Endogenous viruses: insights into viral evolution and impact on host biology

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Endogenous viruses: insights into viral evolution and impact on host biology

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Abstract | Recent studies have uncovered myriad viral sequences that are integrated or ‘endogenized’ in the genomes of various eukaryotes. Surprisingly, it appears that not just retroviruses but almost all types of viruses can become endogenous. We review how these genomic ‘fossils’ offer fresh insights into the origin, evolutionary dynamics and structural evolution of viruses, which are giving rise to the burgeoning field of palaeovirology. We also examine the multitude of ways through which endogenous viruses have influenced, for better or worse, the biology of their hosts. We argue that the conflict between hosts and viruses has led to the invention and diversification of molecular arsenals, which, in turn, promote the cellular co-option of endogenous viruses.

Viruses are the most numerous and diverse genetic entities on earth. They are environmentally ubiquitous and are capable of infecting organisms from all three domains of life, as well as other viruses1. Through various interactions, viruses have profoundly influenced the evolution of cellular life ever since its origin2,3. In addition to the infectious properties of viruses, which enable them to spread horizontally between individuals and across species, many viruses can also become part of the genetic material of their host species, a process that is called endogenization. Such endogenous viral elements (EVEs)4 result from the chromosomal integration of viral DNA (or DNA copies of viral RNA) in the host germ cells, which allows for vertical transmission and potential fixation in the host population. Although endogenization was for a long time thought to be limited to retroviruses, recent studies have shown that all major types of eukaryotic viruses can give rise to EVEs.

The recent availability of large numbers of eukaryotic genome sequences, combined with an enhanced bioinformatics ability to detect and date ancient EVEs, has laid the foundation for the emerging field of palaeovirology5. After describing the discovery of EVEs, we discuss how this burgeoning area has already generated an important shift in our perception of the evolutionary origins and dynamics of several notorious groups of eukaryotic viruses.

Another exciting facet of palaeovirology is that it uncovers how EVEs have contributed to the evolution of the host genome by introducing genetic variation and innovation, which we discuss in sections on mutation of the host genome, influences on host gene expression and domestication into new protein-coding genes with cellular functions. Although other types of invasive, parasitic genetic elements, such as transposable elements, have a similarly broad impact on their host6, EVE sequences may have a greater propensity to be co-opted for certain regulatory and physiological functions. Most transposable elements are engaged in a long-term co-evolutionary relationship with their hosts, pressing them to evolve strategies to minimize their deleterious impact, which, in some cases, may reach a form of commensalism or symbiosis6,7. By contrast, most viruses assume a more destructive parasitic lifestyle, often leading to the death of infected cells, which sets them in an intense and perpetual conflict with their host. This arms race continuously selects for the emergence of genomic adaptations in viruses to manipulate the host cell machinery and to counteract antiviral defences. Ironically, as we will see, such viral weaponry often forms the foundation for the subsequent recruitment of EVEs to cellular functions.

Discoveries of endogenous viruses
Retroviruses are the only known eukaryotic viruses that require chromosomal integration for successful completion of their lifecycle. As such, they encode all of the enzymatic machinery that is necessary to perform this step autonomously. These properties probably explain why endogenous retroviruses (ERVs) were the first EVEs to be characterized and why they still make up the vast majority of EVEs that are recognized today. However, it has been known for some time that other viruses can be endogenized. For example, endogenous...
caulimoviruses (also called pararetroviruses) have been identified in plant genomes since the late 1990s. 

Like retroviruses, caulimoviruses replicate via reverse transcription, and they may have originated from the fusion of a retrotransposon with an envelope (Env)-like gene derived from a distinct virus. Unlike retroviruses, however, plant caulimoviruses possess a double-stranded DNA genome, and they do not normally integrate into the host chromosome. Accordingly, endogenous caulimoviruses are thought to result from fortuitous integration into the genome of germ cells (or meristematic cell progenitors of germ cells), and they generally correspond to partial viral genomes that are incapable of further replication. In rare cases, some of the integrated caulimoviruses appear to have retained the capacity to produce infectious particles but, intriguingly, only under stress conditions. 

Although a few instances of endogenization of various other viruses (shown in brackets) have been known for some time in some plants (Geminiviridae and Potyviridae), arthropods (Baculoviridae and Flaviviridae) and algae (Phycodnaviridae), it is only in the past few years that the pervasiveness of viral endogenization has been appreciated. This progress was largely fuelled by the advent of whole-genome sequencing and bioinformatics, which led to the identification of dozens of non-retroviral EVEs in various eukaryotes (Table 1). As for caulimoviruses, endogenization of these non-retroviral EVEs is thought to result from accidental chromosomal integration events. Little is known about the underlying molecular mechanisms, but several studies have identified sequence signatures at the EVE–host genome junction that point to retroposition events, suggesting involvement of the enzymatic machinery encoded by retrotransposons residing in the host genome. Further, it has been shown that viral DNA can be used to patch double-strand breaks in the host chromosome by non-homologous end joining, which might provide a mechanism for the endogenization of viruses with DNA genomes or DNA replication intermediates (for example, Ref. 19). Finally, it has been proposed that, for some viruses, integration of the viral genome could be facilitated by viral proteins. The discovery of a wide range of non-retroviral EVEs suggests that almost any major type of eukaryotic virus may be endogenized, sometimes in multiple hosts.

