Resident viruses and their interactions with the immune system

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The human body is colonized with a diverse resident microflora that includes viruses. Recent studies of metagenomes have begun to characterize the composition of the human ‘virobiota’ and its associated genes (the ‘virome’), and have fostered the emerging field of host-virobiota interactions. In this Perspective, we explore how resident viruses interact with the immune system. We review recent findings that highlight the role of the immune system in shaping the composition of the virobiota and consider how resident viruses may impact host immunity. Finally, we discuss the implications of virobiota–immune system interactions for human health.

The human body is colonized by commensal microorganisms that encompass diverse phyla from the three domains of life: Eukarya, Archaea and Bacteria. Most of these microorganisms reside at body surfaces that are in direct contact with the environment, including the intestine, skin and respiratory tract. Research efforts over the past two decades have focused primarily on the bacterial component of the human microbiota and its associated genes (the ‘microbiome’). These efforts have yielded a wealth of insight about the composition of human-associated bacterial communities, how these resident bacteria interact with the immune system and how bacteria–immune system interactions are altered in disease1,2.

Recently, it has become apparent that the microbiota of healthy humans also includes viruses, termed the ‘virobiota’3. Much of our current knowledge of the virobiota is derived from metagenomics, studies in which the DNA (and sometimes RNA) content of a microbial community is sequenced4–6. These studies have revealed that the human microbiome includes many viral genes (the ‘virome’). The intestine and the skin, for example, are both associated with viruses that replicate in eukaryotic cells (eukaryotic viruses), and viruses that replicate in bacteria (bacteriophages or phages)5–7. The intestinal microbiota also contains viruses that infect plants, which are likely associated with the host diet5,6,8.

Bacteria that inhabit the intestine and skin are generally regarded as stable residents that confer metabolic and/or immune benefits to their hosts9. It is therefore reasonable to ask whether viruses also can be stably associated with healthy human tissues. In the case of bacteriophages, a persistent, nonpathogenic association seems possible as viral replication occurs in bacterial hosts, which can themselves be stable members of the microbiota. In contrast, the relationship of eukaryotic viruses to their human hosts is much less clear. Such viruses require host cells to replicate and in most cases trigger innate and/or adaptive immune responses after entry into host cells. Thus, the eukaryotic viruses sampled in metagenomic studies could include those from acute, acute recurrent or chronic infections, or newly emerged latent viruses. In the absence of detailed longitudinal metagenomic studies of the virobiota, it is difficult to know at this point whether the human eukaryotic virobiota includes truly resident viruses that are stably associated with healthy host tissues.

In this Perspective we will discuss how the virobiota interacts with the immune system and how this could impact host health. We will focus most of our discussion on viruses associated with tissues that interface with the external environment (for example, intestine, skin and respiratory tract). It is important to recognize that the study of the virobiota is an emerging field and that our understanding of how these viruses interact with the immune system is currently minimal. Thus, some of our discussion will be speculative by necessity. However, our aim is to provide a framework for thinking about virobiota–immune system interactions in mammalian hosts and to motivate experimental studies of these interactions.

Bacteriophages that associate with host tissues

Metagenomic studies of microbiota at various tissue sites have revealed that many of the viruses associated with healthy human tissues are bacteriophages5,6,10. Sequencing of the metagenome from human fecal material has revealed that many phages associate with the gastrointestinal microbiota, which can harbor up to 10^{14} bacterial cells. Phage populations diversify in the intestine as new members of the bacterial community are introduced, suggesting that phage diversity and bacterial diversity are linked11.

Intestinal phages are genetically and morphologically heterogeneous. They include both lysogenic prophages, which are stably integrated into the bacterial chromosome, and lytic phages, which actively infect and lyse bacteria12. Prophages can undergo lytic induction to produce infectious phage particles when bacterial cells encounter environmental stressors or are stimulated by nutrients13,14. Metagenomic sequencing has revealed a few dominant families of intestinal phages that include double- and single-stranded DNA phages5,15. Although there is minimal variation of intestinal phage populations in individuals over time, there is substantial variation.

