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<td>Kawai, Fusako</td>
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<td>Citation</td>
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STUDIES ON
TRANSGLYCOSIDATION TO
VITAMIN B₆ AND PANTOTHENIC ACID
BY MICROORGANISMS

FUSAKO KAWAI

Department of Agricultural Chemistry
Faculty of Agriculture
Kyoto University

1972
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<th>Abbreviation</th>
<th>Full Form</th>
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<tr>
<td>PIN</td>
<td>Pyridoxine</td>
</tr>
<tr>
<td>PAL</td>
<td>Pyridoxal</td>
</tr>
<tr>
<td>PAM</td>
<td>Pyridoxamine</td>
</tr>
<tr>
<td>PAL-P</td>
<td>Pyridoxal-5'-phosphate</td>
</tr>
<tr>
<td>PIN-G</td>
<td>Pyridoxine glucoside</td>
</tr>
<tr>
<td>PIN-4'-G</td>
<td>Pyridoxine-4'-α-D-glucoside</td>
</tr>
<tr>
<td>PIN-5'-G</td>
<td>Pyridoxine-5'-α-D-glucoside</td>
</tr>
<tr>
<td>PaA</td>
<td>Pantothenic acid</td>
</tr>
<tr>
<td>PaA-G</td>
<td>Pantothenic acid glucoside</td>
</tr>
<tr>
<td>PaA-α-G</td>
<td>Pantothenic acid-α-D-glucoside</td>
</tr>
<tr>
<td>PaA-β-G</td>
<td>Pantothenic acid-β-D-glucoside</td>
</tr>
</tbody>
</table>
INTRODUCTION

In 1934, vitamin B₆ was first demonstrated by György as a rat pellagra preventive factor.¹) The apparent and specific chemical properties of vitamin B₆ (PIN) were definitely established by Birch and György.²) Within approximately two years of their publication, five independent groups reported, announcing the isolation of PIN. Lepkovsky,³) Keresztesy and Stevens,⁴) György,⁵) Kuhn and Wendt,⁶) and Ichiba and Michi⁷) had succeeded in isolating PIN as a crystalline form from various natural materials. Subsequently, its chemical constitution was recognized by two independent groups, Stiller et al.⁸) and Kuhn et al.⁹⁻¹¹) The synthesis of the vitamin was accomplished without delay in the same year by Kuhn et al.,¹²) Harris and Folkers,¹³,¹⁴) and Morii and Makino.¹⁵)

It was considered that all the vitamin B₆ activity of natural products could be attributed to PIN before the existence of other forms of vitamin B₆ was recognized by Snell et al.¹⁶) as a result of microbiological assay. These compounds were confirmed as PAL and PAM.¹⁷⁻²¹) Thereafter, it was established that the biological activity of the vitamin B₆ group was displayed by PIN, PAL, and PAM, and their 5′-phosphate esters.²²)

An unidentified phosphorylation product of PAL, which was later identified as PAL-5′-P,²³) was discovered as a coenzyme of tyrosine decarboxylase by Gunsalus et al.²⁴) It has now been made evident that PAL-P is the principal biocatalytically active form of the vitamin as a coenzyme for various enzyme systems, associated almost entirely with
A number of vitamin B₆-dependent enzymes have now been found, and the mechanisms of many PAL-dependent reactions in amino acid metabolism have been interpreted by Braunstein and Shemyakin, and independently by Metzler, Ikawa, and Snell with non-enzymatic model systems.

All vitamin B₆-related compounds owed their vitamin activity to their conversion to PAL-P. The enzymatic interconversion of various forms of vitamin B₆ and their conversion to PAL-P have also been investigated by many workers. Snell et al. investigated the degradation pathway of vitamin B₆ and isolated some degradation products from a culture of certain soil pseudomonads grown on PIN or PAM as the sole source of carbon and nitrogen. Recently, Contractor and Shane have identified a new metabolite of PIN in rat. However, little has been reported on the biosynthetic pathway of vitamin B₆. It has been indicated that common metabolites such as glycerol, aspartate, leucine, and serine are useful for the biosynthesis of the precursors of vitamin B₆, but no immediate precursors have yet been characterized.

Many derivatives of vitamin B₆ have hitherto been chemically synthesized and have also been found to occur naturally. For the purpose of the development of medical drugs or food supplements, many derivatives like vitamin B₆-fatty acid esters, PIN-nicotinic acid esters, PIN-thiamine disulfide, PAL-homocysteine, and vitamin B₆ disulfides have been chemically synthesized. Naturally occurring derivatives of vitamin B₆, such as the PAL-indole.
complex, PIN- or PAL-3-sulfate, PIN glucuronide, and pyridoxilidene-β-phenylethylamine have hitherto been identified as metabolites. However, the glycosides of vitamin B₆ were neither synthesized nor isolated from natural sources until PIN-G was first discovered by Ogata et al. during investigations on the microbial metabolic route of vitamin B₆. In this thesis, the author deals with the microbial formation of PIN-G, the mechanism of its enzymatic formation, and some properties of PIN-G in the following sections.

With regard to sugar derivatives of vitamins, many compounds have already been reported. Thiamine glycosides, the glucose ether of vitamin A, D-glucose pentanicotinate, and vitamin D₂ glucoside tetraacetate are chemically synthesized. Whitby first discovered the formation of riboflavinylglucoside by rat liver in 1950. Katagiri et al. investigated the microbial formation of various riboflavinylglycosides, including riboflavinyl-glucoside, -galactoside, -fructoside, and some oligosaccharides. Suzuki et al. also studied the formation of riboflavinylglycosides by various microorganisms and by various enzyme preparations from plants and microorganisms. It has been suggested that these compounds may play a physiological role in catalyzing the successive transfer of glucosyl groups via the formation of riboflavinyl compounds of oligosaccharides. Recently, Kasai et al. identified riboflavinyl-α-glucoside in cat liver. Miyake and Suzuki have studied the enzymatic formation of the sugar derivative of ascorbic acid. Takagi and Shiraishi have suggested the formation of a sugar derivative of carotenoids in
4'-O-(β-D-Glucopyranosyl)-D-PaA has lately been isolated from tomato juice as a growth factor of certain malo-lactic fermentation bacteria in wine fermentation. The compound showed a microbial activity 100 times as great as that of free D-PaA for a certain lactic acid bacterium. D-PaA is widely distributed in animals, plants, and microorganisms as a component of coenzyme A and plays an important role in many metabolic pathways. It has been clarified by Vagelos et al. and Wakil et al. that 4'-phosphopantetheine is another active form of D-PaA and functions as a prosthetic group in an acyl carrier protein. D-PaA is required by Saccharomyces carlsbergensis 4228 ATCC 9080, Lactobacillus plantarum ATCC 8014, and L. casei ATCC 7469. Recently, bifidus factors, which are growth factors for Bifidobacterium bifidum N4, were isolated from carrot root and identified as 4'-phosphopantetheine-S-sulfonic acid and 3'-dephospho-coenzyme A-S-sulfonic acid by Yoshioka et al. It was suggested that the activity of these derivatives was due to the specific membrane transport of the organisms.

In this thesis, the author describes the microbial formation of PIN-G and PaA-G. In the first chapter, the distribution of PIN-G-forming activity in culture collections and isolated strains of bacteria, yeasts, molds and actinomycetes is discussed. With an isolated strain, Micrococcus sp. No. 431, having the highest activity, several conditions affecting the fermentative production of PIN-G are described. The PIN-G-synthesizing enzyme was also purified from the
organism and the enzymatic properties have been investigated; the biological activity and some properties of PIN-G are discussed in the last section. In the second chapter, the formation of PaA-G was checked in yeasts. PaA-β-G could scarcely be detected at all, but PaA-α-G was found in the culture filtrate of several strains. *Sporobolomyces coralliformis* IFO 1032 showed the highest activity for PaA-α-G formation. PaA-α-G was isolated from the reaction mixture of *S. coralliformis* and characterized.
Chapter I. Microbial Formation of Pyridoxine Glucoside

Section I. Formation of PIN-G by Microorganisms

INTRODUCTION

During the investigation on the metabolism of PIN in microorganisms, PIN-G was first detected in the culture filtrate of Sarcina lutea IFO 3232, grown on a sucrose-peptone medium containing PIN. This compound was isolated from a reaction mixture incubated with intact cells of the organism by paper-pile chromatography, and active charcoal, Dowex 1X2 (Cl\(^-\) type) and Sephadex G-15 column chromatography. The compound was characterized and identified as a mixture of PIN-4'-G and PIN-5'-G by various analytical methods. The chemical structures of PIN-4'-G and PIN-5'-G are shown in Fig. 1.

![Chemical structures of PIN-4'-G and PIN-5'-G.](image)

Fig. 1. Chemical structures of PIN-4'-G and PIN-5'-G.
The distribution of PIN-G-forming activity in microorganisms is an interesting problem in regard to its biological significance. The screening of more producible strains of the compound is useful for the convenient isolation.

This section describes the distribution of PIN-G-forming activity in microorganisms including bacteria, molds, yeasts and actinomycetes. The screening was also performed with isolated strains from soil. An isolated bacterium, No. 431, which had the highest activity for the formation of PIN-G, was identified as Micrococcus sp. No. 431 according to the Bergey's Manual of Determinative Bacteriology etc.100-103)

EXPERIMENTAL

Materials. All chemicals used in this section were obtained from commercial sources.

Microorganisms. Microorganisms preserved in the Laboratory of Physiology of Fermentation and Applied Microbiology, Department of Agricultural Chemistry, Kyoto University (AKU), were used for the screening of PIN-G formation. It was also carried out with the isolated strains from soil in Kyoto Prefecture. All strains used were preserved on a nutrient agar slant by a monthly transfer at 5°C.

Cultivation. The composition of culture media for PIN-G formation by microorganisms is summarized in Table I. All cultivations were carried out in a test tube with 5 ml of the corresponding media. Bacteria except lactic acid bacteria and Clostridium were grown at 28°C for 3 days with shaking. Cultures of Clostridium were grown statically
TABLE I. COMPOSITION OF CULTURE MEDIA FOR PIN-G FORMATION

Glucose was used in place of sucrose or maltose as a blank.

<table>
<thead>
<tr>
<th>Component</th>
<th>Bacteria</th>
<th>Lactic acid bacteria</th>
<th>Yeasts</th>
<th>Molds</th>
<th>Actinomycetes</th>
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<tr>
<td>Sucrose</td>
<td>2.0 %</td>
<td>10.0 %</td>
<td>5.0 %</td>
<td>4.0 %</td>
<td>2.0 %</td>
</tr>
<tr>
<td>Maltose</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>(4.0)</td>
<td>-</td>
</tr>
<tr>
<td>Soluble starch</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.5</td>
</tr>
<tr>
<td>Peptone</td>
<td>1.0</td>
<td>-</td>
<td>0.5</td>
<td>-</td>
<td>2.0</td>
</tr>
<tr>
<td>Polypeptone</td>
<td>-</td>
<td>0.05</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>( \text{NH}_4\text{NO}_3 )</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.1</td>
<td>-</td>
</tr>
<tr>
<td>( \text{NaNO}_3 )</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.1</td>
<td>-</td>
</tr>
<tr>
<td>( \text{KNO}_3 )</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.2</td>
</tr>
<tr>
<td>( (\text{NH}_4)_2\text{SO}_4 )</td>
<td>-</td>
<td>0.03</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>( \text{K}_2\text{HPO}_4 )</td>
<td>0.5</td>
<td>-</td>
<td>0.4</td>
<td>0.1</td>
<td>0.02</td>
</tr>
<tr>
<td>( \text{KH}_2\text{PO}_4 )</td>
<td>0.1</td>
<td>0.5</td>
<td>0.2</td>
<td>-</td>
<td>0.07</td>
</tr>
<tr>
<td>( \text{NaCl} )</td>
<td>0.2</td>
<td>0.1</td>
<td>-</td>
<td>-</td>
<td>0.3</td>
</tr>
<tr>
<td>( \text{KCl} )</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.05</td>
<td>-</td>
</tr>
<tr>
<td>( \text{MgSO}_4 \cdot 7\text{H}_2\text{O} )</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.05</td>
<td>0.01</td>
</tr>
<tr>
<td>Yeast extract</td>
<td>0.2</td>
<td>0.15</td>
<td>0.5</td>
<td>-</td>
<td>0.2</td>
</tr>
<tr>
<td>PIN·HCl</td>
<td>1.22 mg/ml</td>
<td>1.22 mg/ml</td>
<td>1.22 mg/ml</td>
<td>1.22 mg/ml</td>
<td>1.22 mg/ml</td>
</tr>
<tr>
<td>( \text{CaCO}_3 )</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>pH</td>
<td>7.0</td>
<td>7.0</td>
<td>6.0</td>
<td>not adjusted</td>
<td>7.0</td>
</tr>
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</table>

-8-
at 37°C for 4 days. Lactic acid bacteria were cultivated statically at 20-37°C for 2 days. Yeasts were cultivated at 28°C for 2 days with shaking. Molds were cultivated at 28°C for 4 days with shaking and actinomycetes were at 28°C for 3 days with shaking. The cultivations were stopped by immersing culture tubes in boiling water for 5 min. After cooling them in ice-cold water, the cells were removed by centrifugation and the broth was supplied to the assay of PIN-G.

Assay of PIN-G. A suitable amount of the broth was employed for paper chromatography on Toyo filter paper No. 53 with a solvent system of n-butanol:acetic acid:water (4:1:1, v/v). The product on a paper chromatogram was detected by Manaslu lamp (2536 A), cut off and eluted from the paper with a mixture of 2.5 ml of 99% ethanol and 2.5 ml of 25% sodium acetate solution at 37°C for 90 min under occasional stirring in a test tube. To the extract was added 0.2 ml of the diazotized p-aminoacetophenone reagent (104,105) and the absorbancy at 470 nm was measured with a Hitachi Perkin-Elmer spectrophotometer model 139. PIN-G was measured as free PIN with a calibration curve prepared from authentic PIN. The p-aminoacetophenone reagent was prepared by mixing 1 ml of p-aminoacetophenone solution (318 mg of p-aminoacetophenone was dissolved in 4.5 ml of conc. HCl and then diluted to 50 ml with deionized water) and 5 ml of 4.5% sodium nitrite solution.

Identification of bacteria. Classification and identification of an isolated bacterium was performed by standard methods according to the Bergey's Manual of Determinative Bacteriology etc. (100-103)
RESULTS

Distribution of PIN-G-forming Activity in Microorganisms

The screening of PIN-G-forming activity was carried out with 100 strains of bacteria from culture collections. The result is shown in Table II. The high PIN-G-forming activities (more than 10-150 μg/ml of PIN-G as PIN) were distributed specifically in some strains of bacteria belonging to Micrococcus and Sarcina, but the majority of the bacteria accumulated little amounts of PIN-G in their culture media. Pseudomonas fragi formed a PIN-G-like compound in the medium, which was different from PIN-G in Rf values on paper chromatography. This compound still remains to be elucidated.

About 130 strains of yeasts were searched for the accumulation of PIN-G with the screening medium containing sucrose and PIN. As shown in Table III, only several strains among Endomycopsis, Hansenula, Sporobolomyces, Brettanomyces, Kloekera and Trichosporon had the relatively higher activities. The highest activity was found in Torulopsis sake IFO 0435, which accumulated maximally 260 μg/ml of PIN-G as PIN.

It has been reported that mold α-glucosidases readily hydrolyze maltose rather than aryl-α-glucosides, and are not capable of hydrolyzing sucrose. The screening was carried out on a maltose-peptone medium and a sucrose-peptone medium. The results are shown in Tables IV-a and IV-b, respectively. Only 6 strains of molds among 40 strains accumulated 10-150 μg/ml of PIN-G on a sucrose medium, while 14 strains among 34 strains tested accumulated about the same
## TABLE II. DISTRIBUTION OF PIN-G-FORMING ACTIVITY IN BACTERIA

Bacteria were grown on the screening medium described in Table I. The cultivation was carried out as described in the EXPERIMENTAL.

<table>
<thead>
<tr>
<th>Family</th>
<th>Genus</th>
<th>No. of strains tested</th>
<th>PIN-G formed (µg/ml as PIN)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>10&gt;</td>
</tr>
<tr>
<td><strong>Enterobacteriaceae</strong></td>
<td><strong>Escherichia</strong></td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td><strong>Aerobacter</strong></td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td><strong>Erwinia</strong></td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td><strong>Serratia</strong></td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td><strong>Proteus</strong></td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td><strong>Achromobacteriaceae</strong></td>
<td><strong>Alcaligenes</strong></td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td><strong>Achromobacter</strong></td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td><strong>Flavobacterium</strong></td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td><strong>Rhizobiaceae</strong></td>
<td><strong>Agrobacterium</strong></td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><strong>Micrococaceae</strong></td>
<td><strong>Micrococcus</strong></td>
<td>7</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td><strong>Staphylococcus</strong></td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td><strong>Sarcina</strong></td>
<td>5</td>
<td>1</td>
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<tr>
<td><strong>Brevibacteriaceae</strong></td>
<td><strong>Brevibacterium</strong></td>
<td>2</td>
<td>2</td>
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<td><strong>Corynebacteriaceae</strong></td>
<td><strong>Corynebacterium</strong></td>
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<td><strong>Bacillaceae</strong></td>
<td><strong>Bacillus</strong></td>
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<tr>
<td><strong>Pseudomonadaceae</strong></td>
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<td></td>
<td><strong>Xanthomonas</strong></td>
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<td>1</td>
</tr>
<tr>
<td><strong>Lactobacillaceae</strong></td>
<td><strong>Streptococcus</strong></td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td><strong>Leuconostoc</strong></td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td><strong>Lactobacillus</strong></td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td><strong>Pediococcus</strong></td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><strong>Sum total:</strong></td>
<td></td>
<td>100</td>
<td>92</td>
</tr>
</tbody>
</table>
extent of PIN-G when cultured on a maltose medium. However, the amounts of PIN-G formed in a maltose medium were higher than those in a sucrose medium. Especially, *Penicillium rubrum* Stoll IFO 6580 accumulated 210 µg/ml of PIN-G as PIN on a maltose medium. Table V shows that 15 strains out of 45 strains of actinomycetes tested had the ability to form considerable amounts of PIN-G. Among them, *Streptomyces hygroscopicus* IFO 3934 had the highest activity, which accumulated 260 µg/ml of PIN-G as PIN.

Bacteria (324 strains), molds (12 strains) and actinomycetes (6 strains) were isolated from soil in Kyoto Prefecture for a screening of PIN-G formation. The result is shown in Table VI. Neither molds nor actinomycetes had the low activity, but higher activities were found in 6 strains of bacteria, in which a strain No. 431 had the highest activity, accumulating 290 µg/ml of PIN-G as PIN. Thus, the strain No. 431 was selected and used as the PIN-G-forming strain in the following experiment.

*Identification of the Strain No. 431*

The isolated bacterial strain No. 431 described above, was classified and identified as follows, according to the method of the Bergey's Manual of Determinative Bacteriology etc. 100-103)

(a) Morphological: Spheres, 1 micron in diameter in average size, occurring in pairs. Non-motile. Gram-positive.
TABLE III. DISTRIBUTION OF PIN-G-FORMING ACTIVITY IN YEASTS

Yeasts were grown on the screening medium described in Table I. The cultivation was carried out as described in the EXPERIMENTAL.

