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Cryptic inoviruses revealed as pervasive in bacteria and archaea across Earth’s biomes

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Bacteriophages from the Inoviridae family (inoviruses) are characterized by their unique morphology, genome content and infection cycle. One of the most striking features of inoviruses is their ability to establish a chronic infection whereby the viral genome resides within the cell in an exclusively episomal state or integrated into the host chromosome and virions are continuously released without killing the host. To date, a relatively small number of inovirus isolates have been extensively studied, either for biotechnological applications, such as phage display, or because of their effect on the toxicity of known bacterial pathogens including Vibrio cholerae and Neisseria meningitidis. Here, we show that the current 56 members of the Inoviridae family represent a minute fraction of a highly diverse group of inoviruses. Using a machine learning approach leveraging a combination of marker gene and genome features, we identified 10,295 inovirus-like sequences from microbial genomes and metagenomes. Collectively, our results call for reclassification of the current Inoviridae family into a viral order including six distinct proposed families associated with nearly all bacterial phyla across virtually every ecosystem. Putative inoviruses were also detected in several archaeal genomes, suggesting that, collectively, members of this supergroup infect hosts across the domains Bacteria and Archaea. Finally, we identified an expansive diversity of inovirus-encoded toxin–antitoxin and gene expression modulation systems, alongside evidence of both synergistic (CRISPR evasion) and antagonistic (superinfection exclusion) interactions with co-infecting viruses, which we experimentally validated in a Pseudomonas model. Capturing this previously obscured component of the global virosphere may spark new avenues for microbial manipulation approaches and innovative biotechnological applications.

Inoviruses, bacteriophages from the Inoviridae family, exhibit unique morphological and genetic features. While the vast majority of known bacteriophages carry double-stranded DNA (dsDNA) genomes encapsidated into icosahedral capsids, inoviruses are instead characterized by rod-shaped or filamentous virions, circular single-stranded DNA genomes of ~5–15 kb and a chronic infection cycle1–3 (Fig. 1a). Owing to their unique morphology and simple genome amenable to genetic engineering, several inoviruses are widely used for biotechnological applications, including phage display or as drug delivery nanocarriers4–7. Ecologically, cultivated inoviruses are known to infect hosts from only 5 bacterial phyla and 10 genera but can have significant effect on the growth of their hosts when cultivated in the laboratory and can thus easily evade detection. Furthermore, established computational approaches for the detection of virus sequences in whole-genome shotgun sequencing data are not efficient for inoviruses because of their unique and diverse gene content18–20 (Fig. 1b). Finally, inoviruses are probably undersampled in viral metagenomes due to their long, flexible virions with low buoyant density11,22.

Here, we unveil a substantial diversity of 10,295 inovirus sequences, derived from a broad range of bacterial and archaeal hosts, and identified through an exhaustive search of 56,868 microbial genomes and 6,412 shotgun metagenomes using a custom computational approach to identify putative inovirus genomes. These sequences reveal that inoviruses are far more widespread, diverse and ecologically pervasive than previously appreciated, and provide a robust foundation to further characterize their biology across multiple hosts and environments.

**Results**

Inoviruses are highly diverse and globally prevalent. To evaluate the global diversity of inoviruses, an analysis of all publicly available inovirus genomes was first conducted to identify characteristic traits that would enable automatic discovery of divergent inovirus sequences (Supplementary Table 1). Across the 56 known Inoviridae genomes, the gene encoding the morphogenesis (pl) protein, an ATPase of the FtsK–HerA superfamily, represented the only

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Inovirus replication/infection cycle

Host cell entry

Production of new infectious particle (no host cell death)

Viral-encoded extrusion machinery (includes pl)

ssDNA-binding protein

Integrated inovirus

Episomal inovirus (ssDNA)

Propionibacterium phage B5

Inovirus genome diversity

<table>
<thead>
<tr>
<th>Structural proteins</th>
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<tr>
<td>Enterobacteria phage fd</td>
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<tr>
<td>Ralstonia phage HzM3</td>
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<tr>
<td>Vibrio phage VdJphi</td>
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<tr>
<td>Propionibacterium phage B5</td>
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Step 1. pI-like ATPase detection (hmmsearch, score ≥ 30)

Step 2. Extension based on inovirus-like genome context (random forest classifier, score ≥ 0.9)

Step 3 (optional). Genome ends prediction (integration site or circular contig)

Inovirus sequence identification approach

Public (meta)genomes mining

56,868 (draft) genomes (bacteria and archaea)

