Gut microbiota: metabolic programmers as a lead for deciphering Ayurvedic pharmacokinetics

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Gut microbiota has tremendous potential to influence human physiology in terms of health and disease. Factors like improper diet, lifestyle and stress contribute in development of dysbiosis. Ayurveda emphasizes on the importance of mahasrotasa (GIT system) and its homeostasis in maintenance of health. Pharmacokinetic principles of Ayurvedic pharmacology include Vipaka, commonly referred to as ‘postdigestive irreversible process’ which is closely associated with biotransformation of drugs caused by microbiota. There is evidence that the metabolism of polyphenols by the microbiota can influence their bioactivity. Consequently, inter-individual variation in microbial metabolism could have implications for health benefits of phytochemicals.

Keywords: Ayurveda, gut microbiota, pharmacokinetics, metabolic programmes.

As science advances, it opens an arena of novel leads in every field. The seers of Ayurveda have also quoted that in order to obtain thorough knowledge about one shastra (science), one needs to get acquainted with the other scientific literature which complements the former, one should be multifaceted in approach¹.

With the advent of sophisticated instrumentation and omics approaches, microbiome is the most trending field that deals with the mechanism of equilibrium of our body. Recent studies have reported that a host of trillion microbes residing in our body govern the health status. It has been termed as the metabolic organ², wherein the emphasis from the vital organs is now shifting to the gut, especially microflora of human gut for its essential role in the establishment of homeostasis. The microbial ecosystem of our gut comprises native species that are permanently found and those variable ones colonizing transiently. The whole of microorganisms, their genes and metabolites are together termed as ‘microbiome’.

Significance of diet (ahara) and medicine (aushadha) in Ayurveda

Ayurveda, the science of life, bestows certain unique principles in relation to the fundamentals of physiology of our body. Among the three pillars (trayopastambha) quoted by the seers, diet (ahara) is an essential component¹. The significance of ahara has been quoted in four separate chapters (swasthya chatushka) of Charak Samhita⁴. Herein, they have emphasized that the overall maintenance of health is strongly associated with the quantity (matra) of diet (ahara)³. The optimum food consumption further contributes in establishment of balance (bala), luster, radiance (varna), vitality and immunity (oja)⁶. Later, Charak adds that the quantity of ahara to be consumed is closely dependent upon bala (strength) of agni (metabolism)⁷.

The diet to be consumed and the status of agni (energy of metabolism) for its metabolism are interdependent. The strength of agni influences digestion in every season which has been given in the rutucharya (seasonal regimen). Hence, Acharyas have also suggested the diet regimen to be followed during each season (rutu)⁸.

The suitable food for consumption is that which without disturbing the homeostasis of tissues (dhatus) or dosha of the body gets digested as well as metabolized in appropriate time. Thus, it ought to possess individual variations due to variable digestive powers.

Ayurveda has classified the quantity of food (ahara matra) into two types, viz. sarvagraha and parigraha⁹. Sarvagraha refers to the amount as a whole (viz. all food constituents served in a meal together), whereas the individual component portions of the meal, viz. quantity of chapati, daal, etc. is parigraha. Ayurveda also emphasizes upon the pertinent consumption of shadrasa (diet possessing foods with six tastes). Any error in consumption of any individual component of ahara affects the health of tissues which results into ageing or pathology¹⁰.

Modern nutritionists have focused more on the proportions of carbohydrates, fats and proteins in diet, which comes under the parigraha aspect of matra. Ayurvedic
approach to food and dietetics is different from the conventional Western approach. *Charaka Samhita* describes the eight ethics of ahara vidhi, i.e. (1) prakriti (nature of food substances), (2) karana (processing), (3) samyoga (combination), (4) rashi (proportion), (5) desha (region and climate), (6) kala (time), (7) upayogasamstha (rules of consumption), and (8) upayokta (the consumer)\(^{12}\).

**Digestion in ayurveda**

The avasthapaka (digestive process) basically consists of three phases according to Charaka, viz. Madhura, amla and katu\(^{12}\).

**Madhura avasthapaka:** It commences as soon as food comprising shadrasa comes in contact of bodhaka kapha (component for taste perception from salivary secretions) and continues up to amashaya (stomach). Here, some thin and frothy kapha (one of the three bodily doshas responsible for stability, unctuousness, lubrication, immunity and cohesion) is generated. It is often correlated with the breakdown of starch by salivary amylase and partial digestion in fundal part of stomach, where polysaccharides are broken down to monosaccharide.

