Nutrition and pharmacology: general principles and implications for HIV1–4

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ABSTRACT

Food and nutrition play an intimate and inextricable role in all aspects of drug metabolism, safety, and effectiveness. Antiretroviral therapies (ART) have assumed a preeminent position in the prevention, care, and treatment of HIV and its comorbidities. The interaction between food, nutrition, and ART has become an expanding area of interest both in terms of clinical standards of care and as a target for research. Since the original review of this topic by the WHO in 2005, much has been learned (8). This article contains a review of what is known about the general relationships between nutrition and pharmacology, as well as issues specific to ART, with particular attention to their use in low- and middle-resource settings. The importance of food and nutrition on the bioavailability of drugs and vice versa has been an area of historical interest. However, much has been learned about the importance of nutritional status on drug metabolism, distribution, and effectiveness. The impact of traditional therapies (herbal/botanical) is highlighted as an area of clinical concern and one in need of further research. Additional attention is focused on the impact of individual micronutrients on drug pharmacokinetics and pharmacodynamics. Finally, attention is given to the nutritional implications of the metabolic consequences of ART, which include the potential impact of “colliding epidemics” of infection (eg, HIV, tuberculosis) and noncommunicable diseases. Much has been learned, but much remains to be accomplished to ensure the effective integration of nutritional considerations into the effective and safe use of ART. Am J Clin Nutr 2011;94(suppl):1697S–702S.

INTRODUCTION

Nutrition may be defined as the sum total of the processes involved in the taking in and use of food substances through which growth, repair, and maintenance of activities of the body as a whole or in any of its parts are accomplished. The processes of nutrition consist of ingestion, digestion, absorption, metabolism, functional use/activation of dependent systems, and excretion. All these processes are similarly integral to how the body takes in and uses therapeutics/drugs, which include antiretroviral therapies (ART).

Not only do drugs and nutrients share these same processes, their availability and function are also intimately and inextricably entwined. The body’s ability to process foreign substances depends on metabolic systems that rely on essential nutrients (vitamins, minerals, fatty acids, and so forth) obtained through diet. Yang et al (1) offered a teleologic explanation of the synergism between diet and the detoxification of foreign substances, based on the evolutionary change to a complex diet paradoxically rich in essential nutrients but that also contained botanical sources of potentially toxic chemicals. The need to find sources of essential nutrients was linked to the need to develop mechanisms for detoxification of the accompanying toxins. These mechanisms, in turn, became dependent on many of the same essential nutrients, which created interdependence between nutrition and detoxification. In our modern world it is not just exposure to toxins in the environment, but also the response to pharmacologic substances, which, in much the same way as the early botanicals, are being used on a trial-and-error basis to improve the human condition. As is the case with exposures to potentially poisonous herbals/botanicals, exposure to modern medicines can have a healthful or hurtful outcome. The response, either therapeutic or toxic, to any foreign substance is contingent on numerous factors that include stage of development, genetics, general health, and nutritional status.

The general relation between diet, nutrition, and pharmacology is conceptualized in Figure 1. It is discussed in the context of several core concepts, which are outlined in Table 1. Within the context of pharmacokinetics and pharmacodynamics several pathways exist by which nutrition might affect drugs and vice versa (Table 2).

In the context of potential food/nutrient-drug interactions, most of the available information used clinically is focused on factors that pertain to drug pharmacokinetics and, in particular, bioavailability (eg, foods that may affect drug absorption because of physicochemical solubility relationships). There is, however, a historical knowledge base with regard to the role of specific nutrients and pharmacodynamic processes [ie, those Phase I metabolic systems, mixed-function oxygenase/cytochrome P450 (CYP), responsible for the activation, transport, and excretion of...
drugs (2)]. Examples of these types of interactions are listed in Table 3.

In addition, a greater appreciation has emerged for the interaction between genes [which includes genetic polymorphisms in mixed-function oxygenase and related systems (4)] and developmental changes [eg, in infants, pregnancy, and lactation (5, 6)]. Finally, it is also important to note that the disease process (eg, inflammation or response to infection) elicits a unique response to nutrient homeostasis that affects nutrient absorption, availability, and response to treatment (7). This is best exemplified by the anemia of infection and the impact on iron status, and the impact of inflammation via the acute-phase response on the key carriers of iron (eg, ferritin) (8). Thus, a close relation exists between the body’s response to illness, drugs, nutrients, and the requisite systems involved in the functional use of the drugs and nutrients. Many of these concepts are appropriately applied to environmental toxicants as well and have been discussed in greater detail elsewhere (2).

