A Mathematical Model of Tryptophan Metabolism via the Kynurenine Pathway Provides Insights into the Effects of Vitamin B-6 Deficiency, Tryptophan Loading, and Induction of Tryptophan 2,3-Dioxygenase on Tryptophan Metabolites

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Abstract

Vitamin B-6 deficiency is associated with impaired tryptophan metabolism because of the coenzyme role of pyridoxal 5'-phosphate (PLP) for kynureninase and kynurenine aminotransferase. To investigate the underlying mechanism, we developed a mathematical model of tryptophan metabolism via the kynurenine pathway. The model includes mammalian data on enzyme kinetics and tryptophan transport from the intestinal lumen to liver, muscle, and brain. Regulatory mechanisms and inhibition of relevant enzymes were included. We simulated the effects of graded reduction in cellular PLP concentration, tryptophan loads and induction of tryptophan 2,3-dioxygenase (TDO) on metabolite profiles and urinary excretion. The model predictions matched experimental data and provided clarification of the response of metabolites in various extents of vitamin B-6 deficiency. We found that moderate deficiency yielded increased 3-hydroxykynurenine and a decrease in kynurenic acid and anthranilic acid. More severe deficiency also yielded an increase in kynurenine and xanthurenic acid and more pronounced effects on the other metabolites. Tryptophan load simulations with and without vitamin B-6 deficiency showed altered metabolite concentrations consistent with published data. Induction of TDO caused an increase in all metabolites, and TDO induction together with a simulated vitamin B-6 deficiency, as has been reported in oral contraceptive users, yielded increases in kynurenine, 3-hydroxykynurenine, and xanthurenic acid and decreases in kynurenic acid and anthranilic acid. These results show that the model successfully simulated tryptophan metabolism via the kynurenine pathway and can be used to complement experimental investigations.

Introduction

Tryptophan is mainly degraded via the kynurenine (Kyn) pathway. Alterations of the Kyn pathway have been extensively studied and discussed in several contexts including inflammation (2), cancer (3), pregnancy (4,5), oral contraceptive (OC) usage (5–9), and vitamin B-6 deficiency (10–12) in animals and humans. In the liver, tryptophan is first oxidized to N-formylkynurenine by tryptophan 2,3-dioxygenase (TDO) (1,13), whereas in nonhepatic tissues, indoleamine 2,3-dioxygenase (IDO) catalyzes this reaction (14,15). TDO and IDO play an important role in tryptophan homeostasis (16). Whereas TDO has been reported to be induced by tryptophan (16–18), glucocorticoids (18–20), glucagon (21), and estrogens (22), IDO is induced primarily by interferon (IFN)-γ (3,12), and therefore IDO has been extensively studied in the immune response (2,14).

Vitamin B-6 plays an important role in the Kyn pathway because pyridoxal 5'-phosphate (PLP) functions as a coenzyme for the bifunctional enzymes kynureninase (KYN) and kynurenine aminotransferase (KAT; Fig. 1) (23,24). KYN catalyzes the hydrolysis of Kyn to anthranilic acid (AA) and 3-hydroxykynurenine (HK) to 3-hydroxyanthranilic acid (HaA). KAT catalyzes the conversion of kynurenine to kynurenic acid (KA) and HK to xanthurenic acid (XA) (25–27) (Fig. 1).
FIGURE 1 Diagram of the model. The rectangles represent substrates, and the ellipses contain the acronyms of enzymes. The 2 instances of KYN and KAT are labeled differently because they have different substrates and (possibly) different velocities. Transport of tryptophan from the gut to the serum and from the serum to the muscle or liver compartments is by the L- amino acid transporter. Steady state concentrations (μmol/L) are shown: AA, arachidonic acid; AFD, aryl formamidase; gtrp, tryptophan in the gut; HaA, 3-hydroxyanthranilic acid; HAO, 3-hydroxyniolic acid oxygenase; HK, 3-hydroxykynurenine; KA, kynurenic acid; KAT, kynurenine amino-transferase; KMO, kynurenine 3-monoxygenase; Kyn, kynurenine; KYN, kynureninase; Lpool, tryptophan incorporated into proteins in liver; Mpool, tryptophan incorporated into proteins in muscle; Mtrp, tryptophan in muscle; NFK, N-formyl-kynurenine; QA, quinolinic acid; pertrp, tryptophan in the peripheral circulation; PLP, pyridoxal 5'-phosphate; portrp, tryptophan in portal circulation; TDO, tryptophan 2,3-dioxygenase; trp, tryptophan; XA, xanthurenic acid.

effects of vitamin B-6 deficiency on the activity of these 2 enzymes suggest that KYN is more susceptible than KAT (24).

