>. Fig.2 shows that tRNA(76), 1 base shorter, and tRNA(79), 2 bases longer, had no methionine acceptor activity, but tRNA(78), 1 base longer, could accept methionine. The kinetic studies on aminoacylation showed that the differences in aminoacylation activity were reflected only in the  $V_{\text{max}}$  values since the  $K_{\text{m}}$  values of these tRNAs were almost the same. Therefore, both tRNA(77) and tRNA(78) had similar affinities for E. coli methionyl-tRNA synthetase. It appears that the 3'-terminus of tRNA<sup>Met</sup><sub>f</sub> does not influence the affinity of methionyl-tRNA synthetase. Consequently we can assume that both tRNA(76) and tRNA(79) presumably form a complex with the synthetase, but are unable to position the 3'-end adenosine moiety at the catalytic site of the synthetase. One extra base in the 3'-terminus region may not prevent the required contact of the 3'-end adenosine with the catalytic site.

The recognition site of methionyl-tRNA<sub>f</sub><sup>Met</sup> by the transformylase in prokaryotes is still unclear. We have investigated the formylation of the tRNA(78) under the same assay system conditions as for the aminoacylation reaction except for the presence of formyl donor. The formylation ratio for tRNA(78) and tRNA(77) was 0.87 and 0.74, respectively. This may indicate that the addition of 1 extra base in the 3'-terminus region of tRNA<sup>Met</sup> does not affect the shape of the molecule or that the transformylase may recognize not only the 3'-terminal region but also another area as can be seen by comparison of the sequence of tRNA<sup>Met</sup> [22] and tRNA<sup>Met</sup> [23].

## ACKNOWLEDGEMENTS

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## The *in vivo* stability, maturation and aminoacylation of anticodon-substituted *Escherichia coli* initiator methionine tRNAs

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We have constructed eight anticodon-modified *Escherichia coli* initiator methionine (fMet) tRNAs by insertion of synthetic ribotrinucleotides between two fragments ('half molecules') derived from the initiator tRNA. The trinucleotides, namely CAU (the normal anticodon), CAA, CAC, CAG, GAA, GAC, GAG and GAU, were joined to the 5' and 3' tRNA fragments with T4 RNA ligase. The strategy of reconstruction permitted the insertion of radioactive <sup>32</sup>P label between nucleotides 36 and 37. tRNAs were microinjected into the cytoplasm of *Xenopus laevis* oocytes, and the following properties were evaluated: (a) the stability of these eubacterial tRNA variants in the eukaryotic oocytes; (b) the enzymatic modification of the adenosine at position 37 (3' adjacent to the anticodon) and (c) aminoacylation of the chimeric tRNAs by endogenous oocyte aminoacyl-tRNA synthetases.

In contrast to other variants, the two RNAs having CAU and GAU anticodons were stable and underwent quantitative modification at A-37. These results show that the enzyme responsible for the modification of A-37 to N-[N-(9- $\beta$ -D-ribofuranosylpurine-6-yl)carbamoyl]threonine (t<sup>6</sup>A) is present in the cytoplasm of oocytes and is very sensitive to the anticodon environment of the tRNA. Also, these same GAU and CAU anticodon-containing tRNAs are fully aminoacylated with the heterologous oocyte aminoacyl-tRNA synthetases *in vivo*. During the course of this work we developed a generally applicable assay for the aminoacylation of femtomole amounts of labelled tRNAs.

Some time ago a correlation between the anticodon sequence of a tRNA and the identity of neighbouring modified nucleosides was noted [1] (reviewed in [2]). Thus, the hypermodified nucleoside t<sup>6</sup>A, *N*-[*N*-(9- $\beta$ -D-ribofuranosylpurine-6-yl)carbamoyl]threonine, or a derivative thereof is located in position 37 of tRNAs having anticodons terminating in a uridine. One enigmatic exception is the initiator methionine tRNA of *Escherichia coli*, for most other tRNAs including the eubacterial elongator Met-tRNA, which has the same anticodon sequence, and the eukaryotic initiator tRNA contain the A-37 modification [3].

To study the effect of structural modifications in the anticodon loop on the modification of A-37, we have turned to recombinant RNA methods based on T4 RNA ligase. These techniques are particularly well-suited to the preparation of related tRNA chimera, which have substitutions in or near the anticodon, since fragments serving as starting material for the tRNA variant can be readily obtained from controlled

nucleolytic cleavage of tRNA. Several variant tRNA molecules have been constructed in this way from E. coli and yeast tRNA species; these have been useful in evaluating the effect of nucleotide substitution on the interaction of tRNA with aminoacyl-tRNA synthetases, nucleoside modification enzymes and codons during protein synthesis [4-13] (reviewed in [14]). In the present work we prepared eight anticodon-substituted fMet-tRNA chimera. These anticodon variants of the type CAX and GAX (where X is each of the four nucleosides A, C, G and U; CAU is the normal MettRNA anticodon) were synthesized by inserting trinucleotides between 3' and 5' half molecules and analyzed after injection into Xenopus laevis oocytes. Several of these same variants were prepared previously for studies on E. coli methionyltRNA synthetase recognition [8-12]. Using the Xenopus oocyte system [15] we were able to evaluate in vivo the heterologous charging of these variants, the stability of the chimeric tRNA and the structural requirements for nucleoside modification.

## MATERIALS AND METHODS

The ribotrinucleotides CAA, CAC, CAG, CAU, GAA, GAC, GAG and GAU were synthesized by the triester method [16]. Their sequences were determined by complete digestion with RNase T2 and nuclease P1 followed by paper electrophoresis at pH 3.5 [16]. [ $\gamma$ -<sup>32</sup>P]ATP (spec. act. > 3000 Ci/mmol) was obtained from the Amersham International (England). The fMet-tRNA (spec. act. 1400 pmol/ $A_{260}$  unit) was a gener-

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Abbreviations. t<sup>6</sup>A, N-[N-(9- $\beta$ -D-ribofuranosylpurine-6-yl)carbamoyl]threonine; i<sup>6</sup>A,  $N^6$ -isopentenyladenosine; BD-cellulose, benzoylated DEAE-cellulose.

*Enzymes* (IUB Recommendations 1984). Methionyl-tRNA synthetase of L-methionine:tRNA ligase (AMP) (EC 6.1.1.10); T4 RNA ligase (EC 6.5.1.3); T4 polynucleotide kinase (EC 2.7.1.78); alkaline phosphatase (EC 3.1.3.1); pancreatic RNase A (EC 3.1.27.6); RNase T2 (EC 3.1.27.1); nuclease P1 (EC 3.1.30.1); tRNA nucleotidyltransferase (EC 2.7.7.25).

ous gift of Dr D. Söll of Yale University. T4 RNA ligase was purified from *E. coli* strain A19 infected with phage T4 as described [17]. T4 polynucleotide kinase and alkaline phosphatase from *E. coli* strain A19 were obtained from Takara Shuzo Co. (Japan). Pancreatic RNase A, RNase T2 from *Aspergillus orizae* and nuclease P1 from *Penicillium citrinium* were obtained from Boehringer, Mannheim (FRG), Sankyo Co. (Japan) and Yamasa Shoyu Co. (Japan) respectively. Other products were obtained as described earlier [9, 16, 18].

## Preparation of E. coli fMet-tRNA half molecules

E. coli fMet-tRNA was digested by pancreatic RNase A under limiting conditions previously described [9]. The resulting 5' and 3' half-molecule products were purified by 10% polyacrylamide gel electrophoresis under denaturing conditions (7 M urea), and their sequences were determined by mobility-shift analysis [9].

## Ligation, phosphorylation and dephosphorylation

Enzymatic assay conditions for ligation, phosphorylation and dephosphorylation of oligoribonucleotides or tRNA fragments were described previously [16-20]. Trinucleotides, after phosphorylation using T4 polynucleotide kinase, were purified by DEAE-Sephadex A-25 ion-exchange chromatography [19]. First the trinucleotide was ligated to the 5' half molecule. The 3' half molecule of fMet-tRNA was 5' phosphorylated using T4 polynucleotide kinase and  $[\gamma^{-32}P]ATP$ ; this product was then ligated to the 5' half molecule having the attached trinucleotide [9]. The reconstructed <sup>32</sup>P-labelled fMet-tRNA variants were purified by 10% polyacrylamide gel electrophoresis in 7 M urea. The gel mobility of each tRNA was compared to that of a genuine sample of E. coli fMet-tRNA detected by ultraviolet shadowing [52]. The variant tRNAs were eluted from the gel. The solutions were centrifuged to remove small pieces of acrylamide and 2.5 volumes of ethanol were added. After redissolution in buffer, the product was reprecipitated in ethanol.

Complete digestions of RNA molecules with RNase T2 or nuclease P1 were carried out for nearest-neighbour analysis as previously described [18-20].

## Microinjection and fate of tRNA chimera in oocytes

Microinjection into the cytoplasm of X. laevis oocytes was performed as described [15, 18], using 50 nl aqueous labelled tRNA solution at a concentration which gave approximately 200000 cpm (Cerenkov)/nl. This is equivalent to about 1 fmol foreign tRNA/oocyte and corresponds to about 1% of the normal amount of oocyte Met-tRNA. Groups of five oocytes were injected with each sample (reproducibility in tRNA microinjections is about 20%) and then incubated in saline solution [21] at 19°C for the desired period of time (from 1 h to 72 h after injection). At the end of the incubation the oocytes were homogenized in 0.2 ml cold buffer containing 0.2 M sodium acetate (pH 4.5, 4°C), 0.01 M MgCl<sub>2</sub>, 1 mM disodium EDTA and 1% (w/v) of sodium dodecyl sulfate. The nucleic acids were immediately extracted by phenol/ chloroform treatment and ethanol-precipitated as described [18]. Each tRNA precipitate was then redissolved in the same volume of electrophoresis buffer. Given aliquots (the same for each tRNA sample) were then loaded side by side, on a 10% polyacrylamide gel containing 7 M urea. Electrophoresis

was performed at constant voltage for 5-7 h. Location of the  $^{32}P$  (labelled Met-tRNA and the putative degradation products) was performed by autoradiography of the gel, and the band corresponding to intact full-length tRNA was cut out, eluted and its radioactivity counted again.

Identification of the 5'-<sup>32</sup>P-labelled nucleotide at position 37 of the eluted tRNA was carried out by exhaustive nuclease P1 digestion [22] and analysis of the products by two-dimensional chromatography on thin-layer cellulose plates (6.6  $\times$  6.6 cm<sup>2</sup>, see [1]). After detection of the labelled nucleotides on the plates, they were eluted with water and their radioactivity was determined by liquid scintillation.

## Assay of in vivo aminoacylation

tRNA was extracted from oocytes using phenol saturated with 50 mM sodium acetate buffer, pH 4.5, and precipitated with ethanol [23]. It was dissolved in 50 µl 0.1 M triethanolamine/HCl buffer at pH 8.0, and 1 mg phenoxyacetyl ester of *N*-hydroxysuccinimide in 10 µl of anhydrous tetrahydrofuran was added at 0°C [24]. After 10 min the reaction was stopped by dilution with 60 µl cold 20 mM sodium acetate buffer at pH 4.5 containing 20 mM MgCl<sub>2</sub> and 100 mM NaCl. This solution was then applied to a 1-ml column of benzoylated DEAE-cellulose (BD-cellulose), which was pre-equilibrated in 10 mM sodium acetate buffer pH 4.5 containing 10 mM MgCl<sub>2</sub> and 50 mM NaCl. The sample was applied, and the column was washed with 3 ml starting buffer and then with 2 ml of the same buffer containing 1.0 M NaCl. A final wash was composed of the starting buffer containing 1.0 M NaCl and 20% ethanol. All fractions (0.5 ml) were counted in a scintillation counter, and the extent of aminoacylation in the original tRNA preparation was estimated by comparison of the radioactivity of the ethanol fraction to the radioactivity applied to the column.

## RESULTS

## Construction of anticodon-substituted E. coli fMet-tRNA

The enzymatic replacement of the E. coli fMet-tRNA anticodon by several oligoribonucleotides was reported previously [9, 12] and similar procedures have been developed independently by Schulman et al. [10, 11a, b]. In this paper we concentrate on the anticodon variants of the type CAX and GAX (where X is any of the four nucleotides). The reconstruction scheme in Fig. 1 is essentially the same as that previously described [9] except for a simplification of step 3 and the addition of step 7 (see below). Thus, the digestion of fMet-tRNA with pancreatic RNase A generated the two half molecules less the three anticodon nucleotides and the dinucleotide CA of the 3' terminus. The 5' half molecule, composed of pC-1 to Up-33, and the 3' half, A-37 to Cp-75, were separated by 10% polyacrylamide gel electrophoresis under denaturing conditions. The 5' half molecule was then treated by alkaline phosphatase to remove the two terminal phosphates and the product was isolated again by denaturing polyacrylamide gel electrophoresis (step 2). In step 3, synthetic ribonucleotides were joined to the 5' half molecule using RNA ligase. In the earlier procedure [9] this step was performed with synthetic trinucleotides, which were phosphorylated at both ends to prevent self-polymerization. In the present procedure trinucleotides phosphorylated at only the 5' terminus were used; this simplification saves a 3'-dephosphorylation step which would be necessary subsequently to link this

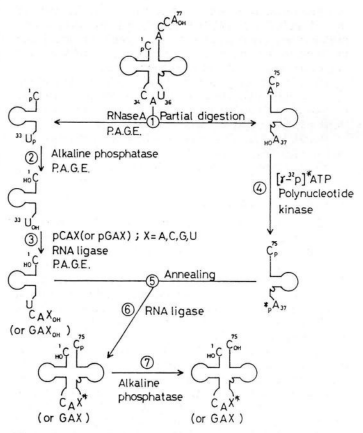


Fig. 1. Synthetic scheme for the preparation of E. coli initiator MettRNA variants. Numbers and names identify the successive steps and molecules used (see text). Each fragment was isolated by electrophoresis on polyacrylamide gel. The asterisk signifies the position of an internal <sup>32</sup>P label introduced in step 4 on the 5'-phosphate of A-37

oligomer to the 3' half molecule. The modification gives yields equivalent to the original procedure, but it is preferable when large numbers of tRNA variants must be prepared.

In step 4 the 3' half molecule was phosphorylated with kinase and  $[\gamma^{-32}P]ATP$ . This radioactive oligomer was then annealed with a ten-times excess of the unlabelled extended 5' half molecule (step 5). The large excess was used to consume all the radioactive fragment and is probably not needed to obtain respectable yields in the ligation step. Ligation of the two molecules in step 6 gave nearly a quantitative yield of the tRNA product, which lacks only the 5' phosphate and the 3' CA terminus. Purification by polyacrylamide gel electrophoresis is shown in Fig. 2. Dephosphorylation by alkaline phosphatase (step 7), a necessity for proper CCA terminal transferase repair in the oocyte [40], was followed by a final gel purification.

