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The healthy human virome: from virus-host symbiosis to disease

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Abstract

Viruses are ubiquitous, essential components of any ecosystem, and of multicellular organism holobionts. Numerous viruses cause acute infection, killing the host or being cleared by immune system. In many other cases, viruses coexist with the host as symbionts, either temporarily or for the duration of the host's life. Apparently, virus-host relationships span the entire range from aggressive parasitism to mutualism. Here we attempt to delineate the healthy human virome, that is, the entirety of viruses that are present in a healthy human body. The bulk of the healthy virome consists of bacteriophages infecting bacteria in the intestine and other locations. However, a variety of viruses, such as anelloviruses and herpesviruses, and the numerous endogenous retroviruses, persist by replicating in human cells, and these are our primary focus. Crucially, the boundary between symbiotic and pathogenic viruses is fluid such that members of the healthy virome can become pathogens under changing conditions.

Introduction

Billions of years of perennial and ubiquitous co-evolution between viruses and cells have produced a broad spectrum of virus-host interaction regimes, ranging from aggressive antagonism to commensalism, whereby viruses coexist with their hosts without harming them, at least, in the short term, and even to mutualism when a virus is beneficial and can be essential to the host [1-4]. The relationship between a particular virus and its host can be rarely, if ever, defined by a single regime. Rather, the mode of virus-host interaction is a function of multiple factors, including the environmental conditions, host and virus population structure, the immunological status of the host and many more.

Virus-host relationships that do not result in the demise of the host cell have been described across the virosphere [3,4]. Hence the concept of the ‘normal’ or ‘healthy’ virome, in principle, is applicable to any organism, including humans. Indeed, although most of the best characterized human viruses cause acute infections and are associated with diseases, a healthy human organism is host to a much greater variety of viruses [5-7]. The substantial majority of these infect bacteria that inhabit the human intestine, but a number of viruses actually reproduce in human cells without causing disease, at least, in the short term.

In this brief review, we systematically discuss the healthy human virome and emphasize that the boundary between “normal” and pathogenic (that is, causally associated with clinically manifested disease) viromes is blurred. Indeed, the same virus can be either a symbiont (with either no perceptible fitness effect on the host, that is, effectively, a commensal, or with a beneficial effect) or a pathogen depending on the conditions such as the health status and developmental stage of the host. We further stress that the current knowledge of symbiotic viruses lags far behind that of the pathogenic viruses.

The healthy human phageome

The human intestine, by far the richest microbial habitat in the body, contains about 10^{14} bacterial and archaeal cells at any given moment [8]. As many microbial communities, the human microbiome hosts a broad variety of viruses, in which tailed bacteriophages (class *Caudoviricetes* within the realm *Duplodnaviria*) comprise the overwhelming majority [8,9], albeit with a considerable contribution by ssDNA phages of the families *Microviridae* [10-12] and *Inoviridae* [13] (realm *Monodnaviria*). Systematic metagenomics surveys of the human phageome identify

thousands of phages but show that relatively few are common components of the healthy gut phageome. Thus, one of the most detailed metagenomics analyses resulted in the identification of only 23 phages that were shared by >50% of the tested individuals [9]. Strikingly, the most prevalent human-associated phage (and most prominent component of the healthy human virome altogether), the crAssphage, has been discovered only through metagenomics [14]. This initial finding has been followed by the characterization of an expansive group of crAss-like phages, also by metagenome analysis [15,16]. Subsequently, some crAss-like phages have been grown in cultures of the respective host bacteria, members of the phylum Bacteroidetes [17,18]. Given the difficulty of growing the human intestine-associated bacteria-phage systems in the laboratory, the actual size and diversity of the healthy human phageome remains to be discovered. Importantly, substantial changes in the human phageome have been associated with various diseases conditions, including infection with human and simian immunodeficiency viruses, and in some cases, the phageome perturbation was, apparently, decoupled from the changes in the microbiome [19-21].

Symbiotic viruses replicating in human cells

Over many decades, diverse viruses have been isolated from healthy humans more or less by chance. In the last few years, systematic surveys of the healthy human virome became possible thanks to the advances of metagenomics [6,7]. Below we briefly discuss the viruses that have been shown, either by traditional virology approaches or by metagenomics (Figure 1a), to be commonly present in healthy humans without causing apparent disease, following the main divisions of the current virus taxonomy [22,23]. The key aspects of the association of these viruses with the human organism are summarized in Figure 1.

