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Protonated Amino Acid Precursor Studies
on Rhodotorulic Acid Biosynthesis
in Deuterium Oxide Media*

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Running title: Biosynthesis of Rhodotorulic Acid

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ABSTRACT: Rhodotorula pilimanae CBS 4479 growing in 99.8% D₂O with protonated sucrose as carbon source was given various additional protonated substrates, the incorporation of which into rhodotorulic acid was examined. It was found that L-ornithine and δ -N-acetyl-L- δ -N-hydroxyornithine are intermediates on the rhodotorulic acid pathway. Indirect evidence was found that O₂ rather than H₂O is the source of the hydroxylamino oxygen. The assembly of the amino acids into the cyclic peptide apparently is similar to the scheme which recently has been established for gramicidin and tyrocidine. Arginine metabolism in R. pilimanae and Saccharomyces cerevisiae appear to be regulated by similar mechanisms.

Introduction

If R. pilimanae and related yeasts are grown in a low-iron medium, large amounts of a diketopiperazine dihydroxamic acid called rhodotorulic acid (RA)¹ are produced (Atkin et al., 1970). Similar but more complex compounds from other organisms have been shown by Emery (1971) to act as iron transporting agents. Two types of these compounds (hydroxamates and phenolates) collectively called siderochromes, have been described in a recent review (Neilands, 1972). Initial studies on the biosynthesis of RA involved the addition of ¹⁴C-labeled amino acids to cultures of R. pilimanae followed by measurement of their incorporation into RA. Labeled glutamate, glutamine, proline, arginine, or ornithine were rapidly converted to RA, and afforded little information on the initial precursor of the molecule. To overcome this problem the metabolic source of the individual hydrogens, rather than the carbon skeleton, was studied. For this purpose R. pilimanae was adapted to grow in 99.8% D₂O on protonated sucrose, and at approximately 1/2 to 2/3 maximum growth (initial phase of RA synthesis) protonated substrates were added to the medium. After about 15 days the RA was purified

¹Abbreviations used: RA, rhodotorulic acid; PMR, proton magnetic resonance; AHO, δ -N-acetyl-L- δ -N-hydroxyornithine.

and the relative protonation determined at each position by proton magnetic resonance (PMR).

Using this technique a variety of intermediates of arginine metabolism were examined as precursors of RA. A number of conclusions were drawn regarding regulation of the metabolism of this amino acid; also, the similarities between arginine metabolism in R. pilimanae and S. cerevisiae are discussed. The initial precursor was shown to be L-ornithine and δ -N-acetyl-L- δ -N-hydroxyornithine (AHO) was demonstrated to be an intermediate on the RA biosynthetic pathway. The similarities between the assembly of the monomer amino acids into RA and the accepted scheme of gramicidin and tyrocidine biosynthesis are noted. The isotopic method has the advantage over conventional radioactive metabolic studies in that the PMR spectrum distinguishes between different hydrogens, thus eliminating the need for chemical degradation of the product to determine the specific activity at each position. Since RA is produced in high yields and is easily crystallized from water it is particularly suited for this type of biosynthetic study.

Methods and Materials

Stock cultures were maintained on malt-agar slants. Crystalline RA was obtained from low-iron cultures of R. pilimanae (Atkin

and Neilands, 1968). D_2O adapted cultures were maintained as liquid cultures and were transferred monthly. 99.8% D_2O (atom % D) was purchased from Bio-Rad Laboratories, Richmond, California. PMR spectra were recorded at 220 MHz on a Varian HR220 spectrometer using 20 mg samples of RA in 0.5 ml of D_6 -dimethyl sulfoxide (Merck, Sharp, and Dohme) and tetramethyl silane as internal standard.

ASKP medium. The medium normally used for growing R. pilimanae contains acetate and citrate, both of which contain carbon-bound hydrogens. Therefore, the following medium was used for all studies performed in D_2O : 2 g ammonium sulfate (enzyme grade), 4 g anhydrous K_2HPO_4 , and 20 g sucrose per l. (designated ASKP medium). Trace metals and thiamine were the same as normally used for R. pilimanae (Atkin and Neilands, 1968) or Ustilago spærogena (Garibaldi and Neilands, 1955). H_2O stock solutions of trace metals and thiamine were evaporated to dryness and then reconstituted with D_2O . D_2O cultures were adjusted to pH 6.8 using concentrated H_3PO_4 . Autoclaving caused a slight precipitation of metal salts.

Determination of Deuterium Concentration. The deuterium concentration was determined spectrophotometrically (Crespi and Katz, 1961) using commercial 99.8% D_2O as a standard. With a hot auto-

clave and tightly cotton-plugged flasks, the deuterium concentration of the medium after autoclaving was 98.5%. At the end of a typical two-three week experiment when the RA was harvested, the deuterium concentration dropped to 97.0 - 96.5%.

Hydroxamate Assay and Hydroxylamine Spray. The standard spectrophotometric hydroxamate assay (Atkin and Neilands, 1968) was employed, utilizing a Beckman DU spectrophotometer. Up to 3 μ moles of hydroxamate were diluted to 3 ml with 5 mM $\text{Fe}(\text{ClO}_4)_2$ -0.1 M HClO_4 . In these conditions AHO was found to have a molar extinction coefficient, A_{505} , of 935 (total AHO determined by a recording titrator) (Neilands and Cannon, 1955). Both the wavelength of maximum absorbance and the intensity of absorption agree with previous reports for monohydroxamates (Seifter et al., 1960). The A_{505} is almost 1/2 and 1/3 of the absorbancy coefficients for RA (Atkin and Neilands, 1968) and ferrichrome (Emery, 1967), respectively. Hydroxylamines were detected by the use of a tetrazolium spray (Snow, 1954).

Adaptation of *R. pilimanae* to D_2O . Two 10 ml cultures of ASKP medium were prepared, one using 100% (99.8%) D_2O and the other 50% D_2O and 50% H_2O ; both were inoculated from a malt-agar slant. Growth was observed in the 50% culture in about four days. One drop of the 50% culture was added daily to the 100% culture starting with day 7. After about 20 days growth was observed in the 100% culture.