**Amla avasthapaka:** Later, the partially digested (pakva and apakva) food comes in contact with amashaya (pyloric part of stomach) with the environment consisting of HCl that facilitates digestion of proteins and the resultant acidified chyme (vidagdha) is propelled into the duo-denum ahead where achha pitta (bile) is produced.

**Katua avasthapaka:** When this food product reaches pak-vashaya (large intestine), it gets further metabolized and hydrolysed by the agni (enzymes) and takes a bolus-form (paripindita) resulting in pungent taste. This stimulates vata dosha. This phase is often equated with the last stage of digestion occurring in the colon along with formation of indole, skatol and excretory products.

The avasthapaka thus can form a part of mechanical and biochemical digestion process in human physiology.

**Difference between vipaka and avasthapaka**

The vipaka (post-digestive metabolism) is the final transformation of food while avasthapaka is the initial phase. Vipaka commences only after the avasthapaka has ceased. This has been mentioned in the Chakrapani commentary\(^{12}\). The term ‘bhinn kala’ (time factor) denotes that these two stages are different. The actual vitiation of doshas and which in turn intimates the end of metabolism (rasa mala vivek) is termed vipaka. The factors that determine vipaka are dosha, mala (faecal matter, urine) and effect on shukra dhatu (tissue responsible for reproduction), which is the last tissue in sequential nutrition as per Ayurveda. Thus, vipaka is closely associated with secretion and absorption functions after the mechanical and chemical digestion. It is quoted as karma nishthaya, i.e. it manifests its properties at the end of complete digestion of food/drug\(^{13}\).

This again exhibits that vipaka is the ultimate transformation of food/drug (dravya) of particular perceptive attribute into final madhura, amla or katu tastes (rasas)\(^{14}\). Thus, it suggests that vipaka is an entity which comes into play after drug/food digestion for tissue nutrition or cellular nutrition.

Unlike conventional medicine, according to Ayurveda any dravya be it medicine (aushadha) or diet (ahara) will have to pass through vipaka for metabolism and manifestation of the action. The difference is of quantity and functions, where ahara is more in amount and results in nutrition, whereas medicines (aushadha) are administered in less amounts and have target actions but not nutritive as a whole to our body. Acharya Kashyapa has termed food (anna) as super medicine (mahabhaishyajya)\(^{15}\).

**Discussion**

Diet composition is a pivotal factor in modulating the gut microbial populations. Several studies have been performed to explore the inter-action between food and gut microbiota in relation to individual health status\(^{16}\). There are references in the treatise wherein a list of foods fit for daily consumption has been enlisted. Besides, the harmful effect of incompatible food (viruddha anna) has also been denoted which again relates with gut microbiota, i.e. incompatible food results in dysbiosis. This can be correlated with dhatvagnimandya that produces systemic effects and sets in pathologies of different diseases. Trillions of bacteria are distributed throughout the gastrointestinal tract from mouth to anus, which facilitate digestion and intestinal homeostasis. The components of diet are highly influential for colonization of microbiota. The variety of flavours (shadrasa) in a meal as stated before often point out the existence of many classes of bio-active compounds, that are necessary for constituting the needed consortium of microflora. Thus, the presence and quality of the consortium decides the fate of metabolism of the ingested food. The probable rationale behind ancient ‘thali’ diet prevalent in many parts of India has been delineated in a recent paper\(^{17}\). Similarly, in Ayurveda, the strength of agni has to be adjudicated on the basis of jarana shakti – capacity of digestion, that ultimately resides over the gut microbiota. Ayurveda considers the derangement of agni, i.e. its pachana karma (metabolism) to be the fundamental etiology for all anomalies. Similarly, unhealthy status of microbiota consortium due to wrong diet results in alteration of metabolism which thereby mirrors the physiopathology of nutritional disorder (closely associated with grahami). The anatomical distribution of gut microbiota resembles the mahasrotasa
Role of gut microbiota on system physiology

The gut microflora is referred to as a key endocrine organ, as it has significant impact on physiology and different human disease states which convert nutritional signals from environment that are hormone-like signals\(^\text{16}\). Gut microbiota aids in routine digestion and metabolizes the substrates producing cytokines, hormones, antigenic responses, activation of cascades, producing neuronal signals, etc. They also produce immune responses on various bodily systems of the host other than the gut.