<table>
<thead>
<tr>
<th>Family</th>
<th>Genus</th>
<th>No. of strains tested</th>
<th>PIN-G formed (µg/ml as PIN)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>10&gt;</td>
</tr>
<tr>
<td><strong>Endomycetaceae</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Saccharomyces</td>
<td>35</td>
<td>35</td>
<td>0</td>
</tr>
<tr>
<td>baker's yeast</td>
<td>2</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>brewer's yeast</td>
<td>3</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>shoyu yeast</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Endomyces</td>
<td>3</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Endomycesis</td>
<td>2</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Schizosaccharomyces</td>
<td>2</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Zygosaccharomyces</td>
<td>3</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Pichia</td>
<td>4</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Hansenula</td>
<td>9</td>
<td>7</td>
<td>2</td>
</tr>
<tr>
<td>Schwanniomyces</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Debaryomyces</td>
<td>2</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Saccharomyces</td>
<td>2</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Hanseniaspora</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Nadsonia</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Lipomyces</td>
<td>2</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td><strong>Sporobolomyceae</strong></td>
<td><strong>Sporobolomyces</strong></td>
<td><strong>5</strong></td>
<td><strong>3</strong></td>
</tr>
<tr>
<td><strong>Cryptococcaeae</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cryptococcus</td>
<td>3</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Torulopsis</td>
<td>7</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>Brettanomyces</td>
<td>3</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Candida</td>
<td>21</td>
<td>21</td>
<td>0</td>
</tr>
<tr>
<td>Kloechera</td>
<td>3</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Wickerhamia</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Rhodotorula</td>
<td>10</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>Trichosporon</td>
<td>3</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td><strong>Sum total:</strong></td>
<td>129</td>
<td>120</td>
<td>8</td>
</tr>
</tbody>
</table>

*) Torulopsis sake IFO 0435 260 µg/ml as PIN
TABLE IV-a. DISTRIBUTION OF PIN-G-FORMING ACTIVITY IN MOLDS

Molds were grown on the sucrose medium described in Table I. The cultivation was carried out as described in the EXPERIMENTAL.

<table>
<thead>
<tr>
<th>Genus</th>
<th>No. of strains tested</th>
<th>PIN-G formed (µg/ml as PIN)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>10&gt;</td>
</tr>
<tr>
<td><em>Mucor</em></td>
<td>9</td>
<td>7</td>
</tr>
<tr>
<td><em>Rhizopus</em></td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td><em>Aspergillus</em></td>
<td>13</td>
<td>12</td>
</tr>
<tr>
<td><em>Penicillium</em></td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td><em>Monascus</em></td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><em>Neurospora</em></td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td><em>Pullularia</em></td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><em>Fusarium</em></td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td><strong>Sum total:</strong></td>
<td><strong>40</strong></td>
<td><strong>34</strong></td>
</tr>
</tbody>
</table>

TABLE IV-b. DISTRIBUTION OF PIN-G-FORMING ACTIVITY IN MOLDS

Molds were grown on the maltose medium described in Table I. The cultivation was carried out as described in the EXPERIMENTAL.

<table>
<thead>
<tr>
<th>Genus</th>
<th>No. of strains tested</th>
<th>PIN-G formed (µg/ml as PIN)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>10&gt;</td>
</tr>
<tr>
<td><em>Mucor</em></td>
<td>8</td>
<td>5</td>
</tr>
<tr>
<td><em>Rhizopus</em></td>
<td>8</td>
<td>7</td>
</tr>
<tr>
<td><em>Aspergillus</em></td>
<td>10</td>
<td>4</td>
</tr>
<tr>
<td><em>Penicillium</em></td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td><em>Neurospora</em></td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td><em>Pullularia</em></td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><em>Fusarium</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><strong>Sum total:</strong></td>
<td><strong>34</strong></td>
<td><strong>20</strong></td>
</tr>
</tbody>
</table>

*) Penicillium rubrum IFO 6580 210 µg/ml as PIN
TABLE V. DISTRIBUTION OF PIN-G-FORMING ACTIVITY IN ACTINOMYCETES

Actinomycetes were grown on the screening medium described in Table I. The cultivation was carried out as described in the EXPERIMENTAL.

<table>
<thead>
<tr>
<th>Genus</th>
<th>No. of strains tested</th>
<th>PIN-G formed (µg/ml as PIN)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>10&gt;</td>
</tr>
<tr>
<td>Mycobacterium</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Nocardia</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>Streptomyces</td>
<td>24</td>
<td>13</td>
</tr>
<tr>
<td>Unidentified</td>
<td>14</td>
<td>11</td>
</tr>
<tr>
<td>Sum total:</td>
<td>45</td>
<td>30</td>
</tr>
</tbody>
</table>

*) Streptomyces hygroscopicus var. angustomyoticus IFO 3934 260 µg/ml as PIN

TABLE VI. DISTRIBUTION OF PIN-G-FORMING ACTIVITY IN MICROORGANISMS ISOLATED FROM SOIL

Microorganisms isolated from soil were grown on the screening medium described in Table I. The cultivation was carried out as described in the EXPERIMENTAL.

<table>
<thead>
<tr>
<th>Microorganisms</th>
<th>No. of strains tested</th>
<th>PIN-G formed (µg/ml as PIN)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>10&gt;</td>
</tr>
<tr>
<td>Bacteria</td>
<td>324</td>
<td>290</td>
</tr>
<tr>
<td>Molds</td>
<td>12</td>
<td>12</td>
</tr>
<tr>
<td>Actinomycetes</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td>Sum total:</td>
<td>342</td>
<td>307</td>
</tr>
</tbody>
</table>

*) Strain No. 431 289 µg/ml as PIN
(b) Cultural:

Gelatin stab: No liquefaction.

Agar slant: Yellow, smooth growth.

Nutrient broth: Turbid, with yellowish sediment.

Potato: Abundant, yellow growth.

(c) Physiological:

Litmus milk: Alkaline, not peptonized and no coagulation; litmus not reduced.

Indole not produced.

Hydrogen sulfide produced.

Ammonia produced from peptone.

Acid from fructose, galactose, xylose, and mannitol. No acid and alkaline from glucose, maltose, dextrin, starch, glycerol, sucrose, arabinose, raffinose, lactose, inulin, mannose and sorbitol. No gas from all carbohydrates.

Citric acid utilized as a sole source of carbon.

Nitrites produced from nitrates.

Catalase positive.

Methyl red test negative.

Voges-Proskauer test negative.

Optimum temperature between 28 and 37°C. Killed at 60°C for 10 min.

Optimum pH between 5.5 and 10.0.

Aerobic.

(d) Source: Isolated from soil in Kyoto Prefecture.
From the data described above, the strain No. 431 may be classified into genus *Micrococcus*. However, the isolated strain was not applicable to any known species in *Micrococcus* described in the manual, and so it was tentatively designated as *Micrococcus* sp. No. 431. The electron micrograph of the organism is shown in Fig. 2.

**DISCUSSION**

The relatively higher activities for PIN-G formation were particularly observed in several strains of bacteria belonging to the genera *Sarcina* and *Micrococcus*. It is an interesting fact that these coccidal strains belonging to the same family, *Micrococcaceae*, have the higher activities. On the other hand, other cocci such as *Staphylococcus*, which is also classified into *Micrococcaceae*, *Leuconostoc*, *Pediococcus* and *Streptococcus* had no activity. Although the trans-glucosidation to riboflavin to form riboflavinyl glucoside has been
demonstrated in various bacterial strains including *Escherichia coli*, *Aerobacter aerogenes*, *Bacillus subtilis*, *Clostridium acetobutylicum*, *Lactobacillus casei* and *Streptococcus lactis*, these strains did not form PIN-G from PIN under the conditions as described in this section. The PIN-G-forming activity was also observed in some strains of yeasts, molds and actinomycetes. When 34 strains of various molds were cultivated on a maltose-peptone medium, 14 strains including *Mucor*, *Rhizopus*, *Aspergillus*, *Penicillium*, and *Fusarium* accumulated 10-150 μg/ml or more than 150 μg/ml of PIN-G. On the other hand, only several strains were found to form a slight amount of PIN-G on a sucrose-peptone medium. These results coincide with the fact that transglucosidase of molds generally hydrolyzes maltose, but not sucrose.

The PIN-G-forming activity in microorganisms which were isolated from soil was distributed mainly among bacteria (34 strains of bacteria, one strain of actinomycetes), in which 7 strains accumulated relatively large amounts of PIN-G (more than 150 μg/ml of PIN-G as PIN). Best of all, an isolated bacterium strain No. 431, which was classified into *Micrococcaceae*, had the highest activity (289 μg/ml of PIN-G as PIN). The various conditions for PIN-G formation by this organism will be discussed in the next section.

**SUMMARY**

The distribution of PIN-G-forming activity was investigated with various kinds of microorganisms under the conditions of their growth on the media containing PIN and sucrose or maltose. The formation of
PIN-G was searched in the culture broth of organisms including bacteria (100 strains), yeasts (129 strains), molds (34 strains) and actinomycetes (45 strains). The screening was also performed with isolated strains of bacteria, molds and actinomycetes (total 342 strains) from soil. The results of screening showed that the higher activities were specifically found in several strains of the bacteria belonging to the genera *Sarcina* and *Micrococcus*. The activity was also shown to be distributed in yeasts, molds and actinomycetes. The activity was found in the isolated strains. A bacterium No. 431, which was identified as *Micrococcus* sp. No. 431 according to the Bergey's Manual *etc.*, showed the highest activity (289 μg/ml of PIN-G as PIN) among the tested organisms and selected to be the most suitable strain for PIN-G formation.
Section II. PIN-G Formation by Micrococcus sp. No. 431

INTRODUCTION

In the preceding section, a bacterium was isolated and identified as Micrococcus sp. No. 431, which accumulated approximately 300 μg/ml of PIN-G as PIN in the culture medium composed of PIN and sucrose. In order to get a large amount of PIN-G, various factors on the formation of PIN-G are studied in this section with the growing cultures of this organism. PIN-G was first detected in the culture filtrate of Sarcina lutea IFO 3232 and was identified as a mixture of PIN-4'-G and PIN-5'-G. In this section, PIN-4'-G and PIN-5'-G were isolated by various separating methods from the culture filtrate of Micrococcus sp. No. 431, which was grown under the optimal conditions. The products were also identified by various analytical methods including IR and NMR spectroscopy. Moreover, PIN-diglucoside (possibly PIN-4', PIN-5', and PIN-4',5'-diglucosides) was isolated from the culture filtrate and identified by the same methods.

EXPERIMENTAL

Materials. Cellulase (Type II, from Aspergillus niger), β-glucosidase (from almonds), maltase (Grade II, from Aspergillus niger) were purchased from Sigma Chem. Co., and α-glucosidase (from yeast) was from Boehringer Mannheim Japan K.K. Other chemicals used in this work were obtained from commercial sources.

Microorganism and cultivation. Micrococcus sp. No. 431, which
was isolated from soil and identified in the preceding section, was used for the formation of PIN-G. The basal medium contained 2.0% sucrose, 1.0% peptone, 0.2% yeast extract, 0.5% $K_2HPO_4$, 0.1% $KH_2PO_4$, 0.2% NaCl, 0.02% MgSO$_4$·7H$_2$O and 1 mg/ml of PIN (pH 7.0). The cultivation was carried out at 28°C for 40 hr with reciprocal shaking.

**Analyses.** After cultivation, the cultural broth was heated for 5 min in a boiling water bath. The supernatant solution obtained by centrifugation was followed to the assay. PIN-G was assayed by paper chromatography as described in the preceding section.

Absorption spectra were recorded on a Shimazu multi-purpose spectrophotometer model MPS-50L or a Hitachi double-beam spectrophotometer type 124.

Paper chromatography of vitamin B$_6$ was carried out with the solvent systems of I. $n$-butanol:acetic acid:water (4:1:1, v/v), II. $n$-amylalcohol:acetone:water (2:1:1, v/v, upper layer), III. water: acetone: $t$-butanol:diethylamine (20:35:40:5, v/v) and IV. water: acetone: $t$-butanol:acetic acid (20:35:40:5, v/v). The diazotized p-aminoacetophenone reagent, a mixture of 1 mL of p-aminoacetophenone solution, 5 mL of 4.5% sodium nitrite and 5 mL of 25% sodium acetate, was sprayed for vitamin B$_6$ detection.

Paper chromatography of sugars was carried out with the solvent systems of I. $n$-butanol:benzene:pyridine:water (5:1:3:3, v/v, upper layer), II. $n$-butanol:pyridine:water (6:4:3, v/v) and III. isopropanol:pyridine:water:acetic acid (8:8:4:1, v/v), and the silver nitrate reagent was sprayed for sugar detection.
A PIN moiety was determined by the diazotized p-aminoacetophenone reagent\(^{104,105}\) as described in the preceding section.

An anthrone method\(^{110}\) was used for the assay of total sugars and a glucostat method\(^{111}\) was used for the assay of a glucose moiety. Reducing sugar was determined by the method of Somogyi.\(^{112}\)

The reaction by several glucosidases was carried out on a reaction mixture containing 212 \(\mu\)g of PIN-G, 20 \(\mu\)moles of 2-mercaptoethanol, 20 \(\mu\)moles of buffer and an appropriate amount of enzyme in a total volume of 1.0 ml. The reaction was carried out at 37\(^\circ\)C overnight. Phosphate buffer (pH 6.5) was used in the incubations with mold maltase and yeast \(\alpha\)-glucosidase, and acetate buffer (pH 4.4) in those with cellulase and \(\beta\)-glucosidase. The reaction mixture was submitted to paper chromatographic analyses for PIN and glucose as described above.

**Measurement of IR and NMR spectra.** IR spectrum of PIN-G was recorded in a micro KBr tablet with a Hitachi infrared spectrometer type EPI-G3.

NMR spectra of PIN-G and PIN-diglucoside were recorded in \(D_2O\) on a Hitachi Perkin-Elmer high resolution NMR spectrometer type R-22 at 90 MHz with DSS as an internal standard.

**RESULTS**

**Cultural Conditions for PIN-G Formation**

**Carbon sources**

The effect of various carbohydrates as the glucosyl donors on
TABLE I. EFFECT OF CARBON SOURCES

*Micrococcus* sp. No. 431 was cultivated on the basal medium described in the EXPERIMENTAL except that various carbohydrates were used as shown in the Table. The cultivation was performed aerobically at 28°C for 40 hr.

<table>
<thead>
<tr>
<th>Carbon sources</th>
<th>Growth</th>
<th>Final pH</th>
<th>PIN-G formed (µg/ml as PIN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glycerin</td>
<td>+++</td>
<td>7.6</td>
<td>0</td>
</tr>
<tr>
<td>Arabinose</td>
<td>++</td>
<td>7.6</td>
<td>0</td>
</tr>
<tr>
<td>Xylose</td>
<td>++</td>
<td>6.6</td>
<td>0</td>
</tr>
<tr>
<td>Glucose</td>
<td>++++</td>
<td>6.4</td>
<td>0</td>
</tr>
<tr>
<td>Fructose</td>
<td>++++</td>
<td>6.4</td>
<td>0</td>
</tr>
<tr>
<td>Galactose</td>
<td>+++</td>
<td>6.4</td>
<td>0</td>
</tr>
<tr>
<td>Mannose</td>
<td>++++</td>
<td>6.0</td>
<td>0</td>
</tr>
<tr>
<td>Mannit</td>
<td>+++</td>
<td>7.0</td>
<td>0</td>
</tr>
<tr>
<td>Methyl-α-D-glucoside</td>
<td>++</td>
<td>8.6</td>
<td>0</td>
</tr>
<tr>
<td>Sucrose</td>
<td>++++</td>
<td>7.0</td>
<td>313</td>
</tr>
<tr>
<td>Maltose</td>
<td>++++</td>
<td>6.6</td>
<td>32</td>
</tr>
<tr>
<td>Trehalose</td>
<td>++</td>
<td>8.8</td>
<td>0</td>
</tr>
<tr>
<td>Lactose</td>
<td>++</td>
<td>8.2</td>
<td>0</td>
</tr>
<tr>
<td>Raffinose</td>
<td>++</td>
<td>7.8</td>
<td>0</td>
</tr>
<tr>
<td>Dextrin</td>
<td>++</td>
<td>8.6</td>
<td>0</td>
</tr>
<tr>
<td>Starch</td>
<td>++</td>
<td>8.6</td>
<td>0</td>
</tr>
<tr>
<td>Inulin</td>
<td>++</td>
<td>8.0</td>
<td>0</td>
</tr>
</tbody>
</table>

The formation of PIN-G was studied with *Micrococcus* sp. No. 431. As shown in Table I, sucrose and maltose, having the α-glucosidic linkage, were effective for the PIN-G formation. Maltose was one-tenth as effective as sucrose, and trehalose was not effective, though it has an α(1→1) glucosidic linkage. Polysaccharides such as dextrin
TABLE II. EFFECT OF NITROGEN SOURCES

*Micrococcus* sp. No. 431 was cultivated on the basal medium described in the EXPERIMENTAL except that various nitrogen sources were used as shown in the Table. The cultivation was performed at 28°C for 40 hr.

<table>
<thead>
<tr>
<th>Nitrogen sources</th>
<th>Growth</th>
<th>Final pH</th>
<th>PIN-G formed (µg/ml as PIN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>+</td>
<td>6.8</td>
<td>83</td>
</tr>
<tr>
<td>Peptone</td>
<td>++++</td>
<td>8.2</td>
<td>269</td>
</tr>
<tr>
<td>Casamino acid</td>
<td>+++</td>
<td>8.2</td>
<td>170</td>
</tr>
<tr>
<td>Urea</td>
<td>+</td>
<td>8.2</td>
<td>41</td>
</tr>
<tr>
<td>(NH₄)₂SO₄</td>
<td>+</td>
<td>6.6</td>
<td>108</td>
</tr>
<tr>
<td>NH₄Cl</td>
<td>++</td>
<td>6.2</td>
<td>104</td>
</tr>
<tr>
<td>NaNO₃</td>
<td>++</td>
<td>7.2</td>
<td>88</td>
</tr>
<tr>
<td>NaNO₂</td>
<td>+</td>
<td>7.2</td>
<td>29</td>
</tr>
<tr>
<td>Asparagine</td>
<td>++</td>
<td>9.2</td>
<td>36</td>
</tr>
<tr>
<td>Glutamate</td>
<td>+++</td>
<td>9.2</td>
<td>91</td>
</tr>
</tbody>
</table>

and starch also had no effect on the formation of PIN-G. No derivative of PIN other than PIN-G was formed with lactose, raffinose and inulin.

*Nitrogen sources*

The incubation was undertaken on a sucrose medium with various inorganic and organic nitrogen compounds. The result is shown in Table II. The organic nitrogen compounds such as peptone and casamino acid were useful for the formation of PIN-G and the growth of the organism. Ammonium sulfate and ammonium chloride were fairly effective on the formation of PIN-G. A considerable amount of PIN-G was formed in a medium containing no nitrogen compounds except 0.2% yeast
extract as a growth factor.

**Surfactants**

The effect of various surfactants on the formation of PIN-G was investigated with the addition of 0.1%, the result being shown in Table III. Nonionic surfactants, Span and Tween series showed a stimulating effect on the formation of PIN-G. Among them, Span 80 was found to be most effective. On the other hand, anionic and cationic surfactants were rather inhibitory on both of the growth and the formation. With the addition of 0.7% Span 80, the amount of PIN-G accumulated reached to a maximum of about 600 µg/ml as PIN from 1 mg/ml of PIN.

The organism was grown on a medium composed of 5.0% sucrose, 1.5% peptone, 0.2% yeast extract, 0.5% K₂HPO₄, 0.1% KH₂PO₄, 0.2% NaCl, 0.02% Na₂MoO₄·2H₂O, 0.1% Tween 60 and 1 mg/ml of PIN (pH 7.0). The cultivation was carried out at 28°C with reciprocal shaking.
**TABLE III. EFFECT OF VARIOUS SURFACTANTS**

*Micrococcus* sp. No. 431 was cultivated on the basal medium described in the EXPERIMENTAL except that 0.1% of a final concentration of various surfactants were added as shown in the Table. The cultivation was performed at 28°C for 40 hr with shaking.