This culture then served as an inoculum for other 100% D₂O cultures. After several transfers deuterated R. pilimanae grew at about 1/2 the rate of protonated R. pilimanae. Cell volume was greater in deuterated cultures than in protonated, owing to an increase in cell size. On a volume basis deuterated cultures produce about 20% of the RA of protonated cultures. Deuterated R. pilimanae often adheres to the sides of the culture flasks. Also, there is a change in some aspect of carotenoid biosynthesis since cultures more than 80% deuterated fail to turn the typical orange-red color of mature protonated cultures but instead remain a dark pink.

Synthesis of δ -N-Acetyl- δ -N-Hydroxyornithine. 10 g of RA in 100 ml of 6 N HCl was sealed in two 2 x 50 cm evacuated Carius tubes and hydrolyzed overnight at 100° to δ -N-L-hydroxyornithine (Atkin and Neilands, 1968). The hydrolysate was flash evaporated at 40° to an oil, dissolved in 100 ml of H₂O and the pH adjusted to 4.3 with pyridine. One equivalent of acetic anhydride (5.5 ml) was added in small portions over one hour while the solution was kept at 50°. Then 0.1 ml portions of acetic anhydride were added until the hydroxamate content reached a maximum as measured at 505 nm in the hydroxamate assay. The reaction mixture was concentrated, acidified with glacial acetic acid, and placed on a 3 x 50 cm Dowex-50 column, which was in the hydrogen form. The column was developed using a

linear gradient between 2 l. of 2 N acetic acid and 2 l. of 2 N pyridinium-acetate, pH 5.0. L-AHO was distinguished from α, δ -N,N-diacetyl- δ -N-hydroxyornithine and δ -N-hydroxyornithine peaks by their elution sequence, electrophoretic mobility (0.1 M pyridinium-acetate, pH 4.8), and ninhydrin, hydroxamate, and tetrazolium tests. The L-AHO was acidified with HCl and the pyridinium-acetate removed by repeated addition and evaporation of water. After passing through 3.5 x 110 cm Sephadex G-10 column with water as eluent, 6.63 g (60% of theoretical) of L-AHO was recovered. The hydrochloride form was crystallized from water by the slow addition of ethanol. The biological activity of AHO was tested using enb mutants of Salmonella typhimurium (Pollack et al., 1970).

Synthesis of DL-AHO. 10 g of RA were hydrolyzed as above and the product racemized by refluxing for 2 hours in a solution containing 100 ml of glacial acetic acid and 50 ml acetic anhydride. The resulting ninhydrin-negative material was decolorized with charcoal and flash evaporated to an oil. An attempt was made to resolve (Greenstein, 1957) a portion of this compound with porcine kidney acylase (Calbiochem); however, no ninhydrin-positive material was produced. The remainder of the diacetyl-DL-hydroxyornithine was hydrolyzed, acetylated, and purified by use of the same procedures applied to L-AHO. The method of Greenstein and Winitz (1961) was

employed for the synthesis of δ -N-acetyl-L-ornithine. Argininosuccinic acid and L- α,γ -diaminobutyric acid \cdot 2HCl were purchased from Sigma Chemical Company. All other chemicals were from regular commercial sources.

Molecular Rotation of L-AHO. 100 mg of AHO in 2 ml of 50% HI were sealed in an evacuated Carius tube and hydrolyzed at 100° for 15 hours. The hydrolysate was flash evaporated at 40° to an oil and the HI and I₂ removed by repeated additions and evaporations of small amounts of 0.1 N HCl. A duplicate sample was prepared to contain 100 mg of a specimen of commercial ornithine which has previously been shown by optical rotatory dispersion to contain greater than 98% of the L-isomer. The samples were diluted to 5.0 ml with 1 N HCl, the ornithine content determined (Chinard, 1952), and the optical rotatory dispersion spectrum recorded with a Cary 60 instrument. No racemization of L-ornithine was observed in this process. The ornithine from L-AHO was determined to be 0.856 L isomer. From these data the molecular rotation for the L isomer of AHO was determined to be $[M]_{400}^{20} = +130^\circ$ and $[M]_{350}^{20} = +172^\circ$ in 1 N HCl. Using these molecular rotations the partially racemized preparation of AHO was found to be 0.670 L isomer.

Purification of Deuterated RA. The D₂O cultures were centrifuged and the medium flash evaporated at 40° to an oil. The oil was dissolved in 50 ml of water and extracted twice with 100 ml of CHCl₃-

phenol (1:1 weight basis). The RA was re-extracted into water after the addition of about 500 ml of ether to the CHCl_3 -phenol. The water phase was concentrated to about 5 ml, acidified with acetic acid, and applied to a 1.5 x 40 cm Dowex-50 column in the hydrogen form. RA was eluted with 2 N acetic acid, and hydroxamate-positive fractions were pooled and the acetic acid removed by repeated additions and evaporation of water. The RA was dissolved in the least possible amount of water and placed on a 3.5 x 110 cm Sephadex G-10 column using water as an eluent. The RA was dried in vacuo over P_2O_5 . Using this procedure quantities as small as 35 mg were recovered.

Recovery of AHO. In the cultures containing exogenous AHO the RA and unconsumed AHO were recovered by placing the acidified, concentrated culture medium directly on a 1.5 x 40 cm Dowex-50 column. The column was developed with a linear gradient between 250 ml of 2 N acetic acid and 250 ml of 2 N pyridinium-acetate, pH 4.8. Two hydroxamate-positive peaks were observed: the first to appear was RA, while AHO was retarded. Both fractions were further purified by removing the pyridinium-acetate by repeated evaporation from water and gel-filtration as described above for RA.

General Procedures. In a typical experiment 100-170 ml of D_2O ASKP medium were inoculated with one to two ml of D_2O -adapted culture and placed on a rotary shaker at 30° . After about four days one ml of the culture was withdrawn and assayed for hydroxamate

concentration. At the same time the protonated supplements were added. Enough supplement was added to make the culture 20 mM except for argininosuccinate, L-AHO, and DL-AHO which were added at a level of 5.0 mM, 13.4 mM and 6.65 mM respectively. After a total incubation period of two to three weeks the RA was purified from the culture. The greatest amount of packed cell volume with the smallest amount of RA produced occurred by day four.