Ayurvedic vis-à-vis conventional ideology of the digestive system

As discussed above, Ayurveda has its own unique way of stating metabolism of diet and drugs in form of vipaka. This metabolic energy is termed ‘agni’ in Ayurveda. Acharya Charak has stated the seat of agni to be grahani, and is said to be located above umbilical region\(^\text{19}\). This coincides with the major seat of gut microbiota. Jatharagni produces the sarakitta vibhajana of ahara (food) and helps in the assimilation of food. This prasada (essence of food) portion is then passed on to bhutagni for further metabolism\(^\text{20}\). This can be compared to various functions that gut microbiota perform in metabolism of carbohydrates, proteins, and fats and other nutrients in the body.

Gut microbiome is a key component in varied metabolic reactions such as digestion of carbohydrates, starch, sugars, oligosaccharides, proteins, non-digestible dietary fibre, transformation and degradation of the conjugated bile salts and oxalate-based complexes. The gut microbiota assist in the enzymatic reaction and stimulate production of various cytokines and other factors that can act either locally or systemically\(^\text{21}\). Commensals like \textit{Bacteroides thetaiotaomicron} and \textit{Lactobacillus innocua} produce antimicrobial proteins such as cathelicidins, (pro) defensins and C-type lectins\(^\text{21}\). The gut microbiota of healthy individual has high-energy metabolic markers such as pyruvic acid, citric acid, fumaric acid and malic acid\(^\text{21}\). The residual products of carbohydrate metabolism from the digestive system are further degraded by organisms (\textit{Bacteroides}, \textit{Roseburia}, \textit{Bifidobacterium}, \textit{Fecalibacterium} and \textit{Enterobacteria} sps) producing SCFA like butyrate, propionate and acetate. These are proven to be high-energy yielding sources\(^\text{21}\) (Table 1). \textit{Bacteroides} \textit{intestinalis}, \textit{Bacteroides} fragile and \textit{Escherichia coli} process bile salts and convert them into secondary products such as deoxycholic and lithocholic acid in the gut, which in turn lead to lipid peroxidation and breakdown of steroids such as cholesterol\(^\text{21,22}\).

Various reports show the possible role of commensals, namely \textit{Oxalobacter formigenes}, \textit{Lactobacillus} species and \textit{Bifidobacterium} species in reducing the chances of oxalate stones formation in the kidney\(^\text{21}\). Pancreatic lipase requires colipase for lipid digestion, commensal \textit{Bacteroides thetaiotaomicron} alleviates the expression of colipase and thus enhances lipid hydrolysis\(^\text{23}\).

Ghrelin release is controlled by cholecystokinin (CKK), glucagon-like peptide-1 (GLP-1), peptide tyrosine tyrosine (PYY), which is eventually modulated by gut microbiota. High intake of lipids and proteins stimulates production of CKK, GLP-1 and PYY pancreatic polypeptides by few intestinal microorganisms. These peptides are responsible for deciding the meal size and long-term energy balance\(^\text{24}\). Gut microbiota metabolize bile salts by generating bile salt species that are distinct from those of the host\(^\text{25}\).

Ayurvedic vis-à-vis conventional ideology of the respiratory system

Each system in Ayurveda has been explained with the term ‘srotasa’, i.e. channels. In case of respiratory system, Pranavaha srotasa has been mentioned. Also, Charak has mentioned the role of ‘mahasrotasa’ which is gastrointestinal system as the main seat of prana related to respiratory system\(^\text{26}\).

The intestinal microbiota influences the respiratory microbiota by direct seeding of microbes from gut to the airway through the gut–lung axis or transferring the metabolites from the gut, short-chain fatty acids\(^\text{27}\). In the intestine, commensals help in production of mucus layer, antimicrobial proteins and IgA, which provides a physical barrier against pathogenic microbes\(^\text{21}\). It has been reported that Paneth cells maintain homeostasis with intestinal microbiota by producing antimicrobial peptides. Reg III (gamma) separates the intestinal and cell wall microbiota; it protects from Gram-positive bacteria which impact pulmonary immunity against fungal pathogens\(^\text{28}\). An experiment with germ free mice showed production of Th17 cells in the intestine after induction of commensal bacteria, which is important for various respiratory diseases\(^\text{29}\) (Table 2).

Ayurvedic vis-à-vis conventional ideology of the cardiovascular system

Heart in Ayurveda is considered under rasavaha (related to heart) and pranavaha srotasa (related to respiratory). It is also considered to be seat of psyche and oja (vitality),
and any derangements in rasa dhatu leads to cardiovascular disease. In physiological aspects, it is clearly stated that for healthy functioning of the heart, an individual must consciously consumes diet favourable for heart and oja function. Thus, in Ayurveda each system has its connections ultimately with gut function.29.  