<table>
<thead>
<tr>
<th>Surfactant</th>
<th>Growth</th>
<th>Final pH</th>
<th>PIN-G formed (µg/ml as PIN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>-</td>
<td>7.4</td>
<td>212</td>
</tr>
<tr>
<td>CTAB</td>
<td>cationic</td>
<td>+</td>
<td>6.4 22</td>
</tr>
<tr>
<td>CPC</td>
<td></td>
<td>+</td>
<td>6.4 33</td>
</tr>
<tr>
<td>Aerosol OT</td>
<td>anionic</td>
<td>+</td>
<td>6.4 19</td>
</tr>
<tr>
<td>Na-Laurate</td>
<td></td>
<td>+++</td>
<td>7.0 92</td>
</tr>
<tr>
<td>Na-Myristate</td>
<td></td>
<td>+</td>
<td>6.8 81</td>
</tr>
<tr>
<td>Na-Palmitate</td>
<td></td>
<td>+++</td>
<td>7.4 248</td>
</tr>
<tr>
<td>Na-Stearate</td>
<td></td>
<td>+++</td>
<td>7.2 243</td>
</tr>
<tr>
<td>Na-Oleate</td>
<td></td>
<td>+</td>
<td>6.6 98</td>
</tr>
<tr>
<td>Tween 20</td>
<td>nonionic</td>
<td>+++</td>
<td>7.2 388</td>
</tr>
<tr>
<td>&quot; 40</td>
<td></td>
<td>+++</td>
<td>7.2 353</td>
</tr>
<tr>
<td>&quot; 60</td>
<td></td>
<td>+++</td>
<td>7.2 376</td>
</tr>
<tr>
<td>&quot; 80</td>
<td></td>
<td>+++</td>
<td>7.0 407</td>
</tr>
<tr>
<td>&quot; 85</td>
<td></td>
<td>+++</td>
<td>6.8 359</td>
</tr>
<tr>
<td>Span 20</td>
<td></td>
<td>+</td>
<td>6.4 94</td>
</tr>
<tr>
<td>&quot; 40</td>
<td></td>
<td>+++</td>
<td>7.0 86</td>
</tr>
<tr>
<td>&quot; 60</td>
<td></td>
<td>+++</td>
<td>7.0 243</td>
</tr>
<tr>
<td>&quot; 80</td>
<td></td>
<td>+++</td>
<td>6.2 524</td>
</tr>
<tr>
<td>&quot; 85</td>
<td></td>
<td>+++</td>
<td>6.4 335</td>
</tr>
</tbody>
</table>

Abbreviations:
CTAB; Cetyltrimethyl ammonium bromide, CPC; Cetylpyridinium chloride, Aerosol OT; Na-dioctylsulfosuccinic acid, Tween 20, 40, 60, 80 and 85; Polyoxyethylene sorbitan monolaurate, monopalmitate, monostearate, monooleate and trioleate, Span 20, 40, 60, 80 and 85; Sorbitan monolaurate, monopalmitate, monostearate, monooleate and trioleate.
TABLE IV. EFFECT OF METAL IONS

Micrococcus sp. No. 431 was grown on the basal medium described in the EXPERIMENTAL except that 5.0% sucrose, 1.5% peptone, 0.7% Span 80, and 0.02% various metal ions were added. The cultivation was carried out at 28°C for 60 hr with shaking.

<table>
<thead>
<tr>
<th>Metal ions</th>
<th>Growth (x50^{-1}, 610 μm)</th>
<th>Final pH</th>
<th>PIN-G formed (μg/ml as PIN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>.193</td>
<td>6.6</td>
<td>695</td>
</tr>
<tr>
<td>EDTA*</td>
<td>.268</td>
<td>6.2</td>
<td>542</td>
</tr>
<tr>
<td>MgSO_4·7H_2O</td>
<td>.308</td>
<td>6.2</td>
<td>528</td>
</tr>
<tr>
<td>Na_2MoO_4·2H_2O</td>
<td>.188</td>
<td>6.4</td>
<td>704</td>
</tr>
<tr>
<td>FeSO_4·7H_2O</td>
<td>.234</td>
<td>6.4</td>
<td>651</td>
</tr>
<tr>
<td>Fe_2(SO_4)_3·xH_2O</td>
<td>.184</td>
<td>6.6</td>
<td>649</td>
</tr>
<tr>
<td>ZnSO_4·7H_2O</td>
<td>.086</td>
<td>7.0</td>
<td>435</td>
</tr>
<tr>
<td>CoCl_2·6H_2O</td>
<td>.122</td>
<td>7.0</td>
<td>386</td>
</tr>
<tr>
<td>MnSO_4·4–6H_2O</td>
<td>.133</td>
<td>8.4</td>
<td>150</td>
</tr>
<tr>
<td>CuSO_4·5H_2O</td>
<td>.188</td>
<td>6.8</td>
<td>297</td>
</tr>
<tr>
<td>AgNO_3</td>
<td>.013</td>
<td>6.4</td>
<td>-</td>
</tr>
<tr>
<td>HgCl_2</td>
<td>.005</td>
<td>6.4</td>
<td>-</td>
</tr>
</tbody>
</table>

*) Ethylenediamine tetra-acetic acid

Incubation time

A time course of PIN-G formation and growth of the organism is shown in Fig. 1. The amount of PIN-G increased in an earlier stage of growth and reached to a maximum in 60 to 80 hr culture. A prolonged cultivation caused a rapid degradation of PIN-G. It may be suggested that the organism has a relatively strong α-glucosidase activity which hydrolyzes PIN-G to liberate PIN.
Fig. 2. Effect of Sucrose and Peptone Concentration.

The organism was grown on a medium composed of various concentrations of sucrose and peptone, 0.2% yeast extract, 0.5% K$_2$HPO$_4$, 0.1% KH$_2$PO$_4$, 0.2% NaCl, 0.7% Span 80 and 1 mg/ml of PIN (pH 7.0). The cultivation was carried out at 28°C for 60 hr with reciprocal shaking.

The effect of metal ions was examined. As shown in Table IV, the presence of metal ions did not affect the formation or excretion of PIN-G by the organism.

*Sucrose and peptone concentrations*

The effect of sucrose and peptone concentrations in culture media was investigated. The cultivation was performed at 28°C for 60 hr with shaking by the addition of 0.7% Span 80. As shown in Fig. 2, the amount of PIN-G increased proportionately to the concentration of sucrose added and reached to a maximum in the presence of 10% sucrose.
The organism was grown on a medium composed of 10% sucrose, 0.5% peptone, 0.2% yeast extract, 0.5% K$_2$HPO$_4$, 0.1% KH$_2$PO$_4$, 0.2% NaCl, 0.7% Span 80 and various concentrations of PIN (pH 7.0). The cultivation was carried out at 28°C for 60 hr with reciprocal shaking.

On the other hand, the concentration of peptone gave no effect on the formation of PIN-G. By the addition of 10% sucrose and 0.5-1.5% peptone, the yield of PIN-G attained to 90% against PIN added.

**PIN Concentration**

The effect of PIN concentration was examined in the presence of 10% sucrose and 0.5% peptone. The incubation was carried out at 28°C for 60 hr with shaking. Figure 3 shows that PIN-G is increased proportionately to the concentration of PIN added and attained to a maximum at...
2 mg/ml of PIN.

**pH of culture**

As shown in Fig. 4, a maximal formation of PIN-G occurred at pH 7.0. At pH 5.0, PIN-G was scarcely formed.

From the results described above, the optimal incubation conditions were determined as shown in Table V.

**TABLE V. OPTIMAL CULTURE CONDITIONS FOR PIN-G FORMATION**

<table>
<thead>
<tr>
<th>Component</th>
<th>Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sucrose</td>
<td>10.0 %</td>
</tr>
<tr>
<td>Peptone</td>
<td>1.5</td>
</tr>
<tr>
<td>Yeast extract</td>
<td>0.2</td>
</tr>
<tr>
<td>K$_2$HPO$_4$</td>
<td>0.5</td>
</tr>
<tr>
<td>KH$_2$PO$_4$</td>
<td>0.1</td>
</tr>
<tr>
<td>NaCl</td>
<td>0.2</td>
</tr>
<tr>
<td>Span 80</td>
<td>0.7</td>
</tr>
<tr>
<td>PIN·HCl</td>
<td>2.44 mg/ml</td>
</tr>
<tr>
<td>pH</td>
<td>7.0</td>
</tr>
<tr>
<td>Water</td>
<td>Tap water</td>
</tr>
</tbody>
</table>

The cultivation was performed aerobically at 28°C for 60 hr.

**Isolation and Identification of PIN-G and PIN-diglucoside**

From the results described above, optimal culture conditions of *Micrococcus* sp. No. 431 for PIN-G formation were settled as shown in Table V. In order to isolate PIN-G on a laboratory scale, the bacterium was cultured with 1000 ml of the optimal medium (each 500 ml of the medium in a 2 liter flask) under the optimal conditions as indi-
cated in Table V. The cultivation was stopped by heating the flask in a boiling water bath for 5 min. After cooling the broth in an ice bath, the cells were removed by centrifugation at 10,000 rpm for 30 min. The supernatant solution was adjusted to pH 2.0 and stored at 4°C overnight. Any material precipitating was removed by filtration. Sixty grams of active carbon was added to the supernatant solution. After stirring the mixture for 1 hr, the charcoal was collected by filtration, washed with 0.01N HCl and water, and then the adsorbed material was eluted with 28% ammonia:99% ethanol:water (5:50:45, v/v). The eluate was concentrated to a small volume in vacuo. The concentrated solution was applied on a Dowex 1X2 (Cl− type, 3 x 10 cm) column, and water was passed through. The effluent was shaken vigorously with chloroform to remove lipophilic impurities. The water layer was concentrated to a small volume to contain 30-50 mg of materials/ml as PIN. Two or three milliliters of the concentrate was repeatedly applied on a Sephadex G-10 (1 x 105 cm), followed by elution with water by 3 ml portions. The absorbancy at 325 μm of each fraction was measured, and an aliquot of the ultraviolet absorbing fractions was chromatographed on Toyo filter paper No. 53 with a solvent system of n-butanol:acetic acid:water (4:1:1, v/v). The elution pattern on a Sephadex column and the paper chromatogram of each fraction are shown in Fig. 5. The fractions (No. 14-16) containing PIN-G were collected, concentrated and applied on a Sephadex column. This procedure was repeated until PIN and unknown compounds were separated from PIN-G fractions. PIN-G fractions were
Fig. 5. Sephadex G-10 Gel Filtration and Paper Chromatogram of Each Fraction.

Column size: 1 x 104 cm, Flow rate: 9 ml/hr, One fraction; 3 ml/hr.
Solvent system: n-butanol:acetic acid:water (4:1:1, v/v)
Fig. 6. Paper Chromatogram of Isolated Compounds.

Solvent system: n-butanol:acetic acid:water (4:1:1, v/v)

The diazotized p-aminoacetophenone reagent was sprayed, followed by respraying 0.2M borate buffer (pH 9.0).

- Yellow spot, orange spot

I. PIN
II. PIN-5'-G
III. PIN-4'-G
IV. PIN-diglucoside (5'-)
V. PIN-diglucoside (4'- and 4',5'-)

followed by paper chromatography with a solvent system of n-butanol: benzene:pyridine:water (5:1:3:3, v/v, upper layer). Two fluorescent zones were detected on the paper chromatogram. Each zone was separately cut off and eluted with 50% ethanol. The first compound with a large $R_f$ value formed a borate complex and was not colored with the diazotized $p$-aminoacetophenone reagent in the presence of borate.
buffer. On the other hand, the second compound with a small $R_f$ value, did not form a borate complex and was colored with the diazotized $p$-aminoacetophenone reagent. It has been reported that 3- and 4-hydroxy groups of PIN combine with borate. \(^{113}\) PIN is colored with the diazotized $p$-aminoacetophenone reagent when 3-hydroxyl group is free. Thus, these two compounds were assumed to be PIN-5'-G and PIN-4'-G, respectively, in which PIN-4'-G was slightly dominant. One of the unknown compounds (I) which were obtained by repeated chromatographies on a Sephadex column was followed to paper chromatography and two fluorescent zones were also confirmed. Each zone was cut off and eluted with 50% ethanol. These compounds were also followed to the identification. Figure 6 shows a paper chromatogram of these compounds.

The identification of PIN-G was carried out with a mixture of

![Absorbance vs. Wave length graph](image)

**Fig. 7. Ultraviolet Absorption Spectra of Isolated PIN-G.**

The absorption spectra were recorded on a Shimazu multi-purpose spectrophotometer model MPS-50L.
PIN-4'-G and PIN-5'-G except NMR spectroscopy. The ultraviolet absorption spectra of the compound at acidic, neutral and alkaline pH were shown in Fig. 7. Absorption maxima of the compound were at 292 μm in 0.1N HCl, at 254 and 327 μm in 0.1M potassium phosphate buffer (pH 7.0), and at 245 and 312 μm in 0.1N NaOH, respectively. These spectra coincided with those of PIN. The compound was hydrolyzed in 0.055N H₂SO₄ at 120°C and 15 lb for 3 hr in an autoclave. As shown in Table VI, the Rf values of a vitamin B₆ component liberated by acid-hydrolysis were consistent with those of PIN in three solvent systems. The Rf values of a sugar component in the hydrolyzate

<table>
<thead>
<tr>
<th>Compound</th>
<th>Solvent 1</th>
<th>Solvent 2</th>
<th>Solvent 3</th>
<th>Solvent 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>PIN-G</td>
<td>0.39</td>
<td>0.07</td>
<td>0.27</td>
<td>0.37</td>
</tr>
<tr>
<td>Acid-hydrolyzate*</td>
<td>0.72</td>
<td>0.40</td>
<td>0.50</td>
<td>0.55</td>
</tr>
<tr>
<td>PIN</td>
<td>0.71</td>
<td>0.41</td>
<td>0.53</td>
<td>0.58</td>
</tr>
<tr>
<td>PAL</td>
<td>0.68</td>
<td>0.39</td>
<td>0.76</td>
<td>0.60</td>
</tr>
<tr>
<td>PAM</td>
<td>0.45</td>
<td>0.35</td>
<td>0.76</td>
<td>0.27</td>
</tr>
</tbody>
</table>

*) hydrolyzed in 0.055N H₂SO₄ at 120°C and 15 lb for 3 hr.

Solvent system:

I. n-butanol:acetic acid:water (4:1:1, v/v)
II. n-amyilalcohol:acetone:water (2:1:1, v/v, upper layer)
III. water:acetone:t-butanol:diethylamine (20:35:40:5, v/v)
IV. water:acetone:t-butanol:acetic acid (20:35:40:5, v/v)

-35-
TABLE VII. PAPER CHROMATOGRAPHY OF ISOLATED PIN-G AND ITS ACID-
HYDROLYZATE (SILVER NITRATE POSITIVE)

<table>
<thead>
<tr>
<th>Compound</th>
<th>Solvent I</th>
<th>Solvent II</th>
<th>Solvent III</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>.Rf values</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PIN-G</td>
<td>.38</td>
<td>.49</td>
<td>.67</td>
</tr>
<tr>
<td>Acid-hydrolyzate*</td>
<td>.23</td>
<td>.34</td>
<td>.54</td>
</tr>
<tr>
<td>Glucose</td>
<td>.23</td>
<td>.34</td>
<td>.54</td>
</tr>
<tr>
<td>Fructose</td>
<td>.28</td>
<td>.37</td>
<td>.56</td>
</tr>
<tr>
<td>Maltose</td>
<td>.13</td>
<td>.22</td>
<td>.47</td>
</tr>
<tr>
<td>Sucrose</td>
<td>.17</td>
<td>.28</td>
<td>.53</td>
</tr>
</tbody>
</table>

*) hydrolyzed in 0.055N H₂SO₄ at 120°C and 15 lb for 3 hr.

Solvent system:
II. n-butanol:pyridine:water (6:4:3, v/v)
III. isopropanol:pyridine:water:acetic acid (8:8:4:1, v/v)

TABLE VIII. MOLAR RATIO OF PIN TO GLUCOSE

<table>
<thead>
<tr>
<th>PIN found (µmole)</th>
<th>Glucose found (µmole)</th>
<th>Molar ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diazotized p-aminoacetophenone method</td>
<td>2.50</td>
<td>-</td>
</tr>
<tr>
<td>Anthrone method</td>
<td>--</td>
<td>2.38</td>
</tr>
<tr>
<td>Glucostat method*</td>
<td>--</td>
<td>2.33</td>
</tr>
</tbody>
</table>

*) PIN-G was hydrolyzed in 0.055N H₂SO₄, at 120°C and 15 lb for 3 hr.

Reducing activity was not found by Somogyi-Nelson method.
TABLE IX. HYDROLYSIS BY α- AND β-GLUCOSIDASES

The isolated compound was hydrolyzed under the conditions described in the EXPERIMENTAL. PIN and glucose moieties were confirmed by paper chromatography.

<table>
<thead>
<tr>
<th>Glucosidases</th>
<th>PIN liberated</th>
<th>Glucose liberated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yeast α-glucosidase</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Mold maltase</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>β-Glucosidase (Emulsin)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Cellulase</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

coincided with those of glucose, as shown in Table VII. The molar ratio of PIN to glucose was confirmed to be 1:1 by the diazotized p-aminacetophenone method,104,105) the anthrone method110) and the glucostat method111)(Table VIII). The reducing activity of the compound was not found with the method of Somogyi.112) Table IX shows the enzymatic hydrolysis of the isolated compound. The compound was hydrolyzed by yeast α-glucosidase and mold maltase, but not by β-glucosidase and cellulase. All these results indicated that the product of Micrococcus sp. No. 431 was PIN-α-glucoside. An IR spectrum of the compound supported this assumption (Fig. 8). As shown in Table X, NMR signals of PIN-G showed no difference between PIN-4'-G and PIN-5'-G.

PIN-diglucoside was also identified by various analytical methods. The ultraviolet absorption spectrum of the compound coincided with that of PIN. On hydrolysis in 0.055N H₂SO₄ at 120°C and 15 lb for 3 hr, the isolated compound gave two components corresponding to glucose
Fig. 8. IR Spectrum of Isolated PIN-G (Micro KBr Tablet).

**TABLE X. NMR SIGNALS OF PIN-4'-'G AND PIN-5'-'G**

<table>
<thead>
<tr>
<th>PIN-4'-'G</th>
<th>PIN-5'-'G</th>
</tr>
</thead>
<tbody>
<tr>
<td>δ 2.49 ppm</td>
<td>δ 2.47 ppm</td>
</tr>
<tr>
<td>δ 3.2-3.9 ppm</td>
<td>δ 3.3-3.9 ppm</td>
</tr>
<tr>
<td>δ 4.87 ppm</td>
<td>δ 4.89 ppm</td>
</tr>
<tr>
<td>δ 5.07 ppm</td>
<td>δ 5.07 ppm</td>
</tr>
<tr>
<td>δ about 5 ppm</td>
<td>δ about 5 ppm</td>
</tr>
<tr>
<td>δ 7.82 ppm</td>
<td>δ 7.85 ppm</td>
</tr>
</tbody>
</table>

(C-2', -CH$_3$, single)  
(ring protons of sugar, multiple)  
(C-5', -CH$_2$, single)  
(C-4', -CH$_2$, single)  
(α-anomeric proton)  
(H-6', single)

NMR spectra were recorded on a Hitachi Perkin-Elmer high resolution NMR spectrometer type R-22 at 90 MHz in D$_2$O with DSS as an internal standard.
Fig. 9. NMR Spectrum of Isolated PIN-diglucoside.