Table 1 | EVEs identified in eukaryotic genomes

<table>
<thead>
<tr>
<th>Group/type</th>
<th>Family or genus</th>
<th>Taxa</th>
<th>Number per haploid genome</th>
<th>Refs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group I/dsDNA</td>
<td>Baculovirus</td>
<td>Insects</td>
<td>Unknown (hybridization data, no sequencing)</td>
<td>124</td>
</tr>
<tr>
<td>Group I/dsDNA</td>
<td>Herpesviridae</td>
<td>Humans</td>
<td>1</td>
<td>125,126</td>
</tr>
<tr>
<td>Group I/dsDNA</td>
<td>Nudiviridae</td>
<td>Parasitic wasps</td>
<td>Several</td>
<td>127</td>
</tr>
<tr>
<td>Group I/dsDNA</td>
<td>Phycodnaviridae</td>
<td>Brown algae</td>
<td>1</td>
<td>128,129</td>
</tr>
<tr>
<td>Group II/ssDNA</td>
<td>Circoviridae</td>
<td>Mammals</td>
<td>1 to 2</td>
<td>4,20,130</td>
</tr>
<tr>
<td>Group II/ssDNA</td>
<td>Geminiviridae</td>
<td>Totenosteae (tobacco and three other species)</td>
<td>5 to 120</td>
<td>130–132</td>
</tr>
<tr>
<td>Group II/ssDNA</td>
<td>Paroviridae</td>
<td>Mammals; shrimp</td>
<td>1 to 3</td>
<td>4,20,89, 133,134</td>
</tr>
<tr>
<td>Group III/dsDNA</td>
<td>Partitiviridae</td>
<td>Plants; arthropods; Protozoa</td>
<td>1 to 4</td>
<td>135</td>
</tr>
<tr>
<td>Group III/dsDNA</td>
<td>Reovirus</td>
<td>Aedes spp. mosquitoes</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>Group III/dsDNA</td>
<td>Totiviridae</td>
<td>Fungi; plants; ticks</td>
<td>1 to 6</td>
<td>16,135,136</td>
</tr>
<tr>
<td>Group IV+/ssRNA</td>
<td>Dicistroviridae</td>
<td>Honeybees</td>
<td>1</td>
<td>137</td>
</tr>
<tr>
<td>Group IV+/ssRNA</td>
<td>Flaviviridae</td>
<td>Medaka fish; mosquitoes</td>
<td>1 to 4</td>
<td>4,21,134, 139</td>
</tr>
<tr>
<td>Group IV+/ssRNA</td>
<td>Pottyviridae</td>
<td>Grapes</td>
<td>Several</td>
<td>140</td>
</tr>
<tr>
<td>Group V−/ssRNA</td>
<td>Bornaviridae</td>
<td>Vertebrates</td>
<td>1 to 17</td>
<td>4,21,17</td>
</tr>
<tr>
<td>Group V−/ssRNA</td>
<td>Bunyaviridae</td>
<td>Ticks</td>
<td>14</td>
<td>4</td>
</tr>
<tr>
<td>Group V−/ssRNA</td>
<td>Filoviridae</td>
<td>Mammals</td>
<td>1 to 13</td>
<td>4,21,141</td>
</tr>
<tr>
<td>Group V−/ssRNA</td>
<td>Nyavirus</td>
<td>Zebrfish</td>
<td>6</td>
<td>21</td>
</tr>
<tr>
<td>Group V−/ssRNA</td>
<td>Orthomyxoviridae</td>
<td>Ticks</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>Group V−/ssRNA</td>
<td>Rhabdoviridae</td>
<td>Insects (ticks and mosquitoes)</td>
<td>1 to 28</td>
<td>4</td>
</tr>
<tr>
<td>Group VI/ssRNA-RT</td>
<td>Retroviridae</td>
<td>Vertebrates</td>
<td>Several hundreds to several hundreds of thousands</td>
<td>36</td>
</tr>
<tr>
<td>Group VII/dsDNA-RT</td>
<td>Hepadnaviridae</td>
<td>Passerine birds</td>
<td>15</td>
<td>4,19</td>
</tr>
<tr>
<td>Group VII/dsDNA-RT</td>
<td>Pararetroviridae</td>
<td>Plants</td>
<td>A dozen to a thousand</td>
<td>8,11</td>
</tr>
</tbody>
</table>

+, positive sense; −, negative sense; RT, reverse transcriptase.
independently and over wide evolutionary periods4 (TABLE 1). These findings also reveal that some viruses have had and may still have a much broader host range than was previously appreciated. For example, endogenous hepadnaviruses19 and filoviruses23 have been found in passerine birds and marsupials, respectively, which are thus far not known to be infected by these viruses. The discovery of EVEs may also be helpful to identify candidate reservoir species of zoonotic viruses. In this regard, the apparent over-representation of filovirus-like EVEs in rodent, insectivore and bat genomes compared with those of other mammals is intriguing given that these taxa rank among the most likely candidate reservoirs for their notorious modern relatives, such as Ebola and Marburg viruses24.

Uncovering the deep evolution of modern viruses EVEs fill in gaps in viral evolution. In spite of the numerical and ecological importance of viruses in the biosphere and their frequently devastating impact on human health, our understanding of viral evolution remains fragmentary. Regarding the early origins of viruses and the nature of their evolutionary relationships with eukaryotes, bacteria and archaea, the emerging consensus posits that viruses may be older than cellular life and may have actually triggered its emergence or at least may have profoundly influenced the early evolution of cells2,3,14. The inferred ancient origin of viruses is based, in part, on the observation that several genes involved in DNA or RNA replication and capsid assembly are structurally more similar among diverse viruses infecting the three domains of life than between viruses and any cellular organisms25,26.

In addition to these inferences on the earliest steps of viral evolution, several facets of the evolution of modern viruses have been extensively investigated. A number of studies have produced robust estimates of the ages, evolutionary rates and epidemiological dynamics of many viruses that currently infect humans, domestic animals and crops23,24. According to these studies, the evolutionary timescale of modern viruses lies within the past million years: most human RNA viruses emerged less than 1,000 years ago28. Hence, a huge gap in our understanding of viral evolution lies between these very recent events and the pre-cellular origins of viruses. The study of EVEs that are closely related to, or even fall within the diversity of, currently circulating viruses offers an opportunity to start filling in this gap. EVEs can provide missing links for deciphering the structural evolution of viral genomes and the origin of new viral genes (BOX 1). In addition, various methods can be used to infer the integration times of EVEs, which, in turn, yield minimum ages for the family of modern viruses to which they belong (BOX 2). In many cases in which EVEs that are closely related to modern viruses could be dated, the minimum age inferred for the most recent ancestor of the viral family has turned out to be far older than was previously estimated using sequence data from circulating viruses. For example, the Circoviridae and Hepadnaviridae families were thought to be <500 and <30,000 years old, respectively25,26, but EVEs from these two families have been dated at >40 million and >19 million years old, respectively20,19. Similarly, EVEs that are clearly related to Lentiviridae and Spumaviridae were traced back to >12 million and >100 million years ago, respectively; far older than had been inferred from sequence comparison of their modern relatives27,28.

