1982

Androgen Biosynthesis in the Testis

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ANDROGEN BIOSYNTHESIS IN THE TESTIS:

I. KINETIC CHARACTERIZATION OF THE STEROIDOGENIC PATHWAYS IN GUINEA PIG, RABBIT, DOG AND RAT TESTIS MICROSOMES.

II. THE EFFECT OF ASCORBIC ACID ON 3-B HYDROXYSTEROID:NAD-OXIDOREDUCTASE ACTIVITY IN THE TESTIS

BY

MARY A. ACCHIARDI

A DISSERTATION SUBMITTED IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE DEGREE OF

DOCTOR OF PHILOSOPHY

IN

BIOLOGICAL SCIENCES
(FOOD SCIENCE & NUTRITION)

UNIVERSITY OF RHODE ISLAND

1982
DOCTOR OF PHILOSOPHY DISSERTATION

OF

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Dissertation Committee

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Dean of the Graduate School

UNIVERSITY OF RHODE ISLAND

1982
The kinetic parameters of testicular steroid synthesizing enzymes involved in the $\Delta_4$ and $\Delta_5$ pathways for testosterone production were examined in four species. Testis microsomes were prepared as a source of steroidogenic enzymes. Radiolabeled steroid-intermediates were used as substrates and products were separated by thin layer chromatography. Michaelis constants, ($k_{0.5}$), and $V_{\text{max}}$ values were determined for 3 $\beta$-Hydroxysteroid oxidoreductase (EC 1.1.1.145), 17$\alpha$-Hydroxylase (EC 1.14.99.9) and $C_{17}-C_{20}$ Lyase reactions using $\Delta_4$ and $\Delta_5$ substrates.

The purpose of the study was to develop a kinetic model for predicting the predominant steroidogenic pathway in mammalian testes. The key step in determining the direction of the pathway is the conversion of pregnenolone to either progesterone ($\Delta_4$) or to 17$\alpha$-hydroxypregnenolone ($\Delta_5$). The most definitive kinetic predictors were found to be 1) the Michaelis constant of the 3BHSOR reaction for PREG and, 2) the $V_{\text{max}}$ of the PREG 17$\alpha$-Hydroxylase reaction. The $\Delta_4$ species demonstrate a low $k_{0.5}$ for PREG in the 3BHSOR reaction and a low $V_{\text{max}}$ for the conversion of PREG$\rightarrow$17$\alpha$PREG. Conversely, $\Delta_5$ species demonstrate a high $k_{0.5}$ for PREG in the 3BHSOR reaction and a high $V_{\text{max}}$ for the hydroxylation. The relative affinity of 3BHSOR and 17$\alpha$-Hydroxylase for PREG was found to be a useful predictor of the predominant pathway. A high ratio, indicating greater affinity of the hydroxylase for PREG, favors
conversion to Δ₅ steroids and a low ratio is indicative of the Δ₄ pathway. Several species were shown to employ mixed Δ₄ and Δ₅ pathways in the production of testosterone.

Differing kinetic parameters and the absence of lyase activity in the opposite, (nonfavored), pathway in the dog, rabbit and guinea pig are evaluated as evidence for the existence of hydroxylase and lyase activities as four distinct proteins. Additionally, a soluble C₁₇-C₂₀ lyase, with dual nucleotide specificity, was identified in the dog. Regulation of testosterone biosynthesis is discussed in relation to hydroxylase and lyase activity in the four species.

Similar procedures were employed to evaluate the role of ascorbic acid in steroid synthesis. Guinea pigs with latent hypovitaminosis C were used as a model for evaluation of the effect of ascorbic acid on 3 BHSOR activity. The enzyme was inhibited 37-76% for the various Δ₅ substrates by the hypovitaminosis C pretreatment. Partial reversal of inhibition could be obtained by adding ascorbate or dehydroascorbate to in vitro incubations.
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ABBREVIATIONS

Lyase - C₁₇-C₂₀ Lyase

3BHSOR - 3B-Hydroxysteroid:NAD Oxidoreductase (EC 1.1.1.145)

17α-Hydroxylase - 17α-Steroid Hydroxylase:NADPH-oxygenoxygenoxidoreductase (EC 1.14.99.9)

17BHSOR - 17B-Hydroxysteroid NADH-Oxidoreductase (EC 1.1.1.64)

PROG - Progesterone; Pregn-4-ene-3,20-dione

PREG - Pregnenolone; 3B-Hydroxy pregn-5-ene-20-one

17αPROG - 17α-Hydroxyprogesterone; 17α-Hydroxypregn-4-ene-3,20-dione

17αPREG - 17α-Hydroxypregnenolone; 3Β, 17αDihydroxypregn-5-ene-20-one

DHEA - Dehydroepiandrosterone; 3B-Hydroxyandrost-5-ene-17-one

Α - Androstenedione; Androst-4-ene-3,17-dione

ADIOL - Androstenediol; Androst-5-ene-3,17-diol

T - Testosterone; 17α-Hydroxyandrost-4-ene-3-one

Cyanoketone - 2α-cyano 4,4,17α-trimethyl-androst-5-ene-17B-3-one

α - alpha

β - beta

Δ - delta (indicates unsaturation)

mg - milligram

µg - microgram

mm - millimeter

mM - millimolar

nm - nanomole or nanometer

nM - nanomolar
$k_{0.5}$ - Michaelis constant
$V_{\text{max}}$ - maximum velocity
min - minute
gm - gram
ml - milliliter
$P_{450}$ - cytochrome P-450
$b_5$ - cytochrome $b_5$
LH - Lutenizing hormone
ACTH - Adrenocorticotrophin hormone
GH - growth hormone
NAD - Nicotinamide adenine dinucleotide
NADH - reduced nicotinamide adenine dinucleotide
NADP - nicotinamide adenine dinucleotide phosphate
NADPH - reduced nicotinamide adenine dinucleotide phosphate
Tris - Tris 2(hydroxymethyl)2-aminomethane
TLC - thin layer chromatography
UV - ultraviolet
Rf - relative migration of steroid on TLC;

$$Rf = \frac{\text{distance steroid moved}}{\text{distance solvent front moved}}$$
PREFACE

The manuscript in this thesis was prepared according to the format of the Journal of Biological Chemistry.

All data presented in this thesis are the original work of the author, except where work by J. Hologittas (Ph.D. dissertation, University of Rhode Island, 1981) is cited and used for purposes of comparison.
I. KINETIC CHARACTERIZATION OF THE STEROIDOGENIC PATHWAYS IN GUINEA PIG, RABBIT, DOG AND RAT TESTIS MICROSONMES
INTRODUCTION

The leydig cell of the testis is the principal site of androgen biosynthesis in the male of dimorphic species. The major androgens produced by the testes, androstendione, dehydroepiandrosterone, testosterone and dihydrotestosterone are synthesized from cholesterol (1,2). In all steroidogenic tissues, the initial step is the sidechain cleavage of cholesterol to form pregnenolone (3-6). This step is considered to be rate limiting in androgen biosynthesis and is modulated by the pituitary gonadotrophin, LH. The mitochondrial sidechain cleavage system is common to all steroid producing tissues and involves two cytochrome P-450 dependent hydroxylations of cholesterol at C20 and C22 followed by cleavage between C20 and C22. The hydroxylations are catalyzed by mixed function oxidases in a reaction requiring NADPH, molecular oxygen and cytochrome P-450. 20,22 dihydroxycholesterol is cleaved by a desmolase to yield C-21 pregnenolone and isocaproic acid.

Subsequent metabolism of pregnenolone to testosterone appears to require translocation of pregnenolone out of the mitochondrion and into the microsomal fraction of the leydig cell. However, the enzyme which converts pregnenolone to progesterone, 3β-Hydroxysteroid oxidoreductase, has been found in the mitochondria in rat adrenal preparations (7),
and in rat testis (8,9). In some species, it is possible that testosterone production may proceed from progesterone of mitochondrial origin, although it is generally presumed that the biosynthesis of testosterone proceeds from pregnenolone via steroidogenic enzymes associated with the smooth endoplasmic reticulum.

Biosynthesis of testosterone from pregnenolone may occur by either of two major pathways, the $\Delta_4$ pathway or the $\Delta_5$ pathway (Figure 1). The $\Delta_5$ pathway proceeds from prenenolone, a $\Delta_5$-6 ene-3β-hydroxy steroid through 17α-hydroxypregnenolone, dehydroepiandrosterone and androstendiol to testosterone. The $\Delta_4$ pathway proceeds from pregnenolone to progesterone, a $\Delta_4$-5 ene-3-keto steroid and, subsequently, through 17α-hydroxy progesterone and androstendione to testosterone. It has been reported that more than one pathway is utilized by the rabbit and the rat (10). These species demonstrate the use of mixed pathways which may proceed from PREG +17PREG + DHEA + A + T or from PREG + 17PREG + DHEA + $\Delta_5$DIOL + T, or PREG + 17PREG + 17PROG A + T. Since differences in testosterone production among species known to exist, it is possible that these differences may be accounted for by preferential use of the $\Delta_4$, $\Delta_5$ or mixed pathways.

Microsomal biosynthesis of testosterone from pregnenolone involves at least five enzymatic activities:
1) oxidation of the C3 hydroxyl group to a ketone,
2) isomerization of the $\Delta_5$-6 double bond to the $\Delta_4$-5 position,
3) 17α-hydroxylation,
4) side chain cleavage between C17 + C19 and
5) reduction of the C17 ketone. The number of enzymes involved in catalysis of these reactions is controversial. Several possibilities are apparent. First, 3-BHSOR, which catalyzes the conversion of Δ5-hydroxysteroids to Δ5 ketone intermediates may exist as a single protein which accepts PREG, 17PREG, DHEA and ADIOL as substrate, or, multiple forms of the enzyme may exist, each with a specific substrate. Four isoenzymes of 3BHSOR have been reported in Pseudomonas testeroni (11) but not much evidence exists for mammalian forms. Evidence for a single 3BHSOR has been reported in the human adrenal (12), sheep adrenal cortex (13) and human testis (14). Secondly, the lyase and hydroxylase activities may exist as:

1) Four separate proteins, PREG lyase, PRO lyase, PREG 17α-Hydroxylase, PROG 17α-Hydroxylase
2) two proteins, one being specific for PREG, the other for PROG, or, 3) one protein with both lyase and hydroxylase activities (with either one or two active sites). Lastly, testis 17-B Hydroxysteroid oxidoreductase is believed to exist as at least two proteins, one which is product stimulated and one which is insensitive to T (15), or, possibly, one for the forward reaction A + T and a second enzyme catalyzing the oxidation of T + A (16). However, pig testis 17BHSOR was found to purify as a homogeneous enzyme (17).
It is necessary to determine the number of enzymes involved on steroid biosynthesis in order to predict the direction of the pathways. As suggested by Nakajin et al. (18), a single 17αHydroxylase and lyase with one active site would not readily release the 17αhydroxy product which is to serve as substrate for the lyase. This would, therefore, limit the use of mixed pathways. It is the purpose of this study to kinetically examine the steroidogenic pathways in two species, the rabbit and guinea pig and compare these data with kinetic data from the rat and dog to develop a kinetic model with which to predict the predominant pathway.
METHODS AND MATERIALS

**Experimental Animals**

New Zealand white male rabbits (age 4-6 months) were obtained from a local source and used immediately. Male Hartley guinea pigs were purchased from Charles River Laboratories, Wilmington, Massachusetts and maintained on rabbit chow (Agway, Kingston, Rhode Island), with an oral supplement of 10 mg. ascorbic acid per day. Water and feed were supplied ad libitum. Guinea pigs were purchased at 10 days of age, (150-200 grams), and raised until they reached 42 days of age, (sexually immature), or to 100 days of age, (700-850 grams). Dog testes for the $C_{17}^-C_{20}$ lyase experiment were obtained from a local animal hospital. The donor was a one-year-old retriever type dog who had been brought to the hospital for orchiectomy.