The D₂O medium in tightly cotton-plugged Erlenmeyer flasks was autoclaved using a preheated autoclave for 10 minutes at 120°. Using this procedure there was little protonation of the D₂O. The supplements were dissolved in about 10 ml of D₂O and the pD adjusted to 6-8 with concentrated H₃PO₄ or KOH (in D₂O) prior to sterilization. Samples of the amino acid supplements were analyzed by electrophoresis at pH 4.8 (0.1 M pyridinium-acetate), pH 7.0 (0.1 M phosphate) and pH 10.0 (0.1 M carbonate) before and after sterilization. Electrophoretograms were developed with ninhydrin to determine if any new amino-containing compounds were produced by the autoclaving procedure. In addition, samples of argininosuccinate before and after autoclaving were chromatographed in phenol:H₂O (100:20,w:w), a system which can resolve argininosuccinate and its anhydride form (Ratner, 1957). No new ninhydrin-positive products were observed.

Calculation of Data from PMR Spectra. PMR peak assignments for RA have been reported previously (Atkin and Neilands, 1968). In

reduced RA (Atkin and Neilands, 1968) FMR spectrum (Figure 1) the β and γ methylene hydrogens are resolved (Figure 1) and have been assigned (Llinás, 1971). Due to low yields in the reduction and limited solubility in dimethyl sulfoxide the spectra of the reduced RA species were not determined. Since the RA samples were purified from water, and dimethyl sulfoxide was used as a solvent, the amide NH and hydroxamate NOH resonances appear as distinct peaks. However, due to the acidic nature of the hydroxamate group its resonance is broadened due to exchange. So, the amide NH area was used as a standard by defining it as one.

As the instrument's integrator was unreliable it was necessary to integrate the area under each peak by cutting it out and weighing the paper. Three expanded scale copies of each spectrum were made and each paper cut-out was weighed three times to the nearest 0.1 mg. Most weight measurements were in the 10-150 mg range. The average of the nine measurements for each peak was divided by the number of hydrogens of that type in RA and normalized to the ring amide proton to give $[H]_{\text{observed}}^i$ ($i = \alpha, \beta + \gamma, \delta$ or CH_3). Using this procedure and a protonated RA sample, numbers within ± 0.015 of the expected value of 1.000 were obtained. The variations in the weight (120 mg) of five 25 cm^2 samples of the paper used for recording the spectra was $\pm 0.8 \text{ mg}$.

When an exogenous substrate was supplied a correction was required for the endogenous production of RA. The fraction (X) of the RA which resulted from exogenous substrate was determined by assuming: 1) The fraction of protonation ([H]) at each carbon for endogenously produced RA is the same as that when no exogenous substrate is supplied (see discussion); and, 2) no H-D exchange for the non- α -hydrogen aliphatic hydrogens of the exogenous substrate, i.e. $[H]_{\text{exogenous}}^i = 1.000$ for $i = \beta + \gamma, \delta, \text{CH}_3$. The experimental X^i values were then calculated:

$$X^i = \frac{[H]_{\text{observed}}^i - [H]_{\text{endogenous}}^i}{1.000 - [H]_{\text{endogenous}}^i}$$

The X^i values were averaged (see Column A, Table I) to give X if the exogenous substrate had the hydrogens of the type considered. The $i = \alpha$ values were not considered in computing X. The $[H]_{\text{exogenous}}^i$ can be computed based on

$$[H]_{\text{exogenous}}^i = \frac{[H]_{\text{observed}}^i - [H]_{\text{endogenous}}^i (1-X)}{X}$$

Results

Table I lists the relative retention (X) of protons from exogenous substrates furnished for the biosynthesis of RA. The percent of RA derived from the exogenous substrate (see methods), when present, is also listed. Corrections have been made for the endogenous

production of RA. The substrate for the L-AHO experiment was actually 85.6% L isomer while that for the DL-AHO experiment was 67.0% L isomer.

The relative protonation of the endogenous production of RA is recorded in Table II. D-alanine, D-ornithine, D-arginine, α -acetyl-L-glutamate, α -acetyl-L-ornithine, L-lysine, and argininosuccinate were also tried as exogenous substrates; however, they were not incorporated, as the relative protonation of the RA recovered in these experiments was similar to the "none" values.

In the experiments involving AHO, the AHO that was not converted into RA was recovered and its isomeric composition determined. The values observed are recorded in Table III. The AHO recovered from the DL-AHO experiment had a $[M]_{400}^{20}$ of -140° indicating that it was completely D isomer. This D isomer was as active or more active on a molar basis than AHO that was 85.6% L isomer in supporting the growth of a number of siderochrome-requiring mutants of Salmonella typhimurium.

Protonated exogenous amino acids were added to aliquots of a protonated ASKP culture initially 0.94 mM in RA. After 48 hours of exposure to the exogenous amino acid the RA concentration was again determined (Figure 2). The amino acid additives for this experiment were prepared in the same manner as the deuterated cultures except that H_2O was used instead of D_2O .

Discussion

Precursors of RA. The data in Table I show that the two δ protons of exogenous arginine, citrulline, and ornithine both appear in the RA produced when these substrates are present, indicating that one of these is the initial substance on the biosynthetic pathway to RA. If proline, glutamine, or glutamate were the initial precursor of RA one of the δ hydrogens from arginine, citrulline, or ornithine would be lost during the formation of glutamic semialdehyde and the second would be eliminated during the synthesis of glutamate.

The regular loss of α protons in the sequence citrulline \rightarrow arginine \rightarrow ornithine indicates that ornithine, rather than any other urea cycle intermediate, is the initial substance in RA biosynthesis. This does not exclude the possibility of α -acetylornithine as the initial RA precursor. However, three lines of evidence indicate that ornithine rather than α -acetyl-ornithine is the initial precursor of RA biosynthesis: 1) The possible distinction made in R. pilimanae between endogenous and exogenous ornithine (see below) with exogenous ornithine not available for arginine biosynthesis (or α -acetylornithine formation); 2) The interference of RA formation by the ornithine analogs, lysine and α, γ -L-diaminobutyrate (Figure 2); and 3) The reversibility of acetyl-ornithine- δ -transaminase (Albrecht and Vogel, 1964), which would labilize one of the δ hydrogens.