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between individuals\(^5,6\), even when those individuals have similar bacterial community structures\(^9\).

Phages likely have a profound impact on the composition and functional properties of the bacterial microbiota, which in turn could shape development and function of the immune system. First, phages may serve as important reservoirs of genetic diversity in the microbiota by acting as vehicles for the horizontal transfer of virulence, antibiotic resistance and metabolic determinants among bacteria\(^16\). Bacterial acquisition of phage genes could modify the functional properties of the microbiota, thereby substantially impacting host metabolism and immunity.

Second, phages may impact the composition of commensal bacterial populations through the predation of susceptible bacterial strains. This was suggested by recent studies of *Enterococcus faecalis* V583, a commensal of the human intestine. When *E. faecalis* V583 colonizes the intestines of mice, it produces lytic phages derived from chromosomally encoded prophage elements\(^14\). Production of the phage particles confers an advantage to *E. faecalis* V583 in competition with closely related enterococcus strains. This indicates that phages have the potential to influence the assembly and composition of intestinal bacterial communities\(^14\). Phage predation may be widespread, as suggested by the presence of clustered regularly interspaced short palindromic repeat (CRISPR) systems in many human commensal bacteria. CRISPRs are genetic modules in which short repeats of foreign DNA are inserted between spacer sequences in the bacterial chromosome. The spacer sequences are transcribed into short complementary RNAs that target invading DNA for destruction. Bacteria of the gastrointestinal tract and oral mucosa have acquired CRISPR-specific spacer sequences homologous to phage DNA\(^6,17\), suggesting that commensal bacteria are preyed upon by lytic phages in the mammalian host.

The ability of phages to shape the genetic diversity and the species diversity of the microbiota is likely to substantially impact the host immune system. It is well established that certain bacterial groups have distinctive effects on maturation of the immune system. For example, individual commensal species in the intestine influence the proportions of lamina propria T lymphocyte subsets that have distinct effector functions\(^18,19\). This in turn has profound effects on whether proinflammatory or anti-inflammatory immunity develops. Furthermore, commensal bacteria specifically limit and promote viral infection and transmission in mammalian hosts\(^20-22\). It is thus interesting to consider whether phage predation of intestinal bacteria could alter community composition in ways that impact function of the immune system and influence the spread of pathogenic viruses.

**Resident eukaryotic viruses of body surface tissues**

The resident viruses of humans also include viruses that infect eukaryotic cells. Like bacteriophages, populations of eukaryotic viruses in humans are heterogeneous, with distinct groups of viruses predominating in different tissues (Fig. 1). For example, the virobiota associated with adult human skin swabs is composed almost exclusively of polyomaviruses, including Merkel cell polyomavirus, a DNA virus associated with cutaneous cancer\(^7,23\). The frequency of human infection by resident polyomaviruses is likely to be high because ~70% of humans test seropositive for antibodies to Merkel cell polyomavirus\(^23\). Human skin also harbors an abundance of human papillomaviruses that are related to the β- and γ-papillomaviruses\(^7\).

A complex viral population is also associated with the mucosal surfaces of the oral cavity and the respiratory tract. Both PCR and metagenome sequencing have been used to characterize the composition of the viral communities associated with the oral mucosa. These studies show a consistent presence of DNA from human herpes viruses and related viruses\(^24,25\). Metagenomic studies of the nasopharynx microbiota also suggest that the respiratory tract is rich in viruses. Analysis of nasopharyngeal aspirates from patients experiencing respiratory-tract infections revealed the presence of many respiratory viruses, including human respiratory syncytial virus, influenza A virus and rhinoviruses. Viral nucleic acid is also present in respiratory aspirates of healthy children and indicates the presence of adenoviruses, picornaviruses and coronaviruses\(^26,27\).