## Stability of the fMet-tRNA chimera

Each of the eight tRNA chimera were microinjected into the cytoplasm of X. *laevis* oocytes. The degradation of these eubacterial variants in the eukaryotic cytoplasm was measured after incubation of the oocytes at 19°C for different periods of time by quantitative analysis of the remaining tRNAs. This analysis consisted of the gel electrophoresis of tRNA samples extracted from oocytes. As shown in Fig. 3, only products having a few less nucleotides than the tRNA are seen: they migrate more quickly than the full-length molecules (compare with the position of the arrow on Fig. 3).

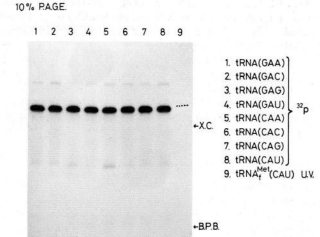


Fig. 2. Autoradiogram of the purification by electrophoresis on polyacrylamide gel (10% in 7 M urea) of eight tRNA chimery. This is the gel after step 6 of Fig. 1. The tRNAs lack the 5' terminus and the  $C^{75}$ - $A^{76}$  of the 3' terminus; therefore, they run correspondingly faster than the authentic fMet-tRNA sample in lane 9 (indicated by the dashed line). The absence of low-molecular-mass fragments reflects a greater than 95% yield of the cyclization reaction (step 6). Migration is from the top to the bottom of the gel as indicated by the bromophenol blue marker (B.P.B.); X.C. corresponds to the position of xylene cyanol

These shorter molecules could represent the tRNA without the 3'CCA terminus while the longer molecules probably correspond to the fMet-tRNA having their 3'CCA end repaired by the tRNA nucleotidyltransferase which is present in the cytoplasm of X. *laevis* oocytes [40, 41]. Interestingly enough, degradation products corresponding to small pieces of tRNA are not apparent on the gel after electrophoresis. This suggests that the very first cleavage in tRNA by oocyte nucleases is the rate-limiting step, after which full degradation of the tRNA into nucleotides occurs rapidly. A similar observation was already made with microinjected yeast tRNA-Asp, where the anticodon loop was cleaved prior to microinjection [18].

The overall rate of degradation of full-length tRNA was evaluated by determination of the amount of radioactivity remaining in the corresponding band on the polyacrylamide gel. These results are summarized in the degradation curves of Fig. 4. The rate of degradation is seen to be, to a first approximation, a simple exponential function, and each tRNA variant has a characteristic half-life in the oocyte. Clearly the fMet-tRNA with a CAU anticodon (the native anticodon) and the GAU anticodon (the anticodon of isoleucine belonging to the same genetic-code quartet) are the most stable having half-lifes of approximately 95 h and 130 h respectively. All other tRNAs had half-lifes on the order of 16-32 h.

## Maturation of tRNA chimera in oocytes

In order to determine the state of modification of the A-37 in the above experiments, the full-length tRNA was extracted from the gels, and each sample was digested by nuclease P1 producing nucleoside 5'-phosphates. Owing to the location of the radioactive phosphate between nucleotides 36 and 37, only the nucleotide in position 36 was labelled, the

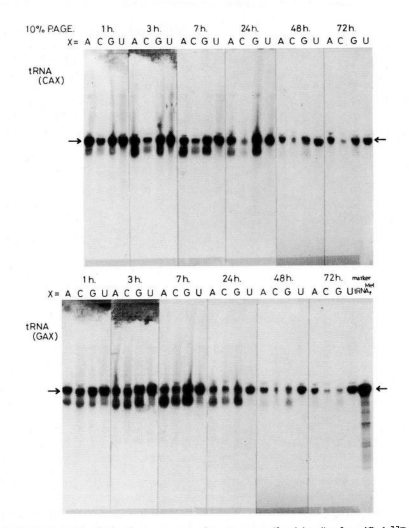


Fig. 3. Stability of microinjected tRNAs in the X. laevis oocytes. A given amount (fmol level) of purified <sup>32</sup>P-labelled tRNA (natural or chimeric) was microinjected into the cytoplasm of a series of five oocytes. At the indicated time after microinjection, total RNA was extracted from the oocytes by phenol/chloroform treatment and ethanol precipitation. RNA samples were then redissolved in electrophoresis buffer and analysed for the presence of degradation products by electrophoresis on a 10% polyacrylamide gel in 7 M urea. The figure shows the resulting autoradiography for *E. coli* fMet-tRNA with anticodon CAX (upper part) or with anticodon GAX (lower part). The last lane (lower part) shows the degradation products of an authentic sample of *E. coli* fMet-tRNA incubated in formamide for 30 min at 100°C (control experiment). The arrows indicate the position of a genuine sample of *E. coli* fMet-tRNA (having its terminal CCA end) detected by ultraviolet shadowing [52]. RNA bands migrating faster than this authentic *E. coli* fMet-tRNA, probably correspond to fMet-tRNA lacking the terminal nucleotides of the amino acid stem (see text)

identity of which could be determined by two-dimensional thin-layer chromatography.

Fig. 5a-h shows representative autoradiograms of these experiments and illustrates clearly that A-37 was rapidly modified to t<sup>6</sup>A in the tRNA variants having the native CAU (Fig. 5a) and GAU (Fig. 5e) anticodon. Trace amounts of the isopentenyl modification of A-37 was detected in the variants having CAA (Fig. 5b) or GAA anticodons (Fig. 5f), and no modification was detected in variants harboring the CAC, CAG, GAC and GAG anticodons (Fig. 5c, g and h) even after incubations as long as 48 h.

Quantification of the radioactivity in the different spots from the thin-layer chromatogram allows the calculation of the kinetics of *in vivo* modification (Fig. 6). The modification of A-37 to t<sup>6</sup>A occurs very rapidly for the CAU and GAU chimera reaching a plateau at 80-90% transformation. The apparent higher rate of the GAU chimera could be due to sequestering of significant quantities of the natural initiator CAU chimera. Isopentenylation of A-37 in the CAA or GAA anticodon variants is considerably slower: only 6-10% conversion after 72 h. Moreover, the rates of conversion are essentially the same whether the tRNA is injected in the oocyte cytoplasm or nucleus (data not shown).

## A microassay of tRNA aminoacylation

With the set of tRNA variants in hand, it was also of interest to determine which, if any, of them could be aminoacylated in the oocyte. Allende and his coworkers [25a, b] were the first to devise a microassay to evaluate the biological activity of small amounts of tRNA microinjected into amphibian oocytes. This procedure was based on the ability of aminoacylated tRNA, but not uncharged tRNA, to bind to *E. coli* elongation factor EF-Tu. Later on, Corbo et al. [26] adapted this procedure in order to quantify small amounts of  ${}^{32}$ P-labelled tRNAs produced after transcription of a tRNA gene in the *Xenopus* oocyte nucleus. These procedures, however, are unwidely at best, since they depend on a Sephadex G-100 separation of charged and uncharged tRNA. Another technique was based on the use of radioactive

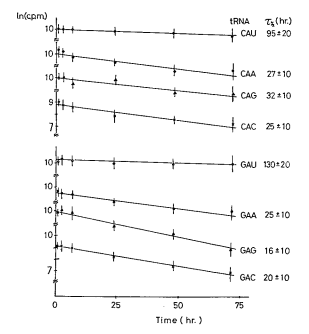


Fig. 4. Stability of microinjected tRNA (natural or chimeric) in the X. laevis oocytes. The results of Fig. 3 were further analyzed by counting the radioactivity recovered from the oocytes in bands corresponding to the full-length tRNA and the tRNA lacking the terminal nucleotides of the amino acid stem (see Fig. 3). These data were then plotted in a semilogarithmic mode as a function of time. In the right-hand column are listed the calculated half-lifes  $t\frac{1}{2}$  of each of the eight tRNA variants assuming an exponential decay law -dN/dt = kN

amino acids [25b] (see also [15]). In all, none of these techniques was particularly well-suited to the determination of variant tRNA aminoacylation for this study; therefore, we set out to establish an alternative method. The technique selected is based on the well-known chromatographic behaviour of phenoxyacetylated aminoacyl-tRNA to be absorbed strongly to BD-cellulose [24]. The derivatized charged tRNA is eluted only in an alcoholic buffer, whereas uncharged and underivatized charged tRNAs are eluted anteriorly in a salt buffer.

## The in vitro aminoacylation of tRNA chimera

Using the above technique we were able to determine the aminoacylation level of each microinjected tRNA variant (Table 1) after various times of incubation in the oocytes. As above, only the natural CAU and the GAU (isoleucine) anticodon-containing tRNAs were aminoacylated to a reasonable extent. Presumably both are aminoacylated with methionine, although the technique based on the detection of <sup>32</sup>P in the RNA does not distinguish between aminoacylating groups. Also, only the fraction of Met-tRNA in which the 3' ends have been fully repaired by the oocyte tRNA nucleotidyltransferase [40] can be charged with the amino acid.

## DISCUSSION

In this paper we have presented evidence that, among the eight E. coli fMet-tRNA anticodon chimera, the GAU and the native CAU variants have the longest half-lifes, are the

only tRNAs where the A-37 is converted to  $t^6A$  and are aminoacylated to a significant degree in amphibian oocytes. This surprising coincidence must be related to the functionality of these tRNAs; tRNAs which can be aminoacylated could be sequestered by proteins and thus be protected from cytoplasmic nucleases (see [15]).

It is indeed remarkable that the A-37 of E. coli fMet-tRNA is modified to t<sup>6</sup>A in the oocyte, since this modification does not take place in E. coli [39]. Thus, there is no intrinsic structural reason, relating to an unusual anticodon conformation for example [27], that this tRNA cannot be modified. The eubacterial modifying enzyme would seem to be more selective in its action. Since the tRNAs variants were injected into the oocyte cytoplasm, we conclude that the t<sup>6</sup>A modifying enzyme is located in the cytoplasm. This result is fully consistent with the fact that this modification has not been found in nuclear tRNA precursors [28]. Considering the cytoplasmic presence of the queuosine (a modified base found at position 34) and the i<sup>6</sup>A-37 modification enzymes shown by the microinjection of yeast tyrosine tRNA [29], it may be that most anticodon-loop-modifying enzymes are cytoplasmic (see also [42]).

Most eukaryotic tRNAs having anticodons ending with U have a t<sup>6</sup>A and those ending with A, a i<sup>6</sup>A, 1-methylguanosine or a Y base at position 37 [1, 30, 31]. A remarkable illustration of these correlations is that mutations resulting in the replacement of C-36 by A-36 or U-36 as well as U-36 by A-36 in glycine or lysine E. coli tRNAs is accompanied by a change in the nature of the modification at A-37 [32-34]. Results of Murgola and coworkers [34, 35], however, point to the fact that the molar yield of such modifications may be quite low, and caution must be used in interpreting the existing correlative data [36]. The low or non-existent conversion of A-37 to the 6-isopentenyl derivative in the CAA and GAA chimera demonstrates that several correlations observed in natural tRNAs do not necessarily hold for synthetic variants. In any case the determinants of nucleotide modification by a given enzyme acting in the anticodon loop are much more complex than the anticodon correlations would lead one to believe. For more discussion, see [36, 42].

The aminoacylation of the E. coli fMet-tRNA variants by the oocyte aminoacyl-tRNA synthetases is also of interest. In previous work it has been unambigiously demonstrated that the anticodon nucleotides of E. coli fMet-tRNA were crucial for recognition by the corresponding homologous enzyme. Indeed, aminoacylation is reduced to levels below experimental detection *in vitro* by base substitution in the wobble position, and lesser, although dramatic, effects result from structural changes at the other two positions (positions 35 and 36) of the anticodon [9-11a]. The mechanism by which just one base substitution in the wobble position reduces the aminoacylation rate of E. coli fMet-tRNA by at least five orders of magnitude remains, however, unclear [11a, 37]. On the other hand, base substitution at position 33 (the constant uridine) or at position 37 (the unmodified adenosine) as well as enlargements of the anticodon by one or two nucleotides at its 5' or the 3' side do not affect drastically aminoacylation of E. coli fMet-tRNA by E. coli methionyl-tRNA synthetase [11a, 12]. From these results it has been concluded that recognition of E. coli fMet-tRNA by its cognate homologous enzyme requires specific interactions of the methionyl-tRNA synthetase with functional groups of the nucleotide base of the anticodon and that the spatial arrangement of those bases in the anticodon loop is less critical than their chemical nature [11a, 37].

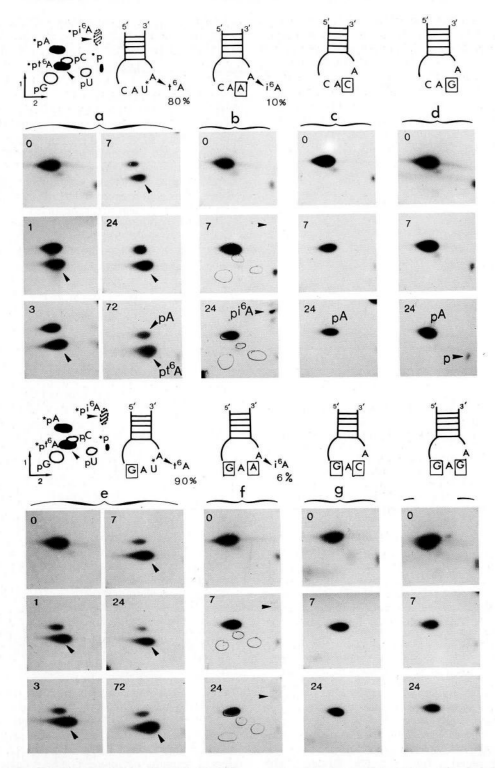


Fig. 5. Autoradiograms of the separation of 5'-nucleosides by chromatography on thin-layer cellulose plates. After isolation from the oocytes and purification by electrophoresis on polyacrylamide gel as shown in Fig. 3, the <sup>32</sup>P-labelled fMet-tRNA recovered from the gel was completely digested into nucleosides with nuclease P1. The resulting nucleoside 5'-monophosphates were separated on thin-layer cellulose plates as described in Materials and Methods. The pattern of nucleosides is given in the upper left-hand corner and each sample is identified by the schematic drawing of its anticodon loop: from (a) to (h) for each of the eight tRNA variants. Nucleotides in boxes are those which differ from the parent *E. coli* fMet-tRNA. Numbers in the corner of each autoradiogram correspond to the time of incubation (h) of the tRNA in the oocytes at 19°C; zero corresponds to tRNA before the microinjection (control experiments). Characteristic spots, which correspond to modified nucleotides pt<sup>6</sup>A and pi<sup>6</sup>A, are identified by arrows. Percentages correspond to molar yield of transformation (mol/ 100 mol) of A-37 to t<sup>6</sup>A-37 or i<sup>6</sup>A-37 in *E. coli* fMet-tRNA after 72 h incubation in the oocytes. Asterisk indicates which compounds are labelled with <sup>32</sup>P in the nuclease hydrolysate of the tRNA

Clearly these rules, which applied to the *E. coli* methionyltRNA synthetase, cannot be extended to the same enzyme from the *X. laevis* oocyte. Indeed, in the experiments reported here the GAU variant of *E. coli* fMet-tRNA (with a base substitution in the wobble position) is at least as active in aminoacylation as the native fMet-tRNA (with normal anticodon CAU): even the reaction rates were so fast *in vivo* that we cannot differentiate between the two tRNAs. It should be pointed out that the non-stoichiometric aminoacylation could result from either the lack of a 5'-phosphate on the tRNA, even though this feature is not thought to effect aminoacyla-