Tlaskalová-Hogenova et al.29 reported that atherosclerosis development was observed in germ free Apo-E deficient mice and not in conventionally reared Apo-E deficient (atherosclerosis model, apolipoprotein E-deficient mice). This denotes that commensals exhibit protective effect on mice health. The microbiota of conventional mice suppressed intestinal expression of fasting-induced adipocyte factor (Fiaf) and a circulating factor 1 (CSF1). Thus, the signalling between neurons and microorganisms for expressing gut hormones (like ghrelin). croorganisms for expressing gut hormones (like ghrelin). 

The gut–brain axis is a two-way communicating channel wherein the development and function of the nervous system is dependent on the gastrointestinal tract and vice versa. Muscularis macrophages (MM) regulate the intestinal motility; it communicates with enteric neurons by providing bone morphogenetic protein 2 (BMP2) signals and these neurons in return facilitates MM homeostasis by macrophage specific factor, viz. Colony stimulating factor 1 (CSF1). Thus, the signalling between neurons and gut is required for GI motility.34. Mucosal enteric glial cells (EGCs) are regulated throughout the intestine by the microbial metabolites. Therefore changes in the functioning of glial cells relate to the gut–brain axis and dysbiosis in the gut.34. As discussed earlier, the neuronal peptides such as CCK, GLP-1 and PYY are stimulated by the gut microbiota protecting it from various neurological diseases (Table 3). 

**Ayurvedic vis-à-vis conventional ideology of the nervous system**

All the functions of the nervous system resemble those of Vata (biophysical energy associated with bodily movements up to cellular level) in Ayurveda. Among five types of vata, samana vata is the enteric nervous system (ENS) which ultimately governs the entire gastrointestinal tract. In real sense, it is the sensory stimulus of ENS (samana vata) that is sent to central nervous system and resulting in vagal stimulation and gastrin feedback mechanism (prana vata) leading to the muscular movements, secretion of gastric juices (vyana vata) for digestion, absorption and discrimination.33.

The gut–brain axis is responsible for the regulation of hypothalamic–pituitary–adrenal (HPA) affecting the behaviour of the host.35. The production of various neurotransmitters such as GABA, serotonin, dopamine and noradrenaline is also done by the commensal thereby regulating the mood, cognition, emotion, motility and various secretions.36–38. Therefore, it can be stated that the central nervous system is controlled by the gut microbiota protecting it from various neurological diseases (Table 3).

**Ayurvedic vis-à-vis conventional ideology of the skeletal muscle system**

In Ayurveda, it is diet and successive products formed that are nutritive to tissues.39. Thus, in case of muscular...
physiology too, diet plays a major role. Malnutrition results in degenerative disorders, whereas over nutrition results in production of ‘ama’ contributing to rheumatic disorders.

The role of gut–muscle axis in the supply of nutrients to muscles and their development has been reported. The gut–muscle axis has been found to be associated with various pathways leading to increasing the metabolic efficiency of myofibre. There can be emergence of new therapies which can reduce muscle wasting and related diseases by inoculation of certain commensal in the host (Table 4).

**Ayurvedic vis-à-vis conventional ideology of the reproductive system**

In the description of healthy spermatogenesis and oogenesis, a role of Shadrasatmak ahara (diet comprising six rasas) has been given prime importance. Thus, proper vipaka can result into formation of respective healthy tissue. In order to have a normal and healthy conception, hita anna sevana (wholesome diet) has been specified in Charak Samhita.

Hormone production in the reproductive system is different for the male and female, which is assisted by different gut microflora. Over the course of pregnancy, the gut microbiota protects the mother and the foetus by producing hormones and signals. The gut impacts the immune system and produces inflammation in the body required in the pregnancy for the protection and development of the foetus. Proteobacteria, *Actinobacteria* and *Lactobacillus* are found in abundance in faecal matter of pregnant women.

The estrogen produced by gut microbiota using dietary estrogens and precursors can reduce metabolic disease during post menopausal period, thereby coping up with the decreased estrogen levels (Table 5).

**Gut microbiota and its association with disease pathogenesis**

The efficiency of gut microbiota in safeguarding human well-being by means of different pathways of metabolism has led to development of numerous associated linkages with the progression of disease owing to dysbiosis. The derangement of gut microbiota consortium has been attributed for development of various diseases as follows.