Presumably, the α protons are lost due to Schiff base formation with pyridoxal enzymes. However, ornithine and arginine decarboxylases would not labilize the α proton (Mandeles et al., 1954). As AHO is a precursor of RA (see below), the α hydrogen data for ornithine and AHO (Table I) indicate that little if any α proton is lost at the ornithine level. A similar observation has been made with $\alpha^{15}\text{-N-}\alpha\text{-}^2\text{H-lysine}$ in intact rats (Clark and Rittenberg, 1951) indicating a non-involvement of α,ω -diamino acids with pyridoxal enzymes. D-ornithine has been reported on several occasions thus implicating an ornithine racemase in a number of organisms (Guinand et al., 1969; Tsuda and Friedmann, 1970). An arginine racemase is known in Pseudo-monas graveolens (Yorifuji et al., 1971) but as this enzyme also catalyzes the racemization of ornithine it probably is not present in R. pilimanae. Initial Schiff base formation of arginine or citrulline with pyridoxal is sufficient to promote exchange of the α proton independently of racemization or transamination (Snell and DiMari, 1970). The hope was to determine the relative retention of the α protons for all the urea cycle intermediates but, unfortunately, exogenously supplied argininosuccinate failed to appear in RA. Using the α proton values for citrulline and arginine it can be determined (exclusive of isotope effects) that a total of 37% of the exogenous citrulline loses the α proton at the citrulline and/or

argininosuccinate level. Similarly, 16% of exogenous arginine α proton is lost before conversion to ornithine. The carbon skeleton of exogenous proline and glutamine reach ornithine via glutamate, at which point they are involved as co-substrates for many transaminases, thereby losing the α proton. Labilization of the α proton might accompany active transport of amino acids; however, this possibility has been excluded by Kessel and Lubin (1965).

As both of ornithine's δ protons are preserved in RA, the possibility of a biosynthetic intermediate unsaturated at the δ carbon can be eliminated. This suggests that O_2 rather than water is the source of the oxygen in the hydroxylamino group, a finding compatible with studies on the biosynthesis of hadacidin (Stevens and Emery, 1966).

As the $\beta + \gamma$, δ , and methyl protons of AHO are incorporated into RA in a 1.00:1.00:1.07 ratio (average of two experiments) it can be concluded that AHO is not hydrolyzed to acetate and hydroxyornithine and then reassembled, but rather the hydroxamic acid containing amino acid (AHO) of RA is synthesized before incorporation into the cyclic dipeptide. If AHO were hydrolyzed and reassembled the acetate would first mingle with endogenously produced deuterated acetate. As most of the RA is protonated (~90% from exogenous AHO) the reassembled AHO would have a lower acetyl to hydroxyornithine protonation ratio than the unhydrolyzed case. Simultaneously, the acetate

would have a higher than endogenous protonation and its condensation with oxaloacetate would eventually lead to ornithine with a higher than normal endogenous protonation at the γ position. Both of these effects would distort the 1:1:1 ratio observed. As more than 50% of the RA was formed from exogenous AHO, RA arises from identical monomers. Interestingly enough, 2,3-dihydroxy-N-benzoyl-L-serine is not a precursor of enterobactin, a representative of the other major type of iron transport compound (Bryce et al., 1971).

Emery (1966) studied the incorporation of AHO labeled with ^{14}C in the acetyl and hydroxyornithine portions, into ferrichrome compounds. He concluded that in ferrichrome, as in RA, the hydroxamic acid portion is formed before the amino acids are cyclized. As Emery has shown that hydroxyornithine is a precursor of ferrichrome, the biosynthesis of RA probably proceeds from ornithine to δ -hydroxyornithine to AHO, which is then assembled into the cyclic dipeptide (Figure 3).

The possibility that δ -N-acetyl-L-ornithine is a precursor of RA and the report of the natural occurrence of this compound (Brown and Fowden, 1966) prompted the investigation of δ -N-acetyl-L-ornithine in this system; however, no incorporation of protons into RA was observed. As this substance is a free amino acid, R. pilimanae should be able to accumulate δ -N-acetyl-L-ornithine. A possible

reason incorporation is not observed is because R. pilimanae does not hydrolyze δ -N-acetyl-L-ornithine or convert it into AHO. Emery (1966) also concluded that this compound is not a precursor of the AHO groups in ferrichrome. He described (1971b) a δ acetylating enzyme which uses δ -N-hydroxyornithine, not ornithine, as a substrate. The amide form was found not to be a precursor of hadacidin (Stevens and Emery, 1966). However, MacDonald (1965) has reported N-hydroxylation of a diketopeperazine precursor of aspergillic acid, and ϵ -N-hydroxylysine does not appear to be an intermediate in the formation of mycobactin P and S (Tateson, 1970). Apparently, two methods of forming hydroxamic acids have evolved.

The lack of incorporation of α -acetylglutamate and α -acetylornithine into RA is not sufficient evidence to indicate that R. pilimanae does not use these intermediates in the biosynthesis of ornithine. Other organisms have been reported which lack the ability to transport these substances but still use them as intermediates (Prozosky, 1967; and references therein). The failure of argininosuccinate to be incorporated may be due to similar reasons, as argininosuccinate lyase mutants of Proteus mirabilis are unable to grow on argininosuccinate (Prozosky, 1967).

The high level of protonation of the δ position from exogenous glutamate, glutamine, or proline is unexplained since glutamate and

glutamine do not originally have δ hydrogens and those of proline are lost in conversion to glutamate. Little if any glutamic semi-aldehyde is converted to ornithine via ornithine- δ -transaminase, an essentially irreversible reaction (Strecker, 1965). Also, the α hydrogen of proline is preserved in this conversion. If R. piliminae is considered to use the accepted acetylated pathway for ornithine synthesis (see Figure 3 and below) one of the hydrogens originates during a transamination and the second from NADPH. Although Dunathan (1970) has observed transfer of the α hydrogen to the amino acceptor during transamination (δ on RA), a more probable source of the δ protonation is from NADPH arising from metabolism of proline, glutamine, or glutamate, as α -ketoglutarate produces RA with more than endogenous protonation (Table II).