6,412 shotgun metagenome assemblies

7,458 hits

1,477 predicted complete

3,098 partial or ‘fuzzy’ prophages

1,818 predicted complete

3,902 partial or ‘fuzzy’ prophages

Fig. 1 | Overview of inovirus infection cycle, diversity and sequence detection process. a, Schematic of the inovirus persistent infection cycle and virion production. Inovirus genomes and particles are not to scale relative to the host cell and genome. ssDNA, single-stranded DNA. b, Comparison of selected inovirus genomes from isolates. The pI-like genes (the most conserved genes) are coloured in red, and sequence similarity between these genes (based on blastp) is indicated with coloured links between genomes. Putative structural proteins that can be identified based on characteristic features (gene length and presence of a TMD) are coloured in grey. Other genes are coloured in grey. c, Representation of the custom inovirus detection approach. The pI-like ATPase gene is coloured in red and other genes are coloured in grey. Dotted arrows indicate the region around pI-like genes that were searched for signs of an inovirus-like genome context and attachment site (see Supplementary Notes). d, Results of the search for inovirus sequences in prokaryote genomes and assembled metagenomes, after exclusion of putative false positives through manual inspection of predicted pI proteins (see Supplementary Notes). Predictions for which genome ends could be identified are indicated in green, while predictions without clear ends (that is, partial genomes or ‘fuzzy’ prophages with no predicted att site) are in blue, adding up to 10,295 curated predictions in total. Sequences for which no inovirus genome could be predicted around the initial pI-like gene are in grey. See also Supplementary Figs. 1–3.

Inovirus sequences were identified in 6% of bacterial and archaeal genomes (3,609 of 56,868) and 35% of metagenomes (2,249 of 6,412). More than half of the species (n = 3,675) were exclusively composed of sequences assembled from metagenomes. These revealed that inoviruses are found in every major microbial habitat whether aquatic, soil or human associated, and throughout the entire globe (Fig. 2 and Supplementary Notes). Hence, inoviruses are much more diverse than previously estimated and globally distributed.

Inoviruses infect a broad diversity of bacterial hosts. To examine the host range of these inoviruses, we focused on the 2,284 inovirus species directly associated with a host, that is, proviruses identified using genome-wide average nucleotide identity (ANI), and only 38 of these included isolate inovirus genomes. About one-third of these species (30%) encoded an ‘atypical’ morphogenesis gene, with an amino-terminal instead of carboxy-terminal TMD (Supplementary Fig. 3). Although this atypical domain organization has been observed in four isolate species currently classified as inoviruses, some of these inovirus-like sequences might eventually be considered as entirely separate groups of viruses. Sequence accumulation curves did not reach saturation, highlighting the large diversity of inoviruses yet to be sampled (Supplementary Fig. 4).

Inovirus sequences were identical using genome-wide average nucleotide identity (ANI), and only 38 of these included isolate inovirus genomes. About one-third of these species (30%) encoded an ‘atypical’ morphogenesis gene, with an amino-terminal instead of carboxy-terminal TMD (Supplementary Fig. 3). Although this atypical domain organization has been observed in four isolate species currently classified as inoviruses, some of these inovirus-like sequences might eventually be considered as entirely separate groups of viruses. Sequence accumulation curves did not reach saturation, highlighting the large diversity of inoviruses yet to be sampled (Supplementary Fig. 4).

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Inoviruses sporadically transferred from bacterial to archaeal hosts. Although no archaea-infecting inoviruses have been reported so far, some inovirus sequences were associated with members of two archaeal phyla (Euryarchaeota and Aenigmarchaeota), which suggests that inoviruses infect hosts across the entire prokaryotic diversity (Fig. 3). These putative archaeal proviruses encoded the full complement of genes expected in an active inovirus (Fig. 4a and Supplementary Notes). Using PCR, we further confirmed the presence of a circular, excised form of the complete inovirus genome for the provirid identified in the Methanobrevibacter smithii MoB genome (Fig. 4b, Supplementary Fig. 6 and Supplementary Notes). This indicates that our predictions in archaeal genomes are probably genuine inoviruses.

Few groups of viruses include both bacteriophages and archaeoviruses. Such evolutionary relationships between viruses infecting hosts from different domains of life might signify either descent from an ancestral virus that infected the common ancestor of bacteria and archaea, or horizontal virus transfer from one host domain to the other26,28. Here, the four archaea-associated inoviruses were clearly distinct from most other inoviruses and clustered only with metagenomic sequences in pI phylogeny (Fig. 4c). In addition, they were classified into two different proposed families (see below) corresponding to the two host groups, reflecting clear differences in their gene content (Fig. 4a,c and Supplementary Notes). The high genetic diversity of these archaea-associated inoviruses, combined with the lack of similarity to bacteria-infecting species, suggest that they are not derived from a recent host switch event.