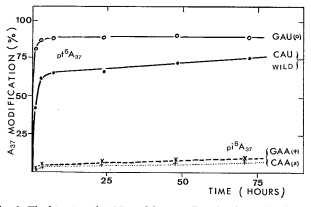


Fig. 6. The kinetics of A-37 modification. Results shown in Fig. 5a - h were analyzed by radioactive counting of the various spots on the thin-layer cellulose plates. From these data were calculated the quantities of pt<sup>6</sup>A and pi<sup>6</sup>A as a function of time, expressed as percentages of the total radioactivity on the thin-layer cellulose plates. The nucleotide sequences of the novel synthetic anticodons in tRNAs corresponding to each curve are given. The kinetics correspond to the post-transcriptional modifications of A-37 in microinjected *E. coli* fMet-tRNA under physiological conditions of a living cell

## Table 1. In vivo aminoacylation level of tRNAs

Activity is expressed as the percentage of radioactivity ( $^{32}$ P) retained on the BD-cellulose column during salt washes (1 M NaCl/sodium acetate buffer at pH 4.5) and eluted in the same salt buffer containing 20% ethanol; the total radioactivity recovered from the column being taken as 100%. These radioactivities correspond to the fraction (%) of Met-tRNA labelled with  $^{32}$ P containing a phenoxyacetyl group, because of the presence of an aminoacyl group at the 3' termini. The error in the evaluations is considered to be 15-20%. The control experiment corresponds to about 15-20 min incubation *in vivo*; it represents the minimal time required to complete the microinjections in one series of five oocytes and to proceed in order to recover the  $^{32}$ P-labelled tRNA from the oocytes by phenol/chloroform extraction as described in Materials and Methods

Anticodon	Aminoacylation after incubation for			
	control	3 h	`12 h	
	%			
CAU	23	× 52 <sup>1</sup>	54	
CAA	3	5	7	
CAC	6	10	10	
CAG	4	8	10	
GAU	24	55	64	
GAA	4	8	9	
GAC	8	6	14	
GAG	5	10	11	

tion (at least with E. coli methionyl-tRNA synthetase) [38], or the incomplete repair of the CCA terminus in the oocvte [40, 41]. Although misacylation of the E. coli fMet-tRNA by an amino acid other than methionine can not be ruled out [11b], generally misacylation rates never reach the proportions of aminoacylation by cognate amino acid [11b, 43, 44]. Also there is no evidence for aminoacylation of a yeast initiator Met-tRNA having an arginine CCU (instead of CAU) anticodon using the homologous partially purified yeast methionyl-tRNA synthetase [45]. Since this variant of yeast Met-tRNA was not aminoacylated by the E. coli enzyme, in contrast to the natural yeast tRNA [45, 46], we conclude that the tRNA recognition by methionyl-tRNA synthetase of eukaryotic cells must be less dependent on the anticodon than is the activity of its E. coli counterpart. Experiments indicating the involvement of the anticodon in tRNA recognition by several aminoacyl-tRNA synthetases are often debated. In several instances it is clear that functional groups of anticodon nucleotides are important in the interaction but this might not be a general rule: see for example [37, 43, 47-49, reviewed in53].

Finally, a method for the determination of femtomole quantities of aminoacylated tRNA is presented. This rapid assay is insensitive to the presence of other (endogenous) unlabelled tRNAs and is therefore particularly well-suited to microinjection techniques. Previous methods had drawbacks relative to the detection limit of radioactive amino acids or tritiated tRNA and to their ease of analysis. The method presented here could be even more easily applied to aromatic amino acids, since their aminoacyl-tRNAs are strongly adsorbed to BD-cellulose without the use of the phenoxyacetyl group [50]. Two cautionary notes must be added, however. First, while the technique used in the assay of microinjected tRNA does determine which fraction of the tRNA is charged, it cannot identify the amino acid involved. The heterologous tRNAs used here could be subject to misaminoacylation, although demonstration of this phenomenon in vivo remains to be made. Also, occasionally a modified nucleotide, present in the labelled tRNA sequence, could react with the phenoxyacetyl group leading to the adsorption of noncharged tRNA to BD-cellulose. This situation could prevail for tRNAs containing 5-carboxymethylaminomethyluridine and 3-(amino-3-carboxypropyl)uridine [51]. In spite of these drawbacks we feel that this method will find many applications in the evaluation of the biological activity of small amounts of tRNAs, especially those produced by recombinant technology.

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#### Received 2 January 1979

#### ABSTRACT

P<sup>1</sup>-Adenosine 5'-P<sup>2</sup>-o-nitrobenzyl pyrophosphate (nbzlppA) has been synthesized as a substrate for T4 RNA ligase catalyzed 3'-phosphorylation. Incubation of oligoribonucleotides and nbzlppA with RNA ligase yielded oligoribonucleotides having a 3'-O-(o-nitrobenzyl)phosphate. Photochemical removal of the o-nitrobenzyl group provided the free 3'-phosphate. Using [P2-32P]nbzlppA, 3'-termini of oligoribonucleotides could be labelled with 32P. This reaction was applied to modify the 3'-end of donor molecules in joining reaction with RNA ligase. A trinucleotide U-A-G was converted to U-A-Gpnbzl and phosphorylated with polynucleotide kinase. pU-A-Gpnbzl was then joined to an acceptor trinucleotide A-U-G to yield A-U-G-U-A-Gp.

#### INTRODUCTION

T4 RNA ligase<sup>2</sup> has been demonstrated to be a useful reagent for joining oligonucleotides.<sup>3-9</sup> For intermolecular reactions the 3'-termini of the 5'-phosphorylated components (donor molecules) have to be modified to prevent self-polymerization and/or cyclization.<sup>10</sup> Several approaches have been reported for preventing these undesired reactions by substitution of various groups at the 2'- or 3'-hydroxyl group. A 2'-substituted mononucleotide can be linked to the 3'-end of oligoribonucleotides by either chemical<sup>8</sup> or polynucleotide phosphorylase catalyzed reactions.<sup>6</sup> Single addition reactions of nucleoside 3',5'-diphosphate catalyzed by RNA ligase have provided 3'-phosphorylated oligonucleotides.<sup>11,12</sup> It was also found that alkyl phosphates and sugar phosphates were recognized by the enzyme if they were linked with adenosine 5'-phosphate<sup>13</sup> in the way found in the active intermediate for the ligase reaction. 5,6,14 In the present paper we wish to report 3'-phosphorylation of oligoribonucleotides using an adenylated photolabile o-nitroVolume 6 Number 2 February 1979

A new method for 3'-labelling of polyribonucleotides by phosphorylation with RNA ligase and its application to the 3'-modification for joining reactions 1

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#### ABSTRACT

p<sup>1</sup>-Adenosine 5'-P<sup>2</sup>-o-nitrobenzyl pyrophosphate (nbzlppA) has been synthesized as a substrate for T4 RNA ligase catalyzed 3'-phosphorylation. Incubation of oligoribonucleotides and nbzlppA with RNA ligase yielded oligoribonucleotides having a 3'-O- (o-nitrobenzyl)phosphate. Photochemical removal of the o-nitrobenzyl group provided the free 3'-phosphate. Using [P2-32P]nbzlppA, 3'-termini of oligoribonucleotides could be labelled with 32P. This reaction was applied to modify the 3'-end of donor molecules in joining reaction with RNA ligase. A trinucleotide U-A-G was converted to U-A-Gpnbzl and phosphorylated with polynucleotide kinase. pU-A-Gpnbzl was then joined to an acceptor trinucleotide A-U-G to yield A-U-G-U-A-Gp.

#### INTRODUCTION

T4 RNA ligase  $^2$  has been demonstrated to be a useful reagent for joining oligonucleotides.<sup>3-9</sup> For intermolecular reactions the 3'-termini of the 5'-phosphorylated components (donor molecules) have to be modified to prevent self-polymerization and/or cyclization.<sup>10</sup> Several approaches have been reported for preventing these undesired reactions by substitution of various groups at the 2'- or 3'-hydroxyl group. A 2'-substituted mononucleotide can be linked to the 3'-end of oligoribonucleotides by either chemical<sup>8</sup> or polynucleotide phosphorylase catalyzed reactions.<sup>6</sup> Single addition reactions of nucleoside 3',5'-diphosphate catalyzed by RNA ligase have provided 3'-phosphorylated oligonucleotides. 11,12 It was also found that alkyl phosphates and sugar phosphates were recognized by the enzyme if they were linked with adenosine 5'-phosphate<sup>13</sup> in the way found in the active intermediate for the ligase reaction. 5,6,14 In the present paper we wish to report 3'-phosphorylation of oligoribonucleotides using an adenylated photolabile o-nitro-

benzyl phosphate ( $P^1$ -adenosine  $5'-P^2$ -o-nitrobenzyl pyrophosphate, nbzlppA) and RNA ligase. Substitution of the alkyl phosphate with  $^{32}P$  enabled us to label the 3'-terminus of oligonucleotides in the same way. The 3'-phosphorylated oligonucleotides could serve as donor molecules after 5'-phosphorylation with polynucleotide kinase. Using these techniques a hexanucleotide was synthesized as a model messenger RNA.

#### MATERIALS AND METHODS

nbzlppA. o-Nitrobenzyl phosphate was synthesized by condensation of o-nitrobenzyl alcohol with phosphoric acid using trichloroacetonitrile<sup>15</sup> and adenylated by treatment with 5'phosphoromorpholidate.<sup>16</sup> Phosphoric acid (85%, 0.116 ml, 1.7 mmol), o-nitrobenzyl alcohol (4.25 g, 27.7 mg) and triethylamine (0.56 ml, 4 mmol) were dissolved in acetonitrile (10 ml) and treated with trichloroacetonitrile (1 ml, 10 mmol) in acetonitrile (1 ml) at 75° for 4 hr in the dark. The extent of reaction was examined by paper electrophoresis. The volatile materials were removed by evaporation and water (20 ml) was added to the residue. The excess of o-nitrobenzyl alcohol was removed with ether (15 ml)x2, and the aqueous layer was concentrated in vacuo, then dissolved in water (14 ml) and applied to a column of Dowex 50X2 (H<sup>+</sup>)(1.7 x 10 cm). o-Nitrobenzyl phosphate was recrystallized from chloroform (20 ml)-acetone (5 ml). The yield was 53%, 0.209 g, 0.89 mmol. Anal. calcd. for C<sub>7</sub>H<sub>p</sub>NO<sub>6</sub>P: C, 36.07; H, 3.56; N, 6.01. Found: C, 35.80; H, 3.54, N, 5.81. o-Nitrobenzyl phosphate (0.2 mmol) was mixed with trin- octylamine (0.2 ml), coevaporated with pyridine three times and allowed to react with adenosine 5'-phosphoromorpholidate (0.2 mmol) in dry DMF (1 ml) at room temperature for 3 days. Paper electrophoresis showed a pyrophosphate which could be converted to ADP by irradiation with UV light. The reaction was stopped after another 3 days by addition of water. DMF was removed and the residue was dissolved in aqueous pyridine. The aqueous pyridine solution was passed through a column (2.7 x 5 cm) of Dowex 50X2 (pyridinium form) to remove tri-n-octylamine. The eluent and washings (200 ml, 10% pyridine) were applied to a column (1.7 x 19 cm) of DEAE-Sephadex A-25 (bicarbonate form). After the column was washed with water (1 L), the product was eluted with a linear gradient of triethylammonium bicarbonate (0 to 0.2 M, total 2 L). The appropriate fractions (0.13 M) were examined by paper electrophoresis and desalted by evaporation. The relative mobility of the product to pA was 0.94. The spectral properties of the product were  $\lambda \max$  (H<sub>2</sub>O) 261,  $\lambda \max$ (H<sup>+</sup>) 256,  $\lambda \max$  (OH<sup>-</sup>) 261 nm. The yield of nbzlppA was 1382 A<sub>260</sub> units, 33%. The pyrophosphate was characterized by enzymic digestion with venom phosphodiesterase to yield o-nitrobenzyl phosphate and pA. The product could be converted to ADP by irradiation with UV light.

<u>nbzl\*ppA</u>. The  $[^{32}P]$  labelled pyrophosphate was synthesized as above except that \*Pi (1 mCi) was diluted with 0.05 mmol of phosphoric acid. The yield was 5.3 µmol, 11%. Enzymes.

RNA ligase was purified according to a procedure of Cranston et al.<sup>17</sup> with a modification using affinity chromatography on ADP-Sepharose.<sup>18</sup> Polynucleotide kinase reactions and other enzymes for characterization of products were described previously.<sup>6,8</sup>

## RNA ligase reaction.

Unless specified otherwise 50 mM HEPES-NaOH (pH 8.3), 10 mM MgCl<sub>2</sub>, 10 mM DTT, 0.1 mM ATP and 0.5-2 µg BSA were used in 10 µl at 25°. Substrate concentration was 0.28-1.5 mM when the enzyme concentration was 140 units/ml. Other methods.

Paper chromatography was performed by the descending technique using solvent systems: A, isopropyl alcohol-concentrated ammonia-water (7 : l : 2, v/v); B, n-propyl alcoholconcentrated ammonia-water (55 : l0 : 35, v/v); C, 0.1 M phosphate (Na, pH 6.8)-ammonium sulfate-n-propyl alcohol (l00 : 60 : 2, v/w/v). Paper electrophoresis was performed using 0.05 M triethylammonium bicarbonate (pH 7.5). Ion-exchange chromatography was performed with DEAE-Sephadex A25 (Pharmacia Co.) and triethylammonium bicarbonate. Homochromatography<sup>19</sup> was performed using Homomix I-V.<sup>20</sup> Partial digestion with Nuclease Pl and venom phosphodiesterase were as described previously.<sup>9</sup>

For the removal of the o-nitrobenzyl group, compounds were

placed in a pyrex tube (1.5 mm thick) and irradiated through a pyrex filter (2 mm thick) inserted in a water jacket. The photolysis apparatus had a 300 W high pressure mercury lamp (Eikosha Model PIH 300) with a quartz water circulating jacket.

## RESULTS

<u>3'-Labelling by phosphorylation with nbzl\*ppA (2) and RNA ligase</u>.

B

A trinucleotide U-A-G (<u>1</u>) was phosphorylated using a two fold excess of nbzl\*ppA and RNA ligase as shown in Chart 1. Chart 1. NO<sub>2</sub> U-A-G + Ap(5')\*pOCH<sub>2</sub>  $\longrightarrow$  U-A-G\*pnbzl  $\xrightarrow{h\nu}$  U-A-G\*p <u>1</u> <u>2</u> <u>4</u>

> Figure 1. Homochromatography of the products after phosphorylation of U-A-G with nbzl\*ppA. Before photoirradiation, U-A-G-\*pnbzl (the slower spot); after photoirradiation, U-A-G\*p (the slowest spot). nbzl\*ppA travels behind the blue marker.