Exogenous acetate appears to a greater extent in the acetyl groups than in the γ position (via citrate synthetase and ornithine synthesis). The acetate consumed in the citric acid cycle maybe segregated from the acetate used in the acetylation step of RA biosynthesis. The high level of protonation of the α position may be due to the formation of protonated NADPH during the metabolism of the exogenous acetate. An examination of the methyl column in Table I shows that there is little perturbation when exogenous substrates are present, i.e., the values are near zero with non-acetyl

containing exogenous substrates. This indicates there is no significant change in the metabolic source of the acetyl groups, justifying the assumption made above (see methods and materials).

In Rhodotorula glutinis it is known that pyruvate will only be utilized after glucose is completely exhausted (Medrano et al., 1969). A similar situation with acetate would explain the low level of incorporation of acetate into RA.

Arginine Metabolism and Regulation in R. pilimanae. The pathways shown in Figure 3 involving arginine metabolism (excluding the δ -N-hydroxyornithine branch) have been partially investigated and thought to exist in a number of fungal genera, namely: Saccharomyces (Middelhoven, 1964; DeDeken, 1962), Aspergillus (Piotrowska et al., 1969), Candida (Middelhoven, 1963), Blastocladiella (Smith and Holmes, 1970), Neurospora (Castañeda et al., 1967), and Ustilago (Priour, P., 1971). They are assumed also to be the major pathways of arginine metabolism in R. pilimanae.

Arginine is known to regulate its own biosynthesis in a number of organisms (Prozosky, 1969; Udaka, 1966; DeDeken, 1962). A similar situation probably exists in R. pilimanae, since when arginine is present in the medium all the RA synthesized comes from exogenous arginine (Table I). This would be the case if exogenous arginine prevented the endogenous synthesis of arginine, as the only source of ornithine would be from the breakdown of exogenous arginine.

When citrulline is present all of the RA is synthesized from the exogenous material (Table I). This could be due to intracellular conversion of exogenous citrulline to arginine, as citrulline does not regulate α -N-acetylglutamic reductase in S. cerevisiae (DeDeken, 1962). The arginine produced from citrulline could now prevent the endogenous production of ornithine from glutamate. This suggests that argininosuccinate synthetase and argininosuccinate lyase are not repressed by arginine, but that the regulation of arginine biosynthesis in R. pilimanae may be by feedback inhibition rather than via the coordinate operon control model. Middelhoven (1969) observed that argininosuccinate lyase in S. cerevisiae was not repressed by exogenous arginine. Unfortunately, argininosuccinate synthetase was not examined.

Decrease in synthesis of RA (Figure 2) when arginine or citrulline is supplied exogenously could be explained by a decrease in the availability of ornithine, as in S. cerevisiae (Middelhoven, 1970) and Aspergillus nidulans (Piotrowska et al., 1969) ornithine transaminase and arginase are coordinately induced by high levels of arginine. Ramos et al. (1970) found that the internal ornithine concentration in S. cerevisiae is lower when arginine is added to their ammonium-containing medium. Citrulline was not tested.

When the ornithine analogs lysine or α, γ -L-diaminobutyrate are added to R. pilimanae cultures (Figure 2) there is a decrease in

the production of RA. In S. cerevisiae high levels of exogenous lysine decrease the internal ornithine level (Ramos et al., 1970), explaining the lower RA production. Another possibility is that lysine and α, γ -L-diaminobutyrate, inhibit the initial hydroxylation enzyme since the acetylation enzyme is known to use either ϵ -hydroxylysine or δ -hydroxyornithine as a substrate (Emery, 1971b).

As there is no loss of α protons at the ornithine level for exogenous ornithine (see above), it can be concluded that a significant portion of the exogenous ornithine is not touring the urea cycle to be reformed into ornithine. If this were the case α protons would be lost at the arginine and citrulline and/or argininosuccinate stage. This could be due to the repression of ornithine transcarbamylase by high levels of exogenous ornithine, which is known to occur in S. cerevisiae (Ramos et al., 1970). Another possible explanation is that for the synthesis of RA a distinction is made between endogenous and exogenous ornithine. Compartmentalization of ornithine is known in Neurospora crassa, as exogenous ornithine is degraded by ornithine transaminase while endogenous ornithine (possibly mitochondrial) is used mainly for the synthesis of arginine (Davis, 1968). A distinction between the two types of ornithine would suggest that the arginine biosynthetic enzymes are segregated from the initial enzyme on the RA pathway.

When 20 mM exogenous ornithine is supplied to deuterated R. pilimanae cultures 88% of the RA produced is derived from the exogenous ornithine (Table I). The remaining RA is derived from endogenous ornithine, whose synthesis was not prevented by this high level of exogenous ornithine. Again, a similar observation has been made in S. cerevisiae, where N-acetylglutamic acid reductase is feedback-inhibited by arginine but not by ornithine. Arginine is known to repress the synthesis of this enzyme in S. cerevisiae, but ornithine was not examined (DeDeken, 1962). In general, the regulation of arginine metabolism appears to be the same in R. pilimanae and S. cerevisiae.

Similarly, since endogenous AHO is produced in the presence of 13.4 mM exogenous AHO it is concluded that the biosynthesis of AHO is not regulated rigidly by AHO. The high level of production of RA from AHO (Figure 2) indicates that the rate-determining step in RA biosynthesis precedes AHO formation. When cultures are shaken vigorously they produce more RA and hydroxylation may be the rate limiting step. Another possibility is that the enzymes in the pathway are sequentially induced and a high level of AHO induces the cyclization enzymes.

The appearance of protons in the acetyl group of RA when α -ketoglutarate was the exogenous substrate (Table II) can be explained

by the reversal of the citric acid cycle. The two protons originally on the γ position of α -ketoglutarate would appear in the methyl group of acetyl-CoA by using the citrate cleavage enzyme (Atkinson, 1969). Since this enzyme exists extramitochondrially (Srere, 1959) and since the acetylating enzyme uses acetyl-CoA as a substrate (Emcry, 1971b), the high incorporation of the protons of α -ketoglutarate into the acetyl portion of RA, and the possibility of two ornithine pools suggests that the biosynthesis of RA does not occur in the mitochondria.