TABLE 1
Core concepts of pharmacology

<table>
<thead>
<tr>
<th>Concept</th>
<th>Description</th>
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<tbody>
<tr>
<td>Pharmacodynamics</td>
<td>The study of the biochemical and physiologic effects of drugs on the body or on microorganisms or parasites within or on the body and the mechanisms of drug action and the relation between drug concentration and effect. A prime example is drug-receptor interactions.</td>
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<tr>
<td>Pharmacokinetics</td>
<td>The action of drugs in the body over a period of time, including the processes of absorption, distribution, localization in tissues, biotransformation, and excretion.</td>
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<tr>
<td>Phases of drug metabolism</td>
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<tr>
<td>Phase I</td>
<td>Oxidation reduction reactions that result in activation, deactivation, or preparation for eventual elimination. These reactions occur primarily in the liver, but also in other tissues (eg, lungs, kidneys, gastrointestinal tract) and use the synergism between 3 primary components:</td>
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<td></td>
<td>• MFO, which include cytochrome P450 enzymes (oxidation)</td>
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<td></td>
<td>• NADPH-P450 reductase (reduction)</td>
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<tr>
<td></td>
<td>• Phospholipid (phosphatidylcholine or lecithin). The phospholipid component provides stability for these membrane-bound enzymes. The dependence on stable membranes introduces the potential for damage due to lipid oxidation, which implies a role for antioxidants in the protection of the integrity of the MFO system.</td>
</tr>
<tr>
<td>Phase II (conjugation reactions)</td>
<td>The attachment of substances, which yields a more polar and water-soluble substance and thereby facilitates elimination.</td>
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$^{1}$ MFO, mixed-function oxygenase.
A more direct conceptualization of the model presented in Figure 1 as it specifically pertains to HIV infection and its treatment is presented in Figure 2. Aside from the mechanisms described above, a number of ART-specific interactions may occur, primarily via the key pathways responsible for drug metabolism. These include drug-drug interactions, drug-botanical/herbal interactions, and drug-nutrient interactions. Whereas the principles may differ, the core mechanisms may often be similar and be mediated via the CYP drug-metabolizing systems, primarily in the liver and gastrointestinal tract.

ART drug-drug interactions were reviewed recently by Fletcher (18), who observed that a specific enzyme, CYP3A4, the most abundant isozyme of the cytochrome system in the human liver, is responsible for the metabolism of ~60% of HIV-related drugs. Most protease inhibitors and nonnucleoside reverse transcription inhibitors are CYP3A4 substrates. Therefore, it is critical to recognize the potential for either inhibition or induction of this enzyme in the use of these drugs. A number of examples are cited in which one drug affects the use of another via this mechanism (18). Although the potential for drug interactions is recognized, what is less well acknowledged is the potential for interactions between ART and other commonly used substances through this same mechanism.

The CYP system has been shown to be the target of a number of other ART interactions with substances commonly used by HIV-infected patients in domestic/US and international settings. Mills et al (12) reported on the impact of African herbal medicines on antiretroviral metabolism and noted specifically a significant inhibition of CYP4A4 by 2 common African herbal remedies (African potato and Sutherlandia).

Flavonoids are a group of substances that occur naturally in fruit (including cocoa), vegetables, beverages (tea, wine), and many dietary supplements (e.g., ginkgo biloba) or herbal remedies. They have been touted as having beneficial effects on a number of health conditions. Of particular relevance to this discussion is evidence that indicates that these compounds significantly affect the activity (induction or inhibition) of CYP isoforms and other related drug-metabolizing enzymes (19, 20).

The historical approach to micronutrients in the context of ART, and more broadly with HIV, is to limit the conversation to...
micronutrient insufficiency (21). However, as evidenced by the discussions above and below, both low and high micronutrient exposure and status must be considered in the context of ART use. With specific regard to nutrients, several have been the focus of investigation in this context.