Realm *Riboviria*

Kingdom *Orthornavira*

The realm *Riboviria* consists of viruses with RNA genomes as well as viruses with DNA genomes that employ reverse transcription in their replication cycles. The kingdom *Orthornavira* encompasses RNA viruses that share homologous RNA-dependent RNA polymerases. Many RNA viruses have been isolated or detected by metatranscriptomics in healthy humans but few can be confidently identified as symbionts replicating in human cells (Figure 1a).

Perhaps, the most notable apparent human symbionts among the orthornaviruses are pegiviruses (family *Flaviviridae*) [24,25]. The prototypical pegivirus has been initially tentatively identified

as hepatitis G virus [26], but no association with hepatitis has been subsequently confirmed [27]. In the family tree of flavivirids, pegiviruses tightly cluster with hepaciviruses which include hepatitis C virus (HCV), a major human pathogen [24,25]. However, unlike HCV, pegiviruses have not been linked to any pathology. Pegivirus infection in humans is common, with the incidence of about 5%, and pegiviruses readily grow in human cell cultures, leaving no doubt that these are bona fide human viruses [25]. Notably, pegivirus infection appears to be associated with benign clinical outcome in AIDS patients [28,29] indicating that these apparent viral symbionts of humans could benefit the host via protection from other viruses.

A recent, intriguing addition to the human virome are statoviruses (for STool-Associated TOmbus-like viruses) that were originally identified in multiple metagenomes from humans, macaques, cows and mice [30], followed by detection in nasal-throat swabs of humans with acute respiratory disease [31]. Although the actual hosts of statoviruses remain unknown, RdRP phylogeny, where statovirus RdRPs cluster with a variety of unclassified tombus-like viruses from invertebrate holobionts (that is, a host together with the entirety of associated symbionts and parasites) [32], suggests that statoviruses are associated with either mammalian diet (e.g., plants) or, more likely, some protist symbionts or parasites.

Another widespread group of putative members of the healthy human virome are members of *Picobirnaviridae*. Picobirnaviruses are commonly detected in mammalian, including human, intestines and have not been convincingly linked to any diseases although are suspected to be associated with diarrhea [33,34]. Picobirnaviruses have never been grown in cell cultures, and their true hosts remain unknown. The presence of highly conserved ribosome-binding sites (Shine-Dalgarno sequences), which are a hallmark of prokaryotic mRNAs, has led to the suggestion that picobirnaviruses infect bacteria in mammalian microbiomes [35].

In addition, certain food-derived plant viruses are present in the human gastrointestinal tract and in feces, sometimes in substantial amounts. The list of such viruses is topped by pepper mild mottle virus (PMMoV), a tobamovirus commonly found in pepper, pepper-derived products and their consumers all over the world [36]. Remarkably, PMMoV retains infectivity after passing through the human alimentary tract. Due to its stability and wide presence in human feces, PMMoV is used as a surrogate marker of fecal contamination of water [37]. However, given that there is no

evidence of plant virus replication in vertebrate cells, any substantial role of plant viruses in human health is an extremely remote possibility.

Kingdom *Pararnavira*

This virus kingdom includes viruses that employ reverse transcription in their replication cycles. Over 3,000 of human endogenous retroviruses (HERVs) are integrated into the host genome, comprising about 8% of human DNA [38]. Accordingly, the HERVs play important and diverse roles in human biology that cannot be discussed in this brief review in any detail (for recent reviews, see [39-42]). Most of the HERVs appear to descend from ancient integration events so that the virus genes are disrupted and rearranged. The legacy of these ancient HERVs are genes encoding virus structural proteins (Gag and Env) that have been recruited for a variety of physiological functions [43], the best known being syncytins, the placental trophoblast receptors [39]. However, some members of the youngest group of HERVs, known as HERV-K or HML2, that are thought to have invaded the human genome less than a million years ago form virus particles, particularly, in early embryogenesis [44,45]. The HERV-K viruses, at least, are bona fide members of the healthy human virome. Many of the other HERVs are expressed as well and are implicated in a variety of functions including modulation of innate immunity, even though functional virus proteins are usually not produced [46]. In particular, it has been reported that expression of one of HERV-K suppresses the spread of invasive melanoma [47]. However, potential associations between HERVs and various diseases have been reported as well. Thus, the relationship between the HERVs and the human host is a typical symbiosis, with both beneficial and potential deleterious effects on the host (Figure 1b).