AHO Racemase. Table III shows that R. pilimanae preferentially consumed the L isomer of the AHO in the biosynthesis of RA. However, since there is not enough L-AHO initially present in the exogenous substrate to account for the amount of RA produced, some D isomer must have been converted into RA. The RA produced under these conditions is possibly an LL or DD diketopiperazine. But, as three chromatography solvent systems (Kopple and Ghazarian, 1968) known to separate DL and LL diketopiperazines failed to resolve the RA from either the L or DL-AHO experiment, R. pilimanae apparently has the capacity to racemize AHO. Ferribactin, from Pseudomonas fluorescens, contains the D form of δ -N-hydroxyornithine (Maurer et al., 1968). The presence of an AHO racemase would explain the observed labilization of the α hydrogen at the monomer level. Because other D-amino

acids were not incorporated into RA, R. pilimanae may not have a specific D-amino acid transporting system. However, D-AHO may enter R. pilimanae as a metal hydroxamate via a transport system that does not specifically examine the configuration at the α carbon, as D-AHO was observed to satisfy a siderochrome requirement of mutant Salmonella typhimurium.

Cyclic Peptide Formation. The incorporation of D-AHO and the labilization of the α hydrogens are consistent with recent developments in the synthesis of the cyclic decapeptides gramicidin and tyrocidine (Kleinkauf et al., 1971; and references therein). The initial enzyme for tyrocidine assembly has been shown to be an ATP-dependent racemase, which can activate either D- or L-phenylalanine. In addition, amino acyl-pantotheine intermediates have been shown to be involved in the assembly process. The labilization of the α hydrogen of thioesters is well known (Bruice and Benkovic, 1966). By analogy to the cyclic decapeptide scheme the RA assembly system should consist of an initial enzyme which is capable of activating L- or D-AHO. On this enzyme the D-AHO is labilized. The second RA assembly enzyme (cyclizer) will also activate AHO and should contain a pantothenic acid moiety that is involved in amide bond formation. The α hydrogens of two AHO monomers which are assembled into a single RA molecule have both been shown to be labile at some

phase in the biosynthesis (Akers and Neilands, manuscript in preparation), an observation consistent with the above predictions, as both monomers are thioesters at one stage of biosynthesis. Work is in progress to isolate the enzyme system involved in the synthesis of RA and the cyclic hexapeptide ferrichrome.

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5093.

TABLE I: Relative Retention of Exogenous Protons into RA^a

Exogenous Substrate	α	$\beta+\gamma$	δ	CH_3	A	%RA from Exogenous Source
L-Glu	0.035 <u>+0.005</u>	1.000 <u>+0.008</u>	0.412 <u>+0.020</u>	0.010 <u>+0.003</u>	$\beta+\gamma$	25
L-Gln	0.050 <u>+0.022</u>	1.000 <u>+0.030</u>	0.290 <u>+0.006</u>	0.006 <u>+0.006</u>	$\beta+\gamma$	23
L-Pro	0.038 <u>+0.009</u>	1.000 <u>+0.038</u>	0.235 <u>+0.049</u>	-0.016 <u>+0.034</u>	$\beta+\gamma$	39
L-Cit	0.344 <u>+0.015</u>	1.010 <u>+0.036</u>	0.989 <u>+0.051</u>	0.061 <u>+0.002</u>	$\beta+\gamma, \delta$	106
L-Arg	0.547 <u>+0.010</u>	1.070 <u>+0.005</u>	0.938 <u>+0.007</u>	0.012 <u>+0.002</u>	$\beta+\gamma, \delta$	100
L-Orn	0.650 <u>+0.009</u>	1.036 <u>+0.031</u>	0.968 <u>+0.024</u>	-0.002 <u>+0.004</u>	$\beta+\gamma, \delta$	88
L-AHO	0.638 <u>+0.012</u>	0.984 <u>+0.013</u>	0.946 <u>+0.009</u>	1.066 <u>+0.079</u>	$\beta+\gamma, \delta, \text{CH}_3$	91
DL-AHO	0.596 <u>+0.029</u>	0.972 <u>+0.005</u>	1.012 <u>+0.011</u>	1.032 <u>+0.013</u>	$\beta+\gamma, \delta, \text{CH}_3$	88

Footnote to Table I

^aThe relative incorporation into RA of protons from the exogenous substrates listed. Values are relative to the proton type (or average of types) listed in column A. The figures have been corrected for endogenous production (Blank values of Table II). The percent of the RA derived from the exogenous material, while present, is also shown. The L-AHO and DL-AHO were actually 85.6% and 67.0% L isomer respectively. The tolerance figures listed are the standard deviation of the mean. These deviations are not meant to indicate the accuracy of the experiments, which is not greater than 1.5% (see methods and material section), but rather to show the precision of the measurements.

TABLE II: Relative Protonation of RA^a

Exogenous Substrate	i =			
	α	$\beta+\gamma$	δ	CH_3
None (Blank)	0.026	0.076	0.085	0.079
	<u>+0.001</u>	<u>+0.002</u>	<u>+0.002</u>	<u>+0.002</u>
Acetate	0.199	0.064	0.078	0.113
α -Keto- glutarate	0.080	0.273	0.161	0.719

^aThe protonation of the RA produced in deuterated cultures, relative to amide NH = 1.000. The deviations listed are the same type as in Table I.

TABLE III: Material Balance for RA Production from AHO in D₂O^a

	AHO Experiment	
	"DL"	"L"
(A) RA present when AHO added	84	81
(B) AHO added (<u>L</u> isomer)	565(378)	1140(976)
(C) Total RA present at end of experiment	578	1367
(D) AHO recovered (<u>L</u> isomer)	87(0)	71(11)
(E) Percent of total RA from exogenous AHO	75.0	79.8
RA from exogenous = (C)(E)	434	1090
AHO consumed = (B)-(D)	478	1069