Substitution rates. Most exogenous viruses are characterized by extremely rapid substitution rates that are often three to six orders of magnitude faster than those of their host30. Typically, viral substitution rates are calculated using samples of modern viruses that have circulated over short periods of time spanning tens or hundreds of years (see the figure in BOX 2). The discovery of EVE sequences that are fossilized in genomes for millions of years but that are still related to and directly alignable to those of modern viruses offers an opportunity to derive viral substitution rates on a much deeper timescale (BOX 2). Surprisingly, such long-term viral substitution rates are considerably slower than short-term rates estimated using only modern viral sequences. For example, in the case of hepadnaviruses and begomoviruses, long-term substitution rates were found to be two to three orders of magnitude slower than short-term rates30,31. At first glance, it is tempting to explain this discrepancy by the fact that the substitution rate of EVEs dramatically plummets following endogenization, as EVE sequences become subject to the much slower mutation rate of the host genome (see the figure in BOX 2). However this ‘mutational freezing’ of the EVE sequence at the time of endogenization has essentially no bearing on the calculation of long-term viral rate because the number of substitutions accumulated at the host rate (indicated by the red triangles in the figure in BOX 2) represents a small fraction of the substitutions observed when the EVE and its closest modern viral relative are compared. That small number of mutations (accumulated at the host rate) can easily be estimated and subtracted in the calculation of long-term viral rate19.

How then can we explain the vast discrepancy between short- and long-term rates of viral evolution? The phenomenon is reminiscent of the so-called time dependency of substitution rates observed in cellular organisms, where rates measured on a shallow timescale (for example, in pedigrees or populations) are invariably faster than rates measured over a deeper, geological timescale27. Several factors have been proposed to explain this incongruity3,14,19,32,33. For instance, it is possible that short-term rates are artificially inflated by a large proportion of slightly deleterious mutations, which are yet to be removed from the population through purifying selection3,14. To assess whether this effect may substantially bias short-term estimates of viral rates, it would be necessary to evaluate the lifespan of deleterious mutations in viral populations. Mutational saturation is a factor that could play in the other direction, leading to an underestimate of the genetic distances between EVEs and circulating viruses and thereby of long-term substitution rates.
As representatives of viral families fossilized at various evolutionary time points, endogenous viral elements (EVEs) can provide insights into the origins of components found in modern viral genomes\(^\text{5,101}\). For example, the origin of the accessory genes of complex retroviruses (such as nef or rev in HIV) has long been a mystery. These genes have diverse functions (for example, regulation of proviral expression, export of retroviral mRNA from the nucleus to the cytoplasm and blocking host restriction factors) and — unlike the structural and enzymatic genes, gag, pol and env, which are common to all retroviruses (shown as grey rectangles in the figure) — they are only found in a subset of retroviral genera (shown as coloured rectangles in the figure).

Until recently, only the vpx gene that is present in HIV-2 and in various simian immunodeficiency viruses (SIVs) had been traced; it appears to have arisen by non-homologous recombination between two different SIVs\(^\text{101-103}\). The discovery of an endogenous lentivirus, PSIV, in the genome of prosimian primates (namely, Malagasy lemurs) helped to shed light on the origin of a second lentiviral accessory gene. This gene, called orf2 (see the figure), shows substantial sequence similarity to the 3’ end of the primate lentiviruses reverse transcriptase domain, suggesting that it might have arisen by partial duplication of this domain, possibly via template jumping during reverse transcription\(^\text{104}\). Interestingly, orf2 is of the same size and is located at the same position (within the pol–env intervening region) as the tat gene of modern lentiviruses, suggesting that it may be a tat orthologue. The orf2 sequence, however, does not share any substantial similarity with known tat genes, which themselves are extremely diverged from each other and are barely alignable. Whether orf2 and tat are truly homologous (that is, whether they have a common ancestor) therefore remains an open question that would need to be addressed by functional studies. Unlike vpx, which is only present in HIV-2 and in a subset of SIVs and must be of very recent origin, tat is present in all primate lentiviruses and in other lentiviruses (such as bovine immunodeficiency virus (BIV) from cows, equine infectious anaemia virus (EIAV) from horses and rabbit endogenous lentivirus type K (RELIK)), which suggests it emerged much earlier than vpx. Thus the link between tat and the 3’ end domain of the lentiviral reverse transcriptase in PSIV might be our best opportunity to trace the origin of tat LTR, long terminal repeat.

The discordance observed between EVE-based studies and modern virus-based studies could also reflect genuine biological differences in the viruses under study. In particular, variations in the replication rate (which could possibly be caused by latency) or in the fidelity of viral polymerases over time\(^\text{19}\) could have a substantial effect; for example, the fidelity of reverse transcriptases from diverse retroviruses is known to vary by 20-fold\(^\text{105}\). Finally, an interesting question is whether viral lineages are best characterized by a single rate with little variation through time or whether their evolution follows a more saltational mode that is typified by rapid changes in substitution rates that are directed by environmental or ecological variables, such as the colonization of new hosts\(^\text{106}\) or the efficiency of the host's immune response. According to the saltational model, elevated short-term viral rates might reflect a transient state of adaptation associated with an intense conflict with a new host, whereas most of the remaining part of a virus evolutionary history would be characterized by a slower rate that is closer to the estimated long-term rate. In line with this scenario, we note that the sample of viruses used to infer short-term rates is strongly biased for zoonotic viruses that may conceivably have been caught in the midst of acute adaptation to their new host. Future efforts should be directed towards developing a simulation approach in order to define the conditions under which a virus could evolve at vastly different rates, depending on the timescale considered, and to assess whether these conditions are realistic or not. It will also be necessary to improve nucleotide substitution models and to continue characterizing new EVEs. Perhaps an ideal prospect to reconcile short- and long-term viral rates would be to identify EVEs with a range of ages spanning a large breadth of the evolutionary history of a viral family and to incorporate these genomic fossils in a phylogenetic analysis of extant viruses.