The tryptophan load test has been used for many years as a functional assessment of vitamin B-6 status (28). Elevation of urinary concentrations of XA, Kyn, and HK after tryptophan loads in vitamin B-6–deficient conditions has been consistently reported in the literature (12,25,29,30). Relatively little is known about the relationship between low cellular concentrations of PLP and the patterns of the other metabolites, including quinolinic acid (QA), HaA, AA, and KA under basal and post–tryptophan load conditions. Moreover, variation in the concentrations of the tryptophan catabolites is found in the published data. For example, Shibata et al. (29) reported an increase in urinary KA and a decrease in KA in vitamin B-6–deficient rats. Bender et al. (30) did not find a change in urinary KA in mice fed a vitamin B-6–deficient diet but reported an increase in XA, Kyn, and HK with HK > XA > Kyn. Yeh and Brown (12) found that, in rats, the percentage of tryptophan load excreted as XA, Kyn, HK, and QA was greater in the vitamin B-6–deficient group than in controls. However, the percentage excreted as KA and HaA did not change in the deficient group. In the same study, humans examined before and after 28 d of a vitamin B-6–deficient diet showed that the percentage of the tryptophan load excreted as XA, Kyn, HK, HaA, KA, and QA all increased in the order HK > Kyn > XA > KA > QA. Similar abnormalities in tryptophan metabolite excretion have been reported in OC users. Miller et al. (31) reported a greater increase in urinary excretion of XA, followed by HK, Kyn, HaA, and KA after a tryptophan load in OC users compared with controls. Donald and Bosse (32) found a higher increase in Kyn followed by XA, HK, and KA in OC users compared with nonusers after a tryptophan load.

The literature is incomplete regarding the basal range of tryptophan metabolites in fasting and fed states and the effect of vitamin B-6 status on their concentrations in plasma and urine. In addition, much of the literature regarding vitamin B-6 and tryptophan metabolism deals with responses to a tryptophan load, rather than with more physiologic dietary intake amounts. Mathematical models constructed by using published experimental kinetic data are excellent tools for the study of nutritional influences on a metabolic system (33). Importantly, such models often can extend, clarify, and provide new insights in metabolism (33,34). Here, we present the development and initial applications of a mathematical model that simulates tryptophan metabolism via the Kyn pathway in the liver. Our model includes tryptophan input in the intestinal lumen, transport of tryptophan from intestinal lumen to circulation by the L-transporter, delivery of tryptophan to the liver, and exchange of circulating tryptophan with muscle and brain. The model was based on Michaelis-Menten constants, V_{max} and K_{m}, for all the enzymes and included provisions for PLP as a coenzyme for KYN and KAT and regulation and inhibition of enzymes.

We report here a mathematical model of whole-body tryptophan metabolism and the results of validation studies that show good agreement of the model with published data. In addition, we report the results of in silico experiments with the model to investigate the effects of various levels of vitamin B-6 deficiency, the effects of TDO induction, and the interaction of vitamin B-6 deficiency.

Materials and Methods

The model. The model was built on the basis of information obtained from published mammalian data on concentrations of tryptophan and its metabolites in the liver, kinetics of the enzymes involved in the Kyn pathway, kinetics of the transporter involved in the transport of tryptophan to tissues, and enzyme regulatory mechanisms (e.g., substrate activation and inhibition of TDO by tryptophan). The kinetic constants used were obtained from different species, including rats, mice, and humans (Table 1). To better approximate whole-body tryptophan metabolism, and to be able to do experiments to examine the effects of different tryptophan inputs (e.g., daily average, oral tryptophan loads), we included 6 compartments in the model: liver, intestinal lumen (gut), portal circulation, peripheral circulation, muscle, and brain. The volume assumptions for the gut and liver compartments were 3 L each and for portal circulation, peripheral circulation, and muscle were 0.33 L, 2.66 L, and 30 L, respectively.

The differential equations (Supplemental Table 1) were derived by using the following approach: If we suppose that tryptophan is being transported from compartment A to compartment B, if V is the rate of decrease in tryptophan concentration in A per unit of time, then the rate of increase of its concentration in B is:

\[
\frac{\text{Vol}(A)}{\text{Vol}(B)} V.
\]

All of the explicit numbers in the differential equations are due to such volume factors. The parameter choices and references for reactions that have Michaelis-Menten kinetics are shown in Table 1 and are in either of the following standard forms:

\[
V = \frac{V_{\text{max}} |S|}{K_{\text{m}} + |S|} = \frac{V_{\text{max}} |S| |S_i|}{(K_{s_i} + |S_i|)(K_{s_j} + |S_j|)}.
\]