**Ayurvedic vis-à-vis conventional pathophysiology of obesity**

In Ayurveda, meda dhatu adipose tissue dysfunction due to improper agni (metabolic energy) has been delineated as a cause for sthauya, i.e. obesity. The vicious circle of hunger to satiety has been linked in etiology of obesity. Excessive consumption of madhura (sweet), sneha (lipid), lack of exercise lead to sthaulya even in upcoming pregnancy. Even treatment peculiarly consists of dietary management along with lifestyle modification and cleansing therapies of gut, viz. vamana (induced emesis), virechan (induced purgation) and Basti (enema).

A potential connection between gut microbiome and development of an obese phenotype has been mentioned by Ley. A proportional decrease in Bacteroidetes with development of an obese phenotype has been mentioned. The estrogen produced by gut microbiota using dietary estrogens and precursors can reduce metabolic disease during post menopausal period, thereby coping up with the decreased estrogen levels.
increase in relative loads of Firmicutes was found to be interrelated with the microbial population of obese mice. This was confirmed in a clinical dietary intervention assessment demonstrating that weight loss in obese individuals was due to increase in the relative proportions of Bacteroidetes. The microbial population within our intestine play a key role in obesity\(^5\). In spite of reports stating that these distinct microbial colonies increase energy harvest which contributes to obesity\(^5\), this has not been found to be totally imperative. It is therefore obvious that there can be multifactorial aspects of microbiota engaged in weight gain etiology and host metabolism. Following weight loss in human subjects, Ley\(^5\) found a decline in weight gain etiology and host metabolism. Following the genesis of urine in Ayurveda consists of moisture components aroused due to imbalances in diet and chronic liver diseases. The domain of bile acid signalling has disclosed the concept of the dynamic and highly regulated gut: liver axis. The etiological factors for liver disorders in Ayurveda have been faulty dietary habits, wrongly given emesis/purgation therapies results in (dhatu vaishamya) malfunctioning in tissue metabolism. These are termed to be santarpan janya (due to faulty lifestyle habits) disorders. The translocation of microbial colonies from the gut to the surrounding tissues with low-grade inflammation is considered to be the underlying mechanism for this disease.\(^6\) Pedersen and other scientists highlighted that the human gut microbiome apparently impinge on insulin resistance coupled with disturbed metabolomics of serum. The contributing species that have been identified for the same are Prevotella copri and Bacteroides vulgates.\(^6\) Complex gene–environment interactions can lead to obesity and its associated metabolic complications. Larsen et al.\(^6\) scientists employed 16S rRNA compositional sequencing in two groups; one of diabetics and other of healthy controls. It was observed that the proportions of clostridia class were reduced while the Bacteroides and Betaproteobacteria were enhanced in a group with T2D compared with control.

**Ayurvedic vis-à-vis conventional pathophysiology of liver diseases**

Type-2 diabetes (T2D) is a rampant global metabolic epidemic often termed as a silent killer. The association of gut microbial composition and progress of T2D are gradually being unveiled\(^5\). The translocation of microbial colonies from the gut to the surrounding tissues with low-grade inflammation is considered to be the underlying mechanism for this disease.\(^6\) Pedersen and other scientists highlighted that the human gut microbiome apparently impinge on insulin resistance coupled with disturbed metabolomics of serum. The contributing species that have been identified for the same are Prevotella copri and Bacteroides vulgates.\(^6\) Complex gene–environment interactions can lead to obesity and its associated metabolic complications. Larsen et al.\(^6\) scientists employed 16S rRNA compositional sequencing in two groups; one of diabetics and other of healthy controls. It was observed that the proportions of clostridia class were reduced while the Bacteroides and Betaproteobacteria were enhanced in a group with T2D compared with control.