A possible scenario would involve an ancestral group of inoviruses infecting the common ancestor of archaea, as postulated for the double-jelly-roll virus lineage28. However, to be confirmed, this hypothesis would require the detection of additional inoviruses in other archaeal clades or an explanation as to why inoviruses were retained only in a handful of archaeal hosts. Instead, on the basis of the current data, a more likely scenario involves ancient and rare events of interdomain inovirus transfer from bacteria to archaea,
including possibly to a *Methanosarcina* host for which substantive horizontal transfers of bacterial genes have already been reported\(^{30}\).

**Gene content classification reveals six distinct inovirus families.** The vast increase of inovirus sequences provided a great opportunity for re-evaluation of the inovirus classification and the development of an expanded taxonomic framework for the large number of inovirus species identified. Similar to other bacterial viruses, especially temperate phages\(^{30}\), inovirus genomes display modular organization and are prone to recombination and horizontal gene transfers\(^{31}\) (Supplementary Fig. 7). Hence, we opted to apply a bipartite network approach, in which genomes are connected to gene families, enabling a representation and clustering of the diversity based on shared gene content. A similar approach has been previously employed for the analysis of DNA and RNA viruses, and was shown to be efficient in cases in which the genomes to be clustered share only a handful of genes\(^{32-34}\). Here, this approach yielded 6 distinct groups of genomes divided into 212 subgroups (Fig. 5a and Supplementary Table 3).

A comparison of marker gene conservation between these groups and established viral taxa suggested that the former *Inoviridae* family should be reclassified as an order, provisionally divided into 6 candidate families and 212 candidate subfamilies, with few shared genes across candidate families (Fig. 5a, Supplementary Fig. 7 and Supplementary Notes). Beyond gene content, these proposed families also displayed clearly distinct host ranges as well as specific genome features, particularly in terms of genome size and coding density (Supplementary Fig. 7). Thus, we propose to establish these as candidate families named ‘Protoinoviridae’, ‘Vespertilinoviridae’, ‘Amplinoviridae’, ‘Paulinoviridae’, ‘Densinoviridae’ and ‘Photinoviridae’, on the basis of their isolate members and characteristics (see Supplementary Notes). If confirmed, and compared with currently recognized inoviruses, the genomes reported here would increase diversity by 3 families and 198 subfamilies.

The host envelope organization seems to play an important role in the evolution of inoviruses, which is reflected in their classification: members of the ‘Protoinoviridae’ and ‘Amplinoviridae’ are associated with dierd hosts—that is, Gram-negative bacteria with an outer membrane—whereas the other candidate families are associated with monoderm hosts or hosts without a cell wall (Supplementary Fig. 7). Conversely, no structuring by biome was observed and all proposed families were broadly detected across multiple types of ecosystems. Hence, we propose here a classification of inovirus diversity into six families based on gene content with coherent host ranges and
specific genomic features, which strongly suggests that they represent ecologically and evolutionarily meaningful units.

**Inovirus genomes encode an extensive functional repertoire.** The extended catalogue of inovirus genomes offers an unprecedented window into the diversity of their genes and predicted functions. Overall, 68,912 proteins were predicted and clustered into 3,439 protein families but only 8,552 singletons (see Methods). A putative function was predicted for 1,133 of the 3,439 iPFs identified as potentially structural based on their size and presence of a TMD (see Methods). Notably, each candidate inovirus family seemed to be associated with a specific set of structural proteins, including distinct major coat iPFs (Supplementary Fig. 8). Conversely, genome replication and integration-associated iPFs were broadly shared across candidate families (Fig. 5b). This confirms that replication-associated and integration-associated iPFs are among the most frequently exchanged among viral genomes and with other mobile genetic elements, especially in small single-stranded DNA viruses.

In addition, 15 distinct sets of iPFs representing potential toxin–antitoxin pairs were identified across 181 inovirus genomes, including 10 unaffiliated iPFs that were predicted as putative antitoxins through co-occurrence with a toxin iPF (Fig. 5b and Supplementary Table 5; see Methods). These genes typically stabilize plasmids or prophages in host cell populations, although alternative roles in stress response and transcription regulation...
have been reported\(^3\). In addition, toxin–antitoxin systems often affect host cell phenotypes, such as motility or biofilm formation\(^1\). Here, similar toxin proteins could be associated with distinct and seemingly unrelated antitoxins and vice versa, suggesting that gene shuffling and lateral transfer occur even within these tightly linked gene pairs (Supplementary Fig. 9). All but one toxin–antitoxin pairs were detected in proteobacteria-associated inoviruses, most likely because of a database bias. Thus, numerous uncharacterized iPFs across other candidate families of inoviruses may also encode previously undescribed toxin–antitoxin systems and, more generally, host manipulation mechanisms.