**Box 1** | EVEs shed light on the structural evolution of viruses

As representatives of viral families fossilized at various evolutionary time points, endogenous viral elements (EVEs) can provide insights into the origins of components found in modern viral genomes\(^\text{5,101}\). For example, the origin of the accessory genes of complex retroviruses (such as nef or rev in HIV) has long been a mystery. These genes have diverse functions (for example, regulation of proviral expression, export of retroviral mRNA from the nucleus to the cytoplasm and blocking host restriction factors) and — unlike the structural and enzymatic genes, gag, pol and env, which are common to all retroviruses (shown as grey rectangles in the figure) — they are only found in a subset of retroviral genera (shown as coloured rectangles in the figure).

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Box 2 | Dating EVEs and inferring long-term viral substitution rates

Various methods can be applied to infer the age of endogenous viral elements (EVEs) and to calibrate the evolutionary timescale of modern virus families. A traditional way of estimating the integration time of endogenous retroviruses (ERVs) is to calculate the divergence between their 5' and 3' long terminal repeats (LTRs) and to divide this distance by the substitution rate of the host genome\textsuperscript{10}. This calculation is based on the fact that the reverse transcription of a given retrovirus inherently generates a provirus that is flanked by identical 5' and 3' LTRs following integration, such that the nucleotide differences that are observed between these LTRs are expected to result from mutations that occurred after integration at the neutral rate of the host genome. This method has been used to infer the age of many ERVs, including most of the ERV families present in the human genome, and various corrections have been proposed that take into account potential biases introduced by gene conversion and heterogeneity in substitution rates between LTRs\textsuperscript{106,107}.

A second method of inferring the age of EVEs relies on the frequency of stop codons in the various endogenous open reading frames\textsuperscript{21}. Finally, EVEs can be dated by screening for the presence or absence of orthologous EVE loci in the genome of species that present a diverse degree of relatedness to the host species in which the EVE was detected\textsuperscript{4,19,20,27,28}. The presence of an EVE locus at the same genomic location in two species indicates that integration predates the divergence of these species, and this provides a lower age boundary. Characterization of an ‘empty site’ (that is, genomic flanking regions without the EVE insertion) in a third more distantly related species can provide an upper age boundary. EVEs that have been dated using this approach can be sequenced in the two more distantly related species in which they were found, and another time estimate can be derived by dividing the distance separating these EVEs by the host neutral substitution rate. In instances in which the host neutral rate is based on a different palaeontological calibration point than the one used for inferring species divergence times, congruence between the two approaches solidifies age estimates\textsuperscript{19}. After the age of EVEs has been estimated, it offers a calibration point to derive a long-term viral substitution rate. An interesting property of EVEs is that, after endogenization, they no longer mutate or evolve at the fast exogenous viral rate. Instead, they are replicated along with their host genome and evolve much more slowly than their exogenous counterparts. Consequently the sequence of an EVE as we see it today is very close to the sequence of the viral genome that became endogenized. It differs from this ancestral sequence by only a few mutations (shown by the red triangles in the figure). Long-term viral substitution rates can therefore be estimated by simply dividing the substitutions (shown by the blue triangles in the figure) between this approximate ancestral sequence (neglecting the few substitutions that occurred at the host rate) and its closest circulating counterpart by the age of the EVE (time of endogenization). To obtain a more accurate estimate of long-term substitution rate, it is possible in some instances to estimate the amount of substitutions accumulated at the host rate after endogenization (shown by the red triangles in the figure) and to remove this amount from the total distance separating EVEs from their closest exogenous counterparts\textsuperscript{19}.
 invisible assault on the host species is hardly possible, but considering that nearly half a million viral insertions have reached fixation in the human genome alone, whereas many more have concurrently colonized other vertebrate lineages, it is easy to predict that viral integration represents a substantial source of natural genomic variation in vertebrate populations. ERVs possess a greater mutagenic potential than other types of EVE, in that each insertion event typically deposits a full-length provirus that is capable of producing new germline or somatic insertions for a long time after the initial endogenization event. Further genomic propagation of ERVs can occur either by reinfection or by intracellular retrotransposition. An ERV may undergo either autonomous replication using its own enzymatic machinery, or it may be complemented by enzymes produced by exogenous viruses or by other retroelements that coexist in the genome. There is evidence that all of these processes have contributed to the genomic expansion of human ERVs over extensive periods of primate evolution. The integrated form of a retrovirus is apparently the most recent ERV wave to have entered our genome, perhaps producing insertions as recently as 150,000 years ago. These data stand in contrast to findings of ERV activity in rodents, where it has engendered extensive genomic variation among inbred strains and at least one heritable de novo insertion with phenotypic effects. Thus, murine rodents offer valuable models for studying the short-term consequences of ERV mutagenesis on genome function and phenotype.

Post-insertional genomic rearrangements. As with other repeated sequences, non-allelic homologous recombination (NAHR) between members of the same EVE family that are located on the same or different chromosomes may lead to genomic rearrangements. Depending on the orientation and location of the elements relative to one another, NAHR may result in deletion, duplication, inversion or translocation events. The frequency and biological impact of NAHR events largely depend on the length, homology, density and distribution properties of the repeats throughout the genome. Generally speaking, families with large numbers of long, highly homogeneous copies are more likely to engage in rearrangements than rare, short and strongly diverged elements. All cases of EVE-mediated rearrangements so far described in humans have implicated ERVs, which is not surprising, given the predominance of these EVEs in our genome. A detailed study of the HERV-K (HML2) subfamily shows that 6 (17%) of the 35 full-length copies examined have undergone some form of NAHR following insertion, despite their fairly recent origin (<20 million years ago). Given that this subfamily accounts for a minuscule fraction (<1%) of human ERVs, these data underscore the potentially profound contribution of ERVs to remodelling genome architecture over a large timescale. ERV-mediated NAHR events can also be pathogenic. For example, a non-reciprocal recombination event between two HERV-I copies on the Y chromosome caused a 792 kb deletion containing the azoospermia factor gene (AZFA; also known as AZFI), apparently causing male infertility. Similarly, HERV-mediated deletion of the eye absent homologue 1 (EYA1) gene was recently linked to branchio-oto-renal syndrome. But somatic rearrangements involving ERVs might be the most frequent and most medically relevant, especially for tumorigenesis, given the long chain of studies linking ERV to cancer (see below). For instance, a recurrent translocation event that creates an oncogenic HERV-K–ETS translocation variant 1 (ETV1) fusion gene has been implicated in prostate cancer. These studies beg for more systematic analyses to assess the importance of ERV-mediated rearrangements in disease and evolution.
Box 3 | The KRAB-ZNF army: a genome defence system against ERVs?