**Ayurvedic vis-à-vis conventional pathophysiology of liver diseases**

In Ayurveda, Madhumeha is often regarded to share pathogenesis and clinical implication with diabetes. Madhumeha is one of the types of prameha, wherein the pathological changes in normal urine have been observed\(^5\). Even some pre-diabetic symptoms stress upon the characteristics of urine\(^5\). The pathogenesis of prameha as given by seers clearly indicate its close correlation with Mutravaha samsthana, i.e. urinary system\(^5\). In Ayurvedic physiology, main function of urine is kleda vahana\(^5\) (moisture components aroused due to imbalances in metabolism.) The genesis of urine in Ayurveda consists of its commencement in pakvashaya\(^5\) (part of colon), which is a unique feature after the assimilation of food occurs in alimentary canal. Ayurveda treatment regimens in diabetes strongly emphasize the establishment of homeostatic environment primely in the gut rightly by emesis and purgation therapies.\(^5\) Also, prescriptions on diet and chronic liver diseases. The domain of bile acid signalling has disclosed the concept of the dynamic and highly regulated gut: liver axis. The etiological factors for liver disorders in Ayurveda have been faulty dietary habits, wrongly given emesis/purgation therapies results in (dhatu vaishamya) malfunctioning in tissue metabolism. These are termed to be santarpan janya (due to faulty lifestyle habits) disorders. The translocation of microbial colonies from the gut to the surrounding tissues with low-grade inflammation is considered to be the underlying mechanism for this disease.\(^6\) Pedersen and other scientists highlighted that the human gut microbiome apparently impinge on insulin resistance coupled with disturbed metabolomics of serum. The contributing species that have been identified for the same are Prevotella copri and Bacteroides vulgates.\(^6\) Complex gene–environment interactions can lead to obesity and its associated metabolic complications. Larsen et al.\(^6\) scientists employed 16S rRNA compositional sequencing in two groups; one of diabetics and other of healthy controls. It was observed that the proportions of clostridia class were reduced while the Bacteroides and Betaproteobacteria were enhanced in a group with T2D compared with control.

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Evidence exhibits the close interlinks of gut microbiota and chronic liver diseases.\(^6\) Factors, namely gut barrier integrity, immune responses of liver to factors derived from gut; potentially contribute to address solutions for chronic liver diseases. The domain of bile acid signalling has disclosed the concept of the dynamic and highly regulated gut: liver axis. Recent facts depict that dysbiosis can lead to the phenomenon of non-alcoholic fatty liver disease (NAFLD) occurring as a sequel to obesity. Patients suffering from this disease have bacterial overgrowth located in small intestine. They also express deranged permeability of intestine that leads to shift of microbiome derived products into the portal circulation.\(^6\) Interference of certain multi-protein structures marks the imbalance of Bacteroidetes: Firmicutes ratio resulting

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**Table 5. Role of gut microbiota related to the reproductive system (after ref. 46)**

<table>
<thead>
<tr>
<th>Organism</th>
<th>Function</th>
</tr>
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<tbody>
<tr>
<td>Clostridia sp., Non Clostridiales sp., Ruminococcaceae sp.</td>
<td>Production of estrogen</td>
</tr>
<tr>
<td>C. scindens</td>
<td>Production of androgen by onverting glucocorticoids to androgens</td>
</tr>
<tr>
<td>Prevotella intermedia</td>
<td>Metabolizes estradiol and progesterone and changes their expression</td>
</tr>
</tbody>
</table>
into hepatic steatosis and inflammatory processes. Alcoholic liver disease expresses similar symptoms proclaiming abundant colonies of Bacteroidetes and Akkermansia, but decreased proportions of Lactobacillus and other similar colonies. Of late, it was explored that patients diagnosed with primary sclerosing cholangitis have distinct gut microbiota. A recent study by Nicolas et al. documented that hepatic gluconeogenesis can be controlled by the gut microbiota. Use of probiotics induces hepatic gluconeogenesis.

Ayurvedic vis-à-vis conventional pathophysiology of inflammatory bowel disease

This can be correlated to ‘pravahika’ in Ayurveda. The management of pravahika totally consists of series of medicated enemas again targeting colon and some wholesome recipes suitable for gut health like medicated porridges and soups.

Recent studies have demonstrated the relation among dysbiosis and gastrointestinal diseases. Imperative changes have been recognized in the microbial population of patients with inflammatory bowel disease (IBD) compared to healthy controls. The ratio of Firmicutes/Bacteroidetes was found to be two times greater than healthy controls in IBD patients.

Ayurvedic vis-à-vis conventional pathophysiology of cardiovascular diseases

Excessive eating, faulty timings of diet, incompatible food ingestion, faulty dietary habits as well as suppression of natural urges lead to formation of ama. This in turn deranges the metabolism in gut and hence hampers healthy functioning of the successive rasa dhatu which causes cardiovascular disorders.

Numerous facets of cardiovascular diseases comprise of plaque formation due to atherosclerotic changes followed by myocardial infarction and lastly failure. All of them are directly interlinked with dysbiosis. The metabolism of dietary phosphatidylcholine is a key example of the metabolic activity of gut microbiota on atherosclerosis development.