In the model, the gut compartment directly receives tryptophan from dietary intake or tryptophan loads. For the purpose of this model, the
steady state concentration of tryptophan in the liver is maintained by not only dietary tryptophan but the tryptophan that comes from proteolysis in the tissues (muscle and liver). We did not consider other sources of amino acids or the effects of plasma albumin on tryptophan concentration. We considered both steady state and variable tryptophan inputs. The daily human intake of tryptophan, which can be calculated by using either catabolism measurements or intake from food, is \(0.5\) g/d (35). This is the equivalent of \(2.5\) mmol/L, or \(100\) μmol/L. The gut compartment in the model has a volume of 3 L, so our steady state input to the gut is \(33.3\) μmol/L per hour. Tryptophan (trp) is removed from the gut and enters the portal circulation by the L-transporter (Vtrpgs) with a \(K_m = 64\) μmol/L (36). We estimated the \(V_{max}\) of the transporter to correspond to \(33.3\) μmol leaving the gut and entering the portal circulation per hour. Because the portal circulation is assumed to have a volume of 0.33 L, and the gut is assumed to have a volume of 3 L, the molar concentration of tryptophan in portal circulation increases by 9

<table>
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<th>Parameter</th>
<th>Units</th>
<th>Model value</th>
<th>Literature values</th>
<th>References</th>
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1. \(V_{max}\) values used in constructing the model and conducting simulations are shown (see text for explanations). AFD, aryl formamidase; HAO, 3-hydroxynalactic acid oxygenase; hk, 3-hydroxykynurenine; KAT, kynurenine aminotransferase; KMO, kynurenine 3-monoxygenase; kyn, kynurenine; KYN, kynureninase; PLP, pyridoxal 5'-phosphate; TDO, tryptophan 2,3- dioxygenase; trpgs, tryptophan from gut to portal circulation; trpl, tryptophan from portal circulation to liver; trpsm, tryptophan from peripheral circulation to muscle.

2. See text for Literature values.

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times the amount that its molar concentration in the gut decreases, which explains the number 9 preceding \( V_{\text{trp}} \) in equation 10.

We assumed that the plasma has a volume of 3 L and is divided into 2 compartments: the portal circulation (0.33 L) and the peripheral circulation, which has a volume of 2.66 L. We divided the plasma volume into these 2 compartments because it is known that the portal concentrations are substantially higher after meals (37,38). During tryptophan loading experiments it is important that much of the tryptophan that is imported from the gut goes first to the portal circulation, then to the liver, and then to the peripheral circulation by mixing of the portal and peripheral circulations (equations 10 and 14). The model also provides for tryptophan to leave the liver and return to the portal circulation (the term \( k_1 \cdot \text{trp} \) in equation 1). Tryptophan is exported from the portal circulation into muscle and brain. The transport into the brain is small (39,40), only \( \sim 3\% \) of the tryptophan whole-body tryptophan utilization, so we represent this simply as a sink for tryptophan indicated by the term \( k_1 \cdot \text{trp} \) in equation 14.

The terms for the mixing of the portal and peripheral circulations are derived as follows. The portal and peripheral circulations are connected as indicated in Figure 1. For any designated time, \( t \), \( \text{pertrp} \) and \( \text{porttrp} \) are the current concentrations of tryptophan in the 2 compartments, respectively. The portal flow of absorbed tryptophan is calculated as follows: In time \( \Delta t \), the portal plasma advances a distance \( \Delta x \) into the peripheral compartment. Because the volume of the peripheral compartment is 2.66 L, then the new peripheral tryptophan concentration (\( \text{newpertrp} \)) is given by the following equation:

\[
\text{newpertrp} = \frac{\Delta x}{2.66} (\text{porttrp}) + (1 - \frac{\Delta x}{2.66}) (\text{pertrp}).
\]

If \( m \) is the velocity of the fluid, then \( m \Delta t = \Delta x \). So rearranging, we get

\[
\frac{\text{newpertrp} - \text{pertrp}}{\Delta t} = m \frac{\text{porttrp} - \text{pertrp}}{2.66}.
\]

Taking the limit as \( \Delta t \to 0 \), we see that

\[
\frac{d(\text{pertrp})}{dt} = 0.375m (\text{porttrp} - \text{pertrp}).
\]

Similarly,

\[
\frac{d(\text{porttrp})}{dt} = 3m (\text{porttrp} - \text{pertrp}).
\]

We choose the mixing constant \( m \) so that the portal and peripheral circulations equilibrate after \( \sim 8 \) h (37).