Realm *Monodnaviria*

Realm *Monodnaviria* includes prokaryotic and eukaryotic viruses which encode homologous replication initiation endonucleases of the HUH superfamily or their inactivated derivatives, as in the case of polyomaviruses and papillomaviruses [22,48]. In addition to phages of the *Microviridae* and *Inoviridae* families mentioned above, several other representatives of *Monodnaviria* have been repeatedly identified as part of the healthy human virome. These include members of the families *Parvoviridae*, *Genomoviridae*, *Smacoviridae*, *Papillomaviridae* and *Polyomaviridae* (Figure 1a). The actual hosts for human-associated genomoviruses [49] and smacoviruses [50] remain unknown but these viruses likely infect human-associated microbes rather than humans directly.

Parvoviruses

Parvoviruses from several genera have been detected in various samples from apparently healthy humans, including human bocaviruses (HBoV; genus *Bocaparvovirus*), adeno-associated viruses (AAVs; genus *Dependoparvovirus*), human parvovirus 4 (PARV4; genus *Tetraparvovirus*), parvovirus B19 (B19V; genus *Erythroparvovirus*) and several protoparvoviruses (genus *Protoparvovirus*) [51-54]. These viruses display highly variable cell tropism and pathogenicity. For instance, AAVs and PARV4 can infect cells from multiple tissue types and are not known to cause any disease. By contrast, HBoV is most commonly found in the respiratory and gastrointestinal tracts as well as in blood [55,56], and is associated with acute respiratory symptoms, especially in children [57]. However, the direct role of HBoV as a pathogen remains unclear. B19V invades red blood cell precursors in the bone marrow and is commonly considered as a human pathogen causing a wide range of pathological conditions, including the fifth disease in children, persistent anemia in immunocompromised patients, transient aplastic crises, hydrops fetalis in pregnant women, and arthropathy [52]. Yet, in healthy adults, B19V infections are largely asymptomatic [58], with the prevalence of up to 25% in healthy human skin biopsies [59]. Thus, B19V is a conditional component of the “healthy” human virome that can turn into a pathogen in response to various factors (Figure 1b).

Polyomaviruses and Papillomaviruses

Monodnaviria includes class *Papovaviricetes* that consists of *Polyomaviridae* and *Papillomaviridae*, two families of viruses with small (5-8 Kbps), circular dsDNA genomes [60,61]. Papovaviruses are thought to have evolved from parvoviruses [48], with polyomaviruses emerging in invertebrates and co-evolving with animals for at least half a billion years [62]. In humans, polyomaviruses lead a low-profile life styles characterized by low propagation levels, evasion of clearance by the immune system and asymptomatic infections in immunocompetent individuals [60]. Apparently, human polyomaviruses have evolved mechanisms to limit their own reproduction levels in order to establish persistent infections [63]. A poster child human polyomavirus is John Cunningham virus (JCV) that establishes life-long latent infections [64]. The seroprevalence of JCV and other human polyomaviruses in adults can reach 90% or higher, with common coinfection by several polyomaviruses that are transmitted through direct contacts between humans or through contaminated objects. Contrary to their name and the oncogenic potential of the large T (tumor) antigen (the early virus protein involved in genome replication) in

experimental settings, most of the human polyomaviruses are not oncogenic. The Merkel cell polyomavirus is the only one that has been convincingly identified as the etiological agent of the eponymous carcinoma, a skin cancer caused by malignant transformation of the skin neuroendocrine cells apparently facilitated by virus genome integration into the host chromosomes [60,65].

Papillomaviruses are the closest, albeit apparently somewhat ‘younger’ relatives of polyomaviruses that likely emerged in vertebrates ~350 mya, but their life style is distinct. The majority of the ~200 known human papillomaviruses (HPVs) belong to Beta and Gamma types (*Betapapillomavirus* and *Gammapapillomavirus* genera, respectively), and cause unapparent productive infections or low-grade disease of the skin or mucosal epithelium (e.g., warts and condylomas) that are normally cleared by the immune system [66,67]. The virus replication cycle is tightly linked to epithelial differentiation and is orchestrated by a regulatory network that involves coordination between the cell cycle, virus DNA replication and transcription, and RNA splicing [67,68]. Most of the infections by the ‘high-risk’, Alpha HPV types (HPV16 and HPV18 being most prevalent; genus *Alphapapillomavirus*) in women result in more prolonged cervical infections, 80-90% of which are asymptomatic and are eventually cleared by the immune system [66,67]. However, a small fraction of such infections, primarily due to defects in the immune response, progress to the formation of persistent papillomas and, in a minority of cases, to cervical cancer. This small fraction of HPV infections, nevertheless, accounts for close to 100% of cervical cancers that affect over 500,000 women annually [69]. On much rarer occasions, the high-risk HPV also cause other types of carcinomas [70]. Carcinogenesis is a dead end for HPV because transformed cells produce no infectious virus. Therefore, despite the carcinogenic potential of high-risk HPVs, by and large, these common components of human virome should be considered symbionts, as demonstrated by the protection from skin cancer caused by immunity to ubiquitous skin-infecting HPVs [71] (Figure 1b).