Several recent studies point to the existence of a previously unrecognized genome surveillance system aimed at repressing the transcription of endogenous retroviruses (ERVs) and possibly other vertebrate retroelements during early embryonic development. A first hint at the system came from a series of elegant studies on the transcriptional silencing of murine leukemia virus in mouse embryonic stem cells. These studies established that silencing occurs through the tethering of the KAP1 (also known as TRIM28) co-repressor complex to proviral DNA by ZFP809, which is a member of the Krüppel-associated box zinc finger (KRAB-ZNF) family of DNA-binding proteins. KAP1 is known to bind several KRAB-ZNF proteins through its amino-terminus, whereas the carboxy-terminus recruits a suite of effectors that promotes the local formation of silenced chromatin (shown in part a of the figure). In early embryonic development, KAP1 is known to bind ESET (also known as SETDB1), which di- or trimethylates histone H3 at lysine 9 (H3K9me2 or H3K9me3, respectively) and to heterochromatin protein 1 (HP1) family members, which ‘read’ H3K9me2 and H3K9me3 marks. KAP1 is also known to interact with another epigenetic silencer, LSD1 (also known as KDM1A), which demethylates lysine 4 on H3 (H3K4me1 or H3K4me2). Finally, KAP1 can also recruit the NURD histone deacetylase (HDAC) complex, which removes acetyl groups on histones. Through each or a combination of these events, it has been shown that KAP1 is a central player for the transcriptional silencing of a wide range of ERV families in mouse embryonic stem cells and in differentiated cells, the epigenetic marks that were deposited by KAP1 in early development may also contribute to the recruitment of DNA methyltransferase 3A (DNMT3A) and DNMT3B, which methylates cytosines at CpG sites (CpGme), reinforcing ERV transcriptional silencing. Together, these data suggest a model in which KAP1 and its associated co-repressor complex can be tethered to ERV DNA in a sequence- or family-specific fashion by a battery of KRAB-ZNF adaptors (shown in part b of the figure). This model is supported by the exceptional expansion, diversification and turnover of KRAB-ZNF genes in tetrapod genomes. These genes are encoded in the hundreds in all tetrapod species examined, but they tend to be poorly conserved in sequence and genomic location even between closely related species. They have been gained and diversified at an extraordinary rate through lineage-specific tandem duplication followed by bouts of positive selection in their ZNF domains, all of which is predicted to change DNA-binding specificity—an evolutionary pattern that is reminiscent of genes that are involved in antiviral immunity (BOX 5). Furthermore, there is a striking positive correlation across species between the number and age of KRAB-ZNF genes and those of their ERV content. These data are consistent with an arms-race model in which the need to silence newly integrated retroviruses, and possibly other retroelements, has driven the duplication and divergence of KRAB-ZNF genes. This model might explain why KRAB-ZNF genes and ERVs tend to cluster together in the genome, and why several KRAB-ZNF genes are themselves under the transcriptional control of ERV-derived LTRs. These fortuitous associations might allow the establishment of a negative-feedback loop to modulate the KRAB-ZNF response. It could also explain the apparent co-option of a subset of ERV-controlled KRAB-ZNF genes for early embryonic development in mice.

Effects on host gene expression

Disruption of gene regulation. Viruses have evolved countless strategies for hijacking and manipulating the machineries of the host to control the expression of their own genes. In particular, the proviral form of retroviruses is adapted to recruit cellular factors to promote their transcription in the context of chromatin. Each of their long terminal repeats (LTRs) contains a basal promoter for RNA polymerase II and enhancers that are responsive to diverse conditions and signals, and the LTRs are bound by many transcription factors (see below). These features enable spatiotemporal control of proviral expression, as well as control of transcription termination and polyadenylation signals. After they have been integrated into the host chromosome, any of these cis-elements has the potential to interfere with the expression of an adjacent host gene (or genes) through myriad mechanisms, including epigenetic effects.

These regulatory activities are likely to augment the deleterious effects that are associated with new EVE insertions and, for those that have reached fixation, to pose a long-lasting burden on the genome. This is reflected in the human genome by a strong statistical depletion of ERVs in the vicinity of promoter regions and near-intron splice sites, presumably because these are functionally sensitive regions in which insertions are rapidly eliminated by purifying selection. The bulk of ERV insertions reaching fixation are transcriptionally silenced in most embryonic and adult tissues through repressive epigenetic marks that are deposited and maintained by what is increasingly recognized to be a dedicated host surveillance system (BOX 5). Failure to maintain or contain these silencing marks would result in the reactivation of dormant ERV insertions, which may trigger new waves of infection or transposition or may perturb local gene expression. This was recently demonstrated by a report showing that derepression of a THE1B ERV in Hodgkin’s lymphoma transcriptionally activates the adjacent colony-stimulating factor 1 receptor (CSF1R) proto-oncogene.
REVIEWS