**Preparation of Microsomes**

Rabbits were sacrificed by injection of a 15cc air embolus into the ear vein. Guinea pigs were sacrificed by dislocation of the cervical vertebrae. Testes were quickly removed, placed on ice in .25M sucrose, decapsulated and weighed. Dog testes were obtained under general anesthesia, placed in .25M sucrose, kept on ice and transported to the laboratory within 1 hour of removal. Dog testes were decapsulated, weighed and minced in .25M sucrose, over ice, before homogenization. Tissues were homogenized in (10% w/v) ice cold .25M sucrose using a motor-driven teflon pestle glass homogenizer.
Microsomes were prepared by centrifugation at 1058Xg for 10 minutes, the supernatant transferred and centrifuged 17,300Xg for 10 minutes and the resultant supernatant centrifuged 105,000Xg for 60 minutes. The microsomal pellet was resuspended in .25M sucrose using a hand-held glass homogenizer with a smooth teflon pestle. Protein was determined by the biuret method of Gornall (19).

Preparation of Radioactive Substrates

Pregnenolone-4-14C, (S.A. 55 mci/mmole), 17α-hydroxypregnenolone-7-3H(N), (S.A. 10 Ci/m mole, and dehydroepiandrosterone-4-14C, (S.A. 55 mci/mmole), were obtained from Amesham Corporation, Arlington Heights, Illinois. Progesterone-4-14C, (S.A. 46 mci/mmole), 17α-hydroxyprogesterone-4-14C, (S.A. 50 mci/mmole) and androstenediol-1,2-3H(N), (S.A. 45 Ci/mmol) were obtained from New England Nuclear, Boston, Massachusetts. Radiochemical purity of isotopes was confirmed by thin layer chromatography.

Non-radioactive steroids were obtained from Steraloids, Wilton, New Hampshire and Sigma Chemical Company, St. Louis, Missouri. Radioactive substrates for enzyme assays were prepared by diluting with a sufficient amount of non-radioactive steroid to achieve specific activities of: 4.44X10^7 dpm/µmole (C14-pregnenolone), 1.08X10^9 dpm/µmole (DHEA), 4.28X10^8 dpm/µmole (androstenediol), 4.06X10^7 dpm/µmole (progesterone) and 3.52X10^7 dpm/mole (17α-hydroxyprogesterone).
**Enzyme Incubations**

Microsomal protein (60-250 µg) was incubated with radioactive substrate (1-40 µM) in 1% propylene glycol, 500 µM NAD, (Sigma Chemical Co., St. Louis, Missouri), or, NADPH, (Schwartz-Mann, Orange, New Jersey), as indicated, .1M Tris, pH 7.4 and 5mM MgCl₂ in a final volume of .1 ml. Δ₅ lyase and hydroxylase reactions were measured in the presence of 5 µg cyanoketone (Sterling-Winthrop Research Institute, Rensselaer, New York), to prevent conversion to Δ₄ products. Protein was added to initiate the reaction. Tubes were incubated 37°C in a reciprocating water bath for 5 minutes, (10 minutes for lyase and hydroxylase reactions, where indicated). The microassay was designed so that the amount of product produced remained well below the Kₐ₅ for the subsequent reaction. In this manner, a single reaction product was measured in most assays.

Enzyme reactions were terminated with .3-.4 ml cold methanol containing appropriate carrier steroids for visualization after thin layer chromatography (50 µg/.4 ml of Δ₄ 3-keto-steroids and 200 µg/.4 ml of Δ₅ 3-hydroxysteroids). Tubes were covered immediately and kept frozen (-5 to -15°C) until separation by TLC.

**Separation and Identification of Steroids**

One half the volume from each incubation tube was placed on a 5x20cm silica gel HLF plate (Analtech, Inc., Newark, Delaware). Each plate was developed in acetone three times,
drying thoroughly between each application, to a point 1.5 cm above the origin. The product and substrate were separated in jar-type TLC tanks lined with Whatman #1 chromatography paper to promote saturation. Elutropic solvents (20) were used for separation as follows:

1. PROG/17αPROG - dichloromethane:butylacetate 9:1 (2 developments)
2. PREG/17αPREG - benzene:acetone 4:1
3. PREG/PROG - chloroform:ethyl acetate 3:1
4. 17αPREG/17αPROG - benzene:acetone 4:1
5. DHEA/A - benzene:acetone 4:1
6. ADIOL/T/A - benzene:acetone 4:1
7. 17αPREG/DHEA - (1)dichloromethane:ethylacetate 9:1 (2)chloroform:diethylether 4:1
8. 17αPROG/A - (1)dichloromethane:ethylacetate 9:1 (2)chloroform:diethylether 4:1

All solvents were chromatographic grade or better. Separated Δ5-3 hydroxysteroids were detected by exposure to iodine vapor and Δ4-3 ketosteroids were detected by UV fluorescence (254 nm). Steroids were identified by Rf value and verified by standards run in the same solvent system.

**Identification of Product**

The product band was scraped from the TLC plate and the silica gel containing radioactive steroid was added directly to a liquid scintillation vial (New England Nuclear, Boston, Mass.), to which 7 ml Bray's Scintillation Cocktail (New England Nuclear, Boston, Mass.) was added. Radioactivity was counted in a Packard TRI-CARB Liquid Scintillation Spectrometer
(Packard, Downer's Grove, Illinois). Efficiency of counting was determined by using the External Standards Ratio method and found to be 48-50% for $^3$H and 83-84% for $^{14}$C.

**Cytochrome P-450 Determination**

Cytochrome P-450 was measured in testis microsomes using a Cary 15 split beam spectrophotometer, (Applied Physics Corp., Monrovia, Calif.), modified with a high intensity light source as described by Menard and Purvis (21). Incubations consisted of 1-4 mg/ml microsomal protein in 0.1M Tris, pH 7.4. Difference spectra were obtained by measuring absorbance of dithionite/CO minus CO/CO at 450nm. The extinction coefficient for changes in absorbance between 450-490 nm is assumed to be $100 \text{ cm}^{-1} \text{ nm}^{-1}$.

**DISCUSSION OF METHODS**

The enzyme assays were designed to measure one product.

1) 3BHSOR reactions were inhibited by cyanoketone for measurement of only $\Delta_5$ product in preg-hydroxylase and lyase reactions.

2) Absence of appropriate cofactor, (NADPH) prevented synthesis of hydroxylated product in PREG $\rightarrow$ PROG reactions.

3) The $\Delta_4$ and $\Delta_5$ hydroxylations were run under conditions such that the product was produced at a rate much below the $k_{0.5}$ for the lyase reactions. While it is known that internally generated substrate may be better utilized than substrate which is added externally, small amounts of product generated in this system did not act as substrate for the lyase reactions.
the lyase reactions. Only two radioactive bands could be detected in the hydroxylase assays corresponding to product and substrate. This may be attributable to the strong inhibition of the lyase by PROG which has been demonstrated by Purvis et al. ⁹

4) The Δ₄ lyase reaction has been shown to produce only androstendione and the Δ₅ lyase consistently produces only DHEA using this assay system. No radiolabeled 17-hydroxy-steroid product bands appear when cold T and ADIOL are added as carrier steroids at the termination of these reactions.

5) The only case in which more than one product is formed from a reaction is the ADIOL + T reaction in which both ADIONE and T are measured to quantitate the total product of the 3BHSOR reaction. Under the assay conditions the production of ADIONE from the product T, (17B-hydroxy-steroidoxidoreductase reaction), remains small.
RESULTS

MICROSOMAL ENZYME ACTIVITY

Kinetic data from rabbit and guinea pig testis microsomes is summarized in TABLE 1. Michaelis constants ($k_{0.5}$) and $V_{\text{max}}$ were estimated by the graphic method of Lineweaver-Burk (22). The $k_{0.5}$ for PREG in the 3BHSOR reaction was found to be three times higher in the rabbit than in the guinea pig (Figure 2). However, $k_{0.5}$ for 17αPREG and DHEA (Figures 3-4) were lower in the rabbit than in the guinea pig. This relative change in affinity for Δ5 substrates is attributable to the decreased $k_{0.5}$ in the rabbit and increase in the $k_{0.5}$ in the guinea pig for 17αPREG + DHEA with respect to $k_{0.5}$ PREG. The $k_{0.5}$ for ADIOL was identical in the rabbit and guinea pig (Figure 5). The $V_{\text{max}}$ of the PREG + PROG reaction was six times greater in the guinea pig than in the rabbit (Figure 2) and similar in the 17αPREG + 17αPROG reaction (Figure 2). The $V_{\text{max}}$ in the guinea pig DHEA → A and ADIOL → T reactions were nearly identical (Figure 4-5) and these velocities were 50% of those obtained with PREG as substrate. The highest 3BHSOR activity in the sexually mature rabbit was obtained with ADIOL (Figure 5), although data from TABLE I would suggest that the 17αPREG: 3BHSOR reaction is the most active. Testis microsomes for the 17αPREG + 17αPROG reaction were obtained from a 4 month old rabbit. Prior to sexual maturation (6 months of age in the rabbit), the amount of interstitial, (Leydig cells), enzyme
is higher on a milligram protein basis due to the absence of fully developed tubules. When the apparent $V_{\text{max}}$ of the $17\alpha\text{PREG} \rightarrow 17\alpha\text{PROG}$ reaction is corrected per nm P-450, the $V_{\text{max}}$ is lower (TABLE III) and the ADIOL $\rightarrow$ T reaction is observed to be the most active.

The ADIOL $\rightarrow$ T reaction is linear in the guinea pig at low protein concentrations (Figure 6). High protein concentrations were necessary to saturate the enzyme before kinetic constants could be determined.

Large differences in the Michaelis constants of the $17\alpha$-hydroxylase reaction for PREG were noted between the rabbit and guinea pig (Figure 7). Although the $k_{0.5}$ for the $\Delta_5$ substrate was greater than the $k_{0.5}$ for $\Delta_4$ progesterone in both instances, the ratio of $k_{0.5}$ PREG/PROG was found to be 20 in the guinea pig as compared to 3.3 in the rabbit. The $V_{\text{max}}$ for the $17$-hydroxylase reaction with $\Delta_5$ and $\Delta_4$ substrates were similar in the rabbit (Figures 7-8), whereas, the velocity with progesterone was 9.6 times greater than with pregnenolone in the guinea pig.

$C_{17-20}$ lyase activity could not be detected in the guinea pig with $17\alpha$PREG as substrate nor in the rabbit with $17\alpha$PROG as substrate. PROG:lyase in the guinea pig was found to have the lowest $k_{0.5}$ of any of the enzymes studied (Figure 9). The $V_{\text{max}}$ in the $17\alpha\text{PROG} \rightarrow A$ reaction in the guinea pig was 25% of the velocity of the preceding $\Delta_4$-hydroxylase reaction. In the rabbit the Michaelis constant of the PREG-lyase reaction (Figure 10) was lower than that of the PREG-
hydroxylase and the $V_{\text{max}}$ values were similar.

**CYTOCHROME P-450 DETERMINATIONS**

TABLE II summarizes cytochrome P-450 levels of testis microsomes. Cytochrome P-450 levels have been shown to reflect the hydroxylase and lyase activities of the microsomes (23). Since the Leydig cell volume and % Leydig cells varies so greatly among species, it is difficult to compare enzymatic activity on a milligram protein basis. Therefore, cytochrome P-450 values have been used as a constant upon which to express relative enzyme activities.