Anelloviruses

Members of the family *Anelloviridae* are among the most enigmatic components of the healthy human virome, both in terms of their evolutionary origin and the impact on human health. Anelloviruses have small icosahedral virions that encapsidate tiny (3 to 3,5 kb) circular ssDNA genomes [72], but unlike the ssDNA viruses in the realm *Monodnaviria*, do not encode recognizable homologs of the rolling circle replication endonuclease. Moreover, no homologs

outside the family have been detected for any of the anellovirus proteins. Hence, anelloviruses are currently not included in the realm *Monodnaviria* and their provenance remains unclear [22,48]. The entire human population is believed to be infected with anelloviruses, and there is no convincing evidence of viral clearance from infected individuals [72,73]. The infections occur at an early age and so far have not been convincingly associated with any disease. The virus load appears to be controlled by the immune system because virus levels increase with the level of host immunosuppression [74,75]. Asymptomatic anellovirus infections are also common in other mammals [73,76-78], suggesting extensive coevolution of anelloviruses with mammalian hosts. Although the potential impact of anelloviruses on human health remains a matter of debate, it has been suggested that they positively influence human physiology by shaping the immunity during early development [72]. Thus, anelloviruses might be the most ‘friendly’, genuinely symbiotic component of the human virome.

Realm Duplodnaviria

The realm *Duplodnaviria* includes viruses with dsDNA genomes that are encapsidated in icosahedral capsids consisting of a distinct type of capsid protein, displaying the HK97 fold (after the first phage for which the capsid protein structure was solved), with the help of the corresponding variety of packaging ATPase, known as the terminase [22]. This realm includes the bulk of the viruses associated with the human microbiome, namely, the numerous tailed bacteriophages, as well as a major component of the virome associated with human cells, the herpesviruses. Some of the human herpesviruses (*Herpesviridae*) infect a variety of cell types and cause life-long latent infections [79]. Of the 9 human herpesviruses identified so far, herpes simplex virus 1 (HSV1), human cytomegalovirus (HCMV), Epstein-Barr virus (EBV) and varicella zoster virus (VZV) are highly prevalent in the human population. Depending on the geographic location, socioeconomic status and, in the case of VZV, vaccination levels, the incidence of these viruses reaches up to 96% [80-82]. The health impact of these viruses (if any) depends on the age and immune system status of the infected person, with ethnicity, gender and genotype being additional significant contributors.

The most stealthy of the human herpes viruses are apparently EBV and HCMV. If acquired in childhood, both of these viruses typically cause asymptomatic infections, mainly, in B-lymphocytes in the case of EBV [83] or in several cell types in the case of HCMV [79]. Such silent infections, however, even if not manifested in disease, cause a range of effects at the molecular,

cellular, tissue and organism levels. In the case of EBV, the gene expression pattern of infected B-cells is reprogrammed [84] and the proportion of plasma cells in blood appears to be increased [7]. Major immunity stimulation effects, a likely result of a balance reached over long virus-host co-evolution process, are well established for HCMV. In particular, ~10% of the CD4⁺ and CD8⁺ T cells in latently infected, otherwise healthy adults are HCMV-specific. Likewise, HCMV causes expansion of adaptive CD57⁺ natural killer (NK) cells targeting virus-infected cells [85]. These powerful arms of the immune system, however, fail to clear the virus due to its armament of immunoevasins, HCMV-encoded proteins involved in modulation of host immunity [86]. Extensive immunity stimulation in HCMV-seropositive individuals appears to enhance responsiveness and protection against heterologous viruses rather than compromise host immunity [86]. These potentially positive effects notwithstanding, damaging consequences of herpesvirus infections appear to be fairly common and vary from mild illness to a variety of life-threatening conditions. Thus, HCMV congenital (*in utero*) infections that occur at up to 2% of childbirths frequently cause neurodevelopmental defects or leukemia [85]. At the other life extremity, in seniors, HCMV seropositivity is associated with increased risks of cancer, cardiovascular disease, and ultimately, mortality.