The spectrum was recorded on a Varian HA-100D spectrometer at 100 MHz in D$_2$O with DSS-D$_6$ as an internal standard.

and PIN, the molar ratio being confirmed to be 2:1 by the glucostat method and the diazotized p-aminoacetophenone method. NMR spectrum showed the existence of PIN and sugar moieties and two $\alpha$-protons (Fig. 9). It was suggested that PIN-4'-diglucoside, PIN-5'-diglucoside and PIN-4',5'-diglucoside might be existent. PIN-4'-diglucoside and PIN-4',5'-diglucoside may be colored with the diazotized p-aminoacetophenone reagent in the presence of borate buffer. On the other hand, PIN-5'-diglucoside may not be colored with the reagent in the presence of borate buffer. The two colored spots on a paper
chromatogram may support this assumption, as shown in Fig. 6.

DISCUSSION

The cultural conditions of the formation of PIN-G were investigated with the isolated strain, *Micrococcus* sp. No. 431, for the convenient isolation of the compound. Sucrose was the most effective glucosyl donor. It seems that the PIN-G-synthesizing enzyme may be a kind of glucosido-invertase like those from yeasts, honey, and some aphids. The amount of PIN-G increased proportionately to the concentration of added sucrose, and was remarkably accelerated by the addition of several nonionic surfactants. The effects of surfactants on the metabolism of microorganisms have been reported in several microbial processes such as glutamic acid fermentation and respiration of yeast. It is not clear whether the stimulative effect of nonionic surfactants on the formation of PIN-G is caused by the acceleration of the permeability of the product or of the enzyme responsible for the transglucosidation reaction. Under the optimal conditions of culture, more than 90% of PIN added was converted to PIN-G by *Micrococcus* sp. No. 431. PIN-G was isolated from the culture broth by the treatment with active carbon, column chromatography on Dowex 1X2 (Cl⁻ type) and Sephadex G-10, and paper chromatography. PIN-4'-G and PIN-5'-G could be separated by paper chromatography with a solvent of n-butanol:benzene:pyridine:water (5:1:3:3, v/v). The two isomers were identified by various analytical methods including absorption spectra, paper chromatography, the color development,
enzymatic hydrolysis, and IR and NMR spectra. Contrary to the case of *Sarcina lutea* IFO 3232, PIN-4'-G was dominant in *Micrococcus*. In addition to PIN-G, PIN-diglucoside was detected in the culture broth, the isolation and identification of the compound being carried out.

It is already reported that riboflavinylglycosides (mono-, di-, tri-, and tetrageglucosides)\(^7\) are formed from riboflavin and sugar by transglucosidation. The mechanism of transglucosidation to PIN to form PIN-G and PIN-diglucoside may probably be similar to the case of riboflavinylglycosides.

**SUMMARY**

With the isolated and identified strain, *Micrococcus* sp. No. 431, several conditions of the formation of PIN-G were investigated by the growing culture. Sucrose and peptone were the most proper compounds as carbon and nitrogen sources. A few nonionic surfactants exhibited a stimulative effect on the formation of PIN-G, in which Span 80 was the most effective. The formation of PIN-G reached to the maximum after 60 to 80 hr of cultivation. Metal ions had no effect. The amount of PIN-G increased proportionately to the concentration of added sucrose. Under the optimal culture conditions, the amount of PIN-G attained to maximally 1800 to 1900 μg/ml (90–95% yield based on PIN added) at 2 mg/ml of PIN. The yield was maximal at pH 7.0. PIN-G was isolated from the culture broth and identified by several methods including IR and NMR analyses. PIN-diglucoside was also produced in
the broth, and the isolation and identification of the compound were performed.
INTRODUCTION

In the previous section, the PIN-G-accumulating activity of bacteria was found to be distributed specifically in genera Sarcina and Micrococcus. An isolated bacterium No. 431, which was classified into a Micrococcus sp., had the highest activity of accumulating PIN-G in the culture broth. Various factors affecting the PIN-G formation were examined with this organism and over than 90% of PIN added was converted to PIN-G under the optimal cultural conditions. However, the enzyme catalyzing the synthesis of PIN-G has not yet been investigated and the mechanism of its formation remains to be elucidated. The present section describes a purification procedure of the enzyme from Micrococcus sp. No. 431, which catalyzes a glucosyl transfer from sucrose to PIN to form PIN-G. The author also reports some properties of PIN-G-synthesizing enzyme purified from Micrococcus sp. No. 431.

EXPERIMENTAL

Materials. Uridine diphosphate glucose was kindly given by the Laboratory of Industrial Microbiology, Department of Food Science and Technology, Kyoto University. Other chemicals used in this section were purchased from commercial sources.

Microorganisms and cultivation. The bacteria except Micrococcus
sp. No. 431 were from laboratory collections. *Micrococcus* sp. No. 431 was isolated from soil in Kyoto Prefecture as reported in the previous section.

The bacteria were grown on a medium containing 3.0% sucrose, 1.0% peptone, 0.2% yeast extract, 0.5% K$_2$HPO$_4$, 0.1% KH$_2$PO$_4$, 0.2% NaCl, and 0.02% MgSO$_4$·7H$_2$O. The pH was adjusted to 7.0. The cultivation was carried out aerobically at 28°C for 24 hr in a 2 liter-shaking flask containing 500 ml of the medium. Cultures were also performed at 28°C for 15 hr in a 20 liter jar fermentor (Marubishi Laboratory Equipment Co., Ltd.) with the rotation of impeller at 225 rpm and the aeration at 15 liters/min.

The cells harvested by centrifugation were washed once with 0.85% saline solution. The washed cells were suspended in 0.01M potassium phosphate buffer (pH 7.0) containing 3 x 10^{-3}M 2-mercaptoethanol and followed to the sonication.

Preparation of cell extract. The washed cells obtained as described above, were disrupted with a Kaijo-Denki 19 Hz ultrasonic oscillator (20 kc) at 0-15°C. The cell debris was removed by centrifugation at 10,000 rpm for 20 min and the supernatant solution was used as cell extract.

Assay of enzyme activity. Through the purification of PIN-G-synthesizing enzyme of *Micrococcus* sp. No. 431, both α-glucosidase and transglucosidase activities were assayed. In the assay of α-glucosidase activity, the standard reaction mixture contained 25 μmoles of sucrose, 100 μmoles of potassium phosphate buffer (pH 8.0),
5 μmoles of 2-mercaptoethanol and a suitable amount of enzyme in a total volume of 2.5 ml. After incubation at 30°C for 30 min, the reaction was stopped by heating the tube in a boiling water bath for 5 min. The α-glucosidase activity was determined by measuring the liberated reducing sugars. The standard reaction mixture for the assay of transglucosidase activity contained 25 μmoles of PIN, 25 μmoles of sucrose, 100 μmoles of potassium phosphate buffer (pH 8.0), 5 μmoles of 2-mercaptoethanol and a suitable amount of enzyme in a total volume of 2.5 ml. After incubation at 30°C for 60 min, the reaction was stopped by heating the tube in a boiling water bath for 5 min. PIN-G was determined by paper chromatography as described in the previous section. One unit of both enzyme activities was defined as the amount of enzyme which produced 1μmole of products per min under the conditions described above. Specific activity was shown as units per E280 μm. Protein was estimated by measuring the optical density at 280 μm with a Hitachi Perkin-Elmer 139 spectrophotometer.

Analytical methods. Reducing sugars were determined by Somogyi-Nelson method. PIN and PIN-G were determined by paper chromatography and the method described in the previous section. For the convenience of time, the ascending development was carried out for 3 hr with running by approximately 10 cm from the spotted line.

Electrophoresis. Polyacrylamide-gel electrophoresis was carried out by a modification of Davis method. Stacking and running gels were polymerized in a Pyrex tube (5 x 65 mm). After the run, the gel was stained with 1% naphthol blue-black, destained electro-
phoretically and stored in 7% acetic acid.

Ultracentrifugal analysis. Sedimentation velocities were measured with the Beckman-Spinco analytical centrifuge model E operating at 59,780 rpm. 120)

Measurement of ultraviolet absorption spectrum. The ultraviolet absorption spectrum was recorded on a Shimazu multi-purpose spectrophotometer model MPS-50L.

RESULTS

Enzyme Activity in Cell Extracts

Five species of Sarcina and Micrococcus were examined for PIN-G-forming activity using cell extracts. The result is shown in Table I. Relatively high transglucosidase activity (PIN-G-forming activity)

TABLE I. PIN-G-SYNTHESIZING ENZYME ACTIVITY IN CELL EXTRACT

Enzyme activities were assayed with the extracts prepared from the cells grown in the basal medium at 28°C for 15 hr on a reciprocal shaker under the standard conditions.

<table>
<thead>
<tr>
<th>Strains</th>
<th>Total absorbancy at 280 μm</th>
<th>α-Glucosidase</th>
<th>Transglucosidase</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>S.A.</td>
<td>T.U.</td>
</tr>
<tr>
<td><em>Micrococcus</em> sp. No. 431</td>
<td>2700</td>
<td>10.2</td>
<td>18300</td>
</tr>
<tr>
<td><em>Micrococcus flavus</em> IFO 3242</td>
<td>2580</td>
<td>4.3</td>
<td>7470</td>
</tr>
<tr>
<td><em>Micrococcus roseus</em> IFO 3764</td>
<td>0</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td><em>Sarcina lutea</em> IFO 1099</td>
<td>762</td>
<td>0.2</td>
<td>100</td>
</tr>
<tr>
<td><em>Sarcina marginata</em> IFO 3066</td>
<td>123</td>
<td>20.1</td>
<td>1650</td>
</tr>
</tbody>
</table>

S.A.: Specific activity (units/E_{280 μm})

T.U.: Total units

-46-
Both α-glucosidase and transglucosidase activities were estimated using cell extract of Micrococcus sp. No. 431 under the standard assay conditions. Incubation was carried out at 30℃.

Ο , Specific activity; ● , Total activity.

was found in two organisms, Sarcina marginata IFO 3066 and Micrococcus sp. No. 431. Total enzyme activity in cell extract of the latter organism was higher than that of the former. Therefore, it seems that Micrococcus sp. No. 431 is an appropriate enzyme source for the purification of PIN-G-synthesizing enzyme.

The effect of culturing time on the formation of both enzymes
of Micrococcus sp. No. 431 is shown in Fig. 1. The specific activities reached to maxima after about 20 hr of incubation. Total activities of α-glucosidase and transglucosidase also reached to maxima after about 20 hr of incubation and remained constant until about 60 hr.

Based on these data, Micrococcus sp. No. 431 was selected to purify PIN-G-synthesizing enzyme.

**Purification of Enzyme**

All operations for the purification of the enzyme were performed at 0-15°C.

Routine enzyme assays in the course of purification were carried out by determining an α-glucosidase activity unless otherwise noted. α-Glucosidase fractions were subjected to the assay of transglucosidase activity.

1) **Sonication.** The washed cells were suspended in 0.01M potassium phosphate buffer (pH 7.0) containing 3 x 10^{-3}M 2-mercaptoethanol, and were disrupted with a Kaijo-Denki 19 Hz ultrasonic oscillator (20 kc) at 0-15°C for 8 hr. Then, the cell debris was centrifuged off at 10,000 rpm for 20 min. The supernatant solution was used as a starting material of the purification.

2) **1st Ammonium sulfate fractionation.** To the cell extract (1425 ml, total Ε_{280} μM = 53,000) was added solid ammonium sulfate to make 0.30 saturation, followed by standing for 30 min. After removal of the resulting precipitate by centrifugation, solid ammonium sulfate was added to the supernatant solution to make 0.80 saturation.
The mixture was left to stand overnight, and the precipitate formed was collected by centrifugation at 10,000 rpm for 20 min and dissolved in 0.01M potassium phosphate buffer (pH 7.0) containing $3 \times 10^{-3}$M 2-mercaptoethanol. The solution was dialyzed overnight against the same buffer.

3) **DEAE-Sephadex column chromatography.** The dialyzed enzyme solution (386 ml, total $E_{280\text{ mu}} = 27,200$) was subjected to DEAE-
Sephadex column chromatography. A DEAE-Sephadex column (5.6 x 53 cm) was preliminarily equilibrated with 0.01M potassium phosphate buffer (pH 7.0) containing 3 × 10^{-3}M 2-mercaptoethanol. The enzyme solution was placed on the column and was washed with the same buffer. Subsequently, it was treated with 0.1M potassium phosphate buffer (pH 7.0) containing 3 × 10^{-3}M 2-mercaptoethanol and 0.2M sodium chloride. Ten milliliters of fractions were collected and the protein was followed by measurement of the absorbancy at 280 nm. Much of the inactive proteins were removed in this step. The enzyme was subsequently eluted with 0.1M potassium phosphate buffer (pH 7.0) containing 3 × 10^{-3}M 2-mercaptoethanol and 0.3M sodium chloride. The elution pattern of the enzyme is shown in Fig. 2. The active fractions (tube No. 424-466) were combined and the enzyme was precipitated by the addition of solid ammonium sulfate (0.65 saturation). The precipitate was collected by centrifugation and dissolved in 0.01M potassium phosphate buffer (pH 7.0) containing 3 × 10^{-3}M 2-mercaptoethanol, and then dialyzed overnight against 2 liters of the same buffer with three changes.

4) 2nd Ammonium sulfate fractionation. The dialyzed enzyme solution was applied to 2nd ammonium sulfate fractionation. The precipitate obtained between 0.45-0.60 saturation of ammonium sulfate was dissolved in 0.01M potassium phosphate buffer (pH 7.0) containing 3 × 10^{-3}M 2-mercaptoethanol and dialyzed for 10 hr against 2 liters of the same buffer with three changes.

5) Hydroxylapatite column chromatography. The dialyzed enzyme solution (24 ml, total $E_{280}$ = 342) was applied to hydroxylapatite
Fig. 3. Hydroxylapatite Column Chromatography.

Column size 3.6 x 13.5 cm, Flow rate 18 ml/hr, One fraction 3 ml.
- O, Protein; •, PIN-G-synthesizing enzyme activity.

column chromatography. The column (3.6 x 13.5 cm) was preliminarily bufferized with 0.01M potassium phosphate buffer (pH 7.0) containing 3 x 10^{-3}M 2-mercaptoethanol. After the enzyme solution was charged, the column was washed with the same buffer and eluted with 0.1M potassium phosphate buffer (pH 7.0) containing 3 x 10^{-3}M 2-mercaptoethanol. The elution pattern is shown in Fig. 3. Active fractions were combined and brought to 0.65 saturation by ammonium sulfate. The precipitate was taken by centrifugation and dissolved in 0.01M
Fig. 4. Second Sephadex G-100 Gel Filtration.

Column size 2.4 x 88 cm, Flow rate 24 ml/hr,
One fraction 3 ml.

O, Protein; ●, PIN-G-synthesizing enzyme activity.

potassium phosphate buffer (pH 7.0) containing $3 \times 10^{-3}$ M 2-mercaptoethanol.

6) Sephadex G-100 column chromatography. The enzyme solution (1 ml, total $E_{280 \text{ nm}} = 60$) was applied to a column of Sephadex G-100 (2.4 x 88 cm). The protein was eluted with 0.01 M potassium phosphate
TABLE II. PURIFICATION OF PIN-G-SYNTHESIZING ENZYME

<table>
<thead>
<tr>
<th>Fraction and step</th>
<th>Total absorbancy at 280 μm</th>
<th>α-Glucosidase</th>
<th>Transglucosidase</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>S.A.</td>
<td>T.U.</td>
</tr>
<tr>
<td>1. Cell extract</td>
<td>53000</td>
<td>8.5</td>
<td>302000</td>
</tr>
<tr>
<td>2. 1st AmSO₄ (30-80%)</td>
<td>27200</td>
<td>15.9</td>
<td>289000</td>
</tr>
<tr>
<td>3. DEAE-Sephadex A-50</td>
<td>342</td>
<td>365</td>
<td>82700</td>
</tr>
<tr>
<td>4. Hydroxylapatite</td>
<td>134</td>
<td>635</td>
<td>56700</td>
</tr>
<tr>
<td>5. 2nd AmSO₄ (45-65%)</td>
<td>60</td>
<td>1040</td>
<td>41800</td>
</tr>
<tr>
<td>6. Sephadex G-100 (I)</td>
<td>23</td>
<td>2450</td>
<td>38000</td>
</tr>
<tr>
<td>7. Sephadex G-100 (II)</td>
<td>11</td>
<td>3020</td>
<td>22500</td>
</tr>
</tbody>
</table>

buffer (pH 7.0) containing $3 \times 10^{-3}$ M 2-mercaptoethanol. The active fractions (fraction No. 46-60) were collected, concentrated by ammonium sulfate at 0.65 saturation, and dissolved in 0.01M potassium phosphate buffer (pH 7.0) containing $3 \times 10^{-3}$ M 2-mercaptoethanol. The enzyme solution was rechromatographed on Sephadex G-100. By the 2nd Sephadex G-100 column chromatography, a single peak was obtained as shown in Fig. 4. The fractions from No. 52 to No. 65 were used as the purified enzyme. The enzyme was purified approximately 354-fold with 7.4% recovery. A summary of the purification procedure is presented in Table II.

Purity and Properties of Enzyme

Disc electrophoresis

Disc electrophoresis (polyacrylamide-gel) was attempted with the purified enzyme. The electrophoretic pattern (Fig. 5) showed a single
band, supporting that the obtained enzyme was homogeneous.

**Ultracentrifugal analysis**

The ultracentrifugal analysis of the purified enzyme was performed with a Beckman-Spinco analytical centrifuge model E. As shown in Fig. 6, the Schlieren patterns indicated the presence of a single component. The sedimentation coefficient, \( s_{20,\text{w}}^0 \), was estimated for the enzyme (\( E_{280\text{~mu}} = 5.4 \)) to be 3.68 S.

These results show that the purified enzyme is a single protein.

**Properties of enzyme**

PIN-G-synthesizing enzyme was purified according to the method.

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**Fig. 5. Acrylamide-gel Electrophoresis of PIN-G-Synthesizing Enzyme.**

The purified enzyme (31.5 μg) was applied to the stacking gel and subjected to the electrophoresis at a current of 2.0 mA for 50 min in Tris-glycine buffer (pH 8.3) containing 3 \( \times \) 10\(^{-3}\)M 2-mercaptoethanol. The direction of migration is from the cathode to the anode.
Fig. 6. Sedimentation Patterns of PIN-G-synthesizing Enzyme.

The experiment was carried out at a concentration of 0.36% enzyme in 0.01M potassium phosphate buffer (pH 7.0) containing 3 x 10^{-3}M 2-mercaptoethanol. The photographs were taken at 26, 36, 51 and 66 min after reaching 59,780 rpm. The sedimentation is left to right.

described above. The first ammonium sulfate fraction (the partially purified sample) and the 2nd Sephadex G-100-treated fraction (the purified sample), homogeneous on ultracentrifugation and polyacrylamide-gel electrophoresis, were used throughout this section.

_Ultraviolet absorption spectrum_

The ultraviolet absorption spectrum of the purified enzyme in 0.01M potassium phosphate buffer (pH 7.0) containing 3 x 10^{-3}M 2-mercaptoethanol was measured with a Shimazu multi-purpose recording spectrophotometer model MPS-50L, as shown in Fig. 7. The spectrum showed a typical protein spectrum and the absorption maximum and minimum were found to be at 280 μm and 253 μm, respectively. $E_{280 \text{ μm}}/E_{260 \text{ μm}} = 1.6$. 