UAG \*NB UAG

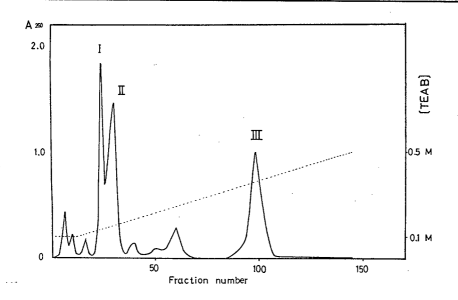


Figure 2. Chromatography of the products in the RNA ligase reaction (605  $\mu$ l) of nbzlppA (1025 nmol) with U-A-G (532 nmol) using RNA ligase (28 units) on a column (0.6 x 15 cm) of DEAE-Sephadex A25. Elution was performed with a linear gradient of triethylammonium bicarbonate (pH 7.5, 0.1 M to 0.5 M, total 220 ml). Fractions of 1.6 ml were collected every 10 min. Peaks: I, pA; II, nbzlppA; III, U-A-Gpnbzl.

The 3'-(o-nitrobenzyl) phosphorylated product (<u>3</u>) was detected by homochromatography. After 4hr, the reaction was almost complete. Fig. 1 shows <u>3</u> as a slower moving compound compared with the pyrophosphate (<u>2</u>) which travels behind the blue marker. Removal of the o-nitrobenzyl group from <u>3</u> was effected by irradiation with UV light of wavelength longer than 280 nm. The 3'-phosphorylated trinucleotide (<u>4</u>) travelled slightly slower than the benzylated starting material (<u>3</u>) as shown in Fig. 1.

Preparation of the 3'-modified trinucleotide and its use in joining reactions.

For joining of pU-A-G to the 3'-end of other oligonucleotides U-A-G was modified at the 3'-hydroxyl group before 5'phosphorylation. U-A-Gpnbzl (5) was prepared in quantity using a two fold excess of unlabelled nbzlppA and RNA ligase and isolated by chromatography on Sephadex as shown in Fig. 2. 5

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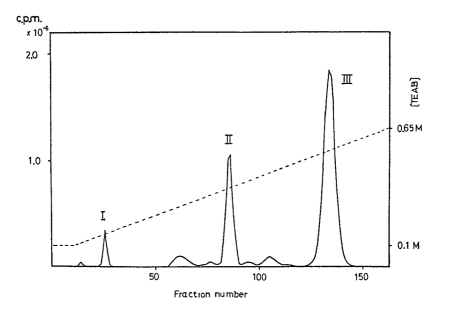


Figure 3. Chromatography of the product in the phosphorylation of U-A-Gpnbzl(292 nmol) with  $[\gamma^{-32P}]$ ATP(480 nmol) and polynucleotide kinase (15 units) on DEAE-Sephadex A25 using a linear gradient of triethylammonium bicarbonate pH 7.5 from 0.1 M to 0.65 M (total, 220 ml). Fractions of 1.6 ml were collected every 10 min. Radioactivity was counted by Cerenkof's method. Peaks: I, \*Pi; II, [ 32P]ATP; III, \*pU-A-Gpnbzl.

was phosphorylated using  $[Y-^{32}P]$ ATP with low specific activity by polynucleotide kinase. The 5'-phosphorylated product (6) was isolated by the similar chromatography (Fig. 3) and joined to A-U-G by treatment with RNA ligase as illustrated in Chart 2. The joined product (7) was analyzed by homochromatography and the reaction mixture was irradiated with UV light to remove the o-nitrobenzyl group. Fig. 4 shows mobilities of the 3'-phosphorylated products before and after photoirradiation. The deblocked hexanucleotide (8) was also isolated by ion-exchange chromatography on DEAE-Sephadex (Fig. 5) and characterized by nearest neighbor analysis (Fig. 6-1) as well as RNase A plus phosphatase treatment (Fig. 6-2). An aliquot of 8 was treated with phosphatase and 9 was isolated by paper chromatography in solvent C. The dephosphorylated hexanucleotide (9) could be an acceptor molecule in a subsequent ligation if the chain is

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Figure 4. Homochromatography of the products in the joining of \*pU-A-Gpnbzl to A-U-G. 1, \*pU-A-Gpnbzl; 2, \*pU-A-Gp; 3, A-U-G\*pU-A-Gp (the slowest spot, after photoirradiation); 4, A-U-G\*pU-A-Gpnbzl (the slowest spot, before photoirradiation).



• B

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to be elongated in the 3'-direction.

Thus U-A-G was joined to A-U-G by successive 3'-modification, 5'-phosphorylation, ligation and 3'-deblocking to yield A-U-G-U-A-G which could serve as a model messenger RNA in protein synthesizing systems.

Chart 2.

 $\begin{array}{ccc} U-A-Gpnbzl &+ [\gamma-{}^{32}P]ATP &\longrightarrow & *pU-A-Gpnbzl & \xrightarrow{A-U-G} \\ & \underline{5} & \underline{6} & & \\ RNA & ligase \\ A-U-G*pU-A-Gpnbzl & \xrightarrow{h\nu} & A-U-G*pU-A-Gp & \longrightarrow & A-U-G*pU-A-G \\ & & & & & 9 \end{array}$ 

The 3'-modification by o-nitrobenzyl phosphorylation of

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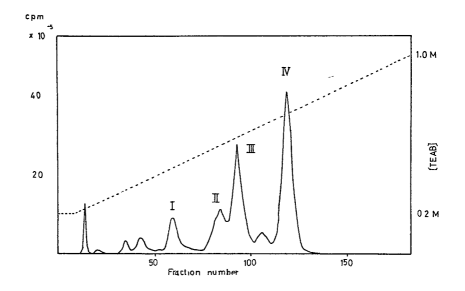


Figure 5. Chromatography of the deblocked product, A-U-G-U-A-Gp after joining of \*pU-A-Gpnbzl (277 nmol) to A-U-G (278 nmol) using RNA ligase (14 units in total volume of 450 µl) on a column of DEAE-Sephadex A25. The conditions for elution were the same as described in Fig. 2. Peaks: I, unidentified, II, mainly dephosphorylated product, A-U-G\*pU-A-G; III, the reaction intermediate, Ap(5')pU-A-Gp; IV, A-U-G\*pU-A-Gp.

C-C-A, A-U-C gave similar results. The conditions for these reactions are summarized in Table I.

#### DISCUSSION

The present study indicates that the adenylated o-nitrobenzyl phosphate (nbzlppA) was recognized by RNA ligase as a donor molecule and that o-nitrobenzyl phosphate was transfered to the 3'-hydroxyl group of ribooligonucleotides. This result is consistent with the previous finding that adenylated cyanoethyl phosphate served as a substrate in RNA ligase reactions.<sup>13</sup> The o-nitrobenzyl group was introduced as a photolabile protecting group for amino acids, carbohydrates and phosphates.<sup>21</sup> Extensive use of this group as protection for the 2'-hydroxyl group of nucleotides in oligonucleotide synthesis has shown no detectable photochemical side reactions during deblocking by irradiation with UV light of wavelength longer than 280nm.<sup>22</sup>

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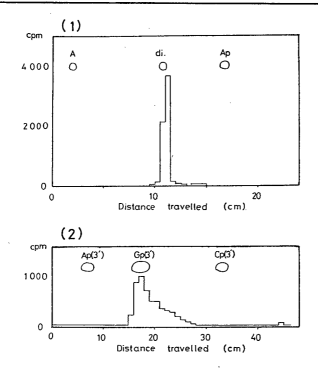


Figure 6. Characterization of A-U-G\*pU-A-G. (1) Paper electrophoresis of the digested product (G\*pU) after RNase A plus phosphatase treatment. A, Ap and dimer indicate markers detected by UV absorption (2) Paper chromatography of the digested product (G\*p) in an RNase M hydrolysis. Ap, Gp and Cp show markers detected by UV absorption.

#### Table I

The 3'-(o-Nitrobenzyl)phosphorylation of Ribotrinucleoside Diphosphate.

Acceptor	(nmol)	nbzlppA (nmol)	RNA ligase (unit)	Total vol. (µl)	Time (hr)	Temp. (C°)	Yield <sup>a</sup> %
C-C-A	14.1	70.4	7.0	50	26	25	82
C-C-A	14.1	70.4	5.3	50	17	37	66
U-A-G	14.1	70.4	7.0	50	26	25	49
U-A-G	139	705	14.0	100	4	25	100
U-A-G	530	1025	28.0	605	19	25	70
A-U-C	13.9	70.5	1.4	10	21	25	97

a, The yield was estimated by counting radioactivity of \*pX-Y-Z and \*pX-Y-Zpnbzl from homochromatogram after the reaction mixture was phosphorylated with  $[\gamma-^{32}P]$ ATP and polynucleotide kinase.

The o-nitrobenzyl group on the 3'-phosphate of oligonucleotides can therefore be removed to give the 3'-phosphomonoester end group. The present procedure provides a new method of 3'phosphorylation of ribooligonucleotides. When the radioactive pyrophosphate (nbzl\*ppA) was used in this reaction, the 3'-end of ribooligonucleotides was labelled with <sup>32</sup>P.

The 3'-o-nitrobenzylphosphorylation of oligonucleotides is also useful as method for modification of the 3'-end. If a 3'-modified oligonucleotide is phosphorylated at the 5'-hydroxyl group, a good donor molecule for RNA ligase reactions is obtained. During 5'-phosphorylation with polynucleotide kinase and ATP, a 3'-phosphate may be removed by an activity which co-chromatographs with polynucleotide kinase.<sup>23</sup> Although 3'phosphatase free kinase<sup>24</sup> may overcome this problem, it is desirable to protect the 3'-phosphate during 5'-kination. This type of reaction was examined in our kination of the trinucleotide (5). The 5'-phosphorylated product (6) was obtained in a high yield (Fig. 3).  $[\chi^{-32}P]$  ATP was used to simplify product analysis in both the phosphorylation and the subsequent RNA ligase reaction. 6 was joined with A-U-G to yield 7 and the protecting group was removed at this stage to yield 8 (Fig. 5). If the chain is to be elongated in the 5'-direction it can be retained for the next 5'-kination reaction. The hexanucleotide with a 3'-phosphomonoester (8) was dephosphorylated by treatment with alkaline phosphatase to obtain 9. By this treatment 9 can be elongated in the 3'-direction.

This 3'-modification by o-nitrobenzylphosphorylation provides a method for facile introduction of a 3'-phosphomonoester which can be removed at later stages by phosphatase treatment if necessary. Once ribotriplets are obtained by whatever means, they can be modified at the 3'-end without the need for addition of mononucleotides and used for RNA ligase reactions in a blockwise fashion. This technique would appear especially useful in synthesis of messenger RNA consisting of trinucleotide codons.

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# Total synthesis of a RNA molecule with sequence identical to that of *Escherichia coli* formylmethionine tRNA<sup>†</sup>

(chemical synthesis of tRNA fragments/RNA ligase/methionyl-tRNA synthetase)

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ABSTRACT A RNA molecule has been synthesized that is identical in sequence to Escherichia coli  $tRNA_f^{Met}$  except that it lacks the base modifications present in the E. coli tRNA. This was achieved by enzymatic joining of chemically synthesized oligonucleotides with chain lengths of 3-10 which were synthesized by the phosphodiester or phosphotriester method. First, guarter molecules of tRNA were constructed by joining of chemically synthesized fragments with RNA ligase. The 5'-quarter molecule (bases 1-20) served as an acceptor in joining reactions with the 3',5'-bisphosphorylated donor molecule (bases 21-34). The 5'-half molecule thus obtained was treated with phosphatase and joined to the 3'-half molecule which was prepared by ligation of the other quarter molecules (bases 35-60, acceptor; bases 61-77, donor) followed by 5'-phosphorylation with polynucleotide kinase. The synthetic tRNA was characterized by oligonucleotide pattern and was partially active in aminoacylation with E. coli methionyl-tRNA synthetase.

Chemical synthesis of nucleic acids has been a challenging problem in organic chemistry since the structure of the nucleic acids was elucidated. Chemical methods to synthesize short ribo- and deoxyribopolynucleotides with defined sequences were established in early 1960s, and those oligonucleotides were important in the elucidation of the genetic code (1). Discovery of DNA ligase allowed the synthesis of bihelical DNAs from chemically synthesized deoxyribopolynucleotides. With this chemical-enzymatic method the genes for yeast alanine tRNA (2) and *Escherichia coli* tyrosine tRNA precursor (3) have been synthesized; the latter was the first synthetic functional DNA molecule. Genes for peptides have also been synthesized by the same approach, and the methods for joining double-stranded DNA pieces with protruding ends have been used in various recently developed reactions for genetic manipulations.

Although tRNAs are the smallest nucleic acids with unique functions, their synthesis has been difficult until recently, mainly because of the lack of good synthetic methods for larger oligoribonucleotides as well as a lack of joining enzymes. After the primary structure of yeast alanine tRNA had been determined (4), the nona- and hexanucleotide corresponding to the terminal sequence of this tRNA were synthesized by phosphodiester block condensation. These fragments in turn were used to form reconstituted molecules with natural tRNA fragments derived by RNase digestions. However, aminoacylation was not possible because the synthetic fragments were too small to form sufficiently stable complexes for recognition by the alanyl-tRNA synthetase (5). The discovery of RNA ligase (6) and its ability to join single-stranded oligoribonucleotides (7) made it possible to elongate synthetic RNA fragments to yield larger molecules such as tRNAs.

The initiator methionine tRNA of prokaryotes has a special role in protein biosynthesis, which manifests itself in several unique properties of that tRNA (8). It was also the subject of detailed modification studies to explain its structure-function relationship (9). Because a RNase T1-digested one-quarter molecule of E. coli tRNA<sup>Met</sup> reconstituted methionine acceptor activity when mixed with the corresponding three-quarter molecules (10), this tRNA seemed an appropriate target for chemical synthesis. The final aim would be to modify systematically the functionally important parts of the molecule.

We began by synthesizing terminal fragments of the tRNA (11-16) and examined the ability of RNA ligase (17-19) to join these fragments. The 5'-terminal icosanucleotide (20), the tetradecanucleotide (bases 21-34) (21), and the 3'-heptadecanucleotide (22) have been obtained by this method. The 5'-quarter molecule here was found to reconstitute methionine acceptor activity when it was combined with the natural RNase T1-generated three-quarter molecule (20). Oligonucleotides corresponding to the rest of the molecule and certain of their analogs have been synthesized either by the phosphodiester method (23, 24) or by the triester method (25, 26).

In this paper we report total synthesis of a RNA molecule with a sequence identical to that of E. *coli* tRNA<sup>Met</sup><sub>f</sub> obtained by the enzymatic joining of chemically synthesized fragments with RNA ligase.

## MATERIALS AND METHODS

**Enzymes.** T4 RNA ligase was purified as described (27). Polynucleotide kinase and *E. coli* alkaline phosphatase were gifts of M. Sugiura. 3'-Phosphatase-free kinase was isolated from *E. coli* infected with T4 PseT1-amN82SP62 as described (20). Other enzymes for characterization of the products were obtained as described (17-20).