The muscle compartment receives tryptophan from the peripheral circulation and exchanges free tryptophan with the tryptophan-incorporated into protein (\( \text{Mpool} \)). We include the muscle compartment in our model because the large amount of protein in muscle is a store of tryptophan and between meals some tryptophan leaves the muscle compartment into the plasma to maintain the plasma concentration. Thus, free tryptophan is used for protein synthesis and, conversely, free tryptophan becomes available from proteolysis. We model these processes as simple reactions whose rates are proportional to concentrations of substrates (tryptophan and protein, respectively). We do not attempt to model the details of protein synthesis, which involves necessarily all the other amino acids. Tryptophan is imported into the muscle compartment by the \( \text{l-transporter} \), \( V_{\text{trp}} \), with \( K_m = 64 \mu\text{mol/L} \) (41). To calculate the molarity of protein in the body, we used the estimate that 20% of the body is protein (42). For a 60-kg person, then 200 g/kg body mass is protein. Assuming a typical protein molecular weight of 50,000 (43), then 200 g is equivalent to 4 mmol, and therefore 200 g/L is \( \sim 4 \mu\text{mol/L} \) or 4000 \( \mu\text{mol/L} \). Our model calculated the steady state concentrations in muscle and liver to be 7232 and 6695 \( \mu\text{mol/L} \), respectively, which is reasonable because muscle and liver should have higher protein concentrations than many other tissues.

To determine the rate of protein synthesis we proceed as follows. In adult rats, \( \sim 5\% - 12\% \) of total protein weight is synthesized each day in the muscle (44-47). Thus, \( \sim 0.05 \times 7000 \mu\text{mol/L} \) should be synthesized in our muscle compartment per day or \( \sim 15 \mu\text{mol/L} \) per hour. Although the rate of protein turnover in rats is higher than in humans (48), the steady state concentration is not affected by the rate of protein synthesis, which generally will be in balance with the rate of proteolysis. Therefore, a higher or lower protein synthesis rate did not affect the predictions of the model. In the model, protein synthesis (\( \text{PRO} \)) was described (depending on available tryptophan) by the following formula:

\[
V_{\text{PRO}} = \frac{V_{\text{max}}(\text{trp})}{(K_m + \text{trp})}
\]

where we choose \( K_m = 25 \mu\text{mol/L} \), a typical cellular tryptophan concentration, and \( V_{\text{max}} = 30 \mu\text{mol/L/hour} \). Because free tryptophan in muscle is \( \sim 25 \mu\text{mol/L} \), the protein synthesis rate in the model is \( \sim 15 \mu\text{mol/L/hour} \).

In the liver, tryptophan is imported from the portal circulation by the \( \text{l-transporter} \), \( V_{\text{trp}} \), and leaves the liver into the peripheral circulation through the term \( k_1 \cdot \text{trp} \). Tryptophan is not just catabolized but is used in a variety of reactions including protein synthesis. Thus, we have reactions representing the use of free tryptophan for protein synthesis and the release of free tryptophan from proteolysis (\( \text{Lpool} \)). This is important because these processes buffer the free tryptophan in the liver. Without these processes, free liver tryptophan would be excessively sensitive to input from the gut. We take these reactions to be the same as in the muscle model discussed above. The velocities of kynurenine 3-monoxygenase (\( V_{\text{MO}} \)), 3-hydroxyanthranilic acid (\( V_{\text{HA}} \)), kynurenine aminotransferase catalyzing Kyn \( \rightarrow \text{KA} \) (\( V_{\text{AKAT}} \)), and kynurenine aminotransferase catalyzing HK \( \rightarrow \text{XA} \) (\( V_{\text{AXAT}} \)) have standard Michaelis-Menten kinetics; constants are given in Table 1.

The enzyme TDO shows both substrate activation and substrate inhibition:

\[
V_{\text{TDO}} = \frac{V_{\text{max}}(\text{trp})}{(K_m + \text{trp} + \frac{\text{trp}}{K_i})} \cdot \frac{2\text{trp}^3}{(20.15)^3 + \text{trp}^3}.
\]

The first term expresses Michaelis-Menten kinetics modified for competitive substrate inhibition (49). We take \( K_m = 43 \mu\text{mol/L} \), from ref 50 and the inhibition constant \( K_i = 170 \mu\text{mol/L} \), from ref 51. The second term requires more discussion. Saimo and Saimo (52) showed that the enzyme TDO exhibits substantial substrate activation. Increasing tryptophan concentration causes the TDO to become more saturated with the iron-porphyrin-containing coenzyme, making the more active holoenzyme. This effect approximately doubles the activity of the enzyme because the tryptophan concentration is increased above normal by a factor of 4. In our model, the normal liver concentration of tryptophan is 20.2 \( \mu\text{mol/L} \), so we chose the last factor in the formula for \( V_{\text{TDO}} \) so that it is equal to 1 when tryptophan is normal and to \( \sim 2 \) when tryptophan increases by a factor of 4.