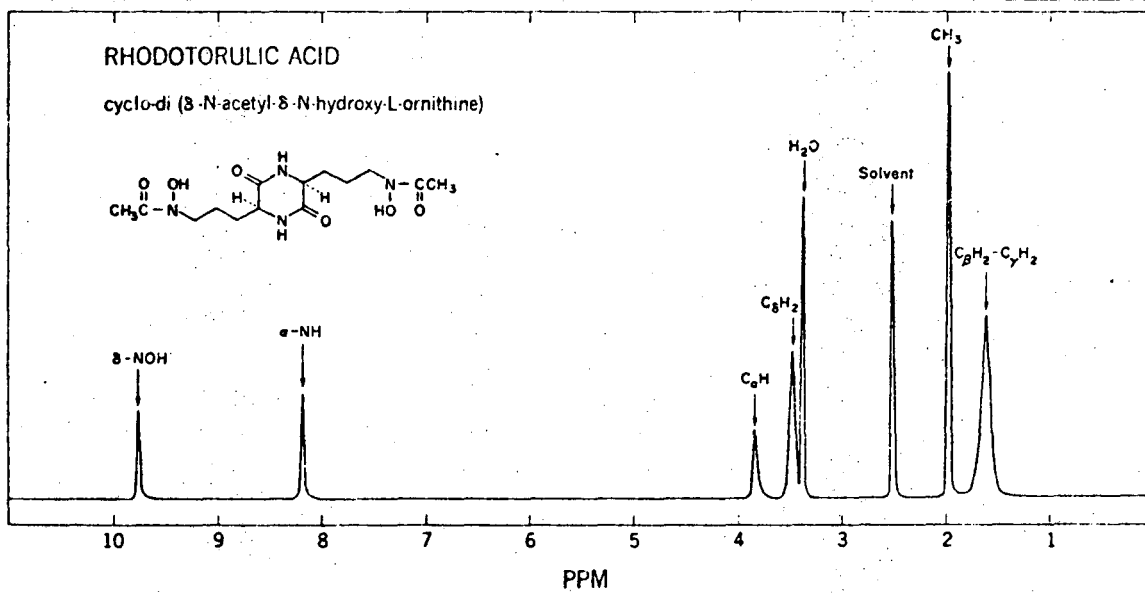
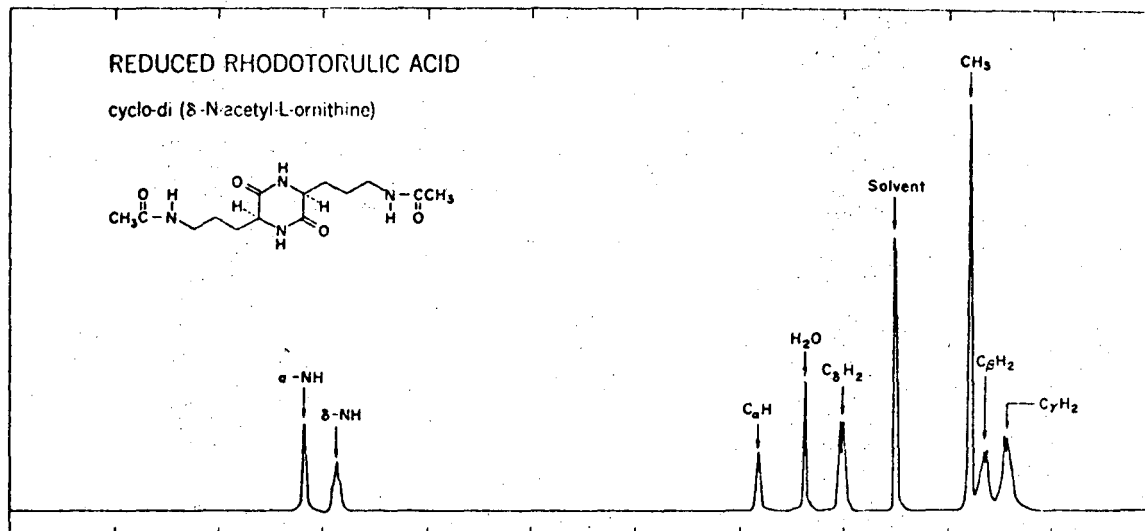
^aThe material balance for RA production from AHO in D₂O. In the experiments involving additions of AHO to deuterated cultures the un-consumed AHO was recovered. Figures in parentheses are amounts of L isomer of AHO. Percentages at Row E are X values (see methods and materials). Figures are μ moles for and μ moles/2 for AHO.