Several studies of the human intestinal virome have suggested that eukaryotic viruses are rare relative to bacterial viruses\(^5,6,8\). However, genetic signatures of eukaryotic single-stranded RNA viruses, single-stranded DNA viruses, double-stranded DNA viruses and retroviruses have been detected in the fecal viromes of healthy individuals\(^5\). Human papillomavirus also has been detected but only in a single subject\(^28\). Genetic signatures of many plant viruses have been detected in metagenomic studies of the human intestinal virome\(^5,8\).

In particular, metagenomic studies of RNA viruses of the human intestine suggest that plant viruses are prevalent in this group and are likely derived from the diet\(^8\). Eukaryotic viral sequences are more prevalent in the intestines of diseased individuals. For example, feces from individuals with diarrheal disease and acute flaccid paralysis harbor a dominant population of DNA and RNA eukaryotic viral sequences derived from picornaviruses and parvoviruses\(^29,30\).

Much remains to be done to fully define the composition of the viral communities associated with human mucosal tissues and skin. For example, most metagenomic studies of human-associated viral populations have focused on analyzing viral DNA content. A more complete characterization of RNA viruses associated with humans will undoubtedly provide a more complete picture of the virobota. Furthermore, as noted above, it remains unclear whether the eukaryotic viruses associated with healthy human mucosal tissues are the result of chronic or latent infections or whether they arise from acute but transient viral infections. Careful longitudinal metagenomic studies of viral populations associated with mucosal tissue will be required to resolve these issues and to identify those viruses that stably inhabit human tissues.
Immune system control of resident viruses

The study of resident viruses is an emerging field, and thus our current understanding of how these viruses interact with the immune system is limited. However, our knowledge of how the bacterial microbiota interacts with the immune system provides a useful framework for our initial consideration of virobiota–immune system interactions. Here we consider key concepts that have emerged from studies of the bacterial microbiota and how they might promote our understanding of host-associated viruses and their interactions with the mammalian immune system.

Studies of commensal bacteria–host interactions have suggested that a key driving force in the evolution of the mammalian immune system has been the need to maintain homeostatic relationships with resident bacteria. In the intestine, for example, the immune system has a central role in controlling the density and the composition of resident bacterial communities. Recent studies in primates suggest that there is a similar relationship between the immune system and the intestinal virobiota. Simian immunodeficiency virus (SIV), a primate virus related to the human immunodeficiency virus (HIV), causes AIDS in rhesus monkeys. SIV infection of rhesus monkeys is associated with damage to the intestinal barrier (enteropathy), which promotes AIDS progression. Until recently, the causes of this enteropathy have not been clear. A metagenomic study of the virome of SIV-infected rhesus monkeys showed that SIV infection is associated with a dramatic expansion of the intestinal virome. This viral expansion was associated with enteropathy, particularly in monkeys that harbored adenoviruses (Fig. 2). These findings suggest that virome expansion is linked to the pathology observed in AIDS and highlights the role of the immune system in controlling virus populations in the intestine.

Viral populations associated with mucosal tissue are also altered in other disease states. In the human lung, cystic fibrosis or acute respiratory infection can result in the expansion of certain viral species. The lungs of patients with cystic fibrosis have many retrotranscribing RNA viruses and human herpes viruses, including Epstein-Barr virus, which has been linked to poor clinical outcomes in children with cystic fibrosis. The lung virome of patients with cystic fibrosis is highly variable as compared to healthy human lungs, where virome composition is relatively constant. It has been proposed that viruses in healthy lungs may cause short-term infections that are rapidly cleared by the host immune system, whereas in the lung of a patient with cystic fibrosis these viruses may be more persistent, perhaps as a result of compromised host immunity. The metabolic profile of the lung of a patient with cystic fibrosis suggests that it is enriched in aromatic amino acids and phosphate. This may influence the outgrowth of bacterial species that can shape the virobiota and possibly alter the severity of disease. It is intriguing to consider that competition among bacteria in the lungs and the success of certain species based on the metabolic environment may modulate the composition of virobiota in the lungs, ultimately leading to progression of disease if host immunity is compromised.