Kinase Treatment, Ligation, and Dephosphorylation. 5'-Phosphorylation by using polynucleotide kinase and  $[\gamma^{-3^2}P]ATP$ was performed as described (20). All 5'-phosphorylations of 3'phosphorylated oligonucleotides were performed by using the 3'-phosphatase-free kinase unless otherwise specified. Ligation was carried out in the presence of a 2-fold excess of ATP with respect to donor molecules in 50 mM Hepes (made pH 8.3 with NaOH)/10 mM dithiothreitol/10 mM MgCl<sub>2</sub>/10% (vol/vol) dimethyl sulfoxide containing bovine serum albumin at 10  $\mu$ g/ ml. The 3'-phosphate was removed by treatment with *E. coli* alkaline phosphatase in 50 mM Tris-HCl (5  $\mu$ l, pH 8.1) at 55°C for 30 min. The enzyme was inactivated by treatment with 1  $\mu$ l

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<sup>&</sup>lt;sup>†</sup>This is paper no. 36 in a series. Paper 35 is ref. 22.

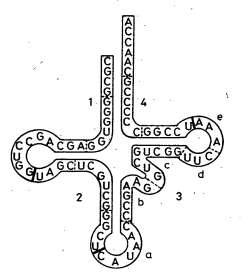


FIG. 1. Structure of E. coli tRNA<sup>fet</sup><sub>f</sub>. The quarter molecules 1, 2, and 4 were obtained by joining chemically synthesized oligonucleotides with RNA ligase. Quarter molecule 3 was joined as shown in Fig. 2.

of 40 mM EDTA at room temperature for 30 min and then at 100°C for 2 min (28) and extracted twice with 2  $\mu$ l of phenol saturated with 50 mM Tris HCl (pH 8.1). The phenol layer was washed twice with water (20  $\mu$ l) and the aqueous phase was subjected to gel filtration on a Sephadex G-50 column (1.1 × 21 cm).

Isolation and Characterization of Joined Products. Paper chromatography was performed with 0.1 M sodium phosphate; pH 6.8/ammonium sulfate/1-propanol, 100:60:2 (vol/wt/vol), as the solvent system. Paper electrophoresis was performed at 900 V/40 cm with 0.05 M triethylammonium bicarbonate (pH 7.5) or 0.2 M morpholinium acetate (pH 3.5). Homochromatography (29) was performed with Homo-mix I-IV (30). Twodimensional chromatography on cellulose plates was as described (31). Polyethylenimine-cellulose plates (Macherey-Nagel, Polygram Cell 300 PEI) were treated as described (32). Polyacrylamide gel electrophoresis was performed on slab gels or on a disc apparatus as described (20).

Nearest-neighbor analysis (17), 3'- and 5'-terminal analysis (20), and partial nuclease P1 digestion (22) for mobility shift analysis were as described. For complete RNase T1 digestion of the product (1 pmol), RNase T1 (1 unit) was used in the presence of phosphatase (180 microunits) in 10 mM Tris HCl (pH 7.5) at 27°C for 4 hr.

Aminoacylation of the Joined Product. The purified E. colimethionyl tRNA synthetase (a gift of J. P. Waller) was used at

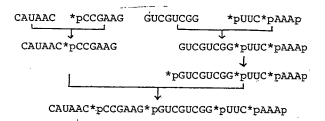


FIG. 2. Synthesis of quarter molecule 3 (bases 35-60).

4  $\mu$ g/ml for 10 pmol of the tRNA in 100 mM Hepes, pH 8.0/ 10 mM Mg(AcO)<sub>2</sub>/10 mM KCl/4 mM ATP/10 mM 2-mercaptoethanol/6  $\mu$ M L-[<sup>14</sup>C]methionine (582 Ci/mol; 1 Ci = 3.7  $\times$  10<sup>10</sup> becquerels) in total volume of 20  $\mu$ l at 37°C for 30 min. Aminoacylation with a crude mixture of E. coli synthetases (33) (0.12 mg/ml) was performed in the presence of 0.1 mM CTP at 37°C for 60 min. The reaction was stopped by addition of 1 M pyridinium acetate (pH 5.0; 5  $\mu$ l) and the mixture was applied to a column  $(0.8 \times 23 \text{ cm})$  of Sephadex G-50 equilibrated with 50 mM pyridinium acetate (pH 5.0). The aminoacylated tRNA was eluted with the equilibration buffer, assayed by Cerenkov's method, desalted by gel filtration on Sephadex G-50 in 0.05 M triethylammonium bicarbonate, and treated with Tris-HCl (pH 9.0) at 37°C for 1 hr to hydrolyze the amino acid. The mixture was applied to a column of Sephadex G-50 to resolve the tRNA and  $[^{14}C]$  methionine. Fractions were assayed using a scintillation counter. An aliquot (10 pmol) of the tRNA was assayed for methionine acceptor activity by acid precipitation as described (20) after <sup>32</sup>P radioactivity became negligible.

## RESULTS

Chemical Synthesis of tRNA Fragments. Most of the synthetic methods used for fragments shown in Fig. 1 have been described previously. The phosphodiester method was used for the synthesis of fragments consisting of bases 1-4 (12), bases 5-10 (13), bases 41-57 (23), bases 58-60 (24), and bases 61-71 (14, 23). The phosphotriester method was applied for synthesis of fragments consisting of bases 11-20 (15), bases 21-34 (unpublished work), bases 35-40 (26), and bases 72-77 (16).

Preparation of Quarter Molecules. Segment 1 (bases 1-20) was prepared by joining three fragments as described (20) and the 3'-terminal phosphate was removed. The next quarter, molecule 2 (bases 21-34), was synthesized by two different approaches (21).

Segment 3 (bases 35-60) was prepared by the joining of five synthetic fragments (Fig. 2). The dodecamer C-A-U-A-A-C-C-C-G-A-A-G (bases 35-46) was synthesized by using 10-fold excess of the acceptor molecule (3a, see Fig. 1) as summarized in

Table 1. Reaction conditions for joining of oligoribonucleotides with RNA ligase

Acceptor, nmol (µM)	Donor, nmol (µM)	АТР, <i>μ</i> М	Enzyme, µg/ml	Temp., °C	Time, hr	Isolated yield, %
CAUAAC	pCCGAAG					
80 (1000)	8 (100)	200	100	25	1	39
GUCGUCGG	pUUCAAAp				-	
15 (150)	10 (100)	200	140	25	1.	36
CAUAACCCGAAG.	pGUCGUCGGUU-				-	
2.1 (150)	CAAp, 1.5 (100)	200	115	25	1	52
1	2				-	~~
0.88 (74)	1.3 (111)	227	535	25	2	31
3	4				_	••
4 (100)	2.5 (63)	200	150	25	2	15
5'-half	p3'-half				-	10
0.14 (4.7)	0.20 (6.7)	100	200	4	17	42

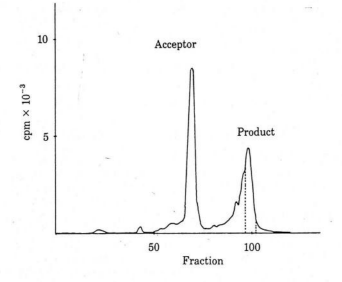


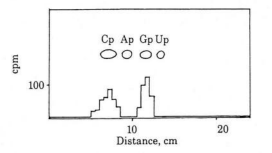
FIG. 3. Disk gel electrophoresis of the 26-nucleotide-long segment  $3(C-A-U-A-A-C^*pC-C-G-A^*pG-U-C-G-U-C-G-U-U-C-A-A-Ap)$  (Fig. 2) on 20% acrylamide. The first peak contained the acceptor (C-A-U-A-A-C^\*pC-C-G-A) and the last peak contained the product.

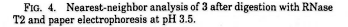
Table 1. The tetradecamer (bases 47–60) was synthesized by joining 3c to the hexanucleotide \*pU-U-C-A-A-Ap which had been obtained by phosphorylation with polynucleotide kinase and  $[\gamma^{-32}P]$ ATP of the joined product from U-U-C (3d) and pA-A-Ap (3e). After 5'-phosphorylation, the tetradecamer was joined to the dodecamer under the condition shown in Table 1. The 26-nucleotide-long segment 3 was isolated by electrophoresis on a 20% acrylamide gel disc as illustrated in Fig. 3. The nearest-neighbor analysis of the product is shown in Fig. 4. The chain length was confirmed by slab gel electrophoresis.

The 3'-phosphorylated heptadecamer 4 was prepared as described (22). The 3'-phosphorylation of C-A-A-C-C-A (16) was done with P1-adenosine-P2-(o-nitrobenzyl) pyrophosphate and RNA ligase (19).

Joining of Quarter Segments to Yield the tRNA Molecule. The 5'-half molecule was synthesized by joining of quarter molecules I and 2. The reaction conditions are summarized in Table I. The mixture was separated by polyacrylamide gel electrophoresis, and the product was detected by autoradiography (Fig. 5A). The extent of the reaction was 73% as measured by assaying gel slices at the appropriate positions. However, the isolated yield after elution from the gel was 31% based on 1.

For the synthesis of the 3'-half molecule, heptadecamer 4 was phosphorylated and joined to the 3 by using the conditions shown in Table 1. The product was isolated as described for the 5'-half molecule in a yield of 20%. It was characterized by nearest-neighbor and terminal analyses. The 3'-half was then 5'-





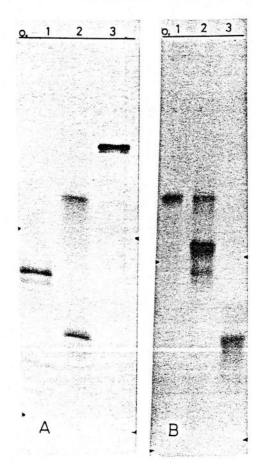


FIG. 5. Polyacrylamide gel electrophoresis of products. (A) Synthesis of the 5'-half molecule from 1 and 2 (lane 2). Lane 1: the 5' quarter molecule, 1. Lane 3: the three-quarter molecule. (B) Synthesis of the 3'-half molecule from bases 35-46 and bases 41-77 (lane 2). Lane 1: bases 35-77. Lane 3: bases 47-60.

phosphorylated with polynucleotide kinase and  $[\gamma^{-32}P]$ ATP and joined to the 5'-half molecule at reduced temperature (Table 1).

The reaction mixture was subjected to gel filtration (Fig. 6A). The joined product was found in peak 1 whereas the acceptor and the donor eluted together in peak 2. This was verified by polyacrylamide gel electrophoresis of the fractions. The molecular weight of the product was estimated to be  $2.6 \times 10^4$  from a plot of logarithm of molecular weight against mobility in polyacrylamide gel electrophoresis. As expected, 5'-end group analysis of the product after <sup>32</sup>P-labeling (with polynucleotide kinase and  $[\gamma^{-32}P]ATP$  and RNase T2 digestion yielded  $[^{32}P]pCp$ . The 3'-end analysis was performed by transferring the labeled 5'phosphate by circularization with RNA ligase followed by hydrolysis with RNase T2. C\*p (instead of A\*p) was identified as the 3'-end by two-dimensional chromatography. This may be due to removal of pAp by a reverse reaction of RNA ligase which has been observed with large excesses of the enzyme (R. I. Gumport and O. C. Uhlenbeck, personal communication). To avoid this side reaction, the 3'-half molecule was prepared (Fig. 5B) and the 3'-phosphate of the 43-unit segment was removed during the 5'-phosphorylation by using polynucleotide kinase with 3'-phosphatase activity. Although the presence of a 3'-OH group on the donor molecule could lead to the donor molecule joining onto itself, it was hoped that the secondary structure would prevent circularization of the 3'-half molecule (donor) during the joining reaction of the halves at the anticodon loop.

Joining of this 43-unit segment to the 5' half was performed under the same conditions as described above, and the product

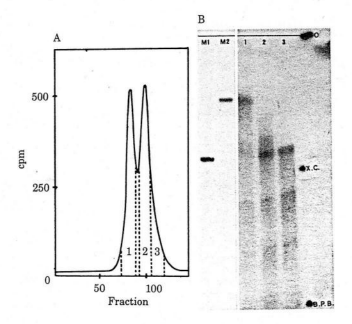


FIG. 6. (A) Gel filtration of products in the synthesis of the total molecule from the 3'- and 5'-half molecules on a column  $(0.7 \times 90 \text{ cm})$  of Sephadex G-200 equilibrated with 50 mM potassium phosphate; pH 7.5/0.1 mM EDTA. Elution was at 50°C with flow rate 1.2 ml/hr; 0.22-ml fractions were collected. Column volume, 34.6 ml: void volume, 11.5 ml. Peak 1 contained the total molecule; the half molecules were eluted in peaks 2 and 3. (B) Polyacrylamide gel electrophoresis of the compound in each peak. M1 and M2 indicate markers of segment *I* and tRNA<sup>Met</sup><sub>f</sub>, respectively. Lanes 1, 2, and 3 correspond to peak 1, 2, and 3.

was isolated in a yield of 17% by gel filtration. The joined product was analyzed as previously and again partial removal of pA from the 3' end was observed. Possibly, dephosphorylation did not go to completion during kination. The whole molecule was then treated with phosphatase to remove the 3'-phosphate residue and then phosphorylated at the 5' end with unlabeled ATP and polynucleotide kinase.

This tRNA molecule was tested for methionine acceptor activity by using L-[<sup>14</sup>C] methionine and purified or crude meth<sub>z</sub> ionyl-tRNA synthetase. Aminoacylation was measured by isolating the aminoacyl-tRNA formed by gel filtration in acidic medium followed by hydrolysis and quantitation of the [<sup>14</sup>C]methionine produced. With the purified enzyme aminoacylation was 6%; with the crude enzyme it was 4%.<sup>‡</sup> The deacylated recovered tRNA was then subjected to RNase T1 digestion for further structural analysis. The T1 fragments were labeled by phosphorylation (28) and mapped by two-dimensional thin-layer chromatography on PEI-cellulose. As shown in Fig. 7, the synthetic tRNA gave essentially the same pattern as the natural tRNA<sup>Met</sup><sub>i</sub>. The 3'-fragment \*pC-A-A-C-C-A (spot 1) was accompanied by \*pC-A-A-C-C (spot 1') in chromatography of the product.