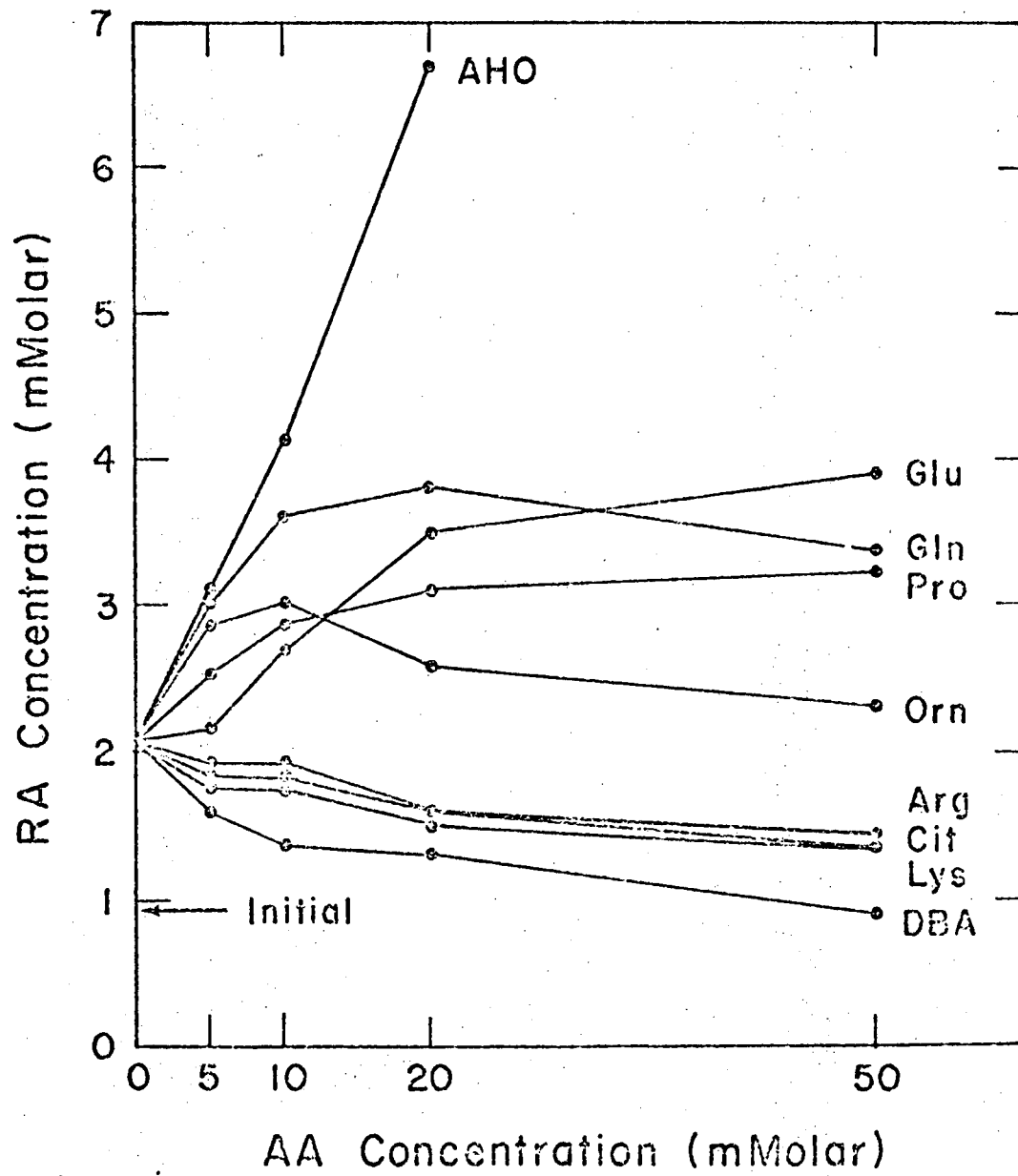
Figure Legends

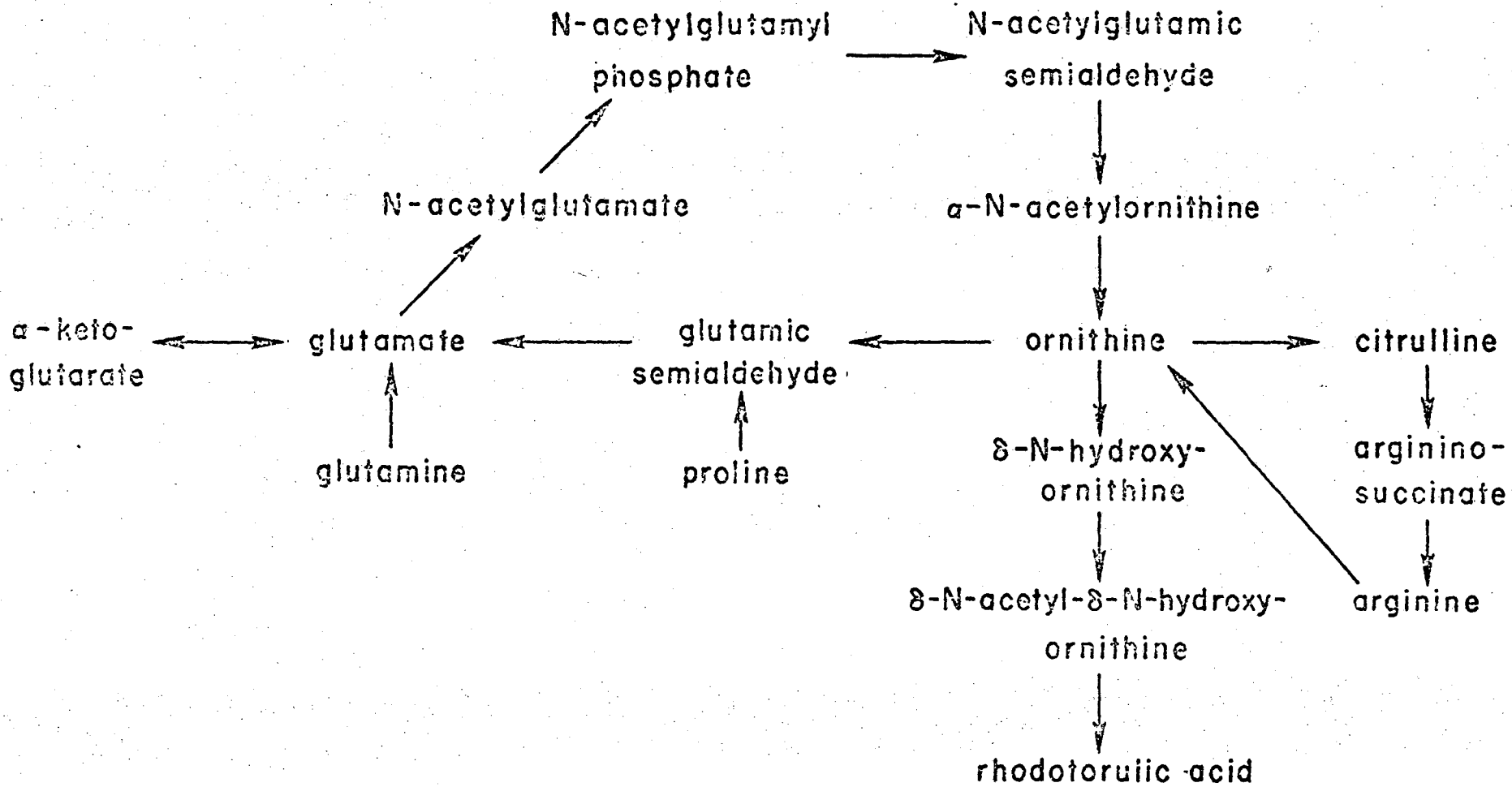
FIGURE 1: The Proton Magnetic Resonance spectra of protonated RA and reduced RA in D_6 -dimethylsulfoxide. The PPM are relative to tetramethylsilane.

FIGURE 2: The effect of exogenous AA on the production of RA. Exogenous AA were added to aliquots of a protonated culture initially 0.94 mM in RA. After 48 hours the RA concentration was again determined. All amino acids were the L isomer except AHO which was 85.6% L isomer. DBA = α,γ -diaminobutyrate.

FIGURE 3: The accepted scheme of arginine metabolism. The proposed RA biosynthetic pathway is shown.







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