Vitamin C has been shown to significantly affect the regulation of several of the key CYP enzymes, which include isoforms of CYP, family 3, and subfamily A (CYP3A), some of which were shown to be decreased by vitamin C deficiency (22). Conversely, the observation of Slain et al (23) is of particular interest in the context of ART use. Their study involved the evaluation of the pharmacokinetics of a specific protease inhibitor (indinavir) in HIV-uninfected subjects who were receiving doses of vitamin C that ranged from 800 to 1000 mg/d. They reported that “concomitant administration of high doses of vitamin C can reduce steady-state indinavir plasma concentrations.” Did the excess vitamin C result in increased CYP450 metabolism of indinavir? The clinical implications of these findings have not been established.

Vitamin D has been highlighted as an important nutrient of concern for the general public (24). Of relevance here, vitamin D has also been identified as a nutrient of concern in the context of ART use. Their study involved the evaluation of the implications of these findings have not been established.

Protein Deficiency:

- Deficiency: ↓ rate of metabolism
- Excess: can ↑ rate of metabolism

Lipids

- Deficiency (or diet high in saturated fatty acids): ↓
- Excess (or diet high in polyunsaturated fatty acids): ↑ activity and induction of MFO enzymes

Carbohydrates

- Excess: ↓

Vitamin C

- Deficiency: ↓
- Excess: ↑ P450 activity

Vitamin B-6

- Deficiency: ↓

Thiamine

- Deficiency: ↑ activity of cytochrome P450
- Excess: ↓ (both reductase and P450)

Riboflavin

- Deficiency: ↓ or ↑ depending on the severity

Vitamin E

- Deficiency: ↓
- Excess: no reported effect

Iron

- Deficiency: ↓ and ↑
- Excess: ↑ in microsomal lipid peroxidation

### TABLE 3
Examples of the impact of specific nutrients on Phase I/MFO metabolism

<table>
<thead>
<tr>
<th>Nutrient</th>
<th>Effect on MFO metabolism</th>
<th>Potential mechanism(s)</th>
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<tbody>
<tr>
<td>Protein</td>
<td>Deficiency: ↓ rate of metabolism</td>
<td>↓ Protein synthesis; ↓ in synthesis of other elements, such as hormones, involved in enzyme induction</td>
</tr>
<tr>
<td>Lipids</td>
<td>Deficiency (or diet high in saturated fatty acids): ↓</td>
<td>↓ Activity of MFO possibly connected to the requirement for polyunsaturated fatty acid in the β-position of phosphatidylcholine (lecithin), which is an essential component of the MFO system</td>
</tr>
<tr>
<td>Carbohydrates</td>
<td>Excess: ↓</td>
<td>Secondary effect due to ↓ protein or possibly inhibition of P450 via ↓ in supporting enzyme components</td>
</tr>
<tr>
<td>Vitamin C</td>
<td>Deficiency: ↓</td>
<td>Alterations in activities of P450 and P450 reductase mediated via either ↑ or ↓ in the expression of specific CYP isoforms in excess or deficiency states</td>
</tr>
<tr>
<td>Vitamin B-6</td>
<td>Deficiency: ↓</td>
<td>↓ Synthesis of heme; possible impairment of protein synthesis</td>
</tr>
<tr>
<td>Thiamine</td>
<td>Deficiency: ↑ activity of cytochrome P450</td>
<td>↑ Activity of specific P450 isoforms and perhaps other enzymes in deficiency by an unknown mechanism.</td>
</tr>
<tr>
<td>Riboflavin</td>
<td>Deficiency: ↓ or ↑ depending on the severity</td>
<td>Excess may be due to ↓ substrate binding</td>
</tr>
<tr>
<td>Vitamin E</td>
<td>Deficiency: ↓</td>
<td>↓ Reductase activity but ↑ P450 activity, such that the metabolism of some drugs will be ↑, whereas others may be ↓</td>
</tr>
<tr>
<td>Iron</td>
<td>Deficiency: ↓ and ↑</td>
<td>Because activities of P450 and reductase are unaffected, it may be due to reduction in antioxidative mechanisms (eg, protection of the lecithin component): lack of effect from excess may be due to rapid metabolic clearance of vitamin E isomers via P450 (3)</td>
</tr>
<tr>
<td></td>
<td>Excess: ↑ in microsomal lipid peroxidation</td>
<td>Differential effects on various components of the MFO system. ↑ Lipid peroxidation could lead to damage to the integrity of the system</td>
</tr>
</tbody>
</table>