\( V_{\text{AFD}} \). The enzyme aryl formamidase (AFD) has a \( K_m = 50 \mu\text{mol/L} \) (53,54) for N-formyl-kynurenine (\( \text{nfk} \)) and shows product inhibition by Kyn with \( K_i = 240 \mu\text{mol/L} \). We take the \( K_i \) of the reaction for Kyn to be 240 \( \mu\text{mol/L} \) and the inhibition constant \( K_m = 20 \mu\text{mol/L} \) from ref 57. PLP is an activator in these reactions. Thus, if \( A \) is an activator of an enzymatic reaction, the most natural way to model its effect is to multiply the \( V_{\text{max}} \) of the reaction by a term of the form

\[
\frac{c[A]}{K + [A]}
\]

This activation function is monotonically increasing and saturates (at level \( c \)), both reasonable assumptions. Then, \( c[K] \) is just the slope of the activation curve for low \([A] \) and \( K \) is the level of the activator that produces half-
maximal activation. The form of this term is exactly the same as if A were a substrate, and thus we believe that it is reasonable to call a K as has been done in the literature. Therefore, our PLP is a cofactor with a KPLP of 0.88 μmol/L (24).

\[
V_{\text{KYN1}} = \frac{V_{\text{max}}(\text{Kyn})(\text{PLP})}{(K_m + \text{Kyn}) (K_{\text{PLP}} + \text{PLP})}
\]

\[
V_{\text{KYN2}} = \frac{V_{\text{max}}(\text{hk})(\text{PLP})}{(K_m + \text{hk}) (K_{\text{PLP}} + \text{PLP})}
\]

Finally, the major downstream metabolites of tryptophan metabolism in the liver, namely Kyn, KA, AA, HK, XA, HaA, and QA, are released from the liver into the circulation and are excreted into the urine in the kidney. We simplify this process by having a linear removal term in each of the differential equations 3–9 of the form d\(y_i\)/dt, where \(y_i\) is the variable. Thus, predicted plasma concentrations will parallel liver concentrations of these metabolites. The coefficients \(d_i\), whose values appear in Table 1, were chosen so these metabolites have their normal concentrations at steady state.

### Results and Discussion

**Predicted steady state concentrations and velocities.** The steady state concentrations of tryptophan and its metabolites are shown in Figure 1. The full names of the variables in the model are indicated in Table 2, and the 15 differential equations formulated for the reactions shown in Figure 1 are provided in Supplemental Table 1. The model prediction for steady state tryptophan concentration in the peripheral circulation was slightly lower than in the portal circulation, as found in refs 37 and 38. Predicted steady state concentrations of tryptophan and tryptophan metabolites in the liver were close to values reported in the literature, which included hepatic tryptophan of 12.3 μmol/L reported in ref 59 and 8.2 μmol/L for Kyn (59). We estimated from other published data the following concentrations for hepatic tryptophan metabolites: 0.06–3 μmol/L for HK (60–61), 0.16–2.5 μmol/L for KA (60,62,63), 1 μmol/L for HaA (60), 0.16 μmol/L for AA (62), and 0.13 μmol/L for XA (62). The velocity split at Kyn corresponds to the velocities of the catalytic reactions catalyzed by kynurenine 3-monooxygenase, kynurenine aminotransferase catalyzing Kyn as has been done in the literature. Therefore, our PLP is a cofactor with a KPLP of 0.88 μmol/L (24).

Model predictions for the fluxes of these reactions were ~80, 10, and 10%, respectively, consistent with the reported dominance of the hydroxylation process forming HK (64,65).