**COMPARISON OF MICROSOMAL ENZYME ACTIVITY IN FOUR SPECIES**

Enzyme activities for four species were expressed as nm product/min/nm P-450. TABLE III summarizes the kinetic data for 3BHSOR activities on the rabbit, guinea pig, dog (24) and rat (24) testis microsomes. The $k_{0.5}$ for PREG in the dog is twenty times higher than the guinea pig enzyme. The rabbit and rat have similar $k_{0.5}$ for PREG and these values are intermediate with respect to the dog (7 to 8 times lower), and guinea pig, (2.5 times higher). The $V_{\text{max}}$ for the PREG + PROG reaction, on a P-450 basis, is the greatest in the rat and lowest in the dog. The $V_{\text{max}}$ for this reaction in the rabbit and guinea pig are intermediate, with the guinea pig value determined to be twice as great as the rabbit and 33% of the rat activity. Michaelis constants of the $17\alpha\text{PREG} + 17\alpha\text{PROG}$ reaction are similar in the rat, rabbit and guinea pig and
high in the dog. The $V_{\text{max}}$ of the $17\alpha\text{PREG} \rightarrow 17 \text{PROG}$ reaction parallel the $V_{\text{max}}$ of $\text{PREG} \rightarrow \text{PROG}$ in the rabbit, guinea pig and rat, although this 3BHSOR activity is the lowest demonstrated in the dog. It is interesting to note that where the $k_{0.5}$ for $17\alpha\text{PREG} + \text{DHEA}$ are decreased in the rabbit with respect to the $k_{0.5}$ for PREG, the values are increased in the guinea pig and remain fairly constant in the rat. The $k_{0.5}$ for DHEA in the dog is the lowest demonstrated for that species, although the $V_{\text{max}}$ of the reaction is low. The $V_{\text{max}}$ of the DHEA $\rightarrow A$ reaction is the highest of the 3BHSOR activities in the rat. The ADIOL $\rightarrow T$ reaction is the most unusual of the 3BHSOR activities. The highest $V_{\text{max}}$ for 3BHSOR in the rabbit and dog is obtained with ADIOL as substrate although the $k_{0.5}$ for ADIOL in both instances is higher than the $k_{0.5}$ for DHEA. The rat $k_{0.5}$ for ADIOL is the lowest of the $\Delta_5$ substrates and the $V_{\text{max}}$ is 23-30% of the values obtained with the other $\Delta_5$ substrates. The $k_{0.5}$ for ADIOL in the guinea pig is similar to $k_{0.5}$ for DHEA and $17\alpha\text{PREG}$ and the $V_{\text{max}}$ is the lowest demonstrated for any of the 3BHSOR substrates.

The kinetic data for the $17\alpha$hydroxylase and $\text{C}_{17}-\text{C}_{20}$ lyase reactions is summarized in TABLE IV. The $k_{0.5}$ for PREG in all species was greater than the $k_{0.5}$ for PROG in the hydroxylase reaction and the $V_{\text{max}}$ of the PROG $\rightarrow 17 \text{PROG}$ reaction was higher than for PREG $\rightarrow 17\alpha\text{PREG}$ in all but the dog. Each species demonstrated both PREG and PROG hydroxylase activity. The rat is the only species which possesses both PREG and PROG lyase activities. In the rabbit and dog microsomal $\Delta_4$ lyase
activity was not detected. In contrast, no Δ₅ lyase activity was detectable in the guinea pig.

Since PROG-lyase activity had been previously reported in the rabbit (25), an experiment was conducted to examine the possibility that a cytosolic Δ₄ lyase existed in dog and rabbit testes. The 105,000Xg supernatant of dog testis was found to have a small but measurable, amount of Δ₄ lyase activity. The enzyme demonstrated a dual specificity for NADH and NADPH although better activity was obtained with NADH. The Michaelis constant of the NADH-cytosolic C₁₇⁻C₂₀ lyase for PROG was 20μM and the $V_{\text{max}}$ of the reaction was determined to be 2 nm/min/nm P-450. No cytosolic Δ₄ lyase activity could be detected in the rabbit 105,000Xg supernatant. It is possible that the failure to detect this activity in the rabbit is attributable to the stage of sexual development, as the animal used in this experiment was four months of age. No microsomal Δ₄ lyase activity could be detected in the presence of 500μM NADH in either the rabbit or the dog.
DISCUSSIONS AND CONCLUSIONS

The synthesis of testosterone from pregnenolone has been shown to proceed by the $\Delta_4$ pathway (26,27,28) and the $\Delta_5$ pathway (29,30,31,32) in several mammalian species. Enzymatic activities of these pathways were determined and are presented in Figure 11. The kinetic data is a useful predictor of the $\Delta_4$ or $\Delta_5$ pathway in testosterone biosynthesis. The key step in determining the direction of the pathway is the conversion of PREG to PROG, ($\Delta_4$), or to 17$\alpha$PREG, ($\Delta_5$). The most definitive kinetic predictors of the $\Delta_4$ or $\Delta_5$ pathway were found to be 1) the Michaelis constant of the 3BHSOR reaction for PREG and, 2) the $V_{max}$ of the PREG-$17\alpha$hydroxylase reaction. The affinity of 3BHSOR for PREG, ($k_{0.5}=2.5 \text{ M}$), in the guinea pig as constrained to the high $k_{0.5}$, (50$\mu$M), in the dog, suggests predominant use of the $\Delta_4$ pathway by the guinea pig and $\Delta_5$ pathway in the dog. The Michaelis constants of the 3BHSOR reaction for PREG on the rat, (6$\mu$M), and rabbit (7$\mu$M) are intermediate and suggest the use of mixed pathways. This prediction is strengthened by the low $V_{max}$ of the PREG-$17\alpha$-hydroxylase reaction in the guinea pig and the extensive activity of this enzyme in the dog. The $V_{max}$ of the $\Delta_5$ 17$\alpha$-hydroxylase reactions in the rat and rabbit are, again, intermediate, suggestive of mixed pathways although favoring the $\Delta_4$ in the rat and $\Delta_5$ in the rabbit.
The ratio of the Michaelis constants for PREG in the 3BHSOR and 17α-hydroxylase reactions (TABLE V) provides further information with which to predict the predominant pathway under conditions of submaximal stimulation where PREG is unlikely to be saturating. In the dog, \(1/2 V_{\text{max}}\) for the hydroxylase reaction would be reached at substrate concentrations 50% lower than the \(k_{0.5}\) for the 3BHSOR reaction. Since the \(V_{\text{max}}\) of the hydroxylase reaction is 5 times greater, most of the PREG would be metabolized to 17αPREG. In the guinea pig, \(1/2 V_{\text{max}}\) will be reached for the 3BHSOR reaction at PREG concentrations which are 10% of the \(k_{0.5}\) for the hydroxylase and metabolism of PREG to PROG would occur at a rate at least 32 times greater than the conversion of PREG → 17 PREG. The Michaelis constants and \(V_{\text{max}}\) for PREG in the 3BHSOR and 17α-hydroxylase reactions are identical in the rabbit and it is probable that 50% of the substrate would be metabolized to PROG and 50% would be converted to 17αPREG. Although the Michaelis constants of the two reactions are similar in the rat, the \(V_{\text{max}}\) is 30 times greater for the 3BHSOR reaction and most of the PREG would be expected to be metabolized to PROG.

Since PROG is a known inhibitor of 3BHSOR activity, it becomes necessary to verify predictions by examining the rate of removal of PROG (TABLE VI). The Michaelis constants for the hydroxylase are of little use for this purpose as small variation on \(k_{0.5}\) for PROG was noted and since the \(k_{0.5}\) for PROG were lower than \(k_{0.5}\) for PREG in each species. The ratio of \(k_{0.5}\) PROG/PREG may be used to demonstrate the relative
affinity of the hydroxylase for $\Delta_4$ and $\Delta_5$ substrates in the guinea pig, rat and rabbit. This ratio is misleading in the dog because the $k_{0.5}$ for PREG is large for both the $\Delta_5$ hydroxylase and the 3BHSOR reactions. However, the $k_{0.5}$ for PROG in the hydroxylase reaction is consistent with data from Eik-Nes (33) demonstrating rapid conversion of PROG to $17\alpha$ PROG in dog testes and the appearance of $17\alpha$PROG as a secretion product in this species. The ratio of hydroxylase activity on a $V_{\text{max}}$ basis is a useful indicator of the relative disappearance of PROG via the $\Delta_4$ pathway and PREG via the $\Delta_5$ pathway. $\Delta_4$ 17-hydroxylase activity is 10 times greater than $\Delta_5$ hydroxylase activity in the guinea pig and 5 times higher in the rat. This comparison strengthens the prediction since 3BHSOR activity in $\Delta_4$ animals is dependent upon an adequate rate of removal of PROG. The ratio of $\Delta_4/\Delta_5$ hydroxylase activity in the dog shows the reaction velocity at saturation to be 8.3 times greater in the $\Delta_5$ pathway.

The kinetic data from the rabbit presents somewhat of a problem. Similar $k_{0.5}$ and $V_{\text{max}}$ values for PREG in the 3BHSOR and hydroxylase reactions and the ratio of $V_{\text{max}}$ PROG/PREG suggest that the rabbit may use either the $\Delta_4$ or the $\Delta_5$ pathway for testosterone production. This appears to be true for $17\alpha$PROG synthesis as the $k_{0.5}$ for the PREG + $17\alpha$PROG reaction is low in the rabbit and the $V_{\text{max}}$ is similar to the PREG-3BHSOR activity but the rabbit testis microsomes do not possess detectable C$_{17}$-C$_{20}$ lyase activity with PROG as substrate.
Therefore, the $\Delta_4$ pathway to T would be blocked in this species. The absence of $\Delta_4$ lyase activity in dog testis microsomes fully supports the prediction that the $\Delta_5$ pathway is the route to T production in the dog. The demonstration of a NADH-linked form of the enzyme is interesting although the high $k_{0.5}$ and low $V_{max}$ of this enzyme would result in the production of very little A for testosterone synthesis. The failure to detect $\Delta_5$ lyase activity in guinea pig testis microsomes may be presented as further evidence that the guinea pig uses the $\Delta_4$ pathway for testosterone synthesis.

These kinetic predictions regarding the direction of the steroidogenic pathway are in agreement with other in vitro data showing that the rat (27) and guinea pig (28) uses the $\Delta_4$ pathway and that the dog (30) uses the $\Delta_5$ pathway for testosterone synthesis. In vivo perfusion techniques developed by Chubb and Ewing (34) and using specific enzyme inhibitors to block multi-directional pathways (10) have demonstrated that the dog and rabbit secrete primarily $\Delta_5$ steroids and the guinea pig (35) and rat secrete mainly $\Delta_4$ steroids. In these studies, no $17\alpha$PREG was detectable in guinea pig testes secretions. This is in agreement with the present finding that the PREG-hydroxylase activity is very low in guinea pig testes microsomes.

An alternate pathway from PREG $\rightarrow 17\alpha$PREG $\rightarrow 17\alpha$PROG $\rightarrow A \rightarrow T$ was proposed for the rat based on secretion studies. This is compatible with the kinetic data for PREG-hydroxylase and $3\beta$HSD activity for PREG in rat testis microsomes. However,
one difference exists between the kinetic data and the perfusion studies. Chubb and Ewing (10) have demonstrated that the rabbit produces a small amount of androstendione in the presence of medrogestone, an inhibitor of 3BHSOR activity. One major difference between the kinetic and the perfusion studies is that the kinetic studies use microsomes from whole testis and perfusion studies use the whole testis. If the rabbit has a soluble lyase such as that seen in the dog, the difference could be explained.

Although the kinetic studies allow for the synthesis of androstendione from DHEA in the rabbit, the presence of inhibitor suggests that androstendione must have been produced from infused 17αPROG in the perfusion study and not from endogenous DHEA. The fact that the testes were under LH stimulation during the perfusion is of little consolation. Presumably, this short term stimulation with high doses of LH maximizes the flux by releasing inhibition on the cholesterol side chain cleavage enzymes. LH stimulation for 4.5 hours would not be expected to stimulate protein synthesis so the appearance of Δ₄ lyase activity cannot be explained on the basis of increased enzyme synthesis. Data from van der Molen et al. (25) suggests one possible explanation for the contrasting data. 17αPROG was shown to be converted to androstendiol in the 800-105,000 x g pellet of a rabbit testis homogenate suggesting that the C₁₇-C₂₀ lyase activity may be associated with the mitochondria which would obviously have been present in this pellet. The present findings can neither confirm nor reject this possibility.
The kinetic data for 17α-hydroxylase and lyase activity in testis microsomes suggests that the Δ5 and Δ4 hydroxylases and lyases are four distinct enzymes. In support of this hypothesis, is the 25 fold difference in Michaelis constants of the hydroxylase reaction for Δ5 and Δ4 substrates in guinea pig testis microsomes. All $k_{0.5}$ for the hydroxylase were shown to be greater for PREG, yet the dog enzyme demonstrated 6.7 times higher activity for the high $k_m$ form of the enzyme. The existence of more than one C17-C20 lyase activity in the testes has been demonstrated by the cytosolic NADH-activated enzyme in the dog. Differing $V_{\text{max}}$ values of the microsomal lyase with respect to PROG and PREG in the rat suggest that these activities are associated with different enzymes. The absence of C17-C20 lyase activity with Δ4 substrate in the dog and rabbit and nondetectable Δ5 lyase activity in the guinea pig should be considered as evidence that two lyases exist and the demonstration of Δ4 hydroxylase activity in the dog and rabbit and weak Δ5 hydroxylase activity in the guinea pig shows that the lyase activity is not necessarily associated with the same protein as the hydroxylase. However, a purified cytochrome P-450 which possesses both hydroxylase and lyase activity has been isolated from testis microsomes in the neonatal pig (36). This purified enzyme was shown to consist of one subunit by electrophoretic and immunochemical criteria. This data is not conclusive since membrane bound enzymes often appear to be homogeneous by these criteria due to the hydrophobic nature of the proteins which causes them to associate and migrate together.
in an electric field. Further evidence (10) was obtained for homogeneity of the neonatal pig cytochrome by temperature pH optimum, denaturation and inhibition studies. The enzyme was shown to possess one heme molecule which suggests one active site. Although the data is consistent with the one active site model of Hochberg et al. (37), this group's own data presents evidence against one active site for lyase and hydroxylase activities. First, the activity of the hydroxylase and lyase exist in a 1:8:1 stoichiometry. Second, the Michaelis constants for the hydroxylase and lyase activities are different, suggesting more than one binding site, and lastly, certain activators which stimulate the hydroxylase activity were unable to stimulate the lyase reaction. Other evidence against one active site was presented by Samuels and Matsumoto (38) who were able to demonstrate an exchange between exogenous 17αPROG and the enzyme bound intermediate generated from PROG, suggesting movement of 17αPROG from one site to another. The presence of one heme is also in contrast to finding in rat testis microsomes (39) that different cytochromes were involved in hydroxylase and lyase activities.