-55-
The purified enzyme was dissolved in 0.01M potassium phosphate buffer (pH 7.0) containing $3 \times 10^{-3}$M 2-mercaptoethanol. The spectrum of the solution was measured by a Shimazu multi-purpose spectrophotometer model MPS-50L.

**Linearity of enzyme activity**

A linear relationship was obtained between enzyme activity and reaction time, up to 30 min for $\alpha$-glucosidase activity and up to 120 min for transglucosidase activity, as shown in Fig. 8. A linearity between enzyme activity and enzyme concentration was observed, up to 150 units for $\alpha$-glucosidase activity and up to 12 units for transglucosidase activity, as shown in Fig. 9.

**Stability of enzyme**

After an aliquot of enzyme solution was kept at various pH at 40°C for 10 min, the remaining enzyme activity was estimated. As shown
Both α-glucosidase and transglucosidase activities were estimated using 1st ammonium sulfate fraction under the standard assay conditions. Incubations were carried out at 30°C.

- α-Glucosidase activity (48.0 units)
- Transglucosidase activity (7.3 units)

Both α-glucosidase and transglucosidase activities were estimated using 1st ammonium sulfate fraction under the standard assay conditions. Incubations were carried out for 30 min (α-glucosidase activity) and 60 min (transglucosidase activity) at 30°C.
An aliquot of enzyme solution (1st ammonium sulfate fraction) was kept at various pH at 40°C for 10 min. After adjustment of pH 8.0, the residual enzyme activity was measured under the standard assay conditions. Incubations were made for 30 min (α-glucosidase activity) and 60 min (transglucosidase activity) at 30°C.

- α-Glucosidase activity (19.0 units)
- Transglucosidase activity (10.4 units)

in Fig. 10, both α-glucosidase and transglucosidase activities were stable at pH 7.0, where 70-80% of initial activities remained.

Figure 11 shows the stability of the enzyme activity against heating. The enzyme was heated at various temperature at pH 8.0 for 15 min. The enzyme was stable in a range from 0°C to 30°C. However,
Both α-glucosidase and transglucosidase activities were estimated using 1st ammonium sulfate fraction under the standard assay conditions. Incubations were carried out for 30 min (α-glucosidase activity) and 60 min (transglucosidase activity) at 30°C at various pH.

- α-Glucosidase activity (26.0 units)
- Transglucosidase activity (20.0 units)

--- Acetate buffer —— Phosphate buffer ——— Tris buffer

It was shown that transglucosidase activity was more stable than α-glucosidase activity against heating; at 80°C, almost all α-glucosidase activity were lost, but 25% of transglucosidase activity remained. It seems that transglucosidase activity is stabilized against heating in the presence of PIN.

**Effect of pH**

The activity curves of the enzyme are shown in Fig. 12. Higher enzyme activities were found at pH of alkaline side, and optimal pH
Fig. 13. Effect of Temperature on Enzyme Activity.

Both α-glucosidase and transglucosidase activities were estimated using 1st ammonium sulfate fraction under the standard assay conditions. Incubations were carried out for 30 min (α-glucosidase activity) and 60 min (transglucosidase activity) at various temperatures.

- α-Glucosidase activity (52.7 units)
- Transglucosidase activity (9.5 units)

was 8.0 with both enzyme activities in phosphate buffer. However, when Tris-HCl buffer was used in place of phosphate buffer, the both activities decreased extremely even at pH 8.0.

Effect of temperature

Optimal temperature on the enzyme activities was observed around 37°C, but the enzyme activities markedly decreased above 40°C under the standard assay conditions, as shown in Fig. 13.

Substrate specificity
TABLE III. EFFECT OF GLUCOSYL DONORS

The reaction was performed using the purified enzyme preparation under the standard reaction conditions except 25 μmoles of glucosyl donor.

<table>
<thead>
<tr>
<th>Glucosyl donor</th>
<th>α-Glucosidase activity (μmole)</th>
<th>Transglucosidase activity (μmole)</th>
</tr>
</thead>
<tbody>
<tr>
<td>D-Glucose</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>D-Fructose</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>Sucrose</td>
<td>362</td>
<td>599</td>
</tr>
<tr>
<td>Maltose</td>
<td>444</td>
<td>892</td>
</tr>
<tr>
<td>Trehalose</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Lactose</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Starch(^a)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Glycogen (^a)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Phenyl-α-glucoside</td>
<td>624</td>
<td>2070</td>
</tr>
<tr>
<td>Methyl-α-glucoside</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Glucose-6-phosphate</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>α-Glucose-1-phosphate</td>
<td>- NaF</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>+ NaF (10^{-2})M</td>
<td>0</td>
</tr>
<tr>
<td>UDPG (^c)</td>
<td>-</td>
<td>0</td>
</tr>
</tbody>
</table>

\(^a\) 10 mg in the standard reaction mixture except 25 μmoles of sucrose.
\(^b\) α-Glucosidase activity 24.1 units, Transglucosidase activity 10.0 units.
\(^c\) Uridine diphosphate glucose.

The effect of various glucosides on the enzyme activity was examined with mono-, di-, and poly-saccharides, methyl- and phenyl-α-D-glucosides, sugar phosphate esters and uridine diphosphate glucose. As shown in Table III, both α-glucosidase and transglucosidase activi-
TABLE IV. EFFECT OF GLUCOSYL ACCEPTORS

The reaction was performed using the purified enzyme preparation (7.6 units) under the standard reaction conditions except 25 μmoles of glucosyl acceptor.

The formation of glucoside was determined by paper chromatography using Toyo filter paper No. 53. The ascending development was carried out in the solvent systems (1), (2) and (3). Glucoside on paper was detected by Manaslu lamp (2536 A filter) and confirmed by spraying with the diazotized p-aminoacetophenone reagent. The ultraviolet absorbing spot of PIN-G was extracted with a mixture of 2.5 ml of 99% ethanol and 2.5 ml of 25% sodium acetate for 90 min at 37°C. The extracted solution was assayed colorimetrically at 470 μm according to the diazotized p-aminoacetophenone method with a Hitachi photoelectric photometer EPO-B.

Solvent system:
(1) acetone:t-butanol:diethylamine:water (35:40:5:20, v/v)
(2) t-butanol:formate:water (70:15:15, v/v)
(3) t-amylalcohol:acetone:diethylamine:water (40:35:5:20, v/v)

<table>
<thead>
<tr>
<th>Glucosyl acceptor</th>
<th>Transglucosidase activity(μmumole)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PIN</td>
<td>453</td>
</tr>
<tr>
<td>PAL</td>
<td>0</td>
</tr>
<tr>
<td>PAM</td>
<td>0</td>
</tr>
<tr>
<td>4-Pyridoxic acid</td>
<td>0</td>
</tr>
</tbody>
</table>

ties were revealed in the presence of sucrose, maltose and phenyl-α-D-glucoside. The reaction rates of enzyme decreased in the following order: phenyl-α-D-glucoside, maltose and sucrose. Essentially no enzyme activity was observed with methyl-α-D-glucoside. α-Glucose-1-phosphate and uridine diphosphate glucose were also inert.
Fig. 14. Effect of Concentrations of Glucosyl Donors 
on Enzyme Activity.

The reactions were carried out at 30°C for 60 min in reaction mixtures 
containing 42.5 units of the enzyme (1st ammonium sulfate fraction), 
25 μmoles of PIN, 100 μmoles of potassium phosphate buffer (pH 8.0), 
5 μmoles of 2-mercaptoethanol and indicated amounts of sucrose, maltose, 
or phenyl-α-D-glucoside as a substrate in a total volume of 2.5 ml. 
Velocity (V) was expressed as μmoles of PIN-G formed/min/unit, and 
substrate concentration (S) as moles per liter.

From these observations, the enzyme seems to belong to one of 
α-glucosidase, but not to α-methylglucosidase, phosphorylase or 
uridine diphosphate glucosyl transferase.

The effect of vitamin B₆ compounds on the enzyme activity was 
examined with PIN, PAL, PAM and 4-pyridoxic acid. As shown in Table 
IV, only PIN served as glucosyl acceptor.

Kinetics

Figure 14 shows the effect of concentrations of sucrose, maltose
Fig. 15. Effect of PIN Concentrations on Enzyme Activity.

The reactions were carried out at 30°C for 60 min in reaction mixtures containing 40.5 units of the enzyme (1st ammonium sulfate fraction), 25 μmoles of sucrose (I), maltose (II), or phenyl-α-D-glucoside (III), 100 μmoles of potassium phosphate buffer (pH 8.0), 5 μmoles of 2-mercaptoethanol and indicated amounts of PIN, in a total volume of 2.5 ml. Velocity (V) was expressed as μmoles of PIN-G formed/min/unit, and substrate concentration (S) as moles per liter.

and phenyl-α-D-glucoside on the formation of PIN-G.

The effect of concentrations of PIN on the formation of PIN-G is shown in Fig. 15.

K\text{ms} for sucrose, maltose and phenyl-α-D-glucoside were calculated from Lineweaver-Burk's plots to be 1.25 \times 10^{-2} M, 1.00 \times 10^{-2} M and 5.26
TABLE V. KINETIC PARAMETERS FOR SUBSTRATES OF ENZYME

<table>
<thead>
<tr>
<th>Substrate</th>
<th>$K_m$ (M)</th>
<th>$v_{max}$ (μmole/min/unit)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sucrose</td>
<td>$1.3 \times 10^{-2}$</td>
<td>2.5</td>
</tr>
<tr>
<td>Maltose</td>
<td>$1.0 \times 10^{-2}$</td>
<td>3.3</td>
</tr>
<tr>
<td>Phenyl-α-D-glucoside</td>
<td>$5.3 \times 10^{-2}$</td>
<td>13.1</td>
</tr>
<tr>
<td>PIN (Sucrose)</td>
<td>$7.1 \times 10^{-3}$</td>
<td>1.8</td>
</tr>
<tr>
<td>(Maltose)</td>
<td>$3.6 \times 10^{-3}$</td>
<td>2.0</td>
</tr>
<tr>
<td>(Phenyl-α-D-glucoside)</td>
<td>$5.7 \times 10^{-3}$</td>
<td>3.3</td>
</tr>
</tbody>
</table>

$x \times 10^{-2}$ M, respectively. $K_m$s for PIN in the coexistence of sucrose, maltose or phenyl-α-D-glucoside were also calculated to be $7.14 \times 10^{-3}$ M, $3.57 \times 10^{-3}$ M and $5.71 \times 10^{-3}$ M, respectively. These $K_m$ values are summarized in Table V.

Effect of various agents

As shown in Table VI, the enzyme activity was affected by sulfhydryl reagents and heavy metal ions. The activity was almost completely inhibited by p-chloromercuric benzoate at a concentration of $10^{-4}$ M. Iodoacetate was a less effective inhibitor at a concentration of $10^{-4}$ M. However, the inhibition was nearly overcome when 2-mercaptoethanol ($10^{-3}$ M) coexisted with p-chloromercuric benzoate ($10^{-4}$ M) or monooiodoacetate ($10^{-4}$ M). Heavy metal ions, which are known as the inhibitors of the sulfhydryl enzymes, also showed an inhibitory effect on the enzyme activity. At a concentration of $10^{-4}$ M of metal ion, the activity was almost or completely inhibited by Cu$^{2+}$, Hg$^{2+}$, Ag$^+$ and Pb$^{2+}$, and moderately inhibited by Zn$^{2+}$. Similar sensitivities to
TABLE VI. EFFECT OF SULFHYDRL INHIBITORS

Reaction mixture contained sucrose 25 μmoles, PIN 25 μmoles, potassium phosphate buffer (pH 8.0) 100 μmoles, the dialyzed enzyme solution (the purified enzyme preparation, 13.8 units) and each reagent in a total volume of 3.0 ml. Incubation was made for 60 min at 30°C.

<table>
<thead>
<tr>
<th>Agent</th>
<th>Concentration (M)</th>
<th>Transglucosidase inhibition (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>PCMB*</td>
<td>10^{-4}</td>
<td>96</td>
</tr>
<tr>
<td>PCMB + 2-Mercaptoethanol</td>
<td>10^{-4} + 2 x 10^{-3}</td>
<td>0</td>
</tr>
<tr>
<td>Monoiodoacetate</td>
<td>10^{-4}</td>
<td>19</td>
</tr>
<tr>
<td>Monoiodoacetate + 2-</td>
<td>10^{-4} + 2 x 10^{-3}</td>
<td>0</td>
</tr>
<tr>
<td>Mercaptoethanol</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AgNO₃</td>
<td>10^{-4}</td>
<td>100</td>
</tr>
<tr>
<td>HgCl₂</td>
<td>10^{-4}</td>
<td>89</td>
</tr>
<tr>
<td>CuSO₄</td>
<td>10^{-4}</td>
<td>99</td>
</tr>
<tr>
<td>2(PbCO₃)Pb(OH)₂</td>
<td>10^{-4}</td>
<td>82</td>
</tr>
<tr>
<td>Zn(CH₃COO)₂</td>
<td>10^{-4}</td>
<td>36</td>
</tr>
</tbody>
</table>

*) p-Chloromercuric benzoate

sulfhydryl reagents or heavy metals have been reported with intestine and yeast α-glucosidases. Thus, the enzyme seems to be a sulfhydryl enzyme.

Table VII shows the effect of various agents on the enzyme activity. The activity was not appreciably affected by Na₂HAsO₄, NaF, KCN, carbonyl reagents and chelating agents. But, Tris caused a 56% inhibition at the concentration of 10^{-2}M, which is a well-known
TABLE VII. EFFECT OF INHIBITORS

Reaction mixture contained sucrose 25 µmoles, PIN 25 µmoles, potassium phosphate buffer (pH 8.0) 100 µmoles, 2-mercaptoethanol 5 µmoles, the enzyme solution (the purified enzyme preparation, 15.0 units) and each reagent in a total volume of 3.0 ml. Incubation was made for 60 min at 30°C.

<table>
<thead>
<tr>
<th>Agent</th>
<th>Concentration (M)</th>
<th>Transglucosidase inhibition (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>NaF</td>
<td>$5 \times 10^{-4}$</td>
<td>17</td>
</tr>
<tr>
<td>KCN</td>
<td>&quot;</td>
<td>9</td>
</tr>
<tr>
<td>$\text{Na}_2\text{HAsO}_4$</td>
<td>&quot;</td>
<td>12</td>
</tr>
<tr>
<td>Phenylhydrazine•HCl</td>
<td>&quot;</td>
<td>13</td>
</tr>
<tr>
<td>Semicarbazide•HCl</td>
<td>&quot;</td>
<td>4</td>
</tr>
<tr>
<td>$\alpha,\alpha'$-Dipyridyl</td>
<td>$10^{-4}$</td>
<td>1</td>
</tr>
<tr>
<td>$\sigma$-Phenanthroline</td>
<td>&quot;</td>
<td>1</td>
</tr>
<tr>
<td>EDTA a)</td>
<td>&quot;</td>
<td>5</td>
</tr>
<tr>
<td>Tris b)</td>
<td>&quot;</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>$10^{-3}$</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>$10^{-2}$</td>
<td>56</td>
</tr>
</tbody>
</table>

a) Ethylenediamine tetra-acetic acid
b) Tris-HCl buffer (pH 8.0)

Effect of metal ions

The effect of metal ions at $10^{-3}$M on the enzyme activity was examined in the presence of 2-mercaptoethanol at $1.7 \times 10^{-3}$M and the
TABLE VIII. EFFECT OF METAL IONS

Reaction mixture contained sucrose 25 μmoles, PIN 25 μmoles, potassium phosphate buffer (pH 8.0) 100 μmoles, 2-mercaptoethanol 5 μmoles, the enzyme solution (the purified enzyme preparation, 5.3 units) and each metal ion (10^{-3}M) in a total volume of 3.0 ml. Incubation was made for 75 min at 30°C.

<table>
<thead>
<tr>
<th>Metal ion</th>
<th>Transglucosidase activity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>100</td>
</tr>
<tr>
<td>NaCl</td>
<td>94</td>
</tr>
<tr>
<td>KCl</td>
<td>100</td>
</tr>
<tr>
<td>LiCl</td>
<td>99</td>
</tr>
<tr>
<td>BaCl₂</td>
<td>113</td>
</tr>
<tr>
<td>CaCl₂</td>
<td>113</td>
</tr>
<tr>
<td>FeSO₄</td>
<td>114</td>
</tr>
<tr>
<td>FeCl₃</td>
<td>106</td>
</tr>
<tr>
<td>MgSO₄</td>
<td>105</td>
</tr>
<tr>
<td>NiSO₄</td>
<td>38</td>
</tr>
<tr>
<td>CoCl₂</td>
<td>61</td>
</tr>
<tr>
<td>MnSO₄</td>
<td>104</td>
</tr>
<tr>
<td>Cr₂(SO₄)₃</td>
<td>104</td>
</tr>
<tr>
<td>Na₂MoO₄</td>
<td>95</td>
</tr>
</tbody>
</table>

results are shown in Table VIII. The enzyme was dialyzed overnight against 0.01M potassium phosphate buffer (pH 7.0) containing 5 x 10^{-3} M 2-mercaptoethanol and then tested. Ni²⁺ caused a 60% inhibition and Co²⁺ a 40% inhibition. The activity was practically unaffected by other metal ions tested.
Fig. 16. Determination of Reaction Products by Two-dimensional Paper Chromatography.

Solvent system:
First development; n-butanol:acetic acid:water (4:1:1, v/v).

Identification of Reaction Products by Two-dimensional Paper Chromatography

The reaction products of PIN-G-synthesizing enzyme were confirmed by two-dimensional paper chromatography. The result is shown in Fig. 16. An appropriate amount of the reaction mixture using the purified enzyme preparation was spotted on Toyo filter paper No. 53. The ascending development was carried out in the direction of one-dimension with running by about 26 cm from the spotted line in the following solvent system; n-butanol:acetic acid:water (4:1:1, v/v).
The ascending development was also carried out in the direction of two-dimension with running by 27 cm from the base line in the following solvent system: n-butanol:pyridine:water (6:4:3, v/v). The products on a paper chromatogram were detected with a Manaslu lamp and qualitatively identified by spraying the diazotized p-aminoacetophenone reagent. PIN-G gave two orange spots. Thereafter 0.2M borate buffer (pH 9.0) was sprayed on a paper, followed by drying at 60°C for 10 min. Then, one of the orange spots changed to yellow. These two compounds were identified as PIN-4'-G and PIN-5'-G, as described in the preceding section. Other two spots with smaller Rf values than those of PIN-G were also detected as shown in Fig. 16. One spot gave an orange color in the presence of borate and another orange spot disappeared in the presence of borate. These compounds also seemed to be PIN-4'-G and PIN-4',5'-diglucoside, and PIN-5'-diglucoside, respectively.

DISCUSSION

Throughout the purification of PIN-G-synthesizing enzyme from Micrococcus sp. No. 431, transglucosidase activity to form PIN-G was always associated with α-glucosidase activity to hydrolyze sucrose. It may be indicated that both activities are due to a single enzyme and that the synthesis of PIN-G is catalyzed by a glucosido-invertase which transfers a glucosyl residue from sucrose to PIN.

A number of α-glucosidases with different substrate specificities have been purified from microorganisms, plants and animals. Most of
invertase preparations from various sources\textsuperscript{125-127} catalyze the transfer of fructosyl residue of sucrose. \(\alpha\)-Glucosidases which catalyze the transglucosidation from sucrose to certain alcoholic groups are found in brewer's yeasts,\textsuperscript{114,128,129} honey,\textsuperscript{115} insects,\textsuperscript{116} and bacteria.\textsuperscript{130} It is of interest to note whether or not these \(\alpha\)-gluco-
sidases catalyze the synthesis of PIN-G from PIN and sucrose.