**Inoviruses can both leverage and restrict co-infecting viruses.** Finally, we investigated potential interactions between persistently infecting inoviruses, other co-infecting viruses, and the host clustered regularly interspaced short palindromic repeat (CRISPR)–CRISPR-associated (Cas) immunity systems. CRISPR–Cas systems typically target bacteriophages, plasmids and other mobile genetic elements\(^3\). We detected 1,150 inovirus-matching CRISPR spacers across 42 bacterial and 1 archaeal families. These spacers were associated with three types and eight subtypes of CRISPR–Cas systems, indicating that inoviruses are broadly targeted by antiviral defences (Fig. 6a, Supplementary Table 6 and Supplementary Notes). Several

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**Fig. 5 | Inovirus genome sequence space and gene content.** a, The bipartite network links genes represented as PCs in squares to proposed subfamilies represented as circles with a size proportional to the number of species in each candidate subfamily (log\(_{10}\) scale), grouped and coloured by proposed family. Proposed subfamilies that include viral isolates are highlighted with a black outline. Candidate subfamilies are connected to PCs when ≥50% of the subfamily members contained this PC or ≥25% for the larger proposed subfamilies (see Methods). b, Distribution of iPFs detected in two or more genomes, associated with genome replication, genome integration and toxin–antitoxin systems (see Supplementary Table 5). The presence of at least one sequence from an iPF (column) in a proposed family (row) is indicated with a grey square. Rolling circle replication (RCR) iPFs include only the RCR endonuclease motif, with the exception of iPF_00203 (highlighted with an asterisk), which also includes the C-terminal 53H motif typical of eukaryotic single-stranded DNA viruses. Transposases used by selfish integrated elements are indistinguishable from transposases domesticated by viral genomes using sequence analysis only; hence, these genes are gathered in a single 'integration or selfish element' category. All toxin–antitoxin pairs were predicted to be of type II, except for Toxin_3 (highlighted with an asterisk), which was predicted to be type IV. S-rec, serine recombinase; Y-rec, tyrosine recombinase. See also Supplementary Figs. 7–9.
host groups, most notably Neisseria meningitidis, were clear outliers, that is, they displayed a particularly high ratio of inovirus-derived spacers suggesting a uniquely high level of spacer acquisition and inovirus infection (Fig. 6a). This is particularly notable because inoviruses were recently suggested to increase Neisseria meningitidis pathogenicity\(^1\) and hints at conflicting host–inovirus interactions in this specific group.

Next, we examined instances of ‘self-targeting’, that is, CRISPR spacers matching an inovirus integrated in the same host genome. Among the 1,429 genomes that included both a CRISPR–Cas system and an inovirus prophage, only 45 displayed a spacer match(es) to a resident prophage (Supplementary Table 6), suggesting that self-targeting of these integrated elements is lethal and strongly counter-selected\(^6\). This was confirmed experimentally using the Pseudomonas aeruginosa strain PA14 harbouring an integrated inovirus prophage (P\(i\)F), for which the introduction of a plasmid carrying P\(i\)-targeting CRISPR spacers was lethal (Supplementary Fig. 10a). In the 45 cases of observed self-targeting, the corresponding CRISPR–Cas system is thus probably non-functional or inhibited via an anti-CRISPR (acr) locus, as recently described in dsDNA phages\(^8\). We first evaluated ten hypothetical proteins, and hence candidate Acr proteins, from self-targeted inoviruses infecting P. aeruginosa; however, none showed Acr activity (Supplementary Notes and Supplementary Fig. 10b). Alternatively, inoviruses could leverage the Acr activity of a co-integrated virus. This hypothesis was further reinforced by the fact that 43 of the 45 self-targeted inoviruses were detected alongside co-infecting dsDNA phages, with 5 of these encoding known acr genes (Supplementary Table 6). We confirmed experimentally cross-protection by trans-acting Acr in the P. aeruginosa PA14 model, and observed that co-infection with an acr-encoding dsDNA bacteriophage rescued the lethality caused by self-targeted inoviruses (Supplementary Notes and Supplementary Fig. 10a).