**Simulation: predicted dependence of steady state metabolite concentrations and metabolic rates on vitamin B-6 status.** Changes as a function of decreasing relative cellular PLP concentrations. Relative values for vitamin B-6 status were 1 to 0.7 (adequate status), 0.7 to 0.3 (moderate deficiency), and 0.3 to 0.01 (deficiency). The results showed little or no change in the metabolites until cellular PLP concentrations reached ~50% of the normal value. It is known that in low vitamin B-6 status an increase in urinary Kyn, XA, and HK occurs. In the model predictions, urinary HK increased in moderate deficiency and markedly increased in more severe deficiency, as reported previously (66). The model also predicted that HK increased to the greatest extent, with the increase of Kyn about half that of HK, whereas XA increased almost as much as Kyn, as reported earlier (10). Data on urinary excretion of KA and AA vary in the literature. The predictions of our model for these 2 metabolites suggested that a slight decrease occurred in moderate vitamin B-6 deficiency, which became more pronounced in severe deficiency. The model also predicted that HaA excretion increased slightly in moderate deficiency and then decreased slightly in more pronounced deficiency, as reported (9). These results can be explained by considering that KYN2 is largely saturated with PLP but KAT2 is not (presumably) at typical cellular PLP concentrations, so the first effect in vitamin B-6 deficiency is a reduction in KAT2 but not in KYN2 activity. Thus, urinary HK and HaA both increased. However, more severe vitamin B-6 deficiency and consequently further cellular PLP depletion also caused the rate of KYN2 to be reduced, which yielded a reduction in HaA production and excretion. Concentrations of urinary tryptophan metabolites excreted in a 24-h period in adequate vitamin B-6 status (steady state) and vitamin B-6 deficiency (PLP concentration decreased to 5% of normal value in the model) were compared and are shown in Supplemental Table 2. HK, Kyn, XA, AA, and QA excretion were 70, 10, 10, 4, and 3% higher and KA and AA were 27 and 30% lower, respectively, in deficiency compared with adequate status. Predictions of urinary excretion rates of total tryptophan catabolites with and without simulated tryptophan load were unchanged by the deficiency, even though a change in the proportion of the metabolites occurred. These predictions showed that overall tryptophan degradation flux via TDO was not affected by vitamin B-6 deficiency (Supplemental Table 2).
tration increased by ~600%, peaking at ~3 h after the load in their study. In our simulation, Kyn excretion (Kynout) peaked at 2.5 to 3 h after the load but increased by only 280%. Møller (68) also measured plasma concentrations of tryptophan and Kyn and found that they peaked at 1.5 and 4 h after the load, respectively. In his study, plasma Kyn peaked at ~10% of the tryptophan maximum, whereas our Kyn peak was ~25% of the peak plasma tryptophan concentration. Finally, the author’s HK in the plasma also peaked at 4 h after the load and at approximately one-sixth of the Kyn concentration, whereas our simulation predicted an HK peak that was 70% of the plasma Kyn peak.

Wolf et al. (69) measured many metabolites in the urine over a 24-h period throughout a tryptophan load of 9800 μmol. Their results showed that excretion of KA increased by 300%, whereas in our model it increased by 300%. Kyn increased by 166%, and in our model it increased by 250%; XA increased by almost 272%, whereas in our model it increased slightly more than a factor of 3 (200%); HK increased by almost 166%, whereas the model predicted an increased HK excretion of 250%; and HaA increased by 93%, which was similar to the model prediction.

In a simulated tryptophan load concurrent with vitamin B-6 deficiency (5% of normal PLP concentration), the main changes were the decrease in KA and the increases in Kyn, XA, and HK (Fig. 4). Urinary KA decreased by 25% with the deficiency, whereas Kyn, XA, and HK were higher by 15, 10, and 70%, respectively, compared with their response after the tryptophan load in adequate vitamin B-6 status. This was also demonstrated by predictions of 24-h urinary excretion calculated as an AUC (Supplemental Table 2).

Simulation: dynamic changes of tryptophan and tryptophan metabolites due to meals. Amino acid concentrations after meals and the effects of the composition of the meals on those concentrations have been a target of study for many years. It is known that the protein content of a meal and the secretion of insulin after a meal influence plasma amino acid concentrations (70). For example, insulin has been shown to increase the uptake of branched-chain amino acids by the muscle decreasing their concentration in plasma (70). A meal with protein increases amino acid concentrations in plasma, whereas a protein-free diet causes a decrease in the concentrations of all amino acids (71,72).
Simulations of tryptophan inputs due to meals and the dynamic changes of tryptophan and its metabolites were conducted over a period of 24 h (Fig. 5). To simulate meals, we introduced into the gut one-fourth of the daily dose of tryptophan between 0700 and 0800 h, one-fourth of the daily dose of tryptophan between 1200 and 1300 h, and one-half of the daily dose of tryptophan from 1800 to 1900 h. The model showed increased tryptophan in plasma after each meal and an increase in its metabolite concentrations due to the increase in its oxidation. The model also showed a decline in plasma tryptophan overnight from 30 to 20 μmol/L, consistent with the decline reported after a meal with no protein (73).

Simulation: effects of induction of tryptophan oxidation and OC use on tryptophan metabolites. The increase of tryptophan oxidation due to an increase in the activity of TDO, the first enzyme in the hepatic Kyn pathway, has been shown by many investigators. Knox and Mehler (17) showed the induction of TDO activity by its substrate tryptophan, and Schütz et al. (74) showed that glucocorticoids increase the activity of the enzyme by increasing the synthesis of new mRNA and TDO protein. Additionally, the effects of estrogen on the activity of TDO were shown by Braidman and Rose (22) when they found that administration of 10 μg estradiol benzoate to female rats yielded an increase of TDO activity by 90%. Thus, it is believed that the use of OC agents exerts a similar induction of TDO.