Excess may be due to reduction in antioxidative mechanisms (eg, protection of the lecithin component): lack of effect from excess may be due to rapid metabolic clearance of vitamin E isomers via P450 (3) Differential effects on various components of the MFO system. ↑ Lipid peroxidation could lead to damage to the integrity of the system
across all segments of the population, including HIV-infected people (27).

Vitamin A continues to hold a position of great interest in the global health dialogue and vitamin A insufficiency continues to be a major concern (28, 29). Again, most of the focus of the public health community, generally and in the context of HIV, has been on insufficiency. Several lines of evidence have indicated that vitamin A supplementation may be an issue of concern, particularly for HIV infection (30, 31).

With specific regard to drug metabolism, evidence similar to that for vitamin D and vitamin C exists for an important role for vitamin A. Vitamin A has 3 active forms (retinol, retinal, and retinoic acid) and a storage form (retinyl ester):

\[
\text{Retinyl ester} \leftrightarrow \text{Retinol} \leftrightarrow \text{Retinal} \rightarrow \text{Retinoic acid (I)}
\]

Investigators have reported that 3 primary forms of vitamin A (9-cis-retinal, 9-cis-retinoic acid, and all-trans-retinoic acid) induce CYP3A expression at messenger RNA, as well as enzyme activity levels in both liver and intestinal cells (32). The study by Chen et al (33) showed that, in their model, retinoids were able to alter drug metabolism through CYP3A induction. These reports reinforce the notion that the use of vitamin A supplements may have implications mediated through this role in drug metabolism, which will require further research and clinical attention.

It is clear that in light of what we know about the processes of nutrition and pharmacology, the view of nutrition and specific nutrients must be expanded beyond the desire to prevent and treat undernutrition. Moreover, the use of dietary supplements, whether in the form of traditional therapies (herbal/botanical) or as nutrient supplements intended to correct presumed insufficiency, must be viewed in a larger context. This context must include the health of the individual [the presence or absence of active disease (either communicable and/or chronic), developmental stage, nutritional status (replete or deplete)] and the potential impact of these bioactive substances on all aspects of pharmacokinetics/pharmacodynamics.

CONCLUSIONS

This article has provided an overview of the potential role of nutrition and specific nutrients and other dietary substances in the safe and effective use of ART. A description of these phenomena is just the first step. In terms of care and treatment, what do we need to do to improve our clinical approach? What does it all mean and what can we do?

To start, we have to look more carefully at potential drug-nutrient interactions in the clinical setting and to ask the correct questions, particularly in settings in which poor nutrition might be anticipated. To support our ability to do that, we need to delineate more clearly the role of nutrients in pharmacology beyond just bioavailability. We also need to look more critically at the nutritional context in which people live and to ask some basic questions at the initial visit, such as “Are you hungry?,” “What are you eating?,” and “Are you using dietary supplements and/or traditional therapies?” From a public health perspective a need exists to examine more closely the “one-size-fits-all” public health approach to nutrition, particularly as it relates to micronutrients and especially in the context of infections such as HIV and their treatment (34). We cannot limit our focus to only amelioration of undernutrition. The provision of additional micronutrients in some scenarios may, in fact, exacerbate rather than ameliorate problems. Our ability to determine when that might be the case will depend on our evolving knowledge about these complex relationships as well as our access to the tools needed to evaluate them (eg, biomarkers) for accurate and reliable assessment of nutritional status.

Much work has been done to address the important synergies between food, nutrition, and the safe and effective implementation of ART use to prevent and treat HIV. However, continued effort and vigilance is needed to ensure that these issues are fully integrated into prevention, care, and treatment programs. Only through such efforts can we achieve the goals of all the global efforts to address this compelling and ongoing global health challenge.

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REFERENCES

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