While this represents an instance of beneficial co-infection for inoviruses, we also uncovered evidence of antagonistic interactions between inoviruses and dsDNA bacteriophages. Specifically, 2 of the 10 inovirus-encoded hypothetical proteins tested strongly limited infection of Pseudomonas cells by different bacteriophages (Fig. 6b, Supplementary Figs. 10c and 12 and Supplementary Notes). This superinfection exclusion effect was found to be host and virus strain dependent, which could drive intricate tripartite coevolutionary dynamics. Thus, these preliminary observations indicate that inoviruses may not only evade CRISPR–Cas immunity by leveraging the Acr activity of co-integrated phages, but also significantly influence the infection dynamics of unrelated co-infecting viruses through superinfection exclusion (Fig. 6c). Multiple effects of virus–virus interactions on host ecology and evolution have been recently highlighted or proposed, and are the main focus.

**Fig. 6 | Interaction of inoviruses with CRISPR–Cas systems and co-infecting viruses.** a. Proportion of the spacers matching an inovirus genome and the corresponding distribution of CRISPR–Cas systems. The proportions are calculated only on hosts with at least one spacer matching an inovirus sequence, with hosts grouped at the family rank (hosts unclassified at this rank were not included). In the boxplot, the lower and upper hinges correspond to the first and third quartiles, respectively, and the whiskers extend no further than ±1.5 times the interquartile range. Outliers identified as values larger than the third quartile plus three times the interquartile range from the complete distribution are highlighted in red. The number of observations is indicated next to each family. b. Instances of superinfection exclusion observed when expressing individual inovirus genes in two P. aeruginosa strains: PAO1 and PA14. From top to bottom: cells were transformed with an empty vector, one expressing gene 2687473927 or one expressing gene 2687473923. For each construct, host cells were challenged with serial dilutions (from left to right) of phages: \(\phi\)JB30 and \(\phi\)DM3m. The formation of plaques (dark circles) indicates successful infection, whereas the absence of plaques indicates superinfection exclusion. Interpretation of infection outcome is indicated to the right of each lane, with successful infection represented by a phage symbol and superinfection exclusion represented by a phage symbol barred by a red cross. Results from additional superinfection exclusion experiments are presented in Supplementary Figs. 10 and 12. All superinfection experiments were conducted twice and produced similar results. c. Schematic representation of the possible mutualistic or antagonistic interactions between inovirus prophages (red) and co-infecting Caulovirales (blue). Mutualistic interactions include suppression of the CRISPR–Cas immunity, especially for integrated inoviruses targeted by the host cell CRISPR–Cas system (‘self-targeting’). Antagonistic interactions primarily involve superinfection exclusion, in which a chronic inovirus infection prevents a secondary infection by an unrelated virus.
of a nascent ‘sociovirology’ field20. Given their broad host range (Fig. 3), frequent detection alongside non-inovirus prophages (Supplementary Fig. 5), extended host cell residence time and the experimental results presented here, inoviruses could be driving many of these interactions and are undeniably important to consider in this framework.

Discussion

Taken together, the results presented here call for a complete re-evaluation of the diversity and role of inoviruses in nature. Collectively, inoviruses are distributed across all biomes and display an extremely broad host range spanning both prokaryotic and eukaryotic hosts. Comparative genomics revealed evidence of long-lasting virus–host co-evolution, leading to strong partitioning of inovirus diversity by host taxonomy, high inovirus prevalence in several microbial groups, including major pathogens, and potential interdomain transfer. Even though small (5–20 kb), their genomes encode a large functional diversity shaped by frequent gene exchange with unrelated genomes of viruses, plasmids and transposable elements. Some of the many uncharacterized inovirus genes probably encode molecular mechanisms at the interface of virus–host and virus–virus interactions, such as modulators of the CRISPR–Cas systems, superinfection exclusion genes or toxin–antitoxin modules. This expanded and restructured catalogue of 5,964 distinct inovirus genomes thus provides a renewed framework for further investigation of the different effects that inoviruses have on microbial ecosystems, and exploration of their unique potential for biotechnological applications and manipulation of microorganisms.