The delicate, life-long EBV-host balance arising from most childhood infections is also fragile: if infection occurs in even non-immunocompromised young adults, it results in infectious mononucleosis ('kissing disease') [87]. Similarly, EBV is etiologically linked to Hodgkin's and Burkitt's lymphomas and nasopharyngeal carcinoma which are common in Southern Chinese and Eskimo people, as well as to a host of other diseases in immunocompromised individuals [83]. A different pathology pattern is characteristic of VZV infections that cause varicella (chickenpox) in children followed by prolonged latency that, in about 15% cases, leads to virus reactivation in elderly people with weakened immune control, causing zoster (postherpetic neuralgia) [88].

The HSV1 infections exhibit another distinct pattern of virus-host interactions. This virus is normally acquired in childhood causing mainly oral infections; its seroprevalence varies from 70% in developed nations to 100% in developing nations [81,89]. Upon infecting epithelial cells, HSV1 is transmitted to axons and establishes life-long latency in dorsal root ganglia that is periodically manifested in reactivation leading to recurrent acute infections or asymptomatic virus shedding. Similar to other human herpesviruses, HSV1 efficiently evades antiviral innate immune responses mediated by Toll-like and other pathogen recognition receptors. The underlying mechanisms of

HSV1 immunoevasion involve multiple virus proteins targeting diverse innate immunity signaling pathways [90].

Thus, human herpesviruses display a striking variety of cell tropisms and infection patterns that emerged over long-term virus-human co-evolution and co-adaptation. Some herpesviruses, in particular HCMV and EBV, can reach a perfect balance with the human host and often persist for the host's lifetime without causing any pathology. However, because of the complexity of virus-host interactions that involve a variety of genetic, environmental and socioeconomic factors, this balance is fragile and can be broken in many situations resulting in morbidity or even mortality (Figure 1b). Therefore, herpesvirus-host co-existence that involves the majority of the human population blurs the very concept of a 'healthy human virome'.

Realm Varidnaviria

This realm includes a broad variety of viruses with dsDNA genomes and icosahedral capsids built of protein unrelated to the capsid proteins of duplodnaviruses [22]. Unlike the members of *Duplodnaviria*, varidnaviruses are minor components of the healthy human virome, at best. Several intriguing reports have appeared on the detection of large and giant viruses of the class *Nucleocytoviricota* in healthy humans [91]. Perhaps, the most notable of these findings is the detection in human blood of several members of *Marseilleviridae* one of which has been reported to grow in T lymphocytes [92]. Additionally, several viruses from both *Marseilleviridae* and *Mimiviridae* have been isolated from human stools or detected in human-associated metagenomes [93]. Furthermore, the presence of mimiviruses in peripheral blood mononuclear cells of patients with atypical pneumonia has been reported, and a role for these viruses in the pneumonia pathogenesis has been suggested [94]. Unexpectedly, DNA of *Acanthocystis turfacea* chlorella virus 1, a member of *Phycodnaviridae*, has been detected in nearly half of the tested oropharyngeal samples from healthy humans, and also has been reported to persist in mouse macrophages [95]. However, so far, many of the reports on the presence of members of *Nucleocytoviricota* in human samples and, especially, their ability to replicate in human cells have been disavowed in follow-up studies [96]. Thus, the status of these viruses as components of the healthy human virome except, perhaps, as occasional contaminants, remains dubious.

Conclusions

The healthy virome of any organism, and especially humans, clearly, is an important component

of the holobiont that makes a major contribution to the health status of the host. However, the very concept of a healthy virome is nebulous and fluid because it is virtually impossible to ascertain that any virus would not cause disease under any conditions. A strong case in point are the herpesviruses that are nearly ubiquitous in the human population, remaining symbionts in most individuals most of the time, but consistently cause disease, in some cases, devastating, in immunocompromised individuals. Conversely, it appears plausible that any virus can become beneficial to the host through protection from other viruses, general stimulation of immunity as in the striking case of β -HPV protecting human hosts from skin cancer, or recruitment of virus genes for host functions. The numerous HERVs integrated in the human genome can be considered the paradigm of virus-host symbiosis. Generally, there is no doubt that many viruses evolved multiple mechanisms to manipulate the host innate and adaptive immunity pathways, ensuring virus persistence and controlling the damage to the host, as most conspicuously exemplified by the latent herpesviruses that are virtually ubiquitous in the human population.