#### DISCUSSION

Examination of tRNA structure-function relationship may lead to an understanding of an interesting example of the specific

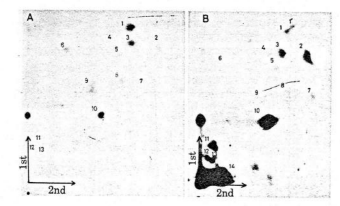


FIG. 7. Two-dimensional thin-layer chromatography of RNase T1 fragments of the natural  $tRNA_f^{Met}$  (A) and of the synthetic nascent molecule (B) on PEI-cellulose plates (20 × 20). The plates were irrigated with 1.4 M lithium formate, pH 3.5/7 M urea (or 10 cm, and then with 2.3 M lithium formate, pH 3.5/7 M urea (1st dimension) and with 0.6 M lithium chloride/20 mM Tris·HCl, pH 8.0/7 M urea (2nd dimension) (33). Spots: 1, pC-A-A-C-C-A; 1', pC-A-A-C-C; 2, ps<sup>4</sup>U-P or pU-G; 3, pC-G; 4, pC-A-G; 5, pA-G; 6, pC-C-C-G; 7, pD-A-G; 8, pU-C-G; 9, pC-U-C-G, pC-U-G, and pA-U-C-G; 10, Pi; 11, pCmU-C-A-U-A-A-C-C-G or pC-U-C-A-U-A-A-C-C-G; 12, pm<sup>7</sup>G-U-C-G; 13, pT- $\Psi$ -C-A-A-A-U-C-C-G or pU-U-C-A-A-A-U-C-C-G; 14, ATP.

recognition of a nucleic acid by a protein. Chemical modifications of tRNAs or genetic approaches to find mutants with base substitutions have been used previously for recognition studies. However, these approaches have certain limitations. Chemical synthesis should provide defined alterations which would be useful in structure-function relationship studies of tRNAs.

The chemical synthesis of oligoribonucleotides that have sequences of E. coli tRNAf and their analogs (e.g. U-G-C-G-G) (25) has provided suitable substrates for the construction of tRNA molecules by joining with RNA ligase. This paper reports the total synthesis of tRNA<sup>Met</sup> from synthetic oligonucleotides with chain lengths 3-10. Oligonucleotides containing modified bases can be joined to other synthetic fragments by methods similar to those described herein. Even though RNA ligase can join short oligonucleotides and is a convenient tool for substituting fragments, it would be desirable to reduce the number of joining steps so as to obtain tRNA molecules in sufficient quantity to provide enough material for biological studies. Chemical synthesis of oligonucleotides as long as quarter molecules of tRNA would yield whole molecules after three ligations. Recently, we synthesized an icosaribonucleotide corresponding to bases 35-54 of the tRNA<sup>Met</sup> by the phosphotriester method (unpublished data) and this fragment will be joined to fragments including modified bases to obtain tRNAs with partial modifications.

The tRNA synthesized in the present work is recognized to a limited extent by E. coli methionyl-tRNA synthetase. We do not know the tertiary structure of this tRNA. If modified nucleotides are necessary for forming the correct conformation required for synthetase recognition, then the low acceptor activity would be explained. Otherwise, a particular nucleotide modification may be required for direct interaction with the enzyme. Further synthetic investigation to identify the levels of modification that increase amino acid acceptor activities should aid in elucidating the mechanisms of these interactions.

We are indebted to Dr. M. Sugiura for suggestions and a gift of polynucleotide kinase and phosphatase. We thank Dr. J.-P. Waller for a gift of the purified methionyl-tRNA synthetase of *E. coli* and Oak Ridge National Laboratory for formylmethionine tRNA from *E. coli* K-12

<sup>&</sup>lt;sup>‡</sup> Acid precipitation of the amino acid carried by the aliquot of the synthetic tRNA (10 pmol) was measured after <sup>32</sup>P radioactivity became negligible with a control (-tRNA<sub>f</sub><sup>Met</sup>, 66 cpm); it was found to be 25/ cpm (0.32 pmol after subtraction of background) when the natural tRNA<sub>f</sub><sup>Met</sup>(10 pmol) accepted methionine (2866 cpm, 4.8 pmol). Thus, the aminoacylation of the synthetic tRNA was 6.7% with respect to the intact tRNA.

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conjugated anti-sheep serum (directed against the globulins in the sheep H3 HA antiserum) and rhodamine-conjugated antihamster serum (directed against SV40 large-tumor antiserum). These cells showed coordinate expression of the HA and largetumor antigen, indicating the specificity of the observed immunofluorescent reactions. Therefore, we conclude that the HA synthesized in HA–SV40-infected AGMK cells is expressed on the cell surface in the absence of influenza virus infection.

## DISCUSSION

We have described the construction of a recombinant viral genome consisting of an SV40 vector and a cloned full-length DNA coding for the HA protein of influenza virus. Infection of AGMK cells with this recombinant virus produced a putative HA polypeptide that was immunoprecipitable with HA antiserum. The polypeptide showed a molecular size corresponding to that of uncleaved influenza HA and it was glycosylated, as shown by incorporation of radioactive labeled sugars. Furthermore, the putative HA product appeared to be functionally active; extracts from cells infected with the HA-SV40 recombinant exhibited specific hemagglutination not seen in control cell extracts. The HA product of HA-SV40 exhibited, in the absence of other influenza viral functions, properties characteristic of a surface glycoprotein. These observations suggest that the HA product of HA-SV40 is similar to the HA synthesized in cells infected with influenza virus. This is expected as our cloned HA DNA contains full-length sequences that code for the entire polypeptide sequence. Analysis of amino acid sequences suggests that the HA molecule includes three functional domains: an NH<sub>2</sub>-terminal prepeptide signal for transport of the polypeptide from the cytoplasm to cell membranes, a COOH-terminal hydrophobic peptide for anchorage of the polypeptide in the cell membrane, and an internal "hinge" peptide region necessary for activation of viral infectivity through specific cleavage (30, 31. Our finding of HA synthesis demonstrates that amino acid sequences encoded by cloned HA DNA are sufficient for expression of the HA on the surface of eukaryotic cells. We did not observe, however, cleavage of the HA<sub>0</sub> into the HA<sub>1</sub> and HA2 subunits during infection with the HA-SV40 viral recombinant. This was not surprising as cleavage of HA<sub>0</sub> occurred at a low level in AGMK cells without added trypsin during a productive infection of these cells with influenza A virus (Fig. 2A).

Only one of the nine HA-SV40 isolates examined synthesized HA. There are several possible explanations for the failure of other isolates to produce the polypeptide. Some of the HA-SV40 recombinants may have sustained deletion of DNA sequences that are required for transcription or translation. Also, some HA-SV40 hybrids may contain HA DNA inserted in an opposite orientation so that the sense (+(HA RNA strand is not transcribed. In the latter case, the (-) HA RNA strand that is present in genomic RNA would be synthesized. Analysis of nucleotide sequences at the junctures of SV40 and HA DNA should help to differentiate between these possibilities.

Our HA DNA recombinant should be useful in elucidating several interesting properties of the influenza HA. Individual domains that specify polypeptide functions can be rigorously tested through introduction of deletions or site-specific mutations. The regions that are associated with cell-receptor binding and, similarly, the separate antibody-binding sites that are defined by classes of monoclonal antibodies can be dissected at the molecular level. Experiments involving phenotypic mixing should answer the question of whether hemagglutinin coded for by cloned DNA is expressed normally on the surface of viral particles. If so, it would then be possible to seek evidence for complementation between HA–SV40 recombinant DNA and influenza viral mutants defective in HA function.

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## CHEMICAL SYNTHESIS OF THE 5'-HALF MOLECULE OF *E.coli* tRNA<sub>2</sub><sup>Giy</sup>

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Abstract—A tritriacontanucleotide which has the sequence of the 5'-half molecule of *E.coli* glycine tRNA<sub>2</sub>, was synthesized by the phosphotriester method involving *p*-anisidate protection for the 3'-phosphate ends. Di- and trinucleotide units were prepared from 5'-dimethoxytrityl-2'-O-tetrahydrofuranyl-3'-O-(o-chlorophenyl)phosphoryl derivatives of uridine, N-benzolycytidine, N-benzolyadenosine and N-iso-butyrylguanosine by condensation with 3',5'-unprotected nucleosides followed by phosphorylation to give 3'-phosphodiester blocks. The 3'-terminal dimers and trimers were synthesized by using 3'-(o-chlorophenyl)phosphoro-*p*-anisidates instead of 3',5'-unprotected nucleosides. The 3'-phosphodiesters of oligonucleotides with a chain length of larger than 5 were obtained by removal of the 3'-phosphoro-*p*-anisidate with isoamyl nitrite. The 5'-dimethoxytrityl group was removed by treatment with zinc bromide under anhydrous conditions. Fragments were designed to use common dimer blocks and to reduce the step for 5'-deblocking of larger fragments. Finally a 3'-phosphodiester\_block with a chain length of 20 was condensed with a 5'-OH component (tridecanucleotide). The fully protected 33 mer was deblocked and purified by chromatography. The structural integrity of the product was confirmed by mobility shift analysis and complete digestion with RNase T2.

Chemical synthesis of ribooligonucleotides is an important subject in organic chemistry. It also provides a useful approach for studies on biological and physico-chemical properties of ribonucleic acids. In the early 1960s short oligomers such as all possible ribotriplets were synthesized by the phosphodiester method and used to elucidate the genetic code.<sup>1</sup> Later phosphodiester ribooligonucleotide blocks were prepared for the synthesis of tRNA fragments.<sup>2</sup> For the synthesis of larger oligonucleotides, the phosphotriester method becomes a method of choice with introduction of phenyl derivatives as protecting groups for internucleotidic phosphates and arenesulfonyl azolides as activating reagents for phosphodiester groups.<sup>3</sup> We have been synthesizing ribooligonucleotides either by the phosphodi- or triester method, and have performed the synthesis of E.coli formyl methionine tRNA by enzymatic joining of the chemically synthesized fragments with RNA ligase.<sup>4</sup> The methodology used in this study can be applied to replacement of functional parts of the tRNA molecule with synthetic oligonucleotides. Synthesis of larger fragments is advantageous in reducing enzymatic joining steps to construct modified tRNAs. Larger quantities of oligonucleotides, which can only be obtained chemically, are very useful for studies on interaction of nucleic acids with proteins, such as aminoacyl-tRNA synthetases.5 Completely chemical synthesis of RNA of the size of tRNA is a challenging subject in chemistry and has its own value. In the present paper we report a synthesis of a tritriacontanucleotide having the sequence of the 5'-half molecule (1-33) of *E.coli* tRNA<sub>2</sub><sup>Gly.6</sup> The synthesis involved phosphotriester block condensations using a combination of tetrahydrofuranyl<sup>7</sup> and dimethoxytrityl<sup>8</sup> groups for the 2'- and 5'-OH functions, respectively.9 The abbreviated scheme of the synthesis is shown in Fig. 1. The 5'-dimethoxytrityl group was shown to be removed selectively in the

presence of the 2'-O-tetrahydrofuranyl group on treatment with zinc bromide,<sup>9</sup> although this reaction required anhydrous conditions, in contrast to the conditions used with deoxyoligonucleotides.<sup>10</sup> In the phosphotriester synthesis, protection of the 2'-OH group is an essential problems and various groups have been used in combination with selectivity removable 5'-O-protecting groups. Tetrahydropyranyl,<sup>8</sup> 4-methoxytetrahydropyranyl,<sup>11</sup> tert-butyldimethylsilyl,<sup>12</sup> and *o*-nitrobenzyl<sup>13</sup> groups have been used for synthesis of larger ribooligonucleotides: oc-tadecamer,<sup>14</sup> nonadecamer<sup>15</sup> and eicosamer.<sup>16</sup> The present tetrahydrofuranyl group has previously been used and shown to have the properties required for a 2'-O-protecting groups, e.g. facile introduction, stability during synthesis and complete removal at the final stage.9,17-1

Preparation of di- and trinucleotide blocks. For the synthesis of larger oligonucleotides, condensation of protected oligonucleotides has obvious advantages. Preparation of oligonucleotides with phosphotriester internucleotidic phosphate requires two kinds of protecting groups for phosphates. One of those has to be removed selectively. A variety of combinations has been reported for the phosphotriester synthesis. Phenyl derivatives,<sup>20</sup> 2-cyanoethyl,<sup>21</sup> phenylethyl derivatives<sup>22</sup> and 5'-chloroquinolinyl<sup>23</sup> have been used for protection for internucleotide phosphates. As removable protecting groups, 2-cyanoethyl,<sup>24</sup> 2,2,2,-trichloroethyl,<sup>25</sup> anilido,<sup>26a</sup> and anisido<sup>16,26b</sup> groups are used at the terminal phosphate. In the synthesis 3'-(o-chlorophenyl)-p-anisido present phosphoryl derivatives (5) were synthesized by phosphorylation of 5'-dimethoxytrityl-2'-O-tetrahydrofuranylnucleosides (2) with o-chlorophenyl panisidophosphorochloridate26b (3) as the phosphorylating reagent followed by removal of the 5'-dimethoxytrityl group with zinc bromide as illustrated in Fig. 2 using the condensations described

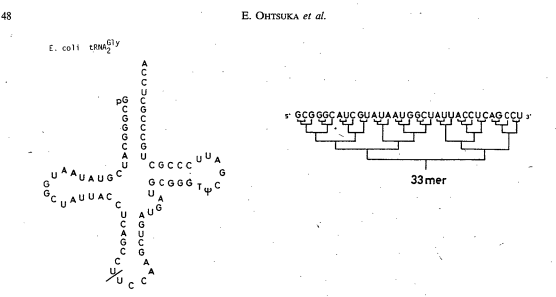
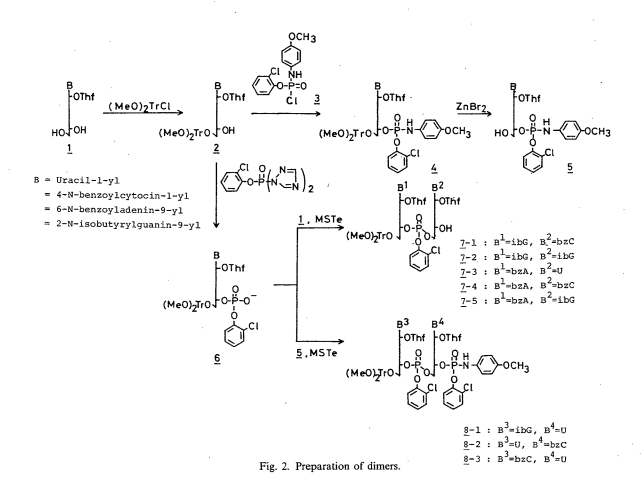


Fig. 1. Structure of the *E.coli* tRNA<sub>2</sub><sup>Giy</sup> and the 5'-half sequence (1-33).

previously.<sup>9</sup> Dimer units (7) were prepared by condensation of 3',5'-unprotected nucleoside (1) with 5'-dimethoxytrityl-2'-O-tetrahydrofuranyl-nucleoside 3'-(o-chlorophenyl)phosphates (6), which in turn were prepared by phosphorylation of 2 with ochlorophenyl bis-(1H-1,2,4-triazol-1-yl) phosphate<sup>27</sup> using mesitylenesulfonyl tetrazolide (MSTe, 1-(2,4,6trimethylbenzenesulfonyl)-1H-tetrazole).<sup>28</sup> For further elongation in the 3'-direction, 7 was phosphorylated. The terminal dimer blocks (8) were obtained by condensation of the 3'-phosphodiesters (6) with the 5'-free nucleotides (5). Reaction conditions for the preparation of dimers are summarized in Table 1. The dimers were isolated by chromatography on silica gel or alkylated silica gel.