During digestion, phosphatidylcholine which is found in primary sources like meat, fish and egg is hydrolysed into trimethylamine by gut bacteria. It is later oxidized in liver in the form of trimethylamine N-oxide.

Ayurvedic vis-à-vis conventional pathophysiology of chronic obstructive pulmonary disease

This is ‘ama janya’ disorder according to Ayurveda, which again proves its link to gut. Existing high-throughput next-generation sequencing (NGS) techniques have reported that the airways harbour varied microbial communities. Microbial diversity is copious in severe chronic obstructive pulmonary disease (COPD) cases. Cameron et al. found remarkable changes in a COPD cohort summed with multiple features of COPD. Those comprised of specific exacerbation phenotypes, treatment differences and altered levels of key serum and sputum markers.

Ayurvedic vis-à-vis conventional pathophysiology of nervous system disorders

In Ayurveda, ahara is known to affect the properties of psyche satva, raja and tama. Accordingly, raja and tama are known to cause imbalance of doshas leading to psychiatric disorders.

The gut microbiota critically contributes to activities of the central nervous system. These frequently encompass immunological, neural and endocrinial pathways. This also consists of direct interface between enteric neurons and gut microbiota. Besides this, synthesis of neurohormones like serotonin, dopamine, etc. with regulation of hypothalamo-pituitary axis are all attributed to gut microbial functions.

Neuropsychiatric disorders often go hand in hand with certain modification in the gut microbial colonies. Examples include anxiety, autism, bipolar disorder, chronic fatigue syndrome, major depressive disorder, stress and schizophrenia. It is also interconnected to neurodegenerative diseases, viz. Parkinson’s disease, Alzheimer’s disease, dementia, and stroke.

Ayurvedic vis-à-vis conventional pathophysiology of cancer

In Ayurveda, cancer can be termed as the bad prognosis of any disease mentioned in classics related to the respective systems/srotas.

Latest studies suggest that perturbations in the gut microbial communities further add to tumorigenesis. The Gram-negative Fusobacterium nucleatum has been most strongly linked to colorectal cancer. The growth in gastric cancers also has an alliance with microbiota. This depends on the concentration of H. pylori and the corresponding changes in other microbial communities in the gut. Likewise, distinct microbial communities are found in case of oesophageal cancer after dysbiosis. Pertaining to liver, it has been stated that patients with hepatocellular carcinoma show profuse quantities of E. coli in faeces, much higher than healthy controls. Some studies have also described that gut microbiota aids in genesis of pancreatic cancer by promoting inflammatory changes, activating the immune response and by disseminating cancer-associated inflammation.
Drug metabolism by gut microbiota

The liver is the chief organ in charge of executing biotransformation and drug metabolism. However, drug metabolism may also partly occur at the intestinal level, which aids to have significant effect on the drug efficacy and safety.

Human gut microbiota influences the pharmacokinetics of many drugs, viz. chemotherapeutics, immunotherapies, anti-inflammatory and antimicrobial drugs and antiretroviral therapies. It is observed that microbiota augment or inhibit the desired therapeutic effect in case of these drugs. They carry out these processes by aiding biotransformation. Many factors, including gut transit time, host genotype, metabolic type, dietary intake and absorption influence microbial drug metabolism. Among them, reduction and hydrolysis occur frequently.

The class of carbohydrates and proteins that are complex and hence tedious to digest form the major substrates at the disposal of the microbiota. These substrates undergo fermentation in the presence of microbiota, which results in the production of a range of metabolites, including SCFA, branched chain fatty acids, ammonia, amines and phenolic compounds. Gases, viz. hydrogen, methane and hydrogen sulphide are also produced. In addition, the role of intestinal microbiota is implicated in the production of vitamins and the activation or inactivation of isoflavonoids.

Gut microbiota participate in drug metabolism in various ways:

- They produce enzymes that degrade drugs directly.
- They compete with drugs over the metabolizing enzymes.
- They alter activity of host drug metabolizing enzymes.
- They produce enzyme-inducing metabolites that alter the drug metabolic activities or pathways.
- The first reported example of drug metabolism by gut microbiota is that of azo reduction of orally administered prontosil and neoprontosil to sulphanilamide. This modification was the first known pro-drug activation by gut bacteria because metabolite sulphanilamide was found to have antibacterial effect. It has been found that reduced numbers of gut microbiota by pre-treatment of antibiotics reduces acetaminophen metabolism in gut. The pharmacokinetic alterations of amiodarone and its active metabolite by treatment with probiotics was witnessed which rendered significant changes in the pharmacological effect of amiodarone, making the active metabolite therapeutically more potent. These are some of the positive pharmacological actions exhibited due to metabolism of drugs by the innate gut microbiota.