The purified PIN-G-synthesizing enzyme utilized phenyl-\(\alpha\)-D-
glucoside and maltose as well as sucrose as glucosyl donors. It showed no transglucosidase activity with \(\alpha\)-glucose-1-phosphate, which is an intermediate in the reaction of sucrose phosphorylase. These results are compatible to the assumption that the enzyme belongs to a glucosido-
invertase (\(\alpha\)-glucosidase).

It is known that yeast \(\alpha\)-glucosidase has a low specificity for glucosides of holoside and heteroside while mold \(\alpha\)-glucosidase has a high one. PIN-G-synthesizing enzyme purified from \textit{Micrococcus} sp. No. 431 had a specificity for both holoside and heteroside.

Among vitamin B\textsubscript{6} compounds tested, only PIN was effective as glucosyl acceptor for the formation of PIN-G. Yet, further investigations are required on the specificity of the enzyme for other glucosyl acceptors. It was also shown that this enzyme examined here, like many other \(\alpha\)-glucosidases, was characteristic of the sulfhydryl nature.

\textbf{SUMMARY}

An enzyme catalyzing the synthesis of PIN-G was demonstrated in cell extracts of \textit{Sarcina} and \textit{Micrococcus} incubated on a medium contai-
ning PIN and sucrose. *Micrococcus* sp. No. 431 was found to have a high activity of the enzyme and the enzyme activity reached to a maximum after 20 hr of cultivation.

The enzyme which synthesized PIN-G from sucrose and PIN by trans-glucosidation was purified from *Micrococcus* sp. No. 431 by means of ammonium sulfate fractionation, and DEAE-Sephadex, hydroxylapatite and Sephadex G-100 column chromatography. The enzyme was purified about 354-fold and confirmed to be homogeneous on polyacrylamide-gel electrophoresis and ultracentrifugation.

Some properties of PIN-G-synthesizing enzyme were investigated with purified enzyme preparations from *Micrococcus* sp. No. 431.

The enzyme was stable at pH 7.0 and between 0°C and 30°C. The maximal activity was obtained at pH 8.0 and 37°C. Besides sucrose, phenyl-α-D-glucoside and maltose served as glucosyl donors. Of vitamin B₆ compounds tested, only PIN served as glucosyl acceptor. The enzyme activity was inhibited by PCMB and heavy metal ions, and the inhibition was prevented by 2-mercaptoethanol, indicating the enzyme would be a sulfhydryl enzyme. The activity was not affected by chelating agents and not activated by metal ions.

The reaction products of PIN-G-synthesizing enzyme were confirmed by two-dimensional paper chromatography. Besides PIN-G (PIN-4'-G and PIN-5'-G), PIN-diglucoside was detected on a paper chromatogram.
Section IV. Stability and Biological Activities of PIN-G

INTRODUCTION

In the previous sections, it has been reported that PIN-G, PIN-4'-G or PIN-5'-G, is synthesized from PIN and glucosyl donors e.g. sucrose, maltose or phenyl-α-D-glucoside by various microorganisms including bacteria, yeasts, molds and actinomycetes. Micrococcus sp. No. 431, having the highest activity for PIN-G formation, accumulated maximally 1800 to 1900 μg/ml of PIN-G (90-95% yield based on PIN added) at 2 mg/ml of PIN. PIN-G-synthesizing enzyme was purified from Micrococcus sp. No. 431 to a homogeneous state. It was suggested that the enzyme might be a glucosido-invertase (α-glucosidase) which catalyzed the transfer of glucosyl residue from sucrose to PIN to form PIN-G.

However, the properties and biological activity of PIN-G have not been investigated as yet. The present paper describes the stability of PIN-G against irradiation of ultraviolet light and heating in comparison with that of PIN and PIN-borate complex. The microbiological activity of PIN-G is also examined in this section. As to the investigations on the transport of vitamin B₆ into human erythrocytes, Yamada et al.¹³¹-¹³³ have reported that free forms of vitamin B₆ are actively transported into the erythrocytes against a concentration gradient, and Suzue et al.¹³⁴ have reported that PAL-P is transported into cells without hydrolysis to PAL. It is of interest to note whether PIN-G is also transported into erythrocytes. In this section, the cellular transport of PIN-G with rabbit erythrocytes was investi-
gated in vitro, comparing with that of PIN as a standard. The various factors affecting the transport of vitamin B₆ compounds were also investigated.

EXPERIMENTAL

Materials. PIN-G was prepared from PIN and sucrose by intact cells of Micrococcus sp. No. 431. A reaction mixture containing 2 g of PIN, 5 g of sucrose, 35 mmoles of potassium phosphate buffer (pH 8.0) and 2.2 g (as dry wt.) of the intact cells in a total volume of 350 ml was incubated aerobically at 28°C for 11 hr. After removal of cells by centrifugation, the supernatant solution was adjusted to pH 2.0, and 50 g of active charcoal was added, and stirred at room temperature for 2 hr. Active charcoal collected by filtration was washed with 100 ml of 0.01N HCl and 200 ml of deionized water, followed by elution with ammoniacal ethanol (99% ethanol 50:28% ammonia 5:water 45, v/v). The eluate containing PIN and PIN-G was concentrated under reduced pressure at 30°C and subjected to Sephadex G-10 column chromatography (1 X 120 cm). PIN-G was eluted with deionized water at a flow rate of 7 ml per hr. The PIN-G fractions were collected and concentrated under reduced pressure at 30°C to about 5 ml and applied to the preparative paper chromatography on Toyo filter paper No. 526 with a solvent system of n-butanol:acetic acid:water (4:1:1, v/v). A small amount of concomitant PIN could be removed from PIN-G in this procedure. The fluorescent zone of PIN-G ($R_f$ = 0.4) was cut off and extracted with deionized water. The extract was concentrated under

-74-
reduced pressure and used in this section. The amount of PIN-G was indicated as that of PIN moiety.

The Alsevers-treated rabbit erythrocytes were purchased from commercial sources. Other materials used in this work were commercial products.

Degradation of vitamin B$_6$ compounds by ultraviolet irradiation and heating. Ultraviolet light by a sterilizing lamp GL-15 (National Electric Co., Ltd.) was irradiated directly upon 5 ml of a sample solution in a test tube (1.5 X 11 cm) from a distance of about 30 cm. Five milliliters of the sample was repeatedly heated in a test tube (1.6 X 160 cm) five times at 120°C and 15 lb for 1 hr in an autoclave.

The degradation of vitamin B$_6$ compounds was followed by measuring the decrease in the absorbancy at 254 µm and 325 µm for PIN-G and PIN, and at 295 µm for PIN-borate complex with a Shimazu multi-purpose spectrophotometer model MPS-50L.

Microbiological assay. Microbiological activity of vitamin B$_6$ was assayed by the method of Atkin et al.,$^{135}$ using Saccharomyces carlsbergensis 4228 ATCC 9080.

Transport of vitamin B$_6$ into rabbit erythrocytes. Experiments were performed according to the method of Yamada et al.$^{131-133)}$

1. Preparation of rabbit erythrocytes. The commercial rabbit erythrocytes were washed three times at room temperature with an ice-cold 0.9% saline solution by centrifugation and resuspension. The fluffy layer of white blood cells was removed as much as possible. The packed cells were finally suspended in an equal volume of 0.9%
saline solution, and equilibrated to the desired temperature in a water bath for 10 min. Aliquots of this suspension were used in the experiment.

2. Incubation and analysis. The incubation mixture contained an aliquot of the erythrocyte suspension having a hematocrit value of 0.50, 6 mM glucose and a suitable amount of vitamin B₆ in total 5 ml of 1/15M Krebs-Ringer phosphate buffer (pH 7.4). The incubation was carried out at 37°C for the indicated time intervals. After incubation, the cells were collected by centrifugation at 3,000 rpm for 10 min, and washed three times with an ice-cold 0.9% saline solution. The cells were then hemolyzed in 8 ml of 1/12N H₂SO₄ to liberate intracellular vitamin B₆ compounds. The hemolysate was hydrolyzed at 120°C and 15 lb for 210 min in an autoclave, and was deproteinized by adding 0.5 ml of 10% sodium tungstate. The supernatant solution was used for determining the total vitamin B₆ by microbioassay with *Saccharomyces carlsbergensis* 4228 ATCC 9080. PIN-G could be assayed as PIN-1 according to this procedure.

3. Preparation of ghosts (erythrocyte membranes). According to the method of Post, 20 ml of erythrocytes were washed three times with 70 ml of an ice-cold 0.9% saline solution at room temperature. All the further procedures were carried out at 2°C. The cells were packed tightly by centrifugation, and the supernatant solution was discarded. Sixty milliliters of distilled water was added to the packed cells by forceful injection from a syringe. The hemolysate mixed by inversion was centrifuged at 15,000 x g for 30 min.
transparent dark red supernatant solution was carefully sucked and discarded. The residue was suspended into 80 ml of $5 \times 10^{-4}$ M histidine-imidazole buffer (pH 7.1), followed by centrifugation at 15,000 x g for 10 min. The treatment was repeated eight times until the supernatant solution became colorless. A large volume of fluffy pink precipitate, called ghosts, was separated by decantation and was stored at pH 7.

4. Determination of vitamin $B_6$ compounds adsorbed on ghost cells. An appropriate amount of PIN or PIN-G was incubated with histidine-imidazole buffer (histidine 7 µmoles and imidazole 13 µmoles, pH 7.1), 5 µmoles of MgCl₂, 200 µmoles of NaCl, 200 µmoles of KCl, 4 µmoles of adenosine triphosphate (ATP), and the ghosts in a final volume of 4.0 ml. The reaction was carried out at 37°C for 60 min. After removal of the ghosts by centrifugation, the amounts of PIN or PIN-G in the supernatant fluid were determined by microbioassay. PIN-G was assayed as PIN after hydrolysis to PIN and glucose by heating at 120°C and 15 lb for 210 min in 0.055N H₂SO₄.

RESULTS

Stability of PIN-G against Ultraviolet Light

The stability of PIN-G against ultraviolet light was investigated in comparison with that of PIN or PIN-borate. PIN-G and PIN were adjusted to pH 6.8 with NaOH. PIN-borate complex was prepared by mixing PIN with borate buffer, pH 6.8, in a final concentration of 0.1M. Changes in absorption spectra of PIN-G, PIN, and PIN-borate
Irradiation changes in ultraviolet absorbancy of vitamin B₆ compounds during irradiation

<table>
<thead>
<tr>
<th>Irradiation</th>
<th>PIN-G</th>
<th>PIN</th>
<th>PIN-borate</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>E₂₅₄ μ</td>
<td>E₃₂₅ μ</td>
<td>E₂₅₄ μ</td>
</tr>
<tr>
<td>0 hr</td>
<td>1.62 (100)</td>
<td>1.18 (100)</td>
<td>1.07 (100)</td>
</tr>
<tr>
<td>6</td>
<td>1.58 (98)</td>
<td>1.02 (86)</td>
<td>0.99 (93)</td>
</tr>
<tr>
<td>9</td>
<td>1.55 (96)</td>
<td>0.97 (82)</td>
<td>0.98 (92)</td>
</tr>
<tr>
<td>22</td>
<td>1.52 (94)</td>
<td>0.81 (69)</td>
<td>0.84 (79)</td>
</tr>
<tr>
<td>47</td>
<td>1.47 (91)</td>
<td>0.64 (54)</td>
<td>0.58 (54)</td>
</tr>
</tbody>
</table>

Complex on irradiation of ultraviolet light are shown in Fig. 1.

As shown in Table I, PIN-borate complex is the most stable against ultraviolet light, for 60% of the initial absorbancy at 295 μ remains even after 47 hr of irradiation. The higher stability of PIN-G was observed, 54% of the original absorbancy at 325 μ being retained after 47 hr of irradiation. On the other hand, the absorbancy of PIN decreased to 30% of the original.

Stability of PIN-G against Heating

PIN and PIN-G neutralized to pH 6.8 with NaOH were heated five times at 120°C and 15 lb for 1 hr in an autoclave, cooled, and followed by standing in an ice box overnight. After centrifugation, the absorbancy of the supernatant solution was measured. The result is shown in Table II. About 90% of the initial absorbancy of PIN-G was retained. PIN-borate complex was not degraded at all, while PIN was decomposed to about 80% of the initial absorbancy.
Fig. 1. Changes in Absorption Spectra of PIN, PIN-G and PIN-borate Complex during Irradiation.

The irradiation of ultraviolet light was performed as described in the text. Ultraviolet absorption spectra were recorded on a Shimazu multi-purpose spectrophotometer.
TABLE II. CHANGES IN ULTRAVIOLET ABSORBANCY OF VITAMIN B<sub>6</sub> COMPOUNDS BY HEATING

Heating was performed as described in the text. Ultraviolet absorbancy was measured by a Shimazu multi-purpose spectrophotometer.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Wave length</th>
<th>Absorbancy Heating time</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0 hr</td>
</tr>
<tr>
<td>PIN-G</td>
<td>E&lt;sub&gt;254&lt;/sub&gt; mμ (%)</td>
<td>1.62 (100)</td>
</tr>
<tr>
<td></td>
<td>E&lt;sub&gt;325&lt;/sub&gt; mμ (%)</td>
<td>1.18 (100)</td>
</tr>
<tr>
<td>PIN</td>
<td>E&lt;sub&gt;254&lt;/sub&gt; mμ (%)</td>
<td>0.77 (100)</td>
</tr>
<tr>
<td></td>
<td>E&lt;sub&gt;325&lt;/sub&gt; mμ (%)</td>
<td>1.40 (100)</td>
</tr>
<tr>
<td>PIN-borate</td>
<td>E&lt;sub&gt;295&lt;/sub&gt; mμ (%)</td>
<td>1.03 (100)</td>
</tr>
</tbody>
</table>

From these results, it seems that PIN-G is more stable than PIN against irradiation of ultraviolet light and heating.

Effect of Heating Time on Hydrolysis of PIN-G

PIN-G could be bioassayed as PIN, after hydrolysis to PIN and glucose, by heating at 120°C and 15 lb in 0.055N H<sub>2</sub>SO<sub>4</sub>. The effect of heating time on hydrolysis of PIN-G was investigated. The result is shown in Fig. 2. Thirty-three per cent of PIN-G was hydrolyzed by heating for 1 hr, and 100% hydrolysis was achieved for 3 hr by the heat treatment.

Microbiological Activity of PIN-G for Saccharomyces carlsbergensis 4228 ATCC 9080

As shown in Fig. 3, PIN-G showed about 20% of microbiological
Fig. 2. Effect of Heating Time on Hydrolysis of PIN-G.

The growth of *S. carlsbergensis* 4228 ATCC 9080 on the Atkin's medium containing each hydrolyzed sample (PIN-G 30.4 mg as PIN) was measured from optical density at 610 μm after 23 hr of incubation.

Fig. 3. Microbiological Activity of PIN-G.

Microbiological activity was assayed as described in the text. Optical density at 610 μm was measured after each time of incubation as shown in the figure.
activity of equivalent mole of PIN after incubation for 24 hr. The microbiological activity of PIN-G was increased with the prolonged incubation and reached to above 50% of activity of equivalent mole of PIN.

**Transport of PIN-G into Rabbit Erythrocytes**

**Effect of incubation time on the transport of PIN and PIN-G into erythrocytes**

The cells were incubated with 5 µg of PIN and PIN-G for 10, 30 and 60 min under the conditions described in the EXPERIMENTAL. The results are given in Table III. The amount of vitamin B₆ in the

**TABLE III. EFFECT OF INCUBATION TIME ON THE TRANSPORT OF PIN AND PIN-G INTO ERYTHROCYTES**

Erythrocytes (0.8 ml of cell suspension) were incubated at 37°C for indicated time intervals in 1/15M phosphate buffer (pH 7.4) containing 6 mM glucose and 5 µg of PIN or PIN-G.

<table>
<thead>
<tr>
<th>Substrate added</th>
<th>Incubation time</th>
<th>Vitamin B₆ concentration</th>
<th>Concentration ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>PIN</td>
<td>min</td>
<td>µg/ml</td>
<td>µg/ml</td>
</tr>
<tr>
<td>PIN-G</td>
<td>10</td>
<td>1150</td>
<td>987</td>
</tr>
<tr>
<td></td>
<td>30</td>
<td>1840</td>
<td>927</td>
</tr>
<tr>
<td></td>
<td>60</td>
<td>2420</td>
<td>877</td>
</tr>
<tr>
<td>PIN</td>
<td>10</td>
<td>265</td>
<td>1060</td>
</tr>
<tr>
<td></td>
<td>30</td>
<td>365</td>
<td>1060</td>
</tr>
<tr>
<td></td>
<td>60</td>
<td>385</td>
<td>1050</td>
</tr>
</tbody>
</table>
erythrocytes incorporated was increased with incubation time. The concentration ratio of intracellular PIN to extracellular PIN was 2.8 after 60 min. However, the amount of transported PIN-G was about 20% of that of transported PIN.

Effect of concentrations of PIN and PIN-G on the transport into erythrocytes

The cells were incubated with 1 to 20 μg of PIN and PIN-G for 60 min. As shown in Fig. 4, the amount of intracellular vitamin B₆

![Graph showing effect of PIN and PIN-G concentrations on intracellular vitamin B₆ levels](image)

Fig. 4. Effect of Concentrations of PIN and PIN-G on the Transport into Erythrocytes.

Erythrocytes (2.0 ml of cell suspension) were incubated at 37°C for 60 min in 1/15M phosphate buffer (pH 7.4) containing 6 mM glucose and various concentrations of PIN or PIN-G

--- PIN, ---- PIN-G

-83-
Fig. 5. Effect of Incubation Temperature on the Transport of PIN and PIN-G into Erythrocytes.

Erythrocytes (1.2 ml of cell suspension) were incubated at various temperature in 1/15M phosphate buffer (pH 7.4) containing 6 mM glucose and 1 µg of PIN or PIN-G.

was increased proportionately to that of PIN or PIN-G added. However, the amount of intracellular vitamin B$_6$ transported from PIN-G was very small, as compared with that from PIN. The result suggests that a barrier may be present for the transport of PIN-G.

Effect of incubation temperature on the transport of PIN and PIN-G into erythrocytes

The cells were incubated with 1 µg of PIN or PIN-G at different temperature. The result is shown in Fig. 5. Neither PIN nor PIN-G
TABLE IV. EFFECT OF GLYCOLYTIC INHIBITORS ON THE TRANSPORT OF PIN AND PIN-G INTO ERYTHROCYTES

Erythrocytes (2.0 ml of cell suspension) were incubated at 37°C for 60 min in 1/15M phosphate buffer (pH 7.4) containing 6 mM glucose and 1 μg of PIN or PIN-G.

<table>
<thead>
<tr>
<th>Inhibitor</th>
<th>Concentration</th>
<th>Intracellular vitamin B₆</th>
</tr>
</thead>
<tbody>
<tr>
<td>PIN</td>
<td>None</td>
<td>611 μg/ml</td>
</tr>
<tr>
<td></td>
<td>NaF</td>
<td>5 x 10⁻³ M</td>
</tr>
<tr>
<td></td>
<td>CH₂ICOOH</td>
<td>&quot;</td>
</tr>
<tr>
<td>PIN-G</td>
<td>None</td>
<td>86</td>
</tr>
<tr>
<td></td>
<td>NaF</td>
<td>5 x 10⁻³ M</td>
</tr>
<tr>
<td></td>
<td>CH₂ICOOH</td>
<td>&quot;</td>
</tr>
</tbody>
</table>

was incorporated into the cells at 5°C. But, the incorporation of vitamin B₆ was accelerated at 28°C and 37°C.