Methods

Construction of an Inoviridae genome reference set. Genome sequences affiliated to Inoviridae and ≥2.5 kb were downloaded from NCBI GenBank and RefSeq on 14 July 2017 (refs. 40–44). These were clustered at 98% ANI to remove duplicates and screened for cloning vectors and partial genomes (Supplementary Table 1). Two of these genomes (Stenotrophomonas phage phiSM99, NC_007189, and Balstonia phage RSS30, NC_021862) presented an unusually long segment (≥1 kb) without any predicted genes, associated with a lack of short genes that are typical of Inoviridae. For these, genes were predicted de novo using Glimmer45 trained on their host genomes (NC_010943 for phiSM99 and NC_003295 for RSS30) with standard genetic code. Similarly, genes for Acholeplasma phage MV-L1 (NC_001341) were predicted de novo using Glimmer with genetic code 4 (Mycoplasma/Spiroplasma) and trained on the host genome (NC_010163), followed by a manual curation step to integrate both RefSeq-annotated genes and these newly predicted CDS.

Protein clusters (PCs) were computed from these genomes from an all-versus-all blastp of predicted CDS (thresholds: e < 0.001, bit score ≥ 30) and clustered with InMap46. Sequences from these PCs were then aligned with MUSCLE47, transformed into an HMM profile and compared with each other using HHSearch44 (cut-offs: probability ≥ 90% and coverage ≥ 50%, or probability ≥ 99%, coverage ≥ 20% and hit length ≥ 100). The larger clusters generated through this second step are designated here as iPFs. Only ten PCs were clustered into larger iPFs, but these were consistent with the functional annotation of these proteins. For instance, one iPF combined two PCs both composed of replication initiation proteins. Marker genes were identified from a bipartite network linking Inoviridae genomes to iPFs (Supplementary Fig. 1). Only the genes encoding the morphogenesis (pl) protein represented good candidates for a universally conserved gene across all members of the Inoviridae, and HMM profiles were built for the three pl iPFs. To optimize these profiles, sequences were first clustered at 90% amino acid identity with cd-hit48, then aligned with MUSCLE47 and the profile generated with hmmbuild49.

These reference genomes were also used to evaluate the detection of the Inoviridae structural proteins based on protein features beyond sequence similarity (see Supplementary Notes). Here, signal peptides were predicted using SignalP inoviridae (Supplementary Table 2) were compared with the reference morphogenesis (pl) proteins with hmmssearch49 (hmmer.org, score ≥ 30 and e < 0.001) for the pl-like iPFs and blastp44 (bit score ≥ 50) for the singleton pl protein (Acholeplasma phage MV-L1). These included 54,405 bacterial genomes, 1,304 archaeal genomes and 1,149 plasmid sequences. A total of 6,819 hits were detected, from which 795 corresponded to complete inovirus genomes. These included 213 circular contigs, that is, likely complete genomes, and 582 integrated prophages with canonical attachment (att) sites, that is, direct repeats of ≥10 bp in a tRNA or outside of an integrase gene. All sequences were manually inspected, and there were plausible inovirus genomes (see Supplementary Notes). The predicted pl proteins from the curated genomes were then added to the references to generate new improved HMM models. Using these improved models, an additional set of 639 putative pl proteins was identified. New models were built from these proteins and used in a third round of searches, which did not yield any additional genuine inovirus sequence after manual inspection.

An automatic classifier was trained on this extended inovirus genome catalogue, that is, the reference genomes and the 795 manually curated genomes, to detect putative inovirus fragments around pl-like genes, based on 10 distinctive features of inovirus genomes (Supplementary Fig. 2 and Supplementary Notes). These 795 manually curated genomes were identified from 17 host phyla (order: Tenericutes/Cyanobacteria) and were later classified into 5 proposed families and 245 proposed subfamilies (see below ‘Gene-content-based clustering of inovirus genomes’). Three types of classifiers were tested: random forest (function ‘randomForest’ from R package randomForest40 using 2,000 trees, other parameters left as default), random forest with conditional inference (function ‘forest’ from R package party47 using 2,000 trees, other parameters left as default) and a generalized linear model with lasso regularization (function ‘glmnet’ from R package glmnet51). The efficiency of classifiers was evaluated using a ten-fold cross-validation in which the input data set was partitioned into ten equal-sized subsamples, with one retained for validation and the other nine used for training through the ten possible permutations. Results were visualized as a ROC curve generated with ggplot2 (refs. 52,53). The importance of features in the random forest classifier was evaluated using the function ‘importance()’ from the package ‘randomForest’.