We first simulated the induction of TDO activity by increasing the TDO V_max to 200% of the basal level (i.e., a doubling). The effects of increased TDO in the urinary excretion of Kyn metabolites in urine are shown in Table 3. Here, the model predictions were consistent with published findings from a study of TDO induction by hydrocortisone, which yielded an increase of urinary Kyn (16). Second, we simulated the increase of TDO activity in the presence of low vitamin B-6 status as has been reported for OC users (Fig. 6) (75,76). In these simulations, the V_max of TDO was increased by 80% and PLP concentration was lowered to 50% of the normal value in the model (moderate deficiency). The model predicted that TDO induction concurrent with vitamin B-6 deficiency augments the elevation of urinary Kyn, HK, and XA and a decrease in the other metabolites that are otherwise associated with vitamin B-6 deficiency. These effects were more pronounced when a simulation of tryptophan load was given in this simulation of OC users, as has been reported in the literature (32).

Further considerations regarding the model. We presented here a mathematical model that simulates tryptophan metabolism via the Kyn pathway. The model is compartmentalized in tissues that are involved in tryptophan metabolism, including intestinal lumen, liver, muscle, and brain. The model uses kinetic constant values and concentrations that are based on values reported in the literature for rodents and humans. We used values reported in rodents when information on metabolite concentrations or enzyme kinetics in humans was not available. This approach has been previously used in the development of other metabolism models in which predictions have been highly satisfactory [e.g., (33,34)]. Although this model uses simplified assumptions (e.g., the metabolite concentrations in plasma are considered to parallel those in liver and the excretion of the metabolites in urine is assumed to be linear, thus skipping the pass of the metabolites through the blood and the uptake by the kidney), we consider that this does not affect the predictions of the model because it successfully reproduces experimental data.

We focused most of our experiments in simulating the effects of induced low vitamin B-6 on the Kyn pathway in the liver, which allowed comparisons of model predictions with the
results of studies in which dietary vitamin B-6 deficiency has been associated with an impaired Kyn pathway. Early studies showed the effects of dietary vitamin B-6 deficiency on tryptophan metabolism by measuring tryptophan metabolite concentrations in urine after tryptophan loads. However, Schaeffer et al. (77) showed that a tryptophan load test of 24.5 μmol (5 mg) of L-tryptophan/100 g body weight was not effective in detecting vitamin B-6 marginal deficiency in adult female Long-Evans rats. Our model was useful in assessing changes in HK, KA, and AA in the marginal (moderate) deficiency range, which shows the usefulness of the model to study the effects of a mild vitamin B-6 deficiency. Indeed, the model predictions agreed well with human data obtained in our laboratory from marginal vitamin B-6–deficient participants in whom plasma HK was significantly increased and KA was significantly decreased without a prior tryptophan load (V. da Silva, L. Rios-Avila, Y. Lamers, M. Ralat, Ø. Midttun, E.P. Quinlivan, T.J. Garrett, B. Coats, M.N. Shankar, S.S. Percival, Y.-Y. Chi, K.E. Muller, P.M. Ueland, P.W. Stacpoole, J.F. Gregory, unpublished results). However, to observe the effects on the other metabolites, the model requires a simulation of a more pronounced deficiency.

On the basis of the structure of the Kyn pathway (Fig. 1) we can predict that the urinary excretion of KA and AA would decline during vitamin B-6 deficiency and that Kyn would increase because the exit processes are being restricted. The cases of XA and HaA in vitamin B-6 deficiency are more interesting because there are 2 conflicting effects. First, HK will clearly increase because the flux in will be higher and KAT2 and KYN2 will be somewhat dysfunctional due to reduced PLP availability. The effects of vitamin B-6 deficiency on XA and HaA are due to the reduced activity of both KAT2 and KYN2. These concentrations are not independent because as KYN2 becomes more dysfunctional, the resulting increase in HK will tend to increase XA. In general, the lack of effect on vitamin B-6 status over a large range is because the K_m values of KYN2 and KAT2 for PLP are very low (0.88 and 0.85 μmol/L, respectively) so that their rates are only weakly affected by a reduction in cellular PLP concentration that is generally well above the K_m values.

We used the model to simulate tryptophan load tests in the presence or absence of vitamin B-6 deficiency. In adequate vitamin B-6 status, the tryptophan load induced an increase in all the tryptophan metabolites in blood and urine as expected. In the presence of vitamin B-6 deficiency, the increase in HK, Kyn, and XA was consistent with the effects of deficiency on the PLP-dependent enzymes. However, our XA response to vitamin B-6 deficiency (under tryptophan load or not) does not seem to be as

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**FIGURE 5** Dynamic changes of tryptophan and tryptophan metabolites after simulation of tryptophan input due to meals in a period of 24 h. The tryptophan inputs were simulated by delivering one-fourth of the daily dose of tryptophan between 0700 and 0800 h, one-fourth of the daily dose of tryptophan between 1200 and 1300 h, and one-half of the daily dose of tryptophan from 1800 to 1900 h. The word "out" denotes urinary metabolites. AA, anthranilic acid; bptrp, tryptophan in peripheral circulation; gut, tryptophan in gut; HaA, 3-hydroxyanthranilic acid; HK, 3-hydroxykynurenine; hptrp, tryptophan in portal circulation; KA, kynurenic acid; Kyn, kynurenine; Ltrp, tryptophan in liver; Mtrp, tryptophan in muscle; QA, quinolinic acid; XA, xanthurenic acid.