*Effect of glycolytic inhibitors on the transport of PIN and PIN-G into erythrocytes.*

Sodium fluoride and monoiidoacetate are known as glycolytic inhibitors which depress the active transport. Effect of sodium fluoride or monoiidoacetate (5 x 10⁻³M) on the transport of PIN and PIN-G is shown in Table IV. The incorporation of PIN and PIN-G was fairly inhibited in the presence of the inhibitors. This result would suggest that the transport of PIN and PIN-G might be dependent upon the energy supplying system.

*Effect of ATP and glucose on the transport of PIN-G into erythro-*

-85-
As shown in Table V, the transport of PIN-G was evidently enhanced by the addition of glucose or ATP. This result leads to the suggestion that the transport of PIN-G may be related to the energy supplying system.

**Adsorption of PIN and PIN-G on ghost cells**

From the results described above, it was shown that PIN and PIN-G was found in erythrocytes after incubation. But it should be noted whether these vitamin B₆ compounds are really transported through erythrocyte membranes or merely combined or adsorbed on membranes (ghost cells). The author examined to clarify whether PIN and PIN-G were combined to ghost cells. As shown in Table VI, the vitamin B₆ compounds are not combined to ghost cells, but remain uncharged in erythrocytes after incubation.

**TABLE V. EFFECT OF ATP AND GLUCOSE ON THE TRANSPORT OF PIN-G INTO ERYTHROCYTES**

Erythrocytes (1.5 ml of cell suspension) was incubated at 37°C for 60 min in 1/15M phosphate buffer (pH 7.4) containing 6 mM glucose, 20 μmoles of ATP and 5 μg of PIN-G.

<table>
<thead>
<tr>
<th>Addition</th>
<th>Intracellular vitamin B₆</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>84 μg/ml</td>
</tr>
<tr>
<td>+ Glucose</td>
<td>99</td>
</tr>
<tr>
<td>+ ATP</td>
<td>132</td>
</tr>
<tr>
<td>+ Glucose, + ATP</td>
<td>144</td>
</tr>
</tbody>
</table>
TABLE VI. ADSORPTION OF PIN AND PIN-G ON GHOST CELLS

The ghost cells (prepared from 0.75 ml of erythrocytes) were incubated at 37°C for 60 min with PIN or PIN-G under the conditions described in the EXPERIMENTAL.

<table>
<thead>
<tr>
<th>Concentration</th>
<th>Incubation 0 min</th>
<th>Incubation 60 min</th>
</tr>
</thead>
<tbody>
<tr>
<td>PIN</td>
<td>40 μg</td>
<td>41.3 μg</td>
</tr>
<tr>
<td></td>
<td>80 &quot;</td>
<td>81.5 &quot;</td>
</tr>
<tr>
<td>PIN-G</td>
<td>50 &quot;</td>
<td>51.7 &quot;</td>
</tr>
<tr>
<td></td>
<td>100 &quot;</td>
<td>100.0 &quot;</td>
</tr>
</tbody>
</table>

the medium. Therefore, it seems to be concluded that a large part of PIN and PIN-G incubated with rabbit erythrocytes was actively transported into the cytoplasm.

*Retention of PIN and PIN-G in the erythrocytes*

After incubation at 37°C for 60 min with various concentrations of PIN and PIN-G, the cells were separated from the suspending medium by centrifugation and washed three times with a 0.9% saline solution. The result is shown in Table VII. A great part of PIN and PIN-G in the first washing is considered to be the contamination of the medium. The amount of PIN and PIN-G eluted in the second and third washings is smaller than the intracellular vitamin B₆, showing the retention of PIN and PIN-G in the erythrocytes.
TABLE VII. RETENTION OF PIN AND PIN-G IN THE ERYTHROCYTES

Erythrocytes (2.0 ml of cell suspension) were incubated at 37°C for 60 min in 1/15M phosphate buffer (pH 7.4) containing 6 mM glucose and the indicated amounts of PIN or PIN-G. After incubation, the cells were separated by centrifugation and washed three times with 5 ml of an ice-cold 0.9% saline solution.

<table>
<thead>
<tr>
<th>Added amount</th>
<th>in Washings</th>
<th>in Erythrocytes</th>
</tr>
</thead>
<tbody>
<tr>
<td>PIN</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 µg</td>
<td>118</td>
<td>99</td>
</tr>
<tr>
<td>5</td>
<td>354</td>
<td>199</td>
</tr>
<tr>
<td>10</td>
<td>1320</td>
<td>682</td>
</tr>
<tr>
<td>PIN-G</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>158</td>
<td>31</td>
</tr>
<tr>
<td>5</td>
<td>410</td>
<td>111</td>
</tr>
<tr>
<td>10</td>
<td>451</td>
<td>260</td>
</tr>
</tbody>
</table>

DISCUSSION

It has been reported that the dipolar form of PIN in a neutral solution is very labile to heat and light. On the other hand, PIN-borate complex is reported to be thermostable. It seems that the stability of PIN-borate complex depends on the monopolar form in which 3' and 4' positions of PIN are masked by borate. It was also reported that 4' or 5' position of PIN was first attacked by photooxidation and that by heating at 120°C for 30 min in a neutral solution, PIN formed dimer or polymer, in which 4' hydroxymethyl group of PIN was combined to nitrogen atom of 1' position of another PIN. In PIN-G, the dipolar form is maintained, but 4' or 5' position of PIN is
masked by glucosyl residue. The higher stability of PIN-G seems to be based on that 4' or 5' position of PIN is masked by glucosyl residue.

PIN-G served as vitamin B₆ (PIN) vitamer, which showed 20% of microbiological activity of equivalent mole of PIN for *S. carlsbergensis* 4228 ATCC 9080. However, the activity of PIN-G was gradually increased accompanied with the prolonged incubation.

It was suggested that PIN-G was hydrolyzed to PIN and glucose by α-glucosidase of *S. carlsbergensis* 4228 ATCC 9080 and then functioned as PIN.

The cellular transport of vitamin B₆ compounds has been investigated by Yamada *et al.*131-133) and Suzue *et al.*134) with human erythrocytes. Yamada *et al.* reported that free forms of vitamin B₆ were actively transported into the cells, while their phosphate esters were hardly transported. They also reported that PAL was most rapidly and abundantly transported into the erythrocytes. Suzue *et al.* found out that PAL-P was also incorporated into red blood cells without being hydrolyzed.

As described in this paper, PIN and PIN-G were transported into rabbit erythrocytes. The concentration of intracellular vitamin B₆ incorporated was 2.8 times as high as that found in the medium of PIN, and 0.4 times in the medium of PIN-G. The cellular transport of PIN and PIN-G was increased with the incubation time and the concentration of them added. At low temperature (5°C), the transfer of PIN and PIN-G into the cells was remarkably depressed. Sodium fluoride and moniodoacetate considerably inhibited the transport of PIN and PIN-G.
By the addition of ATP and glucose, the transport of PIN-G was accelerated. All these results suggest that both of PIN and PIN-G are actively transported through erythrocyte membranes. No PIN and PIN-G was combined to ghost cells, which would show that these vitamins were truly incorporated into erythrocytes. A small amount of PIN and PIN-G was leaked out from the cells into saline washings, but a great part of them were retained in the cells. This result also supports the above suggestion. However, the amount of PIN-G transported was about 10-20% of that of equal mole of PIN. This is probably due to the existence of a permeability barrier for PIN-G. It remains to be investigated whether PIN-G is incorporated directly into the cells or it is incorporated in the form of PIN after PIN-G is degraded by a membrane α-glucosidase.

SUMMARY

PIN-G was found to be as stable as PIN-borate complex against irradiation of ultraviolet light and heating, while PIN was less stable.

Intact PIN-G showed about 20% of microbiological activity of equivalent mole of PIN for *Saccharomyces carlsbergensis* 4228 ATCC 9080. The activity was gradually increased accompanied with the prolonged incubation. It seems that PIN-G is hydrolyzed to PIN and glucose by α-glucosidase of the organism and can be bioassayed as PIN. Hydrolysis of PIN-G by heating in 0.055N H₂SO₄ yielded PIN quantitatively.
PIN and PIN-G are transported into rabbit erythrocytes in vitro. The amounts of vitamin B₆ compounds transported into the erythrocytes were bioassayed with *Saccharomyces carlsbergensis* 4228 ATCC 9080 after hydrolyzing them to free forms of vitamin B₆ by heating for 210 min in 1/12N H₂SO₄. The transport of PIN and PIN-G was affected by incubation time, incubation temperature and concentration of PIN and PIN-G added. The transport of PIN-G increased with the coexistence of ATP or glucose. The transport of PIN and PIN-G was decreased by the addition of sodium fluoride or monoiodoacetate. The adsorption of PIN and PIN-G on ghost cells was not found. Small amounts of transported PIN and PIN-G were leaked out by washings, indicating that large parts of them were retained in the erythrocytes.
Chapter II. Microbial Formation of a New Derivative of Pantothenic Acid: Pantothenic Acid-α-Glucoside

INTRODUCTION

PaA was first recognized from natural sources with the aid of a microbiological assay employing the Gebrüder Mayer strain of yeast Saccharomyces cerevisiae, growing in a synthetic medium. The compound also showed the biological activity for chicks, designated as a "filtrate factor" or a "chick antidermatitis factor". Jukes and Wooley et al. in 1939 simultaneously announced that PaA, isolated in pure form from liver by Williams et al., and the filtrate factor were apparently identical. Lipmann and coworkers demonstrated that coenzyme A contained a bound PaA which could be released effectively by the combined actions of a liver enzyme and alkaline phosphatases.

PaA is widely distributed in animals, plants and microorganisms as a component of coenzyme A. However, other complexes derived from coenzyme A, as well as phosphorylated forms of both PaA and pantethine, have also been demonstrated to occur widely. Many workers have investigated the biosynthetic and degradative metabolisms of PaA. A large number of derivatives of PaA except metabolites described above, have been found in nature and have also been chemically synthesized. Recently a β-glucoside of PaA was isolated from tomato juice as a growth factor for malo-lactic fermentation bacteria in wine fermentation. However, an α-glucoside
of PaA has not been synthesized chemically and biologically.

The present chapter describes the microbial formation of the glucoside of D-PaA. A considerable amount of α-glucoside of D-PaA was formed by intact cells of *Sporobolomyces coralliformis* IFO 1032, when incubated with maltose and D-PaA. The isolation and identification of the compound is presented.

EXPERIMENTAL

**Materials.** DEAE-Cellulose was kindly given by the Green Cross Co., Osaka. Cellulase (Type II, from *Aspergillus niger*), β-glucosidase (from almonds), maltase (Grade II, from *A. niger*) were purchased from Sigma Chem. Co., and α-glucosidase (from yeast) was from Boehringer Mannheim Japan K.K. Takadiastase B was purchased from Sankyo Co., Ltd. Other chemicals used in this work were obtained from commercial sources.

**Microorganisms and cultivation.** Yeasts preserved in the Laboratory of Physiology of Fermentation and Applied Microbiology, Department of Agricultural Chemistry, Kyoto University were used for a screening. All strains used were preserved on a nutrient agar slant by a monthly transfer at 5°C.

Yeasts were grown on a medium composed of 5% glucose, 0.5% peptone, 0.5% yeast extract, 0.5% K₂HPO₄, 0.2% KH₂PO₄, and 0.02% MgSO₄·7H₂O, adjusted to pH 6.0. The cultivation was carried out at 28°C for 48 hr on a reciprocal shaker with 50 ml of the medium in a 300 ml shaking flask. After cultivation, the cells were harvested by centrifugation at 5000 rpm, and washed once with 0.9% saline solution. The washed
cells were suspended in 2-5 ml of 0.01M potassium phosphate buffer (pH 7.0) containing $5 \times 10^{-3}$ M 2-mercaptoethanol and subjected to the reaction.

*Reaction conditions.* The standard reaction mixture contained 50 μmoles of D-PaA, 100 μmoles of maltose or cellobiose, 100 μmoles of potassium phosphate buffer (pH 7.0), 3 μmoles of 2-mercaptoethanol, a final concentration of 0.1% of Span 80, and 0.5 ml of intact cell suspension in a total volume of 1.5 ml. The incubation was performed at 28°C for 24 hr aerobically. Glucose was used as a blank in place of maltose or cellobiose. The reaction was stopped by immersing a tube in a boiling water bath for 5 min. After cooling the reaction mixture in an ice bath, the cells were removed by centrifugation and the supernatant fluid was assayed for a derivative of D-PaA.

*Analyses.* PaA was bioassayed by the paper disc method using *Saccharomyces carlsbergensis* 4228 ATCC 9080. PaA glucoside was identified as follows. A suitable amount of reaction mixture (20-50 μl) was spotted on Toyo filter paper No. 53 (2 cm width) and developed with a solvent system of n-butanol:acetic acid:water (4:1:1, v/v). After development, the dried paper strip was cut off in 0.5 cm widths from the 2 cm width and subjected to bioautography by *S. carlsbergensis* 4228 ATCC 9080 and *Lactobacillus plantarum* ATCC 8014. An appropriate amount of Takadiastase B or mold maltase solution (4 mg/ml of 0.01M acetate buffer, pH 5.0) was daubed on a paper strip with a pipette after it was placed on a culture plate in order to get a clearer active zone of PaA-α-G with a maltose-PaA.
medium. An appropriate amount of cellulase (2 mg/ml of 0.01M acetate buffer, pH 5.0) was similarly used for the detection of PaA-β-G with a celllobiose-PaA medium.

Paper chromatography of PaA-G and its acid-hydrolyzate was carried out on Toyo filter paper No. 53 with the solvent systems as follows; n-butanol:acetic acid:water (4:1:1, v/v), n-propanol:28% ammonia:water (6:3:1, v/v), n-butanol:benzene:pyridine:water (5:1:3:3, v/v, upper layer) and isopropanol:pyridine:water:acetic acid (8:8:4:1, v/v). The PaA moiety was confirmed by bioautography and the glucose moiety was determined by the color development with the silver nitrate reagent.\(^{109}\)

Glucose was determined by the glucostat method\(^{111}\) and reducing sugar was estimated by the Somogyi-Nelson method.\(^{112}\)

The reaction by several glucosidases was carried out on a reaction mixture containing 2.6 μmoles of PaA-G, 10 μmoles of 2-mercaptoethanol, 50 μmoles of buffer and an appropriate amount of enzyme in a total volume of 1.0 ml. The reaction was carried out at 37°C for 3 hr. Phosphate buffer (pH 7.0) was used in the incubation with yeast α-glucosidase and acetate buffer (pH 5.0) in that with mold maltase, β-glucosidase and cellulase. The reaction mixture was submitted to bioautographic and paper-chromatographic analyses for PaA and glucose as described above.

*Measurement of IR and NMR spectra.* IR spectrum of PaA-G was measured in a micro KBr tablet with a Hitachi infrared spectrometer type EPI-G3. NMR spectrum was measured in D₂O on a Varian HA-100D spectrometer at 100 MHz with DSS-D₆ as an internal standard.
RESULTS

PaA-G Formation by Yeasts

PaA-G-forming activity was checked with yeasts. Maltose was used for PaA-α-G formation and cellobiose was used for PaA-β-G formation. Glucose was also used in place of maltose or cellobiose as a blank. A considerable amount of α-glucoside of PaA was found in the reaction mixture of some strains of yeasts belonging to Saccharomyces, and Sporobolomyces coralliformis (Table I). S. coralliformis IFO 1032 seemed to have the highest activity for PaA-α-G formation. A β-glucoside was scarcely detected in all strains. No derivative was found with a glucose medium, as shown in Fig. 1. Thus, S. coralliformis IFO 1032 was selected and used as the PaA-α-G-forming strain in the following experiment. The formation of PaA-α-G at 48 hr of incubation was larger than at 24 hr of incubation.

![Bioautogram of PaA-G.](image)

Solvent system: n-butanol:acetic acid:water (4:1:1, v/v)
I. maltose medium, II. cellobiose medium, III. glucose medium
TABLE I. DISTRIBUTION OF PaA-G-FORMING ACTIVITY IN YEASTS

Intact cells of yeasts were incubated in the reaction mixture described in the EXPERIMENTAL. The incubation was performed at 28°C for 24 hr aerobically.

<table>
<thead>
<tr>
<th>Strains</th>
<th>PaA-α-G formed</th>
<th>PaA-β-G formed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saccharomyces cerevisiae AKU 4100</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&quot; delbrueckii IFO 0285</td>
<td>++</td>
<td>-</td>
</tr>
<tr>
<td>&quot; carlsbergensis IFO 0641</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot; rouxii IAM 4369</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot; sake Kyokai No. 7 AKU 4111</td>
<td>++</td>
<td>-</td>
</tr>
<tr>
<td>&quot; bayanus IFO 0206</td>
<td>-</td>
<td>±</td>
</tr>
<tr>
<td>&quot; sake Fukumusume AKU 4017</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>&quot; rouxii IFO 0686</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot; cerevisiae AKU 4104</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot; sake Unryu AKU 4019</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot; &quot; Shokkomasamune AKU 4015</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot; &quot; Hozan AKU 4013</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Bass Bier Hefe (Burton on Trent No. 1)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>AKU 4002</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kagawakenshi (Shoyu) AKU 4033</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Fukumoto 3 AKU 4034</td>
<td>++</td>
<td>-</td>
</tr>
<tr>
<td>Beer yeast (Nippon) AKU 4038</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Brennerei Hefe Rasse 12 AKU 4010</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Sporobolomyces salmonicolor IFO 0374</td>
<td>-</td>
<td>±</td>
</tr>
<tr>
<td>&quot; &quot; IFO 0375</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot; coprophilus IFO 1422</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot; coralliformis IFO 1032</td>
<td>++</td>
<td>±</td>
</tr>
<tr>
<td>&quot; pararoseus IFO 1103</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot; salmonicolor IFO 1038</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot; odorus IFO 1035</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

++; good, +; fair, ±; slight, -; nil.
Isolation of PaA-α-G

For the purpose of good formation of PaA-α-G-synthesizing enzyme, *S. aoralliformis* was grown on a maltose medium with 500 ml of the medium in a 2 liter shaking flask. The cultivation was carried out at 28°C for 48 hr on a reciprocal shaker. The cells were collected by continuous centrifugation at 5000 rpm, washed once with 0.9% saline solution and suspended in the same solution. Twenty-one grams (calculated as dried cells) of intact cells of the organism were incubated with 40 g of maltose, 4 g of D-PaA, 8 mmoles of potassium phosphate buffer (pH 7.0), and 0.4 g of cetyltrimethyl ammonium bromide (surfactant) in a total volume of 400 ml. The reaction was carried out aerobically at 28°C for 40 hr on a 2 liter shaking flask. The reaction was stopped by immersing the incubation flask (adjusted to pH 7.0) in boiling water for 5 min, and the cells were removed by centrifugation at 10,000 rpm for 20 min. Isolation of a product was made from the supernatant solution as a starting material. The supernatant solution was adjusted to pH 4.0 and left at 4°C overnight. Any material precipitating was removed by filtration. Active charcoal (60 g) was added to the filtrate, followed by stirring for 1 hr. Then, the charcoal was collected by filtration, washed with 0.01N HCl and deionized water and eluted with five volumes of acetone and an equal volume of 50% acetone. The eluate was concentrated by evaporation *in vacuo*. The concentrated solution was submitted to paper chromatography with Toyo filter paper No. 526 in a solvent system of *n*-butanol:acetic acid:water (4:1:1, v/v), followed by bioautography with *Saccharomyces carlsber*
Two different bioactive zones were detected, one of which with a higher \( R_f \) value was identical to D-PaA. The other bioactive zone with a \( R_f \) of 0.5, was cut off and extracted with 50% ethanol. The extract was concentrated in vacuo and desalted with Amberlite CG-50 (\( H^+ \) type). The concentrated effluent was applied to a Sephadex G-10 column (1 x 105 cm), and eluted with water. The active fractions were adsorbed on a DEAE-cellulose column (\( OH^- \) type), and eluted with 0.02M LiCl. The eluted fraction was desalted again with Amberlite CG-50 (\( H^+ \) type), and adjusted to pH 5.0 with HCl. Saturated Ba(OH)\(_2\) solution was added to the fraction until the pH became to 7.0-7.2. Excess Ba(OH)\(_2\) was removed repeatedly by concentration and filtration. The completely dried Ba-salt of the compound was dissolved in methanol, followed by adding ether. A resultant fluffy precipitate was collected and dissolved in water, and subjected to paper chromatography in the solvent system as described above. The active zone was extracted with 50% ethanol, and the extract was evaporated, lyophilized and dried in vacuo on \( P_2O_5 \) and KOH. The lyophilized powder (yield 21.9 mg) was used for the characterization.