On the basis of the inflection point observed on the ROC curves, the random forest was selected as the optimal method as it provided the highest true-positive rate (>92%) for false-positive rates of <1% (Supplementary Fig. 2). This model was then used to classify all putative inovirus fragments that had not been identified as complete genomes previously, using a sliding window approach (up to 30 genes around the putative pl protein), and looking for the fragment with the maximum score in the random forest model (if ≥0.9). For the predicted integrated prophages, putative non-canonical att sites were next searched as direct repeats (10 bp or longer) around the fragment. Overall, 3,908 additional putative inovirus sequences were detected, including 738 prophages flanked by direct repeats. A similar approach was used to search for inovirus sequences in 6,412 metagenome assemblies (Supplementary Table 2). Predicted proteins were compared with the 4 HMM profiles as well as to the Acholeplasma phage MV-L1 singleton sequence, which led to 27,037 putative pl proteins using the same thresholds as for isolate genomes. The final data set of inovirus sequences predicted from these metagenome assemblies consisted of 6,094 sequences, including 922 circular contigs, 44 prophages with canonical att sites (direct repeats of 10 bp or longer in a tRNA or next to an integrase) and 994 prophages with non-canonical att sites (direct repeats of 10 bp or longer).

Clustering of inovirus genomes in putative species. Next, we sought to cluster these putative inovirus genomes along with the previously collected reference genomes to remove duplicated sequences and to select only one representative per species. This clustering was conducted according to the latest guidelines submitted to the International Committee on Taxonomy of Viruses (ICTV) for Inoviridae, that is, “95% DNA sequence identity as the criterion for demarcation of species”54 (https://talk.ictvonline.org/files/ictv_official_taxonomy_updates_since_the_8th_report/m/prokaryote-official/6774/download), and included our 10,295 sequences alongside the 56 reference genomes. Notably, however, predictions spanning multiple tandemly integrated inovirus prophages had to be processed separately, otherwise they would have led to clusters gathering multiple species. To detect these cases of tandem insertions, we searched for and clustered separately all predictions with multiple pl proteins, as this gene is expected to be present in single copy in inoviruses (n = 800 sequences).

These non-tandem seed sequences were then clustered incrementally with priority given to complete genomes over partial genomes as well as fragments identified in microbial genomes over fragments from metagenomes. First, circular contigs and prophages with canonical att sites identified in a microbial genome were clustered, and all other fragments were affiliated to these seed sequences. Next, unaffiliated fragments detected in microbial genomes and with non-canonical att sites (that is, simple direct repeats) were clustered together, and other fragments were affiliated to this second set of seed sequences. Finally, the remaining unaffiliated sequences detected in microbial genomes were clustered together. This allowed us to use the more ‘certain’ predictions (that is, circular sequences and prophages with identified att sites) preferentially as seeds of putative species.

These 795 manually curated genomes were identified from metagenomes, as well as to separately cluster putative tandem fragments, that is, those including multiple pl proteins. All the clustering and affiliation was done with a threshold of 95% ANI on 100% of alignment fraction (according to the ICTV guidelines), with sequence similarity computed using mummer49.
Clustering of predicted proteins from non-redundant inovirus sequences. Predicted proteins from the representative genome of each putative species were next clustered using the same approach as for the reference genomes. A clustering into PCs was first achieved through an all-versus-all Blastp using hits with e ≤ 0.001 and bit score ≥ 50 or bit score ≥ 30 if both proteins are ≤70 amino acids. HMM profiles were constructed for the 5,142 PCs and these were compared all-versus-all using HHSearch, keeping hits with ≥90% probability and ≥50% coverage or ≥99% probability, ≥20% coverage and hit length of ≥100. This resulted in 4,008 protein families (iPFs).

The PCs were subsequently used for taxonomic classification of the inovirus sequences (see below), while iPFs were primarily used for functional affiliation. iPF functional affiliations were based on the alignment of all PCs against Pfam v30 (score ≥ 30), as well as manual inspection of individual iPFs using HHPred57. PCs containing pl-like proteins were also further evaluated to identify potential false positives stemming from a related ATPas encoded by another type of virus or mobile genetic element (see Supplementary Notes). The criteria used to determine genuine inovirus pl-like PCs were: the PC members closest known functional domain was Zot (based on the hmmsearch against PFAM), the proteins contained one or two TMD (either N-terminal or C-terminal), at least half of the sequences encoding this PC also include other genes expected in an inovirus sequence such as replication initiation proteins, and no significant similarity could be identified to any other type of ATPas using HHpred. iPF families were next clustered using the same approach as for the reference genomes. A clustering of predicted proteins from non-redundant inovirus sequences.