The characterization of endogenous viral elements (EVEs) offers a unique opportunity to carry out functional studies of extinct viruses and to shed light on various aspects of the co-evolution between viral pathogens and their hosts. Two groups independently reconstructed an infectious version of human endogenous retrovirus (HERV)-K, the most recently active but apparently now defunct ERV in the human genome. These studies led to a detailed characterization of the HERV-K replication cycle and of its level of infectivity in various cultured cells. The ability of the researchers to generate infectious HERV-K sequences that were built from segments of different endogenous HERV-K loci suggests that human cells could do the same through recombination events between HERV-K genomic loci or transcripts. These results are consistent with occasional observations of HERV-K particles, notably in teratocarcinoma cell lines, and they substantiate the hypotheses according to which HERV-K recombination or retrotransposition could have been involved in several pathologies, including cancer and HIV/AIDS. Reconstruction of full-length or partial genomes of extinct EVEs can also be used to learn about the evolution and activity spectrum of host factors that are known to protect the cell against modern viruses. For example, studies of the resurrected HERV-K (a betaretrovirus) and of partially reconstructed chimpanzee retrovirus 1 and 2 (CERV1 also known as PTERV1) and CERV2, which are both gammaretroviruses, have shown that the replication of these ancient retroviruses was most likely to be restricted to the restriction by tripartite motif-containing 5α (TRIM5α) — a host factor that currently protects primates against various retroviruses but see REF. 119 (BOX 5). By contrast, the replication of HERV-K, CERV1 and CERV2 was strongly affected by apolipoprotein B mRNA editing, enzyme-catalytic, polyprotein-like 3G (APOBEC3G), which may explain, in part, why this host factor has evolved under strong positive selection throughout the primate radiation. In another study, the TRIM5α protein from the ring-tailed lemur was shown to restrict the now-extinct lemur prosimian immunodeficiency virus (PSIV), as well as several extant lentiviruses. Functional studies of ancient viral proteins may also enhance our comprehension of the molecular mechanisms underlying tropism and governing viral host range. For instance, Soll et al. reconstructed and expressed a functional CERV2 env gene and identified the receptor used by this ancient virus to enter human cells. Interestingly, comparative sequence analysis of the receptor unveiled a series of mutational events that have rendered hamster cells resistant to CERV2. In a recent tour de force in palaeo- virology, Goldstone et al. expressed and produced crystal structures for the capsid domain of two prehistoric lentiviruses (namely, PSIV and rabbit endogenous lentivirus type K (RELIK)). Despite low levels of sequence identity, the structures display remarkable similarity to each other and to those of modern lentiviral capsids and, likewise, are bound by cyclophilin A (cypA), which is a host factor that is known to be essential for HIV-1 infectivity.

**ERVs as an abundant source of promoters.** Although the battery of cis-regulatory sequences carried by ERVs poses a hazard to host gene expression, it is also likely to predispose them to occasional co-option or ‘exaptation’ for regulatory function. Examples abound in the literature of ERV-derived sequences that have been incorporated into the ‘normal’ regulation of mammalian genes, most frequently as promoters, enhancers or polyadenylation signals. Promoters acquired from LTRs typically function as alternative, tissue-specific promoters in addition to the broadly used ancestral promoters. Hence, LTR-derived promoters are often co-opted during evolution to spatially increase or restrict gene expression patterns (for a recent example, see REF. 60). Because ERVs have been inserted and co-opted at various time points during evolution, these effects are confined to certain lineages and sometimes to certain individuals, thereby contributing to expression divergence among species and even within species.

High-throughput capture of transcription start sites (TSSs) and ultra-deep sequencing of poly(A)− RNA populations in mouse tissues and human cell lines have uncovered a staggering quantity of ERV-driven unique transcripts (~8,000 in mice and ~50,000 in humans), accounting for ~4% and ~7% of all TSSs mapped in these species, respectively. Around 40% of these transcripts show highly restricted spatial-temporal expression and most are lineage-specific, as only a small fraction of ERV insertions (~10%) are shared between human and mouse genomes. Given that a large number of ERV insertions are not yet fixed in mice, it would be interesting to examine the impact of ERV insertional polymorphism on intraspecific variation in gene expression and the possible phenotypic consequences.

Although some of the ERV-derived transcripts are produced from LTRs that function as traditional sense promoters in the 5′ region of protein-coding genes, most are actually driving the transcription of non-coding RNAs (ncRNAs). This category encompasses so-called cis−natural antisense transcripts that originate from ERVs that have been inserted in introns or in the 3′ region of protein-coding genes, as well as a profusion of other ncRNAs that emanate from intergenic or intronic elements. Furthermore, it is common for ERV LTRs to function as bidirectional promoters, thereby serving not one but two transcriptional units, upstream and downstream of their integration site. Even more surprising, promoter activity is not restricted to LTRs but frequently arises from sequences mapping within the internal, coding region of ERVs. Presumably, most of these internal promoters arose de novo from cryptic motifs that had been activated through post-insertional point mutations or positional effects. Together, these data reveal an extraordinarily complex, tightly regulated and evolutionarily labile ERV-derived ‘subtranscriptional’ framework. It may be an integral part of the ‘flexible RNA scaffolds’ that are increasingly recognized as having a central role in genome regulation.

**Dispersal of novel transcription factor binding sites.** Another common route to EVE co-option is through the donation of enhancers and binding sites for transcription factors. Recently, several landmark studies have provided support to the four-decade-old hypothesis that mobile genetic elements had a pivotal role in the dispersion and turnover of transcription factor binding sites that coordinate gene regulatory networks in mammals. Although any type of interspersed repetitive DNA may become involved in this process, the existing literature points to a disproportionate participation of ERVs in wiring large regulatory circuits. In one study, Wang et al. found that >1,500 binding sites for p53 in the human genome lie within the LTRs of class I ERVs, and that most occurred in just two families (namely, theLTR10 and MER61 families) that invaded an ancestral anthropoid genome 40–63 million years ago. Remarkably, the ERV-derived p53 sites account for approximately one-third of those mapped in human colorectal cancer cells, where p53 is

**Box 4** | ‘Resurrection’ of extinct viruses

The characterization of endogenous viral elements (EVEs) offers a unique opportunity to carry out functional studies of extinct viruses and to shed light on various aspects of the co-evolution between viral pathogens and their hosts. Two groups independently reconstructed an infectious version of human endogenous retrovirus (HERV)-K, the most recently active but apparently now defunct ERV in the human genome. These studies led to a detailed characterization of the HERV-K replication cycle and of its level of infectivity in various cultured cells. The ability of the researchers to generate infectious HERV-K sequences that were built from segments of different endogenous HERV-K loci suggests that human cells could do the same through recombination events between HERV-K genomic loci or transcripts. These results are consistent with occasional observations of HERV-K particles, notably in teratocarcinoma cell lines, and they substantiate the hypotheses according to which HERV-K recombination or retrotransposition could have been involved in several pathologies, including cancer and HIV/AIDS. Reconstruction of full-length or partial genomes of extinct EVEs can also be used to learn about the evolution and activity spectrum of host factors that are known to protect the cell against modern viruses. For example, studies of the resurrected HERV-K (a betaretrovirus) and of partially reconstructed chimpanzee retrovirus 1 and 2 (CERV1 also known as PTERV1) and CERV2, which are both gammaretroviruses, have shown that the replication of these ancient retroviruses was most likely to be restricted to the restriction by tripartite motif-containing 5α (TRIM5α) — a host factor that currently protects primates against various retroviruses. By contrast, the replication of HERV-K, CERV1 and CERV2 was strongly affected by apolipoprotein B mRNA editing, enzyme-catalytic, polyprotein-like 3G (APOBEC3G), which may explain, in part, why this host factor has evolved under strong positive selection throughout the primate radiation. In another study, the TRIM5α protein from the ring-tailed lemur was shown to restrict the now-extinct lemur prosimian immunodeficiency virus (PSIV), as well as several extant lentiviruses. Functional studies of ancient viral proteins may also enhance our comprehension of the molecular mechanisms underlying tropism and governing viral host range. For instance, Soll et al. reconstructed and expressed a functional CERV2 env gene and identified the receptor used by this ancient virus to enter human cells. Interestingly, comparative sequence analysis of the receptor unveiled a series of mutational events that have rendered hamster cells resistant to CERV2. In a recent tour de force in palaeo-virology, Goldstone et al. expressed and produced crystal structures for the capsid domain of two prehistoric lentiviruses (namely, PSIV and rabbit endogenous lentivirus type K (RELIK)). Despite low levels of sequence identity, the structures display remarkable similarity to each other and to those of modern lentiviral capsids and, likewise, are bound by cyclophilin A (cypA), which is a host factor that is known to be essential for HIV-1 infectivity.