- Some negative or adverse effects also have been reported, viz. alleviation of drug-induced enteropathy in case of diclofenac, indomethacin and ketoprofen, via β-glucuronidase inhibition mechanism. More examples include reduction of nitrazepam by microbiota human intestinal tract to 7-aminonitrazepam, which is then further metabolized in the liver to metabolite 7-acetylnitrazepam which is teratogenic.

- Gut microbiota show evidence of a noteworthy action on the metabolism of polyphenols. The metabolism of chlorogenic acid, naringin, and rutin, representative bio-actives of three common families, viz. hydroxycinnamates, flavonones and flavonols respectively was studied by Rechner et al. using in vitro mixed culture model of the human microflora.

- Some examples of the effect of gut microbiota on metabolism of certain medicinal plants are as follows.

  **Yashtimadhu (Glycyrrhiza glabra)**

  There is evidence which denotes that gut microbiota is responsible for conversion of glycyrrhizin to 18β-glycyrrhetic acid completely.

  **Panax ginseng**

  The gut microbiota is reported to metabolize orally administered ginsenoside Rb1 to bioactive compounds such as 20-Oβ-D-glucopyranosyl-20(S)-protopanaxadiol often known as compound K. The same action is inhibited when performed after administration of antibiotics.

  **Pueraria species**

  The effects of metabolites calycosin and daidzein are biologically superior to those of puerarin and daidzin, which are metabolized by gut microbiota.

  **Aloe vera**

  Barbaloin, a C-glycoside present in aloe, is reported to be metabolized in human faecal suspensions to the deglycosylated aloe-emodin anthrone. It exerts cathartic effect which is not exhibited by barbaloin.
**Scutellaria baicalensis**

Microbiota hydrolyse baicalin to baicalein which increase its bioavailability. Some pharmacological properties of baicalein, viz. anti-inflammatory, cytotoxicity for cancerous cells, anti-angiogenic potential are more than baicalein98.

**Citrus fruits**

Rutin is a flavonoid glycoside found in many citrus fruits which is metabolized by the human intestinal microbiota into several metabolites, including quercetin-3-O-glucoside, quercetin, the aglycone and 3,4-dihydroxyphenylacetic acid99. Similarly, hesperidine present in the variety of orange is primarily metabolized by gut microbiota into hesperetin which in turn exhibits enhanced cytotoxic activity in cancerous cells100.

Naringenin from grapes metabolizing in naringin inhibits hydroxymethylglutaryl-CoA (HMG-CoA) reductase. It also induces antioxidant enzymes, such as hepatic superoxide dismutase and glutathione peroxidase, which leads to reduction of hepatic cholesterol and lipid peroxidation101. Therefore, gut microbiota initiated pharmacokinetics and toxicological changes of drugs should be considered in future studies. It is also essential to accomplish the exact characterization of metabolites produced by the gut microbiota because these differ from those produced by hepatic metabolism. Finally, regulation of the gut microbiota may be a good target for enhancing the pharmacological effects of drugs.

**Future scope**

In spite of such extensive research (both *in vitro* and *in vivo*) related to gut microbiota, there are still deficient data regarding the exact mechanism of action of medicinal plants and also dietary components. This is still a big question related to the collective fate of metabolism of diet along with drugs which are often consumed. It can pave the way to unveil the role of ahara (diet) and aushadha (medicine) abiding by the principles of pharmacodynamics and pharmacokinetics of dravya (drugs) in Ayurveda.

The different responses of drugs due to their varied metabolism by gut microbiota explain vividly the need to probe into the fundamental concept of agni, which in turn affects vipaka and its degrees. This ultimately reciprocates on drug utilization and excretion at biota level, which highlights the importance of sara (utilizing of necessary metabolites)—kitta vibhajan (elimination by converting into excretory products at minute level.)

Most studies focus on individual components/constituents may it be of drug or food interaction with individual microbial communities. However, the very focus of Ayurveda resides upon the doctrine of holistic approach. Study designs which consider interface between whole drugs and not the isolates and consortium of gut microbiota will help further discover the fundamentals of the Indian traditional system of medicine.


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Received 18 July 2019; revised accepted 25 May 2020 doi: 10.18520/cs/v119/i3/451-461