Table VII summarizes the ratio of lyase to hydroxylase activity in the rat, guinea pig, dog and rabbit. The ratios suggest independent synthesis of hydroxylase and lyase. It appears that these activities may exist as two subunits that associate to form a 1:1 dimer, a 2:1 dimer, or, as 2:1 dimers plus free hydroxylase. The activity of the free hydroxylase
is assumed to be higher as steric hinderance would exist in the dimeric form. This would easily account for the 1:1.1 stoichiometry demonstrated in the rabbit. Studies on postnatal development of microsomal enzyme activity in rat testes (40) also suggests independent synthesis of hydroxylase and lyase. Between 23-32 days of postnatal development, hydroxylase and lyase activities exist in a 1:1 ratio with maximum C_{17-20} lyase activity occurring at 32 days. At 32 days the ratio of the rate of synthesis of hydroxylase lyase becomes 2.2. In all probability, this increased rate of synthesis of the hydroxylase represents the transition from 1:1 stoichiometry to the 1:2 dimer. Additionally, the hydroxylase activity does not reach maximal until 40 days, demonstrating independent synthesis of this enzyme. Different rates of synthesis under stimulation by gonadotrophic hormones was demonstrated by Latif and Purvis (41). The rate of synthesis of hydroxylase/lyase in the hypophysectomized rat was 2.44 under stimulation by LH, FSH and GH.

The ratio of hydroxylase to lyase activities shows an excess amount of Δ_5 hydroxylase on the dog and an excess amount of Δ_4 hydroxylase in the rabbit and guinea pig. The activity of PROG hydroxylase on a milligram protein basis is fairly constant in the 4 species (.2nm/mg/min in the rat, .24nm/mg/min in the guinea pig, .159nm/mg/min in the rat and .12nm/mg/min in the dog). These values are in agreement with Menard, Stripp and Gillette (42), showing no more than a 2-fold difference in hydroxylase activity. It appears that the excess amount of
hydroxylase is involved in the control of testosterone production by the gonadotrophic hormones. It is well accepted that the rate limiting step in testis steroid synthesis, under low LH concentrations, is the side chain cleavage of cholesterol. Stimulation with high levels of LH for long periods of time is known to stimulate protein synthesis resulting in increased amounts of 17-hydroxylase and C\textsubscript{17}-C\textsubscript{20} lyase. The lyase has been shown to be more sensitive to low concentrations of LH than the hydroxylase (41) and under these conditions one could easily increase testosterone production by stimulating lyase synthesis in the presence of excess hydroxylase.

Whereas high LH levels stimulate the synthesis of the hydroxylase to a greater extent than the lyase (41), the hydroxylase is more susceptible to desensitization by high LH.

Others have suggested that the opposite lyase, the \Delta\textsubscript{5} lyase in the rat (43) and the \Delta\textsubscript{4} lyase in the human (44) is the rate limiting step in testosterone production. The kinetic analysis of the steroidogenic enzymes in the dog, rat, rabbit and guinea pig does not preclude this hypothesis. The human testis is thought to be under periodic LH stimulation. It is possible that under long-term, maximal, LH stimulation the opposite lyase and hydroxylase may be synthesized and, in this manner, release inhibition on testosterone synthesis by increasing the number of alternate pathways.
References


Figure 1. Steroidogenic pathways in mammalian testis microsomes.
Figure 2. Lineweaver-Burk replots of PREG:3BHSOR reactions in rabbit and guinea pig testis microsomes. Incubations run in presence of 0.1M Tris, pH 7.4, 5mM MgCl₂ and 500 µM NAD at 37°C. Protein-65 µg (guinea pig), 185 µg (rabbit). k₀.₅=7µM (rabbit), 2.5 µM (guinea pig). V₅₀=0.12nm/min/mg (rabbit), 0.77 nm/min/mg (guinea pig).
Figure 3. Lineweaver-Burk replots of 17αPREG:3BHSOR reactions in rabbit and guinea pig testis microsomes. Incubations run in presence of 0.1M Tris, pH 7.4, 5mM MgCl$_2$ and 500 µM NAD at 37°C. Protein-175 µg (rabbit), 161 µg (guinea pig). $k_{0.5}$=4 µM (rabbit), 6 µM (guinea pig). $V_{max}$=.79nm/min/mg (rabbit), .70nm/min/mg (guinea pig).
Figure 4. Lineweaver-Burk replots of DHEA:3BHSOR reactions in rabbit and guinea pig testis microsomes. Incubations run in presence of 1M Tris, pH 7.4, 5mM MgCl₂ and 500 µM NAD at 37°C. Protein-336µg (rabbit), 197 µg (guinea pig) kᵦ₅0=2,0µM (rabbit), 4,0 µM (guinea pig). Vₘₐₓ=k₀₅=2.0 µM (rabbit), 4.0 µM (guinea pig). Vₘₐₓ=.1 nm/min/mg (rabbit), .43nm/min/mg (guinea pig).
Figure 5. Lineweaver-Burk replots of ADIOL:3BHSOR reactions in rabbit and guinea pig testis microsomes. Incubations run in presence of .1M Tris, pH 7.4, 5mM MgCl₂ and 500 µM NAD at 37°C. Protein-336 µg (rabbit), 537 µg (guinea pig). 

k₀.₅=5 µM (rabbit), 5 µM (guinea pig).

Vₘₐₓ=.2 nm/min/mg (rabbit), .35nm/min/mg (guinea pig).
Figure 6: Michaelis-Menten equation graph for guinea pig and rabbit.
Figure 6. Michaelis-Menten plot of ADIOL → T reaction in guinea pig testis microsomes. Incubation run in the presence of 0.1M Tris, pH 7.4, 5mM MgCl₂, 500 µM NAD at 37°C. Protein concentration—161 µg/1 ml final volume.
Figure 7. Lineweaver-Burk replots of PREG:17αHydroxylation reactions in rabbit and guinea pig testis microsomes. Incubations run in presence of .1M Tris, pH 7.4, 5mM MgCl₂ and 500 µM NADPH at 37°C. Protein - 185 µg (rabbit), 197 µg (guinea pig) k₀.₅=4.6 µM (rabbit), 33µM (guinea pig). V₅₅₉= .11 nm/min/mg (rabbit), .028 nm/min/mg (guinea pig).
Figure 8. Lineweaver-Burk replots of PROG:17α-Hydroxylase reactions on rabbit and guinea pig testis microsomes. Incubations run in presence of 1M Tris, pH 7.4, 5mM MgCl₂ and 500 µM NADPH at 37°C, Protein-231 µg (rabbit), 182 µg (guinea pig), k₀.₅=2 µM (rabbit), 1.5 µM (guinea pig), Vₘₐₓ=0.16 nm/min/mg (rabbit), 0.20 nm/min/mg (guinea pig).
Figure 9. Lineweaver-Burk replot of C_{17}-C_{20} lyase: (17α-hydroxyprogrenolone dehydroepiandrosterone) in rabbit testis microsomes. Incubations were run in the presence of .1M Tris, pH 7.4, 5mM MgCl_{2}, 500 µM NADPH at 37°C. Protein-374 µg/.1 ml final volume. k_{0.5}=4µM, V_{max}=.12 nm/min/mg.
Figure 10. Lineweaver-Burk replot of C_{17-20} lyase: (17α-hydroxy progesterone + androstendione) in guinea pig testis microsomes. Incubations were run in the presence of 0.1M Tris, pH 7.4, 5mM MgCl₂, 500 µM NADPH at 37°C. Protein-101 µg/.1 ml final volume. k_{0.5}=.63µM. V_{max}=.06 nm/min/mg.
Figure 11. Comparison of steroidogenic pathways for testosterone biosynthesis in rabbit, guinea pig, dog and rat testis microsomes. Michaelis constants ($k_{0.5}$) are indicated in parenthesis on outer perimeter and $V_{max}$ (nm product/min/nm P-450) are indicated inside. Most likely steroidogenic pathways are indicated by darkened arrows. Blocks in pathways are indicated by slashed arrows.
SUMMARY OF KINETIC DATA IN FOUR SPECIES

**RABBIT**

\[
\begin{align*}
PREG & \quad \xrightarrow{(7.0)} \quad 4 \quad 14 \\
17\alpha\text{PREG} & \quad \xrightarrow{(4.0)} \quad 4.3 \quad 3.8 \\
DHEA & \quad \xrightarrow{(2.0)} \quad 3.3 \quad \xrightarrow{(5)} \quad 6.3 \\
\end{align*}
\]

**GUINEA PIG**

\[
\begin{align*}
PREG & \quad \xrightarrow{(2.5)} \quad 17\alpha\text{PROG} \\
17\alpha\text{PREG} & \quad \xrightarrow{(6)} \quad 4.3 \quad 0 \\
DHEA & \quad \xrightarrow{(4.0)} \quad 4.3 \quad \xrightarrow{(5)} \quad 3.5 \\
\end{align*}
\]

**DOG**

\[
\begin{align*}
PREG & \quad \xrightarrow{(50)^a} \quad 1.7 \\
17\alpha\text{PREG} & \quad \xrightarrow{(47)} \quad 0.35 \quad 1.1 \\
DHEA & \quad \xrightarrow{(2.5)^a} \quad 0.5 \quad \xrightarrow{(13)^a} \quad 5.8 \\
\end{align*}
\]

**RAT**

\[
\begin{align*}
PREG & \quad \xrightarrow{(6)^a} \quad 24.0 \\
17\alpha\text{PREG} & \quad \xrightarrow{(8)} \quad 20.0 \quad 1.7 \\
DHEA & \quad \xrightarrow{(6)^a} \quad 26.0 \quad \xrightarrow{(0.8)^a} \quad 6.0 \\
\end{align*}
\]

\[a\text{Data courtesy of Joanne V. Hologittas, Dept. of Biochemistry and Biophysics, University of Rhode Island.}\]
TABLE I
COMPARISON OF KINETIC DATA IN RABBIT AND GUINEA PIG TESTIS MICROSONES

<table>
<thead>
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<th>Rabbit</th>
<th>Guinea Pig</th>
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<tr>
<td></td>
<td>$k_{0.5}^a$</td>
<td>$v_{max}^b$</td>
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<tr>
<td><strong>3β-Hydroxysteroid Oxidoreductase</strong></td>
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<tr>
<td>17αPREG + 17αPROG</td>
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<td>.79*</td>
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<td>DHEA + A</td>
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<td>PROG → 17αPROG</td>
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<td><strong>C_{17}-C_{20} Lyase</strong></td>
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<td>17αPROG → A</td>
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</table>

a - expressed as $\mu$M substrate

b - expressed as nm/min/mg microsomal protein

* - immature animal
### TABLE II

**CYTOCHROME P-450 LEVELS OF TESTIS MICROSOMES IN FOUR SPECIES**

<table>
<thead>
<tr>
<th>Species</th>
<th>P-450 (nm/mg microsomal protein)</th>
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<tr>
<td>RABBIT (immature)</td>
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<tr>
<td>DOG(^a)</td>
<td>.12(^b)</td>
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<tr>
<td>RAT(^a)</td>
<td>.05</td>
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</tbody>
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\(^a\) - Data - courtesy of Joanne V. Hologittas, Department of Biochemistry & Biophysics, University of Rhode Island, Kingston.