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The long-lasting and conflicting interactions between viruses and their eukaryotic hosts have triggered the emergence of diverse antiviral defence strategies. In addition to transcriptional and post-transcriptional mechanisms of viral silencing found in various eukaryotes (BOX 3), a number of restriction factors have been identified in mammals that block the activity of retroviruses (and sometimes of other viruses) at various steps of their replication cycle124. The more taxonomically widespread and best characterized restriction factors are: the apolipoprotein B mRNA-editing, enzyme–catalytic, polypeptide-like (APOBEC) family of cytidine deaminases, which induce hypermutation of viral RNA; and the zinc finger antiviral protein (ZAP), which degrades viral RNA in the cytoplasm; and tetherin, a protein that prevents release of viral particles from the surface of producer cells. A series of elegant evolutionary analyses of these genes have shown that they have been subject to recurrent episodes of positive selection (shown by the purple branches in the figure), reflecting both the ongoing arms race (terminal purple branches in the figure) and the ancient arms race (internal purple branches in the figure), reflecting both the ongoing arms race (terminal purple branches in the figure) and the ancient arms race (internal purple branches in the figure) that take place at the host–virus interface during the evolution of primates and other mammals.

Impact of EVE gene products

Cellular expression of viral genes. Another mechanism by which EVEs can have an impact on host biology is through the cellular expression of viral coding components (but potentially also non-coding sequences). It is well-established that expression of some retroviral Env proteins, including those that are encoded by some ERVs, can modulate the host immune response17,37,39. Although thousands of env genes have been deposited in the human genome, most of them are corrupted by mutations and/or are not expressed in normal tissues or conditions owing to the defence mounted by the host to repress proviral expression39 (BOX 3). However, ERVs can escape from silencing, at least transiently, through genetic or epigenetic changes triggered by certain cellular states, including cancer37, or by environmental factors, such as diet38 or infection by related or unrelated exogenous viruses16,39. For example, the induction of interferon-α in T cells by Epstein–Barr virus (a herpesvirus) infection leads to transcriptional activation of a normally silent HERV-K18 element at the CD48

A APOBEC3G
T TRIM5α
Z ZAP

[Box 5] Antiviral defence and palaeovirology

<table>
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<tr>
<th>Time (million years ago)</th>
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<tr>
<td>40</td>
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<td>Wooly monkey</td>
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<td>Titi</td>
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Current primate lentiviruses
<200,000 years old

ω > 1
ω < 1

Strongly activated. Furthermore, Wang et al.71 were able to confirm that binding of p53 to some of these ERVs correlated with p53-dependent activation of the nearest adjacent gene. Thus ERV-derived p53 sites seem to represent evolutionarily young functional cis-elements with physiological relevance. It is remarkable that such a large subset of the network of human p53 targets has apparently emerged from primate-specific ERVs because p53 is a deeply conserved, pleiotropic regulator of many biological pathways, and it is often coined the ‘guardian of the genome’ because of its tumour suppressor activity72.

No less astounding are the results reported by two studies35,37,38 showing that ERV’s have contributed thousands of binding sites for OCT4 (also known as POU5F1 and Nanog (accounting for 7–15% of all binding sites mapped in human or mouse embryonic stem cells (ESCs)), which are two master regulators of the pluripotency gene network of ESCs. Because these ERVs are specific to either primate or rodent lineages, the data imply a massive emergence of novel binding sites for these two transcription factors in each lineage. Some ERV families have extensively contributed to the wiring of this network. For example, 235 (33.2%) of 767 LTR9B elements are bound by OCT4 in human ESCs—an 82-fold enrichment over the number expected based on the sheer density of LTR9B elements in the genome35. Alignments of the elements that are bound by OCT4 to the LTR9B consensus ancestral sequence suggest that the OCT4 binding sites pre-existed at the time of LTR9B integration in the genome. As for the aforementioned p53 binding sites, which also appear to have pre-existed in some ERV1 LTR families73, it is tempting to speculate that these binding sites were used by the ancestral retroviruses to link their own transcription to certain conditions of the host cell (namely, stress for p53) or to a specific developmental stage (namely, ESCs in the case of OCT4 and Nanog) that is favourable to their propagation. Paradoxically, the molecular manipulations exerted by exogenous viruses on their host may have promoted the subsequent co-option of some of their endogenous descendants for host regulatory functions.
locus\(^8\). Activation of this element results in overexpression of a truncated Env with superantigen activity, which induces an inflammatory cascade that is reminiscent of the onset of several autoimmune diseases. Intriguingly, overexpression of several HERV loci has been linked, with various degrees of consistency, to several autoimmune and/or neurological diseases, such as multiple sclerosis, type 1 diabetes, rheumatoid arthritis and schizophrenia\(^{16,37,55,56}\). Some ERV-encoded Env proteins also possess immunosuppressive activities that can promote tumorigenesis, as indicated by several studies with mouse cancer models\(^7\). Interestingly, the same immunosuppressive properties may be beneficial at a specific developmental stage or in certain tissues: for example, in the placenta syncytiotrophoblast, as described below.