Trimers were prepared from the above dimers by



Chemical synthesis of the 5'-half molecule of E.coli tRNA<sub>2</sub><sup>Gly</sup>

3'-Phospho- diester * Component (mmol)	5'-OH Component (mmol)	MSTe (mmol)	Time (min)	Product	Yield (%)
DT[G]OH (8.00)	HO[C]OH (11.23)	15.90	25	DT[GC]OH (7-1)	62
DT[G]OH (2.81)	HO[G]OH (3.93)	5.60	25	DT[GG]OH ( <u>7</u> -2)	72
DT[A]OH (10.13)	HO[U]OH (13.99)	19.91	25	DT[AU]OH ( <u>7</u> -3)	75
DT[A]OH (2.45)	НО[С]ОН (3.29)	4.71	35	DT[AC]OH ( <u>7</u> -4)	64
DT[A]OH (2.02)	HO[G]OH (2.59)	3.99	35	DT[AG]OH ( <u>7</u> -5)	88
DT[G]OH (2.00)	HO[U] <u>p</u> An (1.70)	3.40	25	DT[GU] <u>p</u> An ( <u>8</u> -1)	78
DT[U]ОН (1.50)	HO[C] <u>p</u> An (1.40)	2.96	25	DT[UC] <u>p</u> An ( <u>8</u> -2)	85
DT[C]OH (0.60)	HO[U]pAn (0.50)	1.01	30	DT[CU] <u>p</u> An ( <u>8</u> -3)	74

Table 1. Reaction conditions for the synthesis of dimers

DT =  $(Me0)_2$ Tr, An = NH $\phi$ -p-OCH<sub>3</sub>, <u>p</u> = o-chlorophenyl phosphate, [] = protected except for 3' and 5' termini, \* : DT[N]OH was converted to 3'-phosphodiester component by

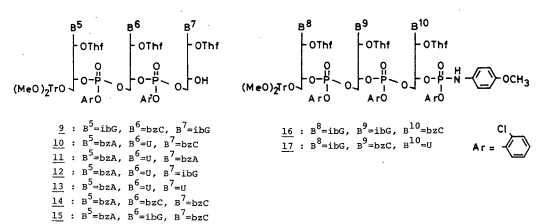
phosphorylation with o-chlorophenyl phosphoroditriazolide followed by treatment with H<sub>2</sub>0.

essentially the same procedure. Structures of the trimers are shown in Fig. 3 and reaction conditions are summarized in Table 2. The trimers were separated by reversed phase chromatography on alkylated silica gel. These dimers and trimers were designed to have purines at the 5'-terminal position, since removal of the dimethoxytrityl group of purine nucleosides was found to be easier.

Synthesis of tritriacontemer by condensation of oligonucleotide blocks. All 3'-diesterified trinucleotide intermediates used in this synthesis were prepared by phosphorylation of trinucleoside diphosphates listed in Table 2. As shown in Fig. 4 the 5'-OH components were derived by removal of the 5'-dimethoxytrityl groups. Yields and conditions of these conversions are summarized in Table 3. Reaction conditions of these nucleotide blocks are given in Table 4. The 3'-phosphoro-p-anisidate of penta- and hexanucleotides were converted to the phosphate by treatment with isoamyl nitrite when elongation was in the 3'-direction. The eicosamer (48) was obtained as the 3'-phosphodiester form by elongating the chain in the 3'-direction. It was activated by the condensing reagent to react with the tridecamer (49). Removal of the 5'-dimethoxytrityl of larger oligonucleotides was avoided as far as possible. Complete removal of the 5'-protecting group of the hexamers and tridecamer was not intended. Unchanged dimethoxytritylated oligonucleotides were recovered when the reaction had been slow. Yields listed in Table 4 varied mainly due to decomposition during chromatography on reversed-phase support. The eicosamer (46) was partially lost by conversion to polar compounds. The fully protected tritriacontamer (50) was not purified by chromatography. It was collected by preparation and isolated after deblocking.

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Deblocking of the product was performed by a procedure similar to that reported previously,<sup>9</sup> by treatment with: (1) isoamyl nitrite in pyridine-acetic acid (5:4), (2) 0.5 M 1,1, 3,3-tetramethylguanidinium *syn*-pyridine-2-carboaldoximate (TMG-PAO),<sup>29</sup> (3) ammonium hydroxide, (4) anion-exchange resin Dowex 50 W  $\times$  2 (pyridinium form), and (5) dilute hydrochloric acid (pH 2). The deblocked product was



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Table	2.	Reaction	conditions	for	the	synthesis	of	trimers	
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3'-Phospho- diester Component*	(mmol)	5'-OH Component	(mmol)	MSTe (mmol)	Time (min)	Product		Yield (%)
DT[GC]OH	(1.72)	НО[G]ОН	(2.43)	3.45	30	DT[GCG]OH	( <u>9</u> )	76
DT[AU]OH	(1.72)	но[с]он	(2.42)	3.23	30	DT[AUC]OH	( <u>10</u> )	72
DT[AU]OH	(1.08)	НО[А]ОН	(1.43)	2.04	30	DT[AUA]OH	( <u>11</u> )	69
DT[AU]OH	(2.92)	H0[G]0H	(4.21)	6.03	25	DT[AUG]OH	( <u>12</u> )	73
DT[AU]OH	(0.66)	но[и]он	(0.93)	1.30	30	DT [AUU]OH	( <u>13</u> )	59
DT[AC]OH	(1.35)	H0[C]0H	(1.89)	2.71	30	DT[ACC]OH	( <u>14</u> )	58
DT[AG]OH	(0.56)	H0[C]0H	(0.79)	1.23	25	DT[AGC]OH	( <u>15</u> )	70
DT[GG]OH	(1.66)	HO[C] <u>p</u> An	(1.67)	3.34	25	DT[GGC]pAn	( <u>16</u> )	73
DT[GC]OH	(3.04)	HO[U] <u>p</u> An	(3.05)	6.02	30	DT[GCU]pAn	( <u>17</u> )	76

DT =  $(Me0)_2$ Tr, An =  $NH\phi$ -p-OCH<sub>3</sub>, <u>p</u> = o-chlorophenyl phosphate, [] = protected except

for 3' and 5' termini, \* : DT[NN]OH was converted to 3'-phosphodiester component by

phosphorylation with o-chlorophenyl phosphoroditriazolide followed by treatment with H<sub>2</sub>0.

separated by gel filtration on Sephadex G-50. Figure 5 shows profiles of gel filtration of the eicosamer (20 mer) and tritriacontamer (33 mer). The product were analyzed by reversed-phase high pressure liquid chromatography (HPLC) and the 33 mer was found to be contaminated with partially protected compounds. Acid treatment for removal of the 2'-O-tetrahydrofuranyl group was repeated and the product was fractionated by reversed-phase HPLC after gel filtration as shown in Fig. 6. The fractionated products were found to be homogeneous (Fig. 7).

The 20 and 33 mer were identified by analysis of the chain length (Fig. 8) and mobility shift method (Fig. 9). Figure 8 shows a radio-autograph of a 20%

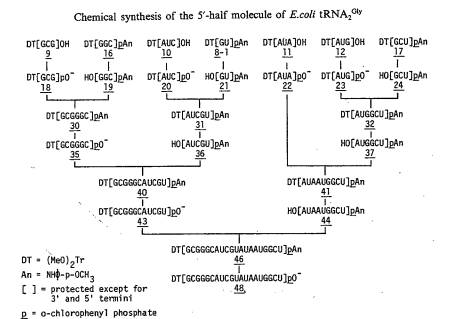
polyacrylamide gel electrophoresis<sup>30</sup> of the 5'-labeled products. For mobility shift analysis<sup>31</sup> and polymers were labeled at either end with polynucleotide kinase<sup>32</sup> and  $[\gamma^{-32}P]ATP$  or with RNA ligase plus 5'-labeled pCp.<sup>33</sup>

Complete removal of the protecting groups and maintaining of the 3'-5' internucleotide linkages were confirmed by complete digestion with RNase  $T2^{34}$  followed by labeling with polynucleotide kinase and  $[\gamma^{-32}p]$  ATP. The results of two dimensional thin layer chromatography on cellulose<sup>35</sup> of the mixture of 5'-labeled pNp are shown in Fig. 10. This test indicated that the 20 mer and 33 mer were digested with RNase T2 to give nucleoside 3'-phosphates.

Subst (mr	rate 101)	1M ZnBr2*	Time (min)	Product	Yield (%)
16	(1.195)	40	2.5	H0[GGC] <u>p</u> An ( <u>19</u> )	67
<u>8</u> -1	(1.289)	40	3	HO[GU] <u>p</u> An (21)	84
<u>17</u>	(2.250)	70	2.5	HO[GCU] <u>p</u> An ( <u>24</u> )	68
<u>8</u> -2	(1.146)	30	20	HO[UC] <u>p</u> An ( <u>27</u> )	73
<u>8</u> -3	(0.354)	10	12	HO[CU] <u>p</u> An (29)	76
<u>31</u>	(0.791)	35	2	HO[AUCGU] <u>p</u> An ( <u>36</u> )	70
<u>32</u>	(1.129)	50	1	HO[AUGGCU]pAn ( <u>37</u> )	57
<u>33</u>	(0.603)	30	5	HO[ACCUC] <u>p</u> An ( <u>38</u> )	- 81
<u>34</u>	(0.221)	11	5	H0[AGCCU] <u>p</u> An ( <u>39</u> )	67
<u>41</u>	(0.380)	25	2	HO[AUAAUGGCU] <u>p</u> An ( <u>44</u> )	56
<u>47</u>	(0.064)	10	2	HO[AUUACCUCAGCCU]pAn (49)	66

Table 3. Removal of the 5'-dimethoxytrityl group

An = NH $\phi$ -p-OCH<sub>3</sub>, <u>p</u> = o-chlorophenyl phospate, [] = protected except for 3' and 5' termini, \*; CH<sub>2</sub>Cl<sub>2</sub>:iso-PrOH = 85:15.



DT[CU]pAn DT[AGC]OH DT[AUU]OH DT[ACC]OH DT[UC]pAn <u>8</u>-3 15 8-2 14 13 HO[CU]pAn DT[AGC]p0 HO[UC]pAn DT[AUU]p0 DT[ACC]<u>p</u>0 28 25 26 DT[AGCCU]pAn DT[ACCUC]pAn <u>34</u> 33 HO[AGCCU]pAn HO[ACCUC]pAn 39 38 DT[AUUACCUC]pAn 42 DT[AUUACCUC]p0 45 DT[AUUACCUCAGCCU]pAn 47 HO[AUUACCUCAGCCU]pAn 49 DT[GCGGGCAUCGUAUAAUGGCUAUUACCUCAGCCU]pAn <u>50</u> deblocking GCGGGCAUCGUAUAAUGGCUAUUACCUCAGCCUp 51

Fig. 4. Synthesis of the 33 mer.

#### EXPERIMENTAL

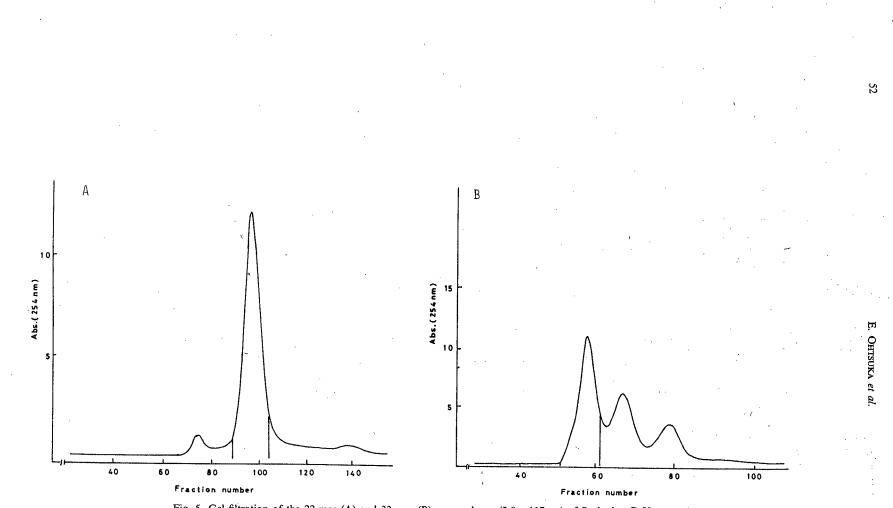
TLC was performed on plates of silica gel (Kieselgel 60 HF<sub>254</sub>, Merck) using a mixtures of CHCl<sub>3</sub> and MeOH. For reversed phase TLC (RPTLC), silanized silica gel (Kieselgel 60 HF<sub>250</sub> Silanisiert, Merck) was used with a mixture of acetone-water. For columns, silica gel (type 60 or 60 H, Merck) was used with a mixtures of CHCl<sub>3</sub>-MeOH. For preparative reversed phase chro-

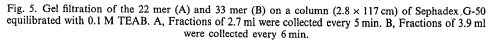
matography, alkylated silica gel (C-18,  $35-105 \mu$ , Waters) was packed with 60-70% acetone and elution was performed with a gradient of acetone (60-80%) in 0.2% aqueous pyridine. HPLC was carried out on an Altex 332 MP apparatus using a reversed phase column (TSK-LS410, Toyosoda).

Two dimensional homochromatography<sup>31</sup> was performed as described previously.<sup>36</sup>

Triethylammonium bicarbonate (TEAB) buffer (pH 7.5)

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Chemical synthesis of the 5'-half molecule of E.coli tRNA2 Gly

3'-Phos diester Componer	oho- nt (mmol)	5'-OH Compone	ent (mmol)	MSTe (mmol)	Time (min)	Product (Chain length)	Yield (%)
<u>9</u> a)	(0.903)	<u>19</u>	(0.791)	1.988	40	<u>30</u> (6 mer)	59
<u>10</u> a)	(1.061)	<u>21</u>	(1.060)	2.620	40	<u>31</u> (5 mer)	79
<u>12</u> a)	(1.702)	24	(1.492)	3.702	35	<u>32</u> (6 mer)	77
<u>14</u> a)	(0.778)	<u>27</u>	(0.794)	1.660	25	<u>33</u> (5 mer)	84
<u>15</u> a)	(0.365)	<u>29</u>	(0.277)	0.650	35	<u>34</u> (5 mer)	82
<u>30</u> b)	(0.456)	36	(0.466)	1.008	40	<u>40</u> (11 mer)	49
<u>11</u> a)	(0.675)	<u>37</u>	(0.646)	1.693	40	<u>41</u> (9 mer)	74
<u>13</u> a)	(0.352)	38	(0.356)	1.062	55	<u>42</u> (8 mer)	64
<u>40</u> b).	(0.222)	<u>44</u>	(0.183)	0.720	50	<u>46</u> (20 mer)	22
<u>45</u>	(0.155)	<u>39</u>	(0.146)	0.464	55	<u>47</u> (13 mer)	63
<u>46</u> b)	(0.020)	<u>49</u>	(0.020)	0.187	90	<u>50</u> (33 mer)	

Table 4. Conditions for block condensation

a) : DT[NNN]OH was converted to 3'-phosphodiester component by phosphorylation with

o-chlorophenyl phosphoroditriazolide followed by treatment with H<sub>2</sub>O.

b) : Fully protected ribooligonucleotide was converted to 3'-phosphodiester component

by treatment with isoamyl nitrite.

was used to wash organic layers containing protected nucleotides.