Does the virobiota shape host immunity?

Early comparisons of germ-free and conventionally colonized mice revealed a profound effect of the microbiota on the formation of lymphoid tissue and subsequent development of the immune system. Since then, many studies have established the importance of resident bacteria in promoting normal development and function of the immune system. Commensal bacteria also can prime immune responses that provide cross-protection for pathogenic infections. Given the presence of diverse families of eukaryotic and prokaryotic viruses at multiple human body sites, it is likely that viruses associated with mucosal tissue also have a pronounced influence on development and function of the immune system and could impact the host’s ability to fight pathogenic viral infections.

The eukaryotic virobiota and the immune system

One way that the virobiota could impact host immunity is by triggering immune responses that protect against pathogenic viral and/or bacterial infections. It has been estimated that the average human has ~8–12 chronic viral infections at any given time. These infections are caused by a variety of different viruses, including papillomaviruses, herpes viruses and polyomaviruses, that often reside in tissues of healthy individuals. Certain resident eukaryotic viruses can cause acute or latent infections that transition to chronic infections, where viral particles are shed throughout life. For example, herpes viruses infect most people during childhood, and latent herpes infection can lead to shedding of viral particles throughout the lifetime of an individual. Although latency and reactivation of herpes virus infection is considered pathogenic, chronic herpes virus infection has been shown to protect the host from viral and bacterial infections. Protection is conferred by chronic infection with select types of herpes viruses, including γ-herpes viruses, which increase basal interferon γ (IFN-γ) expression and facilitate activation of macrophages. This in turn limits the spread of other infecting bacteria and viruses.

Some chronic viral infections lead to a decrease in host immunity that promotes immune suppression and greater susceptibility to infection. As discussed above, SIV can trigger AIDS in some primate species, which results in an expansion of the intestinal virome. In another example, chronic infection by lymphocytic choriomeningitis virus hinders the expression of type I interferons and promotes the opportunistic invasion of murine cytomegalovirus.

These studies highlight a fine balance between viral symbiosis and pathogenesis during chronic infections. As discussed above, the
community of resident human associated viruses is not yet clearly defined, but viruses that maintain chronic infections may represent bona fide stable residents of the human virobota. The nature of the association of particular viruses with the host (chronic or transient) will likely have a profound impact on the immune response to viral and bacterial pathogen challenge. This suggests that when analyzing host immune status, composition of the resident viral community and the potential existence of chronically associated viruses must be considered.

**Bacteriophages and the immune system**

Bacteriophages are a long-overlooked component of the virobota that have the potential to shape mammalian immunity. Several studies have shown that phages can trigger a host immune response or can modulate host immunity. It is not surprising that humoral antibody responses can be mounted to phages that have been used to immunize animals at high titers. Phages can also inhibit activation and proliferation of human T cells in vitro through an unknown mechanism. Finally, phage and their nucleic acids alter the expression of innate immune genes in mouse tissues. Despite these intriguing findings, virtually nothing is known about whether phages can influence innate and adaptive immunity during natural associations with mammals.

There are several possible routes by which phages could become exposed to mammalian tissues and trigger immune responses. For example, it is known that oral administration of phages to animals results in the translocation of phages to systemic tissues. This suggests that mammals have mechanisms for uptake and delivery of phage that may allow intestinal phages to elicit innate and adaptive immune responses. One possible uptake route involves dendritic cells, which are known to sample intestinal luminal contents and can actively phagocytize phage particles in culture. Phage engulfment renders dendritic cells incapable of further phagocytosis, suggesting that phage sampling by dendritic cells could negatively impact their proinflammatory functions. Alternatively, phage antigens could be presented by dendritic cells to T cells, resulting in the development of cell-mediated immunity and cytokine release. Intestinal phages associate with the intestinal mucosal surface in larger numbers in patients afflicted with Crohn’s disease, raising the question of whether phages may influence the development of intestinal inflammatory diseases.