\(^b\) - Average value, mixed species, long-legged dogs.
### TABLE III

TESTIS MICROSONAL

3B-HYDROXYSTEROID OXIDOREDUCTASE ACTIVITY* in four species: *k*<sub>0.5</sub> and *V*<sub>max</sub>

<table>
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<th>Substrate</th>
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<th>Dog</th>
<th>Rat</th>
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<td><em>V</em>&lt;sub&gt;max&lt;/sub&gt;</td>
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<td><em>V</em>&lt;sub&gt;max&lt;/sub&gt;</td>
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<td>4.3</td>
<td>6.0</td>
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<td>ADIOL</td>
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<td>6.3</td>
<td>5.0</td>
<td>3.5</td>
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- enzyme assays were conducted at pH 7.4, 37°C in the presence of 1M Tris, 5mM MgCl<sub>2</sub>, 500µM NAD, 1% propylene glycol.
- data courtesy of Joanne V. Hoboggitas, Dept. of Biochemistry and Biophysics, University of Rhode Island, Kingston.
- *k*<sub>0.5</sub> - expressed as µM concentration substrate.
- *V*<sub>max</sub> - expressed as nm product min/nm P-450.
<table>
<thead>
<tr>
<th></th>
<th>RABBIT</th>
<th></th>
<th>GUINEA PIG</th>
<th></th>
<th>DOG</th>
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<td>$V_{max}$</td>
<td>$k_{0.5}$</td>
<td>$V_{max}$</td>
<td>$k_{0.5}$</td>
<td>$V_{max}$</td>
<td>$k_{0.5}$</td>
<td>$V_{max}$</td>
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<tr>
<td><strong>17 Hydroxylase</strong></td>
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<tr>
<td>PREG</td>
<td>7.0</td>
<td>4.0</td>
<td>33.0</td>
<td>0.28</td>
<td>25.0</td>
<td>6.7</td>
<td>10.0</td>
<td>0.8</td>
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<tr>
<td>PROG</td>
<td>2.1</td>
<td>5.3</td>
<td>1.5</td>
<td>2.0</td>
<td>3.0$^d$</td>
<td>1.6$^d$</td>
<td>2.0$^d$</td>
<td>4.0$^d$</td>
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<tr>
<td><strong>C_{17-C20} Lyase</strong></td>
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<td>$17\alpha$PREG</td>
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<td>3.8</td>
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<td>4.0</td>
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<td>$17\alpha$PROG</td>
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<td>0</td>
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<td>0.63</td>
<td>0</td>
<td>0</td>
<td>3.0$^d$</td>
<td>1.7$^d$</td>
</tr>
</tbody>
</table>

a - enzyme assays were conducted at pH 7.4, 37°C in the presence of .1M Tris, 5mM MgCl$_2$, 500µM NAD, 1% propylene glycol.

b - $k_{0.5}$ - expressed as µM concentration of substrate.

c - $V_{max}$ - expressed as nm product/min/nm P-450.

d - data courtesy of Joanne V. Holloggitas, Dept. of Biochemistry and Biophysics, University of Rhode Island, Kingston.
TABLE V

MICHAELIS CONSTANTS ($k_{0.5}$)\textsuperscript{a} FOR PREGnenolone

IN 3BHSOR AND HYDROXYLASE REACTIONS

<table>
<thead>
<tr>
<th></th>
<th>3BHSOR</th>
<th>HYDROXYLASE</th>
<th>3BHSOR/H</th>
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</thead>
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<tr>
<td>GUINEA PIG</td>
<td>2.5</td>
<td>25.0</td>
<td>0.1</td>
</tr>
<tr>
<td>RAT</td>
<td>6.0</td>
<td>10.0</td>
<td>0.6</td>
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<tr>
<td>RABBIT</td>
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<tr>
<td>DOG</td>
<td>50.0</td>
<td>25.0</td>
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</tr>
</tbody>
</table>

\textsuperscript{a} - $k_{0.5}$ expressed in µM PREG.
### TABLE VI

RELATIONSHIP OF 17 HYDROXYLASE$^{a,b}$ ACTIVITIES IN FOUR SPECIES

<table>
<thead>
<tr>
<th>Species</th>
<th>$k_{0.5}^{\text{a}}$</th>
<th>$v_{\text{max}}^{\text{b}}$</th>
<th>$k_{0.5}$ Ratio</th>
<th>$v_{\text{max}}$ Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Guinea Pig</td>
<td>1.5</td>
<td>25.0</td>
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<tr>
<td>Rat</td>
<td>2.0</td>
<td>10.0</td>
<td>4.0</td>
<td>0.8</td>
</tr>
<tr>
<td>Rabbit</td>
<td>2.1</td>
<td>7.0</td>
<td>5.3</td>
<td>4.0</td>
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<tr>
<td>Dog</td>
<td>3.0</td>
<td>25.0</td>
<td>1.0</td>
<td>6.7</td>
</tr>
</tbody>
</table>

$^a$ $k_{0.5}$, expressed as $\mu$M substrate.

$^b$ $v_{\text{max}}$, expressed as $\mu$M product/min/nm P-450.
TABLE VII
RATIO OF LYASE/HYDROXYLASE ACTIVITY\textsuperscript{a}
FOR PREGNENOLONE AND PROGESTERONE
IN FOUR SPECIES

<table>
<thead>
<tr>
<th></th>
<th>PROG</th>
<th>PREG</th>
</tr>
</thead>
<tbody>
<tr>
<td>RAT</td>
<td>1:2.3</td>
<td>1:1.1</td>
</tr>
<tr>
<td>GUINEA PIG</td>
<td>1:3.2</td>
<td>0</td>
</tr>
<tr>
<td>DOG</td>
<td>0</td>
<td>1:6</td>
</tr>
<tr>
<td>RABBIT</td>
<td>0</td>
<td>1:1.1</td>
</tr>
</tbody>
</table>

\textsuperscript{a} on the basis of $V_{\text{max}}$, expressed as nm prod/min/nm P-450.
II. THE EFFECT OF ASCORBIC ACID ON 3-B HYDROXYSTEROID: NAD-OXIDOREDUCTASE ACTIVITY IN THE TESTIS

Efforts to elucidate the role of ascorbic acid in steroid metabolism have failed to reveal a clear-cut effect on adrenal hydroxylations in the microsomal (1, 2, 3) and mitochondrial (4, 5, 6) fractions of steroid producing tissues, suggestive of a "cooperating" role in the control of steroid synthesis. Slight stimulation of adrenal steroid synthesis in guinea pigs during the early stages of scurvy (9) and decreased (8) levels in the testes of scurvy guinea pigs (4) suggest release of inhibition on the cholesterol side chain cleavage enzymes as ascorbic acid levels are lowered.

Other studies have demonstrated a stimulatory effect of ascorbic acid on the conversion of pregnenolone to progesterone (10). 3-B hydroxy-5-androstan-17-one. NAD-oxidoreductase (EC 1.1.1.34) activity was found to be inhibited in vitamin A deficient rats (11). The vitamin A deficient rat is unable to synthesize ascorbic acid and addition of ascorbic acid to the diet promotes restoration of NAD activity.

The mechanism by which ascorbic acid stimulates adrenal activity is uncertain. It has been proposed that the vitamin may release NADH inhibition toward a steroid-3-hydroxysteroid-dehydrogenase oxidation of the reduced pyridine nucleotide (12, 13, 14). Two distinct enzymes have been reported to be involved in the transport of electrons from NADH to 3-hydroxy-5-
INTRODUCTION

Efforts to elucidate the role of ascorbic acid in steroidogenesis have been both controversial and contradictory. Ascorbic acid has been shown to exert an inhibitory effect on steroid hydroxylations in the mitochondria (1,2,3) and microsomes (4,5,6,7) of steroid producing tissues, suggestive of a "braking" role in the control of steroid synthesis. Slight stimulation of adrenal steroid synthesis in guinea pigs during the early stages of scurvy (8) and decreased cholesterol levels in the testes of scorbutic guinea pigs (9) suggest release of inhibition on the cholesterol side chain cleavage enzymes as ascorbic acid levels are lowered.

Other studies have demonstrated a stimulatory effect of ascorbic acid on the conversion of pregnenolone to progesterone (10). 3B-hydroxysteroid:NAD-oxidoreductase (EC 1.1.1.145) activity was found to be inhibited in vitamin A deficient rats (11). The vitamin A deficient rat is unable to synthesize ascorbic acid and addition of ascorbic acid to the diet promotes restoration of 3BHSOR activity.

The mechanism by which ascorbic acid stimulates 3BHSOR activity is uncertain. It has been proposed that the vitamin may release NADH inhibition through a monodehydroascorbate-dependent oxidation of the reduced pyridine nucleotide (12, 13,14). Two distinct enzymes have been reported to be involved in the transport of electrons from NADH to monodehydro-
ascorbate (15). Ninety percent of the electron transport from NADH to monodehydroascorbate in microsomes goes via NADH-monodehydroascorbate oxidoreductase, (EC 1.6.5.4), and the remaining 10% through cytochrome b5 via ascorbate:ferricytochrome b5 oxidoreductase (EC 1.10.2.1). The physiological function of these proteins is unclear. It is not known whether they operate exclusively as a conservation mechanism for ascorbic acid by reducing free radical formation, or, function to generate oxidized cofactor for steroid synthesis.

This mechanism cannot totally explain the stimulation of 3BHSOR by ascorbic acid. Staudinger (13) demonstrated that the oxidation of NADH was strictly specific for reduced ascorbic acid. However, dehydroascorbic acid has been shown to stimulate 3BHSOR activity in toad testes (16). Depressed NAD/NADH ratios were reported in the tissues of ascorbic acid deficient guinea pigs although no inhibition of the ascorbate linked oxidase was apparent (17).

This study was undertaken to examine the effect of ascorbic acid on 3BHSOR in the testis using the guinea pig with latent hypovitaminosis C as a model. Microsomal 3BHSOR activity was examined in deficient and control animals with four known substrates for the enzyme using a sensitive microassay.
METHODS AND MATERIALS

Experimental Animals

Twelve, all male, Hartley guinea pigs were obtained from Charles River Laboratories, Wilmington, Massachusetts. Animals were purchased at 10 days of age (150-200 grams) and were maintained on rabbit chow (Agway, Kingston, Rhode Island) and supplemented with 10 mg L-ascorbic acid/day, administered by oral intubation. Feed and water were supplied ad libitum and animals were raised for 4 weeks prior to commencement of feeding experiment.

Feeding Experiment

Animals were divided into two groups at 35 days of age. Mean weights were 360 gm (Group I) and 376 gm (Group II). Group I served as the control and received rabbit chow plus 10 mg L-ascorbic acid (ICN Biochemicals, Plainview, N.Y.) in .5 ml distilled water, daily, by oral intubation. L-ascorbic acid solutions were prepared immediately prior to use. Group II was maintained on rabbit chow and received .5 ml distilled water by oral intubation. Vitamin C content of the diet was determined to be 43.75 g per gram of diet. The diet supplied an ascorbic acid intake of approximately 1.3 mg/day for the hypovitaminosis C group, (average intake=30 gm/day), and diet plus oral supplement supplied 11.3 mg/day ascorbic acid in the control. Animals were housed individually and feed and water were supplied ad libitum. Animals were weighed
weekly to monitor onset of Vitamin C deficiency. Control animals were maintained for 9 weeks and the hypervitaminosis C group was maintained until animals began to lose weight (9-13 weeks).