The healthy virome is obviously heterogeneous and consists of 3 distinct components (Figure 1a): i) viruses that systematically enter the human organism, primarily, with food, but do not replicate in humans, ii) viruses infecting prokaryotes and, possibly, unicellular eukaryotes that comprise the healthy human microbiome, and iii) viruses that actually replicate and persist in human cells. With the advances of metagenomics, the human “microbiovirome” has become a subject of intense studies that continue bringing discoveries of new bacteriophage groups. In contrast, the “true” healthy human virome is poorly understood, with many questionable sightings of diverse viruses but little solid evidence on persistence mechanisms. On the whole, and in contrast to the disease-associated virome, the healthy human virome appears to be dominated by DNA viruses, in particular, anelloviruses and herpesviruses, that are substantially more common than RNA viruses in healthy humans. A thorough investigation of this component of the healthy virome can be expected to enhance our understanding of virus-host interactions and have major implications for human health.

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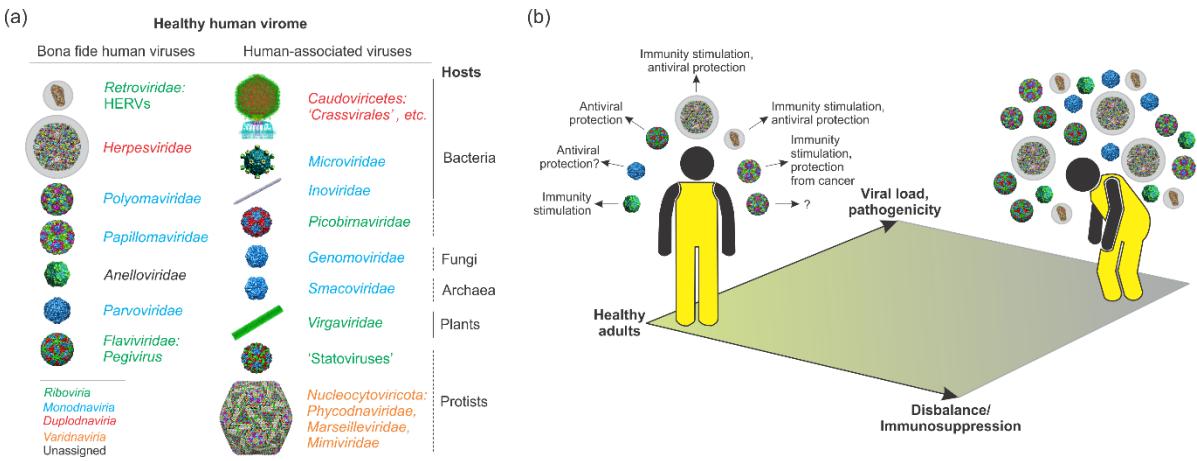
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629 **Figure 1. The healthy human virome.**

630 **(a)** Taxa of viruses found in healthy humans. Taxa of viruses replicating in human cells are shown
631 on the left, whereas those of viruses infecting human-associated microbes or associated with food
632 sources are shown on the right (the actual or suspected hosts are listed next to the corresponding
633 taxa, with the uncertain assignments indicated with broken lines). Depicted virus taxa are
634 represented with virion structures which were retrieved from VIPERdb (viperdb.scripps.edu) or
635 the Electron Microscopy Data Bank (<https://www.ebi.ac.uk/pdbe/emdb/>). When the structure of a
636 virus representing a particular taxon was not available, a structurally related member was chosen
637 instead. Virus taxa are colored according to their realm affiliation (the key is provided at the
638 bottom).

639 **(b)** Fluidity of the healthy human virome. The (potential) beneficial effects of the healthy human
640 virome are indicated next to the corresponding virus structures. Question marks denote
641 uncertainty. Upon changes in the health status/immunosuppression, viruses that cause
642 asymptomatic infections or are beneficial in healthy individuals proliferate and can cause diseases
643 including severe ones. The figure illustrates the general tendency of increased virus load under
644 immunosuppression/disease conditions and should not be interpreted as a quantitative
645 representation of the changes for any depicted group of viruses.