Genome-context-based clustering of inovirus genomes. A bipartite network was built in which genomes and PCs (as nodes) are connected by an edge when a predicted protein from the genome is a member of the PC. This network was then used to classify inovirus sequences as done previously for dsDNA viruses5. PCs were retained in this network if they offer a higher pl-like protein score than any other type of ATPase using HHpred57. PCs containing pl-like proteins were also further evaluated to identify potential false positives stemming from a related ATPas encoded by another type of virus or mobile genetic element (see Supplementary Notes). The criteria used to determine genuine inovirus pl-like PCs were: the PC members closest known functional domain was Zot (based on the hmmsearch against PFAM), the proteins contained one or two TMD (either N-terminal or C-terminal), at least half of the sequences encoding this PC also include other genes expected in an inovirus sequence such as replication initiation proteins, and no significant similarity could be identified to any other type of ATPas using HHpred.

Gene-content-based clustering of inovirus genomes. A bipartite network was built in which genomes and PCs (as nodes) are connected by an edge when a predicted protein from the genome is a member of the PC. This network was then used to classify inovirus sequences as done previously for dsDNA viruses5. PCs were retained in this network if they offer a higher pl-like protein score than any other type of ATPase using HHpred57. PCs containing pl-like proteins were also further evaluated to identify potential false positives stemming from a related ATPas encoded by another type of virus or mobile genetic element (see Supplementary Notes). The criteria used to determine genuine inovirus pl-like PCs were: the PC members closest known functional domain was Zot (based on the hmmsearch against PFAM), the proteins contained one or two TMD (either N-terminal or C-terminal), at least half of the sequences encoding this PC also include other genes expected in an inovirus sequence such as replication initiation proteins, and no significant similarity could be identified to any other type of ATPas using HHpred.

Phylogenetic trees of inovirus sequences. Phylogenies of inovirus sequences were based on multiple alignment of pl protein sequences. To obtain informative multiple alignments, an all-versus-all Blastp search of all pl protein sequences was performed to identify the nearest neighbours of sequences of interest. For sequences detected in archaeal genomes, an additional 10 most closely related sequences with e ≤ 0.001, bit score ≥ 30 and a blast hit covering ≥50% of the query sequence were recruited for each archaea-associated sequence to help populate the tree. A similar approach was used for the tree based on the integrase genes from archaea-associated inoviruses: the protein sequences for the three integrase genes were compared with the NCBI nr database with blastp (bit score ≥ 50, e ≤ 0.001) to gather their closest neighbours across archaeal and bacterial genomes.

Resulting data sets were first filtered for partial sequences as follows: the average sequence length was calculated excluding the top and bottom 10% and all sequences shorter than half of this average were excluded. These protein sequences were next aligned with MUSCLE (v3.8.1751)43, automatically trimmed with trimAL (v1.4.rev15)50 (option gappytout), and trees were constructed using IQ-TREE (v1.5.5) with an automatic detection of optimal model and displayed using iTOL5. The optimal substitution model, selected based on the Bayesian information criterion, was VT + F + R5 for the pI phylogeny of archaeal inoviruses, and LG + R4 for the integrase phylogeny of archaeal inoviruses. Annotated trees are available at http://itol.embl.de/shared/Sirojx (project ‘Inovirus’).

Functional affiliation of iPFs. An automatic functional affiliation of all iPFs was performed by searching the iPFs against PFAM (data extracted from the IMG). To refine these annotations for functions of interest, namely, replication initiation proteins, integration proteins, DNA methylases and toxin–antitoxin systems, individual iPF alignments were submitted to the HHPred website7, and the alignments were visually inspected for conserved residues and/or motifs (Supplementary Table 5, motifs extracted from refs. 72,73 and 74) and used to identify the nearest neighbours of sequences of interest. For sequences detected in archaeal genomes, an additional 10 most closely related sequences with e ≤ 0.001, bit score ≥ 30 and a blast hit covering ≥50% of the query sequence were recruited for each archaea-associated sequence to help populate the tree. A similar approach was used for the tree based on the integrase genes from archaea-associated inoviruses: the protein sequences for the three integrase genes were compared with the NCBI nr database with blastp (bit score ≥ 50, e ≤ 0.001) to gather their closest neighbours across archaeal and bacterial genomes.

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explored to identify the ones with a clear associated CRISPR–Cas system and to affiliate these systems to the different types described. Only spacers for which a can gene could be identified in a region of ±10kb were retained. The CRISPR–Cas system affiliations were based on the set of candidates identified around the spacer and performed following the guidelines from ref. 12.

For host genomes with a self-targeting spacer, additional (that is, non-inovirus) prophages were detected using VirSorter6. The number of distinct prophages was also estimated using the detection of large terminus subunits (hmmsearch against PFAM database, score ≥ 30