Animals were sacrificed by cervical dislocation. Plasma samples were immediately drawn by cardiac-puncture into heparinized syringes and placed on ice prior to centrifugation at 4°C. Testes were quickly removed and placed on ice in 0.25M sucrose. Testes were homogenized in 0.25M sucrose (10% w/v) as described in Manuscript I. Reduced ascorbic acid in plasma and testes homogenates were determined by the micro-method of Zannoni et al. (18).

**Preparation of Testis Microsomes**

This procedure was conducted as previously described in Manuscript I.

**Enzyme Incubations**

Assay of 3BHSOR activity in testes microsomes was performed as described in Manuscript I with the exception that L-ascorbic acid and L-dehydroascorbic acid were added at concentrations of 2.0 mM where indicated. Microsomes from control and hypovitaminosis C animals were pooled for the enzyme incubations.

**Vitamin C Content of Diet**

Reduced ascorbic acid content of the diet was determined by the method of Zannoni et al. (19), with the following
modification; ascorbic acid was extracted from 1 gram of diet into 5 ml of 5% trichloroacetic acid. The extract was filtered and rinsed several times with 5% TCA solution and adjusted to a final volume of 10 ml. An internal standard was used to check for completeness of extraction. Ascorbic acid content was verified by using the AOAC 2,6-dichloroindophenol method (19).
RESULTS

PLASMA AND MICROSONAL ASCORBIC ACID VALUES

The results of ascorbic acid analysis of plasma and testes microsomes from experimental and control guinea pigs are summarized in TABLE I. The plasma ascorbic acid levels were significantly \( p < .05 \) lowered in the hypovitaminosis C group after 9-13 weeks on the diet supplying 1.3 mg ascorbic acid/day. Testis microsomal ascorbate levels were also significantly lower in the hypovitaminosis C animals.

BODY AND TESTES WEIGHTS

Mean body weight of the hypovitaminosis C group was significantly lower \( p < .05 \) after nine weeks on the experimental diet (TABLE II). The wide range in body weights in this group represents the animals that had not begun to lose weight at 9 weeks. No significant difference was seen between mean testes weights. Surprisingly, the microsomal protein was found to be increased in the hypovitaminous C group.

3BHSOR ACTIVITY IN GUINEA PIG TESTIS MICROSONES

Michaelis constants and \( V_{\text{max}} \) values for the 3BHSOR reaction using PREG, 17\( \alpha \)PREG, DHEA and ADIOL as substrate were estimated by the graphic method of Lineweaver-Burk (20) and are summarized in TABLE III. The \( k_{0.5} \) for NAD in the 3BHSOR reaction was determined to be 35\( \mu \)M (Figure 1) and all enzyme
incubations were carried out at greater than saturation levels of the coenzyme. The $k_{0.5}$ for PREG in the 3BHSOR reaction was found to be similar in both control and hypovitaminosis C guinea pig testis microsomes (Figure 2), although the $V_{\text{max}}$ was inhibited 64% by the hypovitaminous C pretreatment. The Michaelis constants of the 3BHSOR reaction for 17αPREG, DHEA and ADIOL (Figures 3-5) were found to be slightly lower in the testis microsomes from the hypovitaminosis C animals but these differences are considered to be negligible. The most severe inhibition of the $V_{\text{max}}$ by hypovitaminosis C pretreatment was seen in the 17αPREG + 17αPROG reaction where inhibition at saturation reached 76%. Lesser inhibition was seen in the DHEA $\rightarrow$ A and ADIOL $\rightarrow$ T reactions, 49% and 37%, respectively.

Since ascorbic acid levels in whole testes of the normal guinea pig were reported to be $36.5 \pm 1.1$ mg/100 gm (21), representing an ascorbic acid concentration of approximately 2mM, ascorbic acid and dehydroascorbic acid were added back to in vitro incubations of the hypovitaminosis testis microsomes at this level. Results are summarized in TABLE IV. Both ascorbic acid and dehydroascorbic acid stimulated 3BHSOR activity at intermediate levels of substrate. No stimulation by ascorbic acid was seen at low substrate concentrations nor at very high substrate concentrations and at no time was activity restored to the level of the control animals. The stimulation of 3BHSOR activity with ascorbate is at least partially protein dependent. The greatest effect was seen
with high protein concentrations where product formation reached 165% and 155% of the reference value in the presence of ascorbic acid and dehydroascorbic acid at 5\textmu M substrate. Similar stimulation of 3BHSOR activity is seen with high protein concentrations at the 10\textmu M substrate level. The dependence of this stimulatory effect upon protein and substrate concentrations suggests that ascorbic acid may be acting to reverse product inhibition. The two products of the reaction, NADH and progesterone are known to be inhibitors of 3BHSOR activity. Since NADH is a competitive inhibitor of NAD it is not likely to be inhibitory with low product appearance in the presence of 500\textmu M NAD. Progesterone inhibition of the 3BHSOR is noncompetitive (22) with an estimated $k_i$ of 2.5\textmu M (Figure 6) in guinea pig testes microsomes. It appeared possible that the ascorbic acid may be reversing product inhibition by progesterone. Microsomes were incubated in the presence of 2\textmu M and 5\textmu M progesterone, 2 to 20\textmu M substrate, with and without the addition of 2mM ascorbate or dehydroascorbate. Neither ascorbic acid nor dehydroascorbic acid were able to reverse progesterone inhibition of 3BHSOR activity.
DISCUSSION AND CONCLUSIONS

The increase in the amount of microsomal protein isolated from the testes of the hypovitaminosis C guinea pigs bears review. It is clear that this increase in protein is not the result of testicular hypertrophy since the testes weights were similar in both groups. The discussion of increased microsomal protein synthesis in the ascorbic acid deficient guinea pig is not within the scope of this study. Since only 30-40% of the microsomes are isolated by the preparation procedure, the increase in microsomal protein may simply be due to an increased yield of microsomes. To answer this question a study would need to be conducted employing a microsomal marker in the whole testes homogenate to quantitate the amount of microsomes present in order to determine % yield. No decrease in the amount of cytochrome P-450 could be detected in the testis microsomes of the hypovitaminosis C pretreated guinea pigs. The activity of PROG-17α-hydroxylase was used as a marker for the P-450 as this enzyme has been shown to exhibit a constant activity/nm P-450 in testis microsomes (23). $V_{\text{max}}$ for the reaction in the control testis microsomes was previously determined to be 0.24 nm/min/mg and was found to be 0.19 nm/min/mg in deficient microsomes. As the amount of P-450 in microsomal preparations is inversely related to the amount of protein isolated, this activity must be corrected for the 20% increase in microsomal protein. On this basis, the activities
of the 17α-hydroxylase in the control and hypovitaminosis C microsomes are identical. This data is in contrast to Degwitz et al. (24) who demonstrated a 30% decrease in liver cytochrome P-450 synthesis following 28 day depletion of ascorbic acid. It is apparent that tissue differences may exist with respect to the protoheme synthesis for P-450 and also probable that chronic hypovitaminosis C may not affect cytochrome P-450 levels to the extent that is seen in the scorbutic animal.

3BHSOR activity was found to be inhibited in testis microsomes isolated from guinea pigs with hypovitaminosis C. This inhibition was present with all four substrates for the enzyme, although the % inhibition was greatest in the reactions having the highest rates of product formation. In vitro additions of ascorbic acid and dehydroascorbic acid were most effective in stimulating 3BHSOR activity with PREG as substrate under conditions where the reaction rate was maximal (i.e., saturating substrate and high protein concentrations). It appears that ascorbic acid may function in some manner which releases product inhibition on the enzyme. The reversal of NADH inhibition of 3BHSOR activity by ascorbic acid has been demonstrated in rat adrenal preparations (12), although the ascorbic acid concentration used was 40mM which is far greater than physiological levels in the testes, and the NADH concentration was 300µM. The incubation procedure used in this study produces NADH at a maximum velocity of .77 nm/min/ng in controls and .28 nm/min/ng in the deficient group.
This would allow for an average NADH concentration from .7-1.0 µM under the assay conditions employed. Since NADH is continually being oxidized in testes microsomes via cytochrome b5, the average concentration is probably much lower than predicted. Inhibition of 3BHSOR activity in testes microsomes of hypovitaminous C guinea pigs and stimulation of enzyme activity with ascorbate and dehydroascorbate in the presence of 500 M NAD can not be explained on the basis of NADH oxidation alone. The monodehydroascorbate-NADH oxidoreductase demonstrated by Staudinger (13) is strictly specific for reduced ascorbic acid. Therefore, the oxidase could not explain the stimulation of 3BHSOR activity seen with dehydroascorbate in this study and in toad testes (16). NADH has been found to inhibit 3BHSOR noncompetitively with respect to pregnenolone in the 0-40% (NH4)2SO4 fraction of human testis preparations (22) but the k_i for NADH was estimated to be 150 M. Since this concentration could never be attained under the assay conditions of the present study, this model can not explain the release of inhibition on 3BHSOR by ascorbic acid. The data suggest that ascorbic acid does not function in the reversal of product inhibition on 3BHSOR since NADH inhibition is unlikely and progesterone inhibition cannot be reversed by the ascorbate addition.

The conversion of pregnenolone to progesterone requires two enzymatic conversions: 1) the NAD-dependent oxidation of the hydroxyl group at the C-3 position to a ketone by 3BHSOR and 2) the isomerization of the Δ5-6 double band to
the Δ₄-₅ position. Although the reaction catalyzed by 3BHSOR is known to be the rate limiting step in the coupled reaction (22), it is possible that ascorbic acid exerts its effect on the isomerase. Since the velocity of the 3BHSOR reaction depends upon the rapid removal of the Δ₅ 3-ketone intermediate, it is feasible that inhibition of the isomerase would result in the decreased conversion of PREG + PROG seen with hypovitaminosis C.
Figure 1. Michaelis Menton plot of 3BHSOR reaction in guinea pig testis microsomes with varying NAD. Incubations run in the presence of 10µM PREG, 0.1M Tris, pH 7.4, 5mM MgCl₂, 1% propylene glycol at 37°C for 5 minutes. Protein-60 g/1 ml final volume. NAD varied from 25-500µM. k₀.5 NAD=35µM.
Figure 2. Lineweaver-Burk replots of 3BHSOR with PREG as substrate in control and hypovitaminosis C guinea pig testis microsomes. Incubations were run in the presence of .1M Tris, pH 7.4, 5mM MgCl$_2$, 500µM NAD, 1% propylene glycol at 37°C for 5 minutes. Protein-65 µg (control), 109 ± 182 µg (deficient) $k_{0.5}$=2.75µM (control), 3.3µM (deficient) $V_{max}$=.79 nm/min/mg (control), .28 nm/min/mg (deficient).
Figure 3. Lineweaver-Burk replots of 3BHSOR with 17αPREG as substrate in control and hypovitaminosis C guinea pig testis microsomes. Incubations were run in the presence of .1M Tris, pH 7.4, 5mM MgCl₂, 500µM NAD, 1% propylene glycol at 37°C for 5 minutes.
Protein - 161µg (control), 162µg (deficient)
K_{0.5} = 2.75µM (control), 3.3µM (deficient)
V_{max} = .77 nm/min/mg (control) .28 nm/min/mg (deficient)
Figure 4. Lineweaver-Burk replots of 3BHSOR with DHEA as substrate in control and hypovitaminosis C guinea pig testis microsomes. Incubations were run in the presence of .1M Tris, pH 7.4, 5 MgCl$_2$, 500µM NAD, 1% propylene glycol at 37°C for 5 minutes.

Protein-197µg (control) 162µg (deficient)

$K_{0.5} = 4.0$µM (control), 1.3µM (deficient)

$V_{max} = .43$ nm/min/mg (control), .22 nm/min/mg (deficient).
Figure 5. Lineweaver-Burk replots of 3BHSOR with ADIOL as substrate in control + hypovitaminosis C guinea pig testes microsomes. Incubations were run in the presence of .1M Tris, pH 7.4, 5mM MgCl₂, 500 M NAD, 1% propylene glycol at 37°C for 5 minutes.