It is also interesting to consider whether bacteriophages might elicit antiviral innate immune responses. Mammalian cells are endowed with the ability to detect viral nucleic acids through several pattern-recognition receptors that are positioned to detect viral entry into cells. These pattern-recognition receptors include viral RNA sensors, such as the endosomal Toll-like receptors TLR7 and TLR8, and RIG-1, a cytoplasmic double-stranded RNA helicase. There are also DNA sensors, including TLR9, which detects endosomal DNA, and the cytoplasmic DNA sensor cyclic-GMP-AMP (cGAMP) synthase. RIG-I can also indirectly sense viral DNA with the aid of RNA polymerase III, which converts cytoplasmic DNA into 5’-phosphorylated RNA that can be recognized by RIG-I (ref. 54). Each of these sensors initiates signaling cascades that activate expression of type I interferon, inflammatory cytokines such as interleukin 6 (IL-6) and IL-1β and chemokines including IL-8 and CXCL-10. If resident bacteriophages can enter host cells, it is possible that they would activate one or more of these pathways, as their nucleic acids would likely be exposed to either the endosomal or cytoplasmic viral nucleic acid sensors. This could induce a tonic stimulation of antiviral immunity that might confer an advantage to the host by protecting against pathogenic viral infections.

Another way in which phages could become exposed to the mammalian immune system is through their association with commensal bacteria. Many commensal bacteria harbor prophages in their genomes and can induce these elements to produce phage particles in vivo. Under certain conditions, such as host immunodeficiency, commensal bacteria from the intestine can enter epithelial cells or become engulfed by phagocytic cells. Phage particles associated with these bacteria could be exposed to the cytoplasm or to endocytic compartments of cells and could thus prime innate or adaptive immune responses. In this way, bacteria could be delivery vehicles for phages that are sensed by the immune system.

Human intestinal phages have diversity-generating mechanisms that may facilitate evasion of host adaptive immunity. Sequencing of the metagenome of the human intestinal virome revealed that phage genomes are rich in regions of hypervariability. Most of these hypervariable regions map to phage tail-fiber genes that are diversified to...
allow the encoded proteins to bind variable phase receptors. Other hypervariable regions have been observed in genes encoding predicted immunoglobulin superfamily proteins, suggesting that such proteins could act as scaffolds for the presentation of diversified phase peptide sequences. The phase diversity-generating mechanism relies on error-prone reverse transcription. Although the physiological relevance of these hypervariable regions is not yet clear, such a diversity-generating mechanism could allow phages to evade antibodies that target the phage particles.

**Future perspectives**

Recent advances in high-throughput sequencing have led to the discovery that viruses are an important component of the microbial communities that inhabit healthy human tissues. Emerging information about the interactions between resident viruses and the immune system suggests that these interactions are as intricate and as coevolved as those involving the bacterial microbiota.

Going forward, it is imperative that resident host-associated viruses be incorporated into models of host-microbiota interactions. It will be important to perform careful longitudinal studies of the viral populations associated with healthy human tissues to determine the stability of these populations and how they may evolve over time. There is a need for further studies aimed at understanding how resident prokaryotic phages and eukaryotic viruses shape commensal bacterial communities and how this impacts host immunity. Future work should also focus on exploring the extent to which resident phages and eukaryotic viruses activate host innate and adaptive immunity. Examination of how resident viruses interact with host cells may reveal new host immune factors that lie outside of currently known canonical pathways of antiviral immunity.

Future experimental explorations of virobiota–immune system interactions will undoubtedly be challenging and will require new tools and approaches. However, such studies hold the promise of exciting insight into the complex relationships between humans and their associated microbial communities and may yield new strategies for enhancing human health.

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