Dinucleoside monophosphates 7; Table 1, Example (7-1)

General methods for phosphorylation and condensation. 5'-Dimethoxytrityl-2'-O-tetrahydrofuranyl-N-isobutyrylguanosine (2, B = ibG; 5.805 g, 7.998 mmol) was dried by evaporation of pyridine and dissolved in pyridine (3 ml).

o-Chlorophenyl phosphoroditriazolide (12.0 mmol in 40 ml of dioxane) was added and the mixture was shaken for 20 min at 30°. TLC and RPTLC showed disappearance of the starting material. After 25 min, 0.1 M TEAB (150 ml) and pyridine (70 ml) were added. The product (6) was extracted with CHCl<sub>3</sub> (150 ml). The aqueous phase was reextracted with CHCl<sub>3</sub>-pyridine (3:1, 40 ml) and the combined organic layer was washed twice with TEAB (150 ml).

After evaporation of organic solvents the residue was dried by evaporation of pyridine 3 times and mixed with 2 (B = bzC) (higher isomer, 4.686 g, 10.15 mmol). The mixture was dried as above and treated with MSTe (4.012 g, 15.90 mmol) in pyridine (40 ml) at 30° for 15 min. Completion of the reaction was confirmed by TLC and RPTLC. After 25 min, water (3 ml) was added and evaporated. The residue was dissolved in CHCl<sub>3</sub> and washed with sat NaHCO<sub>3</sub> aq. The product (7-1) was separated by chromatography on silica gel (Kieselgel 60 H, 150 g,  $\phi$ 10 × 5.4 cm) using a gradient MeOH in CHCl<sub>3</sub>, and precipated with hexane from its soln in CHCl<sub>3</sub>, yield was 62%, 6.512 g, 4.95 mmol.

Dinucleotide 8. Table 1, Example (8-2). Compound 2 (B = U) (0.927 g, 1.50 mmol) was phosphorylated as described above and 6 (B = U) was condensed with 5 (B = BzC) (0.996 g, 1.40 mmol) in the presence of MSTe

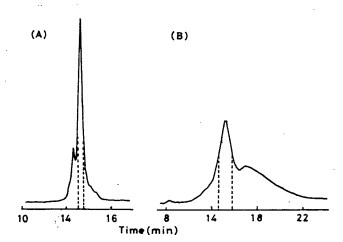
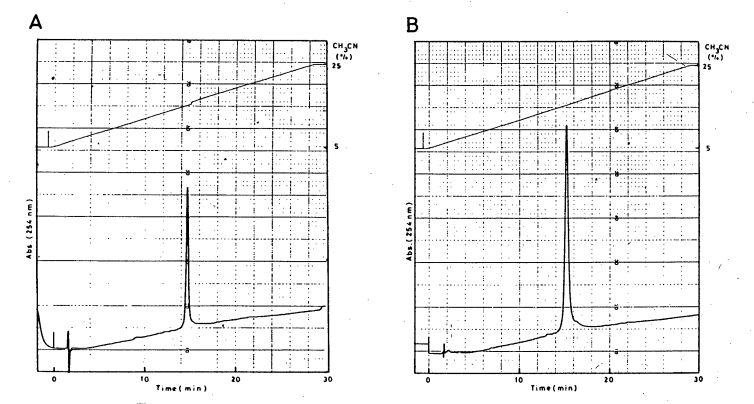
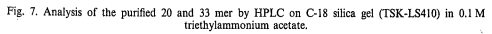


Fig. 6. Purification of the 22 mer (A) and 33 mer (B) by HPLC on silica gel (TSK-LS410) with a flow rate of 2 ml/min. A, a linear gradient of acetonitrile (5-25% during 30 min) in 0.1 M triethylammonium acetate. B, a linear gradient of acetonitrile (11-15% during 03 min) in 0.1 M triethylammonium acetate.

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Chemical synthesis of the 5'-half molecule of E.coli tRNA2 Gly

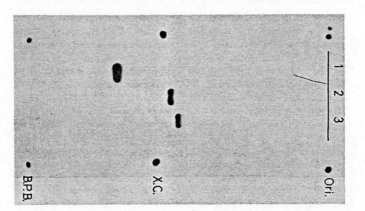


Fig. 8. Gel electrophoresis of the 20 mer (1), 33 mer (2) and a marker (3, 34 mer from the *Ecoli* tRNA<sub>f</sub><sup>Met</sup>) on 10% polyacrylamide.

(0.746 g, 2.96 mmol) at 30° for 15 min. The reaction was checked by TLC and RPTLC. After 25 min water (2 ml) was added and the mixture was concentrated. The residue was dissolved in CHCl<sub>3</sub> and washed twice with sat NaHCO<sub>3</sub> aq (70 ml). The product (8-2) was isolated by reversed phase chromatography on C-18 silica gel ( $\phi$  4 × 9.5 cm) using a gradient of acetone in 0.2% pyridine and precipated with hexane from its soln in CHCl<sub>3</sub>. The yield was 85%, 1.789 g, 1.193 mmol.

Synthesis of the trimers (9–17) Table 2. Trimers (9–15) were synthesized from 7 by phosphorylation as described for the preparation of 6 followed by condensation N, 2'-protected nucleosides (1). For the synthesis of trinucleotides (16, 17), nucleotides (5) were used. Synthetic procedures were the same described for dimers (7, 8) and trimers were isolated by reversed phase chromatography on C-18 silica gel ( $\phi 4 \times 9$  cm) using a gradient of acetone in 0.2% pyridine.

Removal of the 5'-dimethoxytrityl group (Table 3). The trinucleotide (protected GCUp, 17) (4.723 g, 2.25 mmol) was dried by evaporation three times with pyridine, three times with toluene and shaken with 70 ml of 1 M ZnBr<sub>2</sub> in CH<sub>2</sub>Cl<sub>2</sub>-PrOH (85:15, v/v) for 2.5 min at room temp. A soln of 1 M ammonium acetate (200 ml) was added with shaking and the product was extracted with CH<sub>2</sub>Cl<sub>2</sub> (100 ml). The organic soln was washed twice with 1 M ammonium acetate (100 ml) and concentrated. The residue was applied to a column ( $\phi$  5 × 5.5 cm) of Kieselgel 60 H (40 g). The oligonucleotide was eluted with a gradient of MeOH in CHCl<sub>3</sub> and precipated with hexane from its soln in CHCl<sub>3</sub>. The yield was 68%, 2.728 g, 1.521 mmol.

#### Condensation of oligonucleotide blocks

(1) The nonanucleotide (41). The trimer 11 (1.245 g, 0.675 mmol) was phosphorylated by the procedure described for the synthesis of 6, and condensed with 37, (3.254 gm 0.646 mmol) using MSTe (4.27 mg, 1.69 mmol) at 30° for 25 min. Completion of the reaction was checked by TLC and RPTLC and the product was isolated by reversed phase chromatography as described for the trimers.

(2) The undecamer 40 (Removal of the anisido group). The protected 30 (1.817 g, 0.456 mmol) was treated with isoamyl nitrite (3.1 ml, 23 mmol) in pyridine-AcOH (5:4, 14 ml) at 30° for 5 hr. The mixture was added with 0.2 M TEAB (100 ml), pyridine (75 ml), ether-pentane (1:1, 100 ml) on the aqueous phase was washed with ether-pentane (1:1, 100 ml). The product (35) was extracted with CHCl, (140 ml), washed three times with 0.2 M TEAB (100 ml), applied to a column ( $\phi 4 \times 7$  cm) of C-18 silica gel and eluted with a gradient of acetone in 0.2% pyridine. The

hexamer 35 was collected, dried by evaporation with pyridine and condensed with 36 (1.348 g, 0.466 mmol) using MSTe (254 mg, 1.01 mmol) at 30° for 40 min. The product was isolated by reversed phase chromatography as above and precipated with pentane. The yield was 49% 1.505 g, 0.223 mmol. The *R*, values in TLC (10:1) and RPTLC (7:3) were 0.29 and 0.23, respectively.

(3) The tridecamer (47). The octamer 45 (0.756 g, 0.155 mmol) and 39 (0.437 g, 0.146 mmol) were condensed using MSTe (0.117 g, 0.464 mmol) at 30° for 55 min and the product was isolated by reversed phase chromatography on a column ( $\phi$  3 × 8 cm) of C-18 silica gel. The  $R_f$  values in TLC (10:1) and RPTLC (8:2) were 0.43 and 0.70, respectively.

(4) The eicosamer (46). The undecamer 43 and 39 were condensed using conditions shown in Table 4 and 46 was isolated first by reversed phase chromatography on a column ( $\phi 4 \times 7 \text{ cm}$ ) of C-18 silica gel. The product 46 was purified by chromatography on a column ( $\phi 3 \times 2.8 \text{ cm}$ ) of silica gel (Kieselgel 60 H, 7 g) using a gradient of MeOH in CHCl<sub>3</sub>. The yield was 22%, 0.481 g, 0.041 mmol. The  $R_f$  values in TLC (10:1) and RPTLC (8:2) were 0.35 and 0.62, respectively.

(5) The tritriacotamer (50). The eicosamer 46 (239 mg, 0.020 mmol) was treated with isoamyl nitrite (0.15 ml, 1.11 mmol) in pyridine-AcOH (5:4, 1.5 ml) at 30° for 5.5 hr. Completion of the reaction was checked by TLC and RPTLC. The eicosamer 48 was extracted with CHCl<sub>3</sub>pyridine (2:1, 45 ml), washed 4 times with 0.2 M TEAB (50 ml) and precipated with ether-pentane (1:4, 50 ml) from its soln in CHCl<sub>3</sub> (2.5 ml). The eicosamer 48 was reprecipated, dried by evaporation with pyridine and condensed with 49 (150 mg, 0.020 mmol) in pyridine (0.5 ml) using MSTe (27 mg, 0.11 mmol) at 30° for 50 min. Starting materials were detected after 50 min in TLC and RPTLC. The mixture was treated with MSTe (20 mg, 0.079 mmol) for 40 min and added with water (0.5 ml). The product was extracted with CHCl<sub>3</sub>-pyridine (5:2, 70 ml), washed twice with 0.1 M TEAB (50 ml) and precipated with pentane from its soln in CHCl3. The yield of the crude product was 463 mg.

The deblocked 20 mer. The protected 46 (25 mg, 0.002 mmol) was treated with isoamyl nitrite (0.027 ml, 0.2 ml) in pyridine-AcOH (5:4, 0.3 ml) at 30° for 5 hr and the product was extracted with CHCl<sub>3</sub>-pyridine (2:1, 30 ml). The eicosamer was precipated with ether-pentane (1:4, 50 ml), treated with 1 M TMG-PAO in dioxane (2 ml) and water (2 ml) at 30° for 60 hr. The volatile materials were removed and the residue was dissolved in pyridine (2 ml). The mixture was treated with conc ammonia (10 ml) at 55°

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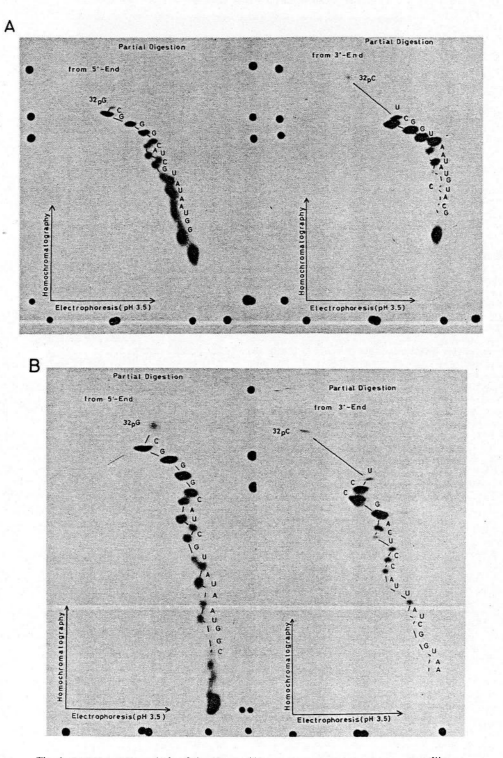


Fig. 9. Mobility shift analysis of the 20 mer (A) and 33 mer (B) using Homo-mix L<sup>31b</sup>

for 6 hr and concentrated. The product was dissolved in aqueous pyridine (30%, 20 ml) and passed through a column (5 ml) of Dowex 50 W  $\times$  2 (pyridinium form). The column was washed with 30% pyridine (100 ml) and combined solns were concentrated. The residue was dissolved in 0.1 M TEAB (50 ml), washed twice with ether (40 ml), evaporated

three times with added toluene and mixed with 0.1 N HCl (15 m). The solution was adjusted to pH 2 with 0.1 N HCl, kept at 25° for 9 hr, neutralized with 0.1 M ammonium hydroxide, washed twice with ether (40 ml) and concentrated. The residue was applied to a column of Sephadex G-50 (Fig. 5a) and the product in peak I (272  $A_{260}$ ) was

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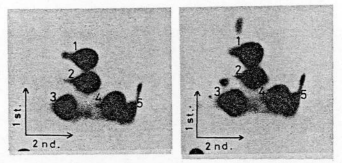


Fig. 10. Two dimensional TLC of mononucleotides obtained by digestion with RNase T2 of the 20 mer (A) and 33 mer (B) Spots: 1, Ap, 2, Cp; 3, Gp; 4, Up; 5, Pi. Solvents: 1st dimension, isobutyric acid-0.5 M ammonium hydroxide (5:3, v/v); 2nd dimension, isopropanol-conc. hydrochloric acid-water (70:15:15, v/v).

collected. An aliquot (6.8  $A_{260}$ ) was subjected to HPLC (TSK LS 410) (Fig. 6) and 1.7  $A_{260}$  units of the pure eicosamer was obtained. The estimated yield from 46 was 16% assuming  $\epsilon$  of the eicosamer being 20 × 10<sup>4</sup>.

The deblocked 33 mer (51). The protected 33 mer 50 (crude, 77 mg) was treated with isoamyl nitrite (0.018 ml, 0.6 mmol) in pyridine-AcOH (1:1, 1 ml) at 30° for 6 hr and treated with 0.5 M TMG-PAO (16 ml) using procedures described for the deblocking of the 20 mer. The 33 mer was then treated with conc ammonia (20 ml) at 55° for 6 hr, concentrated and passed through a column of Dowex 50 W  $\times$  2 (10 ml, pyridinium form). Acid treatment was performed as described for the 20 mer using 0.1 N HCl (30 ml) and 0.01 N HCl. The product was applied to gel filtration (Fig. 5b) and fractions containing the 33 mer were combined (234 A<sub>260</sub>). HPLC analysis showed incomplete removal of protecting groups. The product (220 A260) was retreated at pH in HCl (15 ml) at 25° for 8 hr and neutralized. The 33 mer was subjected to gel filtration and a part of the product (fraction No. 65, 18 A260) was further purified by HPLC (TSK-LS 410) as shown in Fig. 6.

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