Protein-537µg (control), 504µg (deficient)

k⁰.₅=5.0µM (control), 3.8µM (deficient)

Vₘₐₓ = .35 nm/min/mg (control, .22 nm/min/mg (deficient).
Figure 6. Inhibition of 3BHSOR by Progesterone, Replot of Lineweaver-Burk (I vs. Slope). Incubations were run in the presence of .1M Tris, pH 7.4, 5mM MgCl₂, 500μM NAD, 1% propylene glycol for 5 minutes at 37°C, 2-10μM PREG, Protein-173μg. $k_1=2.5\text{ M.}$
The text is not fully legible due to the quality of the image. However, it appears to discuss plasma and tissue levels of acetylsalicylic acid and includes a graph with labeled axes and annotations.
<table>
<thead>
<tr>
<th></th>
<th>PLASMA ASCORBIC ACID (mg/100ml)</th>
<th>MICROSMAL ASCORBIC ACID g/mg protein</th>
<th>g/2 Testes</th>
</tr>
</thead>
<tbody>
<tr>
<td>CONTROL (6)</td>
<td>$0.252 \pm 0.142^b$</td>
<td>$0.323 \pm 0.07^c$</td>
<td>$6.69 \pm 1.71^d$</td>
</tr>
<tr>
<td>HYPOVITAMINOSIS C (6)</td>
<td>$0.152 \pm 0.101^b$</td>
<td>$0.072 \pm 0.07^c$</td>
<td>$1.84 \pm 1.10^d$</td>
</tr>
</tbody>
</table>

a - reduced ascorbic acid
b - significantly different as determined by the 1 Tail T-Test $p < .05$.
c - significantly different as determined by the 1 Tail T-Test $p < .001$.
d - significantly different as determined by the 1 Tail T-Test $p < .001$

The numbers in parenthesis represent number of animals in each group.
TABLE II

MEAN BODY WEIGHT, TESTES WEIGHT AND MICROsomAL PROTEIN IN CONTROL AND HYPOVITAMINOSIS C GUINEA PIGS

<table>
<thead>
<tr>
<th></th>
<th>BODY WEIGHT(^2) (GM)</th>
<th>WEIGHT/2 TESTES (GM)</th>
<th>MICROsomAL PROTEIN (MG/2 TESTES)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CONTROL (6)</td>
<td>714 ± 81(^b)</td>
<td>4.25 ± .55(^c)</td>
<td>20.63 ± 1.22(^d)</td>
</tr>
<tr>
<td>HYPOVITAMINOSIS (6)</td>
<td>617 ± 134(^b)</td>
<td>4.02 ± .62(^c)</td>
<td>25.77 ± 1.85(^d)</td>
</tr>
</tbody>
</table>

\(a\) - mean body weight after 9 weeks on experimental and control diets

\(b\) - values are significantly different as determined by the 1 Tail T-Test \(p < .05\)

\(c\) - values are not significantly different

\(d\) - values are significantly different as determined by the 2 Tail T-Test \(p < .001\)

The numbers in parenthesis represent the number of animals in each group.
<table>
<thead>
<tr>
<th>Reaction</th>
<th>( k_{0.5} )</th>
<th>( V_{\text{max}} )</th>
<th>( k_{0.5} )</th>
<th>( V_{\text{max}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>PREG + PROG</td>
<td>2.75</td>
<td>.77</td>
<td>3.3</td>
<td>.28</td>
</tr>
<tr>
<td>17\alpha\text{PREG} + 17\alpha\text{PROG}</td>
<td>5.5</td>
<td>.67</td>
<td>4.9</td>
<td>.16</td>
</tr>
<tr>
<td>DHEA ( \rightarrow ) A</td>
<td>4.0</td>
<td>.43</td>
<td>1.3</td>
<td>.22</td>
</tr>
<tr>
<td>ADIOL ( \rightarrow ) T</td>
<td>5.0</td>
<td>.35</td>
<td>3.8</td>
<td>.22</td>
</tr>
</tbody>
</table>

\( k_{0.5} \) expressed as \( \mu \text{M} \) substrate

\( V_{\text{max}} \) expressed as nm product/min/mg protein
TABLE IV

THE EFFECT IN VITRO ADDITION OF ASCORBIC ACID\(^a\) AND DEHYDROASCORBIC ACID ON 3BHSOR ACTIVITY\(^b\) IN TESTIS MICROSONOMES OF GUINEA PIGS WITH HYPOVITAMINOSIS C

<table>
<thead>
<tr>
<th>Substrate Concentration (mM)</th>
<th>+ Ascorbic Acid 108 µg Protein</th>
<th>+ Dehydroascorbic Acid</th>
<th>Dehydroascorbic Acid 297 µg Protein</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0</td>
<td>100</td>
<td>100</td>
<td>-</td>
</tr>
<tr>
<td>2.0</td>
<td>134</td>
<td>107</td>
<td>147</td>
</tr>
<tr>
<td>5.0</td>
<td>123</td>
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</tr>
<tr>
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<td>156</td>
<td>147</td>
</tr>
<tr>
<td>15.0</td>
<td>100</td>
<td>103</td>
<td>-</td>
</tr>
</tbody>
</table>

\(^a\) - final concentration, 2mM Ascorbic Acid or Dehydroascorbic Acid

\(^b\) - PREG used as substrate

\(^c\) - Reference - pooled microsomes from hypovitaminosis C guinea pigs - no addition
References


I. ANDROGEN BIOSYNTHESIS IN THE TESTIS

Synthesis of testosterone from acetate was shown to involve cholesterol, pregnenolone, progesterone, 17α-hydroxyprogesterone and androstendione (1). Two pathways for the synthesis of testosterone from cholesterol have been subsequently identified. Slaunwhite and Samuels (2) demonstrated a pathway in rat testes homogenates in which progesterone was hydroxylated at the 17 position followed by side chain cleavage to produce androstendione which is reduced to form testosterone. Neher and Wettstein (3) found dehydroepiandrosterone to be an intermediate in testosterone synthesis in porcine testes homogenates and suggested a pathway in which pregnenolone is converted to 17α-hydroxypregnenolone and then to DHEA. DHEA was presumed to be converted to androstendiol and then reduced to testosterone. Eik-Nes and Kekre (4) confirmed DHEA to be an intermediate in dog testes. Later, Δ5-androstendiol was identified as a probable intermediate in the conversion of DHEA to testosterone in rabbit (5) and rat (6) testis microsomes. Hagen and Eik-Nes (7) demonstrated the conversion of 17α-hydroxypregnenolone to 17α-hydroxyprogesterone, the first alternate route to testosterone from pregnenolone.

The specific enzyme reactions have been further elucidated. The first reaction in adrenal steroid synthesis and
of the \( \Delta_4 \) pathway in the testis is the 3B oxidation of pregnenolone, catalyzed by 3B hydroxysteroid oxidoreductase/\( \Delta_5 \)-3-ketosteroid isomerase. The reaction involves an NAD dependent oxidation at C3 to yield a \( \Delta_5 \)-3-ketosteroid intermediate which is isomerized to progesterone, a \( \Delta_4 \)-3-ketosteroid. The oxidation is rate limiting in the reaction (8) and the overall reaction rate is determined by the rate of conversion of the \( \Delta_5 \)-3-hydroxysteroid to the \( \Delta_5 \)-3-ketosteroid. The reaction is largely irreversible. This is probably attributable to the isomerization step in the coupled reaction, although the conversion was shown to be reversible in the rabbit (9) and ovine adrenal microsomes (10).

The next reaction on either the \( \Delta_4 \) or the \( \Delta_5 \) pathway is the 17α-hydroxylation of PROG or PREG which is catalyzed by a cytochrome-P-450 dependent hydroxylase. The enzyme is of the mixed-function oxidase type and requires molecular oxygen and NADPH. Purvis et al. have demonstrated the P-450 dependence in hypophysectomized rats (11). In the testes microsomes of the hypophysectomized animal, the decay rate of the 17α-hydroxylase activity, \( (t\,1/2=2.3\,\text{days}) \), paralleled the decay of cytochrome P-450, \( (t\,1/2=3.3\,\text{days}) \). Conversely, the hydroxylase was shown to maintain a constant activity per nanomole of cytochrome P-450 when the level of P-450 was increased under HCG stimulation. Further evidence that the enzyme requires cytochrome P-450 is inhibition of the 17α-hydroxylase activity by carbon monoxide (10,11) and
which are known to be inhibitors of cytochrome P-450.

Metabolism of 17α-hydroxysteroids to testosterone involves a second cytochrome-P-450 dependent conversion, catalyzed by C\textsubscript{17}-C\textsubscript{20} lyase. The reaction requires NADPH and molecular oxygen. As with the P-450 dependent steroid 17α-hydroxylase, C\textsubscript{17}-C\textsubscript{20} lyase levels parallel the cytochrome P-450 content of the testes microsomes (11). The reaction has been found to be irreversible (15). Products of C\textsubscript{17}-C\textsubscript{20} lyase reaction are C\textsubscript{19}-ketosteroids which are reduced at C\textsubscript{19} to testosterone. The reaction is catalyzed by 17B-hydroxysteroid oxidoreductase. The enzyme requires NADPH but does not involve cytochrome P-450. The decay rate of 17BHSOR was found to be slower in testis microsomes of hypophysectomized rats, (t\textsubscript{1/2}=4.5 days), than either cytochrome P-450 or the P-450 dependent enzymes (11) and enzyme activity is not inhibited by spironolactone (14). The reaction is capable of proceeding in both forward and reverse directions, however, there is some evidence that the oxidation of T\textsuperscript{+}A and the reduction of A\textsuperscript{-}T are catalyzed by two distinct enzymes (16,17,18).

Although, the \textgreek{Delta}\textsubscript{4} and \textgreek{Delta}\textsubscript{5} pathways for steroid biosynthesis are fairly well defined, the number of enzymes involved in the catalysis of the reactions remains controversial. It has been shown that 3-(17)BHSOR purified from Pseudomonas testeroni yields 4 distinct bands upon electrophoretic
separation, each possessing both 3B and 17BHSOR activity (19). Two subunits were demonstrated and the enzyme was believed to exist as four tetrameric species. However, Shikita and Talalay reported all 3BHSOR activity to be associated with a single band upon electrophoretic separation of the purified *P. testeroni* enzyme (30). No evidence exists for the presence of multiple forms in mammalian species. 3BHSOR/Δ5-3-
ketoisomerase has been isolated from sheep (21) and human (22) adrenal cortex microsomes and found to have similar activity with either DHEA or pregnenolone as substrate. Yates and Deshplande (22) noted that when both substrates were incubated simultaneously, the $k_m$ value decreased 50% and the $V_{max}$ decreased suggesting that the two substrates were competing for the same enzyme. In human testis microsome (23), the $k_m$ for DHEA and androstenediol were found to be similar and the two substrates were shown to be competitive inhibitors of one another providing further evidence that the various substrates of 3BHSOR all compete for the same enzymatic site.

The existence of two C17-C20 lyases, or, two catalytic sites for $17\alpha$REG and $17\alpha$PROG was demonstrated in human testes microsomes (24). $k_m$ values were found to be different for $17\alpha$PROG, (1.7x10^{-5} M), and $17\alpha$PREG, (5.9x10^{-7} M). Additionally, testosterone was shown to be a competitive inhibitor with $17\alpha$PROG as substrate and uncompetitive with respect to $17\alpha$PREG, indicating that T binds to the active site
on the PROG enzyme and at other than the active site on the
PREG enzyme. However, this evidence cannot be considered
definitive as 17αPROG and 17αPREG were found to act as com-
petitive inhibitors of one another. Recently, a homogenous
cytochrome P-450 was purified from neonatal pig testis micro-
somes (25,26). Both lyase and 17αhydroxylase activities were
found to be associated with this enzyme. The $k_m$ values were
similar for both 17αPROG and PROG although the $V_{max}$ for the
hydroxylation was twice as great as the lyase. Both activi-
ties were reported to be inhibited to the same extent by
P-450 inhibitors. These findings are in contrast to those of
Betz (27) who demonstrated that the enzyme activities are as-
sociated with two different proteins in rat testis microsomes.

The question as to the number of enzymes is crucial to
understanding the varying pathways for testosterone synthe-
sis. Two major pathways, the $\Delta_4$ pathway from pregnenolone
to progesterone and the $\Delta_5$ pathway from pregnenolone to 17α
hydroxyprogesterone have been widely accepted. It appears,
however, that some species use mixed, (or alternate), path-
ways to T. The existence of a single lyase and steroid hy-
droxylase would limit the use of mixed pathways since the 17
hydroxy product of the first $\Delta_5$ reaction would likely be
enzyme-bound and not available for the alternate pathways
prior to the lyase reaction.