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**TABLE 3** Model prediction of the effect of extent of induction of TDO activity on urinary excretion of tryptophan metabolites

<table>
<thead>
<tr>
<th>TDO percentage of basal activity</th>
<th>Trp metabolites in urine</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>100 (baseline)</td>
</tr>
<tr>
<td>Kyn</td>
<td>2.00</td>
</tr>
<tr>
<td>HK</td>
<td>1.43</td>
</tr>
<tr>
<td>KA</td>
<td>1.40</td>
</tr>
<tr>
<td>HaA</td>
<td>3.44</td>
</tr>
<tr>
<td>AA</td>
<td>3.21</td>
</tr>
<tr>
<td>QA</td>
<td>3.37</td>
</tr>
</tbody>
</table>

1 AA, anthranilic acid; HaA, 3-hydroxyanthranilic acid; HK, 3-hydroxykynurenine; KA, kynurenic acid; Kyn, kynurenine; QA, quinolinic acid; TDO, tryptophan 2,3-dioxygenase; Trp, tryptophan; XA, xanthurenic acid.
but rather responded to the varying PLP concentrations. which the activities of the enzymes were not completely reduced increase in Kyn, HK, XA, and HaA and a decrease in KA and AA reduction in the enzyme activities. Our model predicted an normal. In that model, Kyn activity was not affected by the it decreased when KAT activity was reduced and KYN was kept different results for each metabolite analyzed. For example, HaA, KA, and XA after simulations in which both KYN and reduction in the activity of each enzyme separately showed the coenzyme-availability. Our model is also useful to predict effects of induction or inhibition of enzyme activities. For example, our model was used to simulate the induction of the activity of TDO as reported for glucocorticoids and sex hormones (18,22) and the induction of TDO activity concurrent with low plasma concentrations of PLP as has been reported in OC users (22,76,79). Inhibitory effects of enzymes also can be simulated, thus providing useful information to interpret abnormalities in the metabolites under different conditions including dietary and genetic disorders.

A computer simulation of tryptophan metabolism via the Kyn pathway in the liver was previously published (80). However, this model did not account for the distribution of metabolites in other tissues nor for enzyme regulation. This previous model predicted the effects of vitamin B-6 adequacy and deficiency (incorrectly designated by that author as “pyridoxal” deficiency) on metabolite concentrations in conditions of presence or absence of the PLP coenzyme. Changes in enzyme activities were simulated by eliminating the coenzyme-dependent rate constants. The results showed a decrease in HaA, KA, and XA after simulations in which both KYN and KAT activities were lost. However, because the enzymes are not equally affected by vitamin B-6 deficiency, simulation of the reduction in the activity of each enzyme separately showed different results for each metabolite analyzed. For example, HaA increased when KYN sensitivity for the coenzyme was lost and the activity of KYN aminotransferase was kept normal, whereas it decreased when KAT activity was reduced and KYN was kept normal. In that model, Kyn activity was not affected by the reduction in the enzyme activities. Our model predicted an increase in Kyn, HK, XA, and HaA and a decrease in KA and AA as a result of the simulation of a general vitamin B-6 deficiency in which the activities of the enzymes were not completely reduced but rather responded to the varying PLP concentrations.

In summary, the results from the model simulations reliably reproduce experimental data and confirm the validation of the model to be used as a tool to study tryptophan metabolism. Moreover, the model predictions confirmed the increases in Kyn, HK, and XA and the decreases in AA and KA in vitamin B-6 deficiency. A further application of the model will be its use as a platform to develop a model to study extrahepatic tryptophan metabolism, the specific role of IDO, and effects of potential redistribution of PLP in inflammatory tissues (81).

Acknowledgments
J.F.G., H.F.N., and M.C.R. designed the study; H.F.N. and M.C.R. constructed the model; H.F.N., M.C.R., and L.R.-A. conducted simulation studies; H.F.N., M.C.R., L.R.-A., and J.F.G. wrote the manuscript; J.F.G had the primary responsibility for the final content; and all authors contributed data on which the model is based. All authors read and approved the final manuscript.

Literature Cited