**Identification of the Compound**

The \( R_f \) values of the isolated compound on bioautography in several solvent systems were examined, comparing with known metabolites of PaA. The result is shown in Table II. The \( R_f \) values of the isolated compound differed from PaA, \( \beta \)-alanine and phospho-PaA. Pantenol, pantethine and pantoyl lactone had no biological response for *Saccharomyces*
carlsbergensis 4228 ATCC 9080. The compound was also responsible for Lactobacillus plantarum ATCC 8014. The organism utilizes PaA, phospho-
PaA, and panthethine, but not pantenol, β-alanine and pantoyl lactone. The activity for both strains increased with the prolonged incubation. It seems that the compound is hydrolyzed to D-PaA and glucose by α-
glucosidases of S. carlsbergensis and L. plantarum and can be bioassayed as PaA.

The isolated compound had no reducing activity for Somogyi-Nelson reagent112). However, on hydrolysis in 0.055N H₂SO₄ at 120°C and 15 lb for 3 hr, or on treatment with yeast α-glucosidase at 30°C and pH

<table>
<thead>
<tr>
<th>Compounds</th>
<th>I</th>
<th>II</th>
<th>Solvent III</th>
<th>IV</th>
<th>V</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isolated compound</td>
<td>.49</td>
<td>.36</td>
<td>.63</td>
<td>.52</td>
<td>.48</td>
</tr>
<tr>
<td>D-PaA</td>
<td>.76</td>
<td>.50</td>
<td>.76</td>
<td>.85</td>
<td>.69</td>
</tr>
<tr>
<td>β-Alanine</td>
<td>.29</td>
<td>.13</td>
<td>.54</td>
<td>.39</td>
<td>.62</td>
</tr>
<tr>
<td>4'-Phospho-PaA160)</td>
<td>.45</td>
<td>.19</td>
<td>.28</td>
<td>.40</td>
<td>.38</td>
</tr>
<tr>
<td>Panthenol</td>
<td>no response</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pantethine</td>
<td>no response</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pantoyl lactone</td>
<td>no response</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Solvent system:
I. n-butanol:acetic acid:water (4:1:1, v/v)
II. n-butanol:pyridine:water (6:4:3, v/v)
III. n-propanol:28% ammonia:water (6:3:1, v/v)
IV. n-butanol:ethanol:water (5:1:4, v/v, upper layer)
V. isobutyric acid:0.5N ammonia:0.1M EDTA (100:80:1.6, v/v)
TABLE XII. IDENTIFICATION OF PaA AND GLUCOSE MOIETIES OF THE ISOLATED COMPOUND

The PaA moiety was confirmed by bioautography using *S. carlsbergensis* 4228 ATCC 9080 and the glucose moiety was identified by the color development with the silver nitrate reagent.109)

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Solvent I</th>
<th>Solvent II</th>
<th>Solvent III</th>
<th>Solvent IV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isolated compound</td>
<td>.46</td>
<td>.58</td>
<td>.24</td>
<td>.73</td>
</tr>
<tr>
<td>Acid hydrolyzate*</td>
<td>.80</td>
<td>.76</td>
<td>.33</td>
<td>.57</td>
</tr>
<tr>
<td>Enzymatic hydrolyzate**</td>
<td>.80</td>
<td>.73</td>
<td>.31</td>
<td>.55</td>
</tr>
<tr>
<td>PaA</td>
<td>.79</td>
<td>.75</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>β-Alanine</td>
<td>.32</td>
<td>.55</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Glucose</td>
<td>-</td>
<td>-</td>
<td>.33</td>
<td>.57</td>
</tr>
<tr>
<td>Maltose</td>
<td>-</td>
<td>-</td>
<td>.22</td>
<td>.49</td>
</tr>
</tbody>
</table>

* hydrolyzed in 0.055N H₂SO₄ at 120°C and 15 lb for 3 hr.
** hydrolyzed by yeast α-glucosidase.

Solvent system:

I. *n*-butanol:acetic acid:water (4:1:1, v/v)
II. *n*-propanol:28% ammonia:water (6:3:1, v/v)
III. *n*-butanol:benzene:pyridine:water (5:1:3:3, v/v, upper layer)
IV. isopropanol:pyridine:water:acetic acid (8:8:4:1, v/v)

7.0, the isolated compound gave two components corresponding to glucose and PaA. The identification of the components by paper chromatography and bioautography is shown in Table III. The molar ratio of glucose and PaA was determined to be 1:1 by the glucostat method and bioassay (Table IV). Enzymatic hydrolysis of the isolated compound by various
TABLE IV. MOLAR RATIO OF PaA AND GLUCOSE

<table>
<thead>
<tr>
<th>Method</th>
<th>PaA found (μmole)</th>
<th>Glucose found (μmole)</th>
<th>Molar ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microbioassay by paper disc method* (S. carlsbergensis)</td>
<td>195</td>
<td>-</td>
<td>1.00</td>
</tr>
<tr>
<td>Glucostat method*</td>
<td>-</td>
<td>211</td>
<td>1.08</td>
</tr>
</tbody>
</table>

*) The isolated compound was hydrolyzed at 37°C overnight by yeast α-glucosidase.

Reducing activity of the compound was not found by Somogyi-Nelson method.

Glucosidases was investigated. Figure 2 shows that the compound is hydrolyzed to glucose and PaA by yeast α-glucosidase and mold maltase, but not by cellulase and β-glucosidase. The data suggest that glucose and PaA are bound through an α-glucosidic linkage in the isolated compound, that is, the compound is α-glucopyranosyl-D-PaA. The IR spectrum of the compound showed that the compound contained PaA (chelating with Ba$^{2+}$) and sugar moiety (Fig. 3). The NMR spectrum of the compound is shown in Fig. 4. The signals shows the presence of two methyl groups (C-3', δ 0.98 and 0.99 ppm, 6H, single), a methylene group (C-2, δ 2.42 ppm, 2H, triplet, $j$=13 Hz), a methylene group (C-3, δ 3.45 ppm, 2H, triplet, $j$=11.5 Hz), a methylene group of C-4' and ring protons of the sugar moiety (δ 3.55-3.90 ppm, multiple), a methyne group (C-2', δ 4.12 ppm, 1H, single), and one anomeric proton of the α-glucosidic linkage (δ 4.89 ppm, 1H, doublet, $j$=3.5 Hz). The comparison of this spectrum with that of authentic PaA suggests that a PaA
Fig. 2. Enzymatic Hydrolysis of the Isolated Compound by Glucosidases.

The PaA moiety was confirmed by bioautography and the glucose moiety was determined by the color development with the silver nitrate reagent.

Solvent system:
I. n-butanol:acetic acid:water (4:1:1, v/v)
II. n-butanol:benzene:pyridine:water (5:1:3:3, v/v)

a. mold maltase
b. yeast α-glucosidase
c. β-glucosidase (emulsin)
d. cellulase
e. isolated compound
moiety is contained in this compound. The ratio of the numbers of protons attributed to two methyl groups in PaA moiety and glucose residue is approximately 1:1. The proposed structures of the isolated compound are shown in Fig. 5. Comparing the methyne signal (δ 4.12 ppm) of the proposed α-D-glucopyranosyl-D-PaA with that of 2'-O-(β-D-glucopyranosyl)-D-PaA and that of 4'-O-(β-D-glucopyranosyl)-D-PaA which were reported by Amachi et al., 86-88) it is suggested that the glucosidic linkage of the compound may be present at C-4' position. From the results described above, it is concluded that the isolated compound is possibly 4'-O-(α-D-glucopyranosyl)-D-PaA.
Fig. 4. NMR Spectrum of the Isolated Compound.

The spectrum was recorded on a Varian HA-100D spectrometer at 100 MHz in D$_2$O with DSS-D$_6$ as an internal standard.

$4'\text{-O-(\alpha-D-glucopyranosyl)-D-pantothenic acid}$

$2'\text{-O-(\alpha-D-glucopyranosyl)-D-pantothenic acid}$

Fig. 5. Proposed Structures for \alpha-Glucosyl-D-pantothenic Acid.
DISCUSSION

PaA-β-G has recently been isolated from tomato juice as a growth factor of malo-lactic fermentation bacteria in wine fermentation. In this chapter, the author has first described the formation of PaA-α-G, a new sugar derivative of PaA, by yeast Sporobolomyces coralliformis IFO 1032.

Many sugar derivatives of vitamins have been isolated from nature, and synthesized by enzymatic and chemical methods. Since riboflavinyl-α-glucoside has been found in rat liver by Whitby in 1950, the formation of various riboflavinylglycosides containing glucoside, galactoside, fructoside and oligosaccharides is reported in microorganisms, plants, and animals by many investigators. A sugar derivative of ascorbic acid has been prepared enzymatically by Miyake et al. PIN-G, the first sugar derivative of PIN to be recognized, has been isolated by Ogata et al. from bacterial cultures during investigations on the metabolism of vitamin B₆ by microorganisms. Microbial distribution of PIN-G-forming activity, purification and properties of the PIN-G-synthesizing enzyme of Micrococcus sp. No. 431 and some physico-chemical and biological properties of PIN-G were investigated in detail by the author in the preceding chapter.

In a preliminary experiment on microbial formation of PaA-G, the author examined the distribution of PaA-G-forming activity in microorganisms under cultural conditions. The cultivations were performed on a PaA-maltose medium and a PaA-cellobiose medium in expect of the formation of PaA-α-G and PaA-β-G in the respective media. Very few
organisms produced an appreciable amount of PaA-β-G in the PaA-cellobiose medium, but some strains of yeast accumulated a considerable amount of PaA-α-G in the PaA-maltose medium. *Sporobolomyces corallicformis* IFO 1032 had a relatively high activity for the formation of PaA-α-G among those tested. It was also found that a large amount of PaA-α-G was formed when PaA and maltose were incubated with intact cells of *S. corallicformis* grown on a maltose medium. Therefore, the large scale reaction with the intact cells was carried out to isolate and identify PaA-α-G for the simplicity of reaction system.

The author isolated a new compound from a reaction mixture of the organism by the treatment with active charcoal, Amberlite CG-50, Sephadex G-10 and DEAE-cellulose column chromatography and paper chromatography. The compound was characterized as α-glucopyranosyl-D-PaA. It was suggested that the glucosidic linkage of the compound might be present at C'-4 from NMR spectrum and the facilitiy of reactivity of C'-2 or C'-4 alcoholic groups. The compound showed the slight microbiological activity for *Saccharomyces carlsbergensis* and *Lactobacillus plantarum*. The activity increased with the incubation period. It seemed that the compound might be gradually hydrolyzed by α-glycosidases of these strains and could be utilized as PaA. The transglucosidation seems to be catalyzed by α-glycosidase, reacting on maltose. However, *Micrococcus* and *Sarcina*, having the high activity of PIN-G formation did not form the compound. These α-glycosidases differ from each other on the substrate specificity for glucosyl acceptor.
SUMMARY

The formation of PaA-G was checked with intact cells of yeasts. PaA-β-G-forming activity was scarcely found, but PaA-α-G-forming activity was found with several strains. Among them, Sporobolomyces aoralliformis IFO 1032 showed the highest activity for PaA-α-G formation. The compound was isolated from a reaction mixture of the organism incubated on a medium containing maltose and D-PaA, by the treatment with active charcoal, Amberlite CG-50, Sephadex G-10 and DEAE-cellulose column chromatography and paper chromatography. The isolated compound was characterized as 4'(2')-O-(α-D-glucopyranosyl)-D-PaA by various analytical methods including bioautography, paper chromatography, and IR and NMR spectroscopy.

The intact compound showed the slight microbiological activity for Saccharomyces carlsbergensis 4228 ATCC 9080 and Lactobacillus plantarum ATCC 8014, which are used as strains for bioassay of PaA. The activity increased gradually with the prolonged incubation.
CONCLUSION

In this thesis, the author has investigated the formation of PIN-G and PaA-G by microorganisms.

In the first chapter, the distribution of PIN-G-forming activity was investigated in various microorganisms. Various factors influencing on the formation, the enzymatic metabolism, the stability and the biological activity of PIN-G were also examined. The distribution of PIN-G-forming activity was searched in culture collections of bacteria, yeasts, molds and actinomycetes together with isolated strains. A relatively high activity of PIN-G formation was found in bacteria belonging to *Micrococcus* and *Sarcina*. The activity was also widely distributed in yeasts, molds and actinomycetes. A bacterium strain No. 431, which was isolated from soil and identified as *Micrococcus* sp. No. 431, had the highest activity among the tested organisms and selected as the most suitable strain for PIN-G formation. With this strain, several conditions on the formation of PIN-G were investigated by the growing culture. Sucrose and peptone were the most excellent substrates as carbon and nitrogen sources. A few nonionic surfactants showed the stimulative effect on the formation of PIN-G, in which Span 80 was the most effective. Metal ions had no effect. The accumulation of PIN-G reached to a maximum after 60 to 80 hr of cultivation and decreased after then. The formation of PIN-G increased proportionately with the concentrations of sucrose and PIN. Optimal pH of the culture was 7.0. Under the optimal conditions, the amount of PIN-G reached to a maximum of 1800 to 1900 μg/ml as PIN moiety (yield over 90%) at 2
mg/ml of PIN added.

PIN-G was isolated from the culture filtrate of *Micrococcus* sp. No. 431 grown under the optimal conditions, and identified by various analytical methods including IR and NMR spectroscopy. PIN-4'-G and PIN-5'-G were separately confirmed by paper chromatography. PIN-diglucoside was also found in the filtrate, which was isolated and identified by the same methods as PIN-G.

The enzyme activity to synthesize PIN-G from PIN and sucrose was demonstrated in intact cells and cell extracts of *Sarcina* and *Micrococcus*. *Micrococcus* sp. No. 431, an isolated bacterium, was found to have the high activity of this enzyme in its cell extract and used as the enzyme source. The enzyme activity of the microorganism reached to a maximum after 20 hr of cultivation. The enzyme which catalyzes the synthesis of PIN-G via transglucosidation from sucrose to PIN was purified from the organism by means of ammonium sulfate fractionation, and DEAE-Sephadex, hydroxylapatite and Sephadex G-100 column chromatography. The enzyme was purified about 354-fold and confirmed to be homogeneous on polyacrylamide-gel electrophoresis and ultracentrifugation.

Some properties of PIN-G-synthesizing enzyme of the organism were studied using the partially and highly purified enzyme preparations. The enzyme was stable at pH 7.0 and between 0°C and 30°C. The maximal activity was obtained at pH 8.0 and 37°C. Besides sucrose, phenyl-α-D-glucoside and maltose served as glucosyl donor. Of vitamin B₅ compounds tested, only PIN served as glucosyl acceptor. The enzyme acti-
vity was inhibited by PCMB and heavy metal ions, and the inhibition was prevented by 2-mercaptoethanol, indicating that the enzyme was a sulfhydryl enzyme like many α-glucosidases. The activity was not affected by chelating agents and not activated by metal ions.

Intact PIN-G showed about 20% of microbiological activity of equivalent mole of PIN for *Saccharomyces carlsbergensis* 4228 ATCC 9080. The biological activity increased gradually accompanied with the prolonged incubation. It seems that PIN-G is hydrolyzed to PIN and glucose by α-glucosidase of the yeast and can be bioassayed as PIN. Hydrolysis of PIN-G by heating in 0.055N H₂SO₄ at 120°C and 15 lb for 3 hr yielded PIN quantitatively.

PIN-G was transported into rabbit erythrocytes *in vitro*. The cellular transport was affected by the incubation time, the incubation temperature and the concentration of PIN-G added. The transport was increased with the coexistense of ATP and glucose and was decreased by the addition of sodium fluoride or monooiodoacetate. These results suggest that PIN-G is actively transported into cells through erythrocyte membranes. Adsorption of PIN-G on ghost cells was not found. This fact shows that PIN-G is truly incorporated into erythrocytes. A small amount of PIN-G was leaked out from the cells in the saline washings, but a great part of it was retained in the cells. This result also supports the above suggestion. However, the amount of PIN-G incorporated was about 10 to 20% of that of equal mole of PIN, showing the existence of a permeability barrier for PIN-G. It remains to be investigated whether PIN-G is incorporated into the cells with
the whole form of PIN-G or with the form of PIN after PIN-G is hydrolyzed by a membrane α-glucosidase of erythrocyte cells.

PIN-G was found to be as stable as PIN-borate complex against irradiation of ultraviolet light and heating, while PIN was less stable.

In the second chapter, the formation of the glucoside of D-PaA was investigated.

The formation of the glucoside of D-PaA was checked with intact cells of yeasts incubated on a medium containing maltose or celllobiose and D-PaA. A β-glucoside of D-PaA was scarcely formed, but an α-glucoside of D-PaA was found with several strains. *Sporobolomyces coralliformis* IFO 1032 showed the highest activity. Then, this new compound was isolated from a reaction mixture of the strain incubated in a medium containing maltose and D-PaA by the treatment with active charcoal, Amberlite CG-50, Sephadex G-10 and DEAE-cellulose column chromatography and paper chromatography. The isolated compound was characterized as 4'(2')-O-(α-D-glucopyranosyl)-D-PaA by various analytical methods including bioautography, paper chromatography, and IR and NMR spectroscopy.

The intact compound showed the slight microbiological activities for *Saccharomyces carlsbergensis* 4228 ATCC 9080 and *Lactobacillus plantarum* ATCC 8014. The activity increased gradually with the prolonged incubation. It seems that the compound is hydrolyzed to D-PaA and glucose by α-glucosidases of *S. carlsbergensis* and *L. plantarum* and can be bioassayed as D-PaA.
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REFERENCES


20) S. A. Harris, D. Heyl and K. Folkers, ibid., 154, 315 (1944).
35) E. E. Snell, "Proceedings of the Fifth Internatl. Congress of


48) S. F. Contractor and B. Shane, Biochem. Biophys. Res. Commun., 39,


72) K. Shirai, Vitamins (Japan), 45, 344 (1972).
75) idem, ibid., 5, 8 (1959).
76) idem, ibid., 5, 298 (1959).
77) idem, ibid., 4, 126 (1958).
81) idem, Vitamins (Japan), 41, 338 (1970).
82) H. Katagiri and S. Tachibana, ibid., 14, 178 (1957).
91) *idem, ibid.*, 53, 410 (1965).
97) *idem, ibid.*, 19, 186 (1971).
99) K. Ogata, Y. Uchida, N. Kurihara, Y. Tani and T. Tochikura, *ibid.*, -119-


132) idem, ibid., 37, 472 (1968).
133) idem, J. Vitaminol., 14, 282 (1968); idem, ibid., 16, 237 (1970).
140) R. J. Williams, C. M. Lyman, G. H. Goodyear, J. H. Truesdail and D. Holaday, ibid., 55, 2912 (1933).
141) T. H. Jukes, ibid., 61, 975 (1939).