The mixed pathways in rat and rabbit testes have been
shown using an in vivo perfusion technique (28,29,30). The
most probable pathways in the rat were predicted to be mixed:

\[ \text{PREG} \xrightarrow{\Delta_5} 17\alpha\text{PREG} + 17\text{PROG} \rightarrow A + T, \text{ or, purely } \Delta_4: \text{PREG} \rightarrow 17\alpha\text{PROG} + A + T. \]

The rabbit is capable of using mixed pathways: PREG + 17\alpha\text{PREG} + 17\alpha\text{PROG} + A + T or \[ \Delta_5: \text{PREG} \rightarrow 17\alpha\text{PREG} + \text{DHEA} + \text{ADIOL} + T \]

but was predicted to favor the \[ \Delta_5 \]
pathway due to low conversion of 17\alpha\text{PROG} + A.

Testosterone biosynthesis on the human has been assumed to rely primarily on the \[ \Delta_5 \]
pathway from PREG + 17\alpha\text{PREG} + \text{DHEA} + \text{ADIOL} + T (31,32). Very low conversion of \text{DHEA} + A was seen in comparison to the large conversion of \text{DHEA} + \text{ADIOL} so the possible use of a mixed pathway from PREG \[ \xrightarrow{\Delta_5} 17\alpha\text{PREG} \xrightarrow{\Delta_5} \text{DHEA} + A \xrightarrow{\Delta_4} T, \]
was not considered to be favorable. The \[ \Delta_5 \]
pathway was found to predominate in the dog by in vivo (33,34) and in vitro (35) methods. The alternate pathway from \text{DHEA} + A + T is unlikely in the dog since \text{DHEA} + 3\text{BHSOR} activity was found to be only 15% that of \text{DHEA-17BHSOR} (35).

Although differences in total steroid secretion among species under maximum stimulation may be accounted for on a basis of testes weight (36) or leydig cell mass (30), differences in testosterone secretion are not related to differences in these two values. It has been suggested by Zirkin et al. (37), that quantitative differences in testosterone production by maximally stimulated testes on the hamster, rat, rabbit, dog, and guinea pig may be explained on the basis of leydig cell ultrastructure. A linear relationship was shown between the amount of leydig cell smooth endoplasmic reticulum and testosterone production. As most of the
steroidogenic enzymes are localized in the SER it appears reasonable that the total amount of this organelle would reflect the steroid synthesizing capacity of the organism. However, the amount of SER enzymes have not been quantitated and it appears plausible that differences in testosterone synthesis among species are related to enzymatic activities of the $\Delta_4$, $\Delta_5$ or mixed pathways and the efficiency of feedback and feedforward controls on these pathways.

II. THE EFFECT OF ASCORBIC ACID ON 3B-HYDROXysteroid oxidoreductase IN THE TESTIS.

Administration of large doses of ascorbic acid has been linked to abortion in animals (38,39,40) and, possibly, humans (40). It has been proposed that the inability to maintain a pregnancy after ingestion of massive doses of ascorbic acid is due to a disruption in steroid metabolism leading to alterations of steroid hormone levels in the blood. The importance of progesterone, in the maintenance of pregnancy has long been recognized. Progestogens are frequently used in the maintenance of pregnancies in females with histories of repeated abortions.

The role of ascorbic acid in steroidogenesis is poorly understood. High levels of ascorbic acid have been shown to inhibit the conversion of cholesterol to pregnenolone; this is considered to be the rate limiting step in the biosynthesis of steroids. Conversion of the C-27 steroid, cholesterol, to C-21$\Delta_5$-pregnenolone involves two mitochondrial
hydroxylations at C-20 and C-22, followed by side chain cleavage between C-20 and C-22 (41). In vitro studies, using levels of ascorbic acid comparable to those of resting ovary and adrenal cortex, have demonstrated ascorbate inhibition of mitochondrial side chain cleavage in rat ovary (42,43) and bovine adrenal cortex (44). Stimulation of steroidogenesis in the ovary by LH and in the adrenal cortex by ACTH results in decreased ascorbic acid concentrations in these tissues before steroid synthesis begins (45). Presumably, this releases the inhibition on the conversion of cholesterol to pregnenolone. Steroid synthesis proceeds in one of two directions, via the Δ₄ pathway which involves a conversion of the Δ₅ 3-hydroxysteroid, pregnenolone, to the Δ₄ 3-ketosteroid, progesterone, by action of the enzymes 3β-Hydroxysteroid oxidoreductase and Δ₄ 3-ketosteroid isomerase, or, by use of the Δ₅ pathway through 17α-hydroxyprogrenenolone. Depressed conversion of Δ₅-hydroxysteroids to Δ₄-ketosteroids has recently been observed in vitamin C deficient guinea pigs (46). Decreased ability to convert 3β-hydroxysteroids to 3-ketosteroids has been seen in Vitamin A deficient rats (47). The Vitamin A deficient rat is unable to synthesize ascorbic acid. When animals were depleted of Vitamin A and Vitamin C was removed from the diet, decreased 3βHSOR activity was seen. Normal activity was achieved by restoration of ascorbic acid. Dehydroascorbate was
shown to stimulate the conversion of pregnenolone to progesterone in toad testes (48).

The results of these studies suggest that ascorbic acid should favor the Δ_4_ pathway. However, in vitro incubations of guinea pig ovary and testis homogenates have shown that the addition of 0.5 mM ascorbic acid stimulated the conversion of Δ_5_ to Δ_4_ steroids from cholesterol to a greater extent than additions at 1.0 or 1.5 mM levels and that the stimulatory effect was absent at 2 mM ascorbate (49). No significant increase in the synthesis of Δ_4_ steroids could be seen when pregnenolone was used as substrate in the presence of 0, 0.5, 1.0, 1.5 and 2.0 mM ascorbate (49). All levels of ascorbic acid were shown to inhibit 17 hydroxylation (49). Ascorbic acid has been shown to inhibit 17α-hydroxy-corticosteroid production in the presence of ACTH in beef adrenal cortex preparations (50) and to favor the production of 17-β-oxycorticosteroids in pig adrenal cortex microsomes (51). Microsomal 21-hydroxylase activity was inhibited by ascorbic acid in bovine adrenal cortex (52, 53).

It has been proposed that ascorbic acid may release NADH inhibition of 3BHSOR in the conversion of pregnenolone to progesterone (54). The reversal of NADH inhibition is presumed to involve an ascorbate dependent NADH oxidation. Staudinger (55) has demonstrated a microsomal NADH-oxidase in rat liver which is strictly specific for ascorbic acid.

The following reaction pathway was proposed:
The flavoprotein was found to be cyanide insensitive and therefore was not NADP:ferricytochrome b₅ oxidoreductase. The protein was later identified to be ascorbate:ferricytochrome b₅ oxidoreductase (EC 1.10.2.1) (56). Existing evidence suggests that most ascorbic acid oxidations follow the two-step process described by Michaelis (57) involving the formation of free radical intermediates. Monodehydroascorbate may be formed by the action of l-ascorbate:oxygen oxidoreductase (EC 1.10.3.3), or, by the comproportionating reaction as shown below:

\[
\begin{align*}
\text{Ascorbate} & \quad \text{Dehydroascorbate} \quad \text{Monodehydroascorbate}
\end{align*}
\]

The monodehydroascorbate free radical formed by comproportionation is equally effective as substrate for NADH linked electron transport as the free radical formed on the enzyme catalyzed reaction. However, the removal of electrons from NADH via monodehydroascorbate need not necessarily be linked to cytochrome b₅. In liver microsomes only 10% of the electrons transported from NADH to monodehydroascorbate utilize cytochrome b₅; 90% of the electrons are transferred by NADH:
monodehydrascorbate oxidoreductase (EC 1.6.5.4) (54).

Heath and Fiddick (58) have shown that NADH oxidation in bovine ocular tissue microsomes is stimulated by ascorbic acid and that dehydroascorbic acid could not substitute for the reduced form. Devine and Rivers (59) found depressed ratios of NAD/NADH in tissues of ascorbic acid deficient guinea pigs which could be attributed to increased levels of NADH, although the activity of the ascorbic acid dependent NADH oxidase was not impaired on the vitamin C deficient guinea pig.

The physiological role of the NADH-monodehydroascorbate oxidoreductases is unclear. It was presumed that this mechanism may conserve ascorbic acid by reducing formation of the free radical, or, may function in steroid synthesis by maintaining adequate levels of oxidized cofactor.

Other evidence suggests that the role of ascorbic acid in steroid synthesis may be related to synthesis of the ferricytochromes, P-450 and b5. Liver cytochrome P-450 and b5 levels were found to decrease in ascorbic acid deficient guinea pigs (60). Twenty-eight days after withdrawal of ascorbic acid, cytochrome P-450 levels were 30% of control and b5 decreased to 45% of control levels. Liver ascorbic acid was found to drop rapidly during the initial 14 days of depletion and then to level off. P-450 had decreased to 45% of control and b5 to 70% at 14 days but continued to fall after ascorbic acid levels had stabilized. Similar,
but less dramatic, results were seen in the adrenal (60). Cytochrome P-450 levels could be normalized with ascorbic acid administration but this normalization was abolished by inhibitors of protein synthesis unless \( \delta \)-amino levulonic acid was administered concurrently. Additionally, phenobarbital or 3-methylcholanthrene, known stimulators of heme synthesis, were able to restore P-450 levels in vitamin C deficient animals (61). This evidence suggests that heme synthesis is impaired in ascorbic acid deficiency.

Testes levels of ascorbic acid have been determined in the rat (62), guinea pig (63,64) and human (65). The normal level in the human testis is the lowest cited (3 mg/100 gm tissue), whereas, higher levels have been reported in the guinea pig (24-36 mg/100 gm tissue), and rat, (25-30 mg/100 gm tissue). Testes ascorbic acid concentrations decrease in scorbutic animals with the degree of depletion apparently being related to the stage of scurvy. Levels of 6 mg/100 gm tissue in early scurvy were decreased to 3 mg/100 gm tissue in late scurvy (63). Gombe et al. (64) reported testes levels of 8.4 mg/100 gm tissue after depletion for three weeks but found no corresponding decrease in plasma or testes testosterone levels. This data was in contrast to that of Kocen and Cavazos (66) who reported arrest of spermatogenesis in ascorbic acid deficient guinea pigs and in agreement with Jones et al. (67) who found no inhibition of steroid synthesis in the adrenals of scorbutic guinea pigs. Cholesterol
levels in the testes were significantly decreased in the testes of scorbutic guinea pigs (64) suggesting release of inhibition on the cholesterol side chain cleavage enzymes by decreased ascorbic acid levels. This evidence supports the inhibitory role of ascorbic acid in steroid synthesis proposed by Sulimovici (42,43) and others (44,45,52,53).
References


57. Michaelis, L. (1932) J. Biol. Chem. 96, 703.
Figure 1. Activity of 5342A activity with varying 
acetic acid concentrations in guinea pig stools 
microsome incubations were run in the 
presence of 5% dextran. As expected, no activity 
was observed at an ionic pH 7.4 at 37°C for 1 
MINUTE. Substrate was 100 mM HCl and protein 
concentration was varied from 1 to 300 mg/ml 
at final volume.

APPENDIX
Figure 1. Linearity of 3BHSOR activity with varying protein concentrations in guinea pig testis microsomes. Incubations were run in the presence of 500µM NAD, 5mM MgCl₂, 1% propylene glycol in .1M Tris, pH 7.4 at 37°C for 5 minutes. Substrate was 10µM PREG and protein concentration was varied from 16 to 200 µg/.1 ml final volume.
Figure 2. Linearity of 3BHSOR activity with time in guinea pig testis microsomes. Incubations were run in the presence of 500 µM NAD, 5mM MgCl₂, 1% propylene glycol in .1M Tris, pH 7.4 at 37°C. Protein concentration was 66 µg/.1 ml final volume. Reactions were incubated for 1, 5 and 10 minutes.


Michaelis, L. (1932) J. Biol. Chem. 96, 703.