### Domestication of viral genes by the host

Despite the potentially disastrous consequences of expressing cellular genes of viral origin, there are circumstances in which viral gene products have been usefully recruited by the host. One common functional theme of domestication is ‘fighting fire with fire’: an EVE produces a protein that offers immunity against an exogenous virus. A classic example is endogenous Jaagsieke sheep retrovirus (enJSRV), which protects the sheep genital tract from exogenous JSRV infection at two levels\(^8\). First, enJSRVs encode an Env that binds a subset of the cellular receptors used by JSRV, decreasing their availability to exogenous particles and leading to a substantial reduction in viral entry\(^7\). Second, enJSRVs express a misfolded form of Gag protein, which co-assembles with JSRV Gag to form chimeric viral particles that are targeted for degradation by the proteasome\(^8\). Another well-known example is the Friend virus susceptibility 1 (Fv1) gene of mice, which derives from the gag gene of an extinct spumavirus-like ERV that infiltrated an ancient mouse genome about 7 million years ago\(^8\) and now restricts murine leukaemia virus (MLV), a very distantly related retrovirus. Although the molecular mechanism by which Fv1 defeats MLV remains poorly understood, recent results indicate that it requires direct interaction between FV1 and the MLV capsid\(^8\). Following co-option, enJSRVs and Fv1 have undergone rapid evolution by positive (or diversifying) selection\(^14,15\). This is a tell-tale signature of many antiviral host factors (BOX 5) that is thought to reflect a molecular arms race arising from conflicting interactions with exogenous retroviruses\(^8\). In the case of sheep enJSRV and JSRV, the arms race is illustrated by the recent emergence (<200 years ago) of JSRV variants that are capable of escaping enJSRV-mediated restriction\(^8\). A function in viral defence has also been hypothesized for several other EVE-derived genes that appear to have evolved, at least transiently, under selective constraint\(^16,21,27,87-91\).
Concluding remarks and perspectives

The bounty of EVEs that has recently been unearthed from eukaryotic genomes is providing a treasure trove of information on the vast period separating the early origin of primitive viruses and the evolutionary dynamics of contemporary viruses. Notably, these genomic fossils are revealing that many modern viral families have much deeper evolutionary roots than was previously anticipated. In turn, the dating of these ancient viral sequences yields new calibration points enabling estimates of viral substitution rates across a macro-evolutionary timescale. These long-term rates turn out to be dramatically slower than those inferred from the analysis of modern viruses. Although this finding might seem surprising, it is inescapable, as one would predict that viral sequences that have diverged from their modern relatives for dozens of millions of years would be unrecognizable over such extended periods if they had evolved under the substitution rates inferred for circulating viruses. Whether the differences between short-term and long-term rates of viral evolution have biological underpinnings or whether they can merely be accounted for by methodological issues remains an open question. Improving our understanding of viral substitution rates is important for making accurate inferences on virus origin and evolution, but it is also crucial for the better monitoring and prediction of the epidemiological trajectory of clinically relevant viruses.

Although the impact that EVEs have on eukaryotic genome evolution has long been appreciated, until recently, most studies in this area were limited to one or a few candidate loci. The development of high-throughput technologies is beginning to provide a genome-wide perspective on the contribution of EVEs to cellular function, and the magnitude of this contribution is exceeding even the boldest predictions. Currently, most of our knowledge of the impact of EVEs on the host genome is based on the study of ERVs, in part because of their numerical dominance, but we can anticipate that further analysis of non-retroviral EVEs will yield new insights. For example, it will be interesting to evaluate whether EVEs can generate antiviral immunity through other mechanisms than those known for ERVs and, in particular, through an RNA-mediated response 97. Furthermore, reconstruction and expression of ancient viruses, as well as the study of their interactions with host factors, could be extended to non-retroviral families using EVEs as template sequences. Thus far, such studies have dealt with the interaction of ancient ‘resurrected’ retroviral sequences with contemporary host factors (BOX 4). However, to delineate the history and processes of host–virus co-evolution more finely, it would be necessary to study the interaction of ancestral sequences that have been reconstructed for both virus and host factors within a robust phylogenetic framework 98.

Undoubtedly, the discovery of new EVEs will continue to widen our knowledge of viral evolution and of the impact that viruses have on their host beyond their immediate pathogenic effects. In addition, the study of EVEs offers an alternative source of information on viruses that is often directly related to those plaguing humans, crops or domestic animals, which has great potential to reveal new targets and avenues for the development of innovative antiviral strategies.

4. Katsourakis, A. & Gifford, R. J. Endogenous viral elements in animal genomes. PLoS Genet. 6, e1001191 (2010). This paper presents a systematic in silico mining of EVEs in animal genomes (that were available at the time), revealing that all major types of eukaryotic viruses can be endogenized.
Along with reference 45, this study blurs the boundary between retrotransposons and retroviruses and suggests an evolutionary continuum between the two.


This was one of the first reports of non-retroviral EVEs in mammalian genomes and an experimental demonstration that Borna disease virus DNA can spontaneously integrate in the genome of human infected cells.


Belyi, V. A., Levina, A. J. & Skalka, A. M. Sequences from ancestral single-stranded DNA viruses in vertebrate genomes: the parvoviridae and circoviridae are more than 40 to 50 million years old. J. Virol. 84, 12458–12462 (2010).


The authors provide an excellent review of the concepts used and methods to delineate the epidemiological dynamics of clinically relevant viruses.


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This paper provides a comprehensive characterization of the various events that took place during the recent common ancestor arm race between domesticated endogenous betaretroviruses and their exogenous counterparts in sheep.


References 108–111 unveil fundamental discoveries in proviral silencing in mammalian ESCs.


The above two references reconstitute an infectious progenitor for a human endogenous retrovirus.


This paper provides a vivid demonstration of the power of evolutionary sequence analysis to shed crucial insight into the interaction of host restriction factors with their viral targets.


Lin, C. L. et al. Persistent Hz-1 virus infection in insect cells: evidence for insertion of viral DNA into host chromosomes and viral infection in a latent status. J. Virol. 73, 128–139 (1999).


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Competing interests statement
The authors declare no competing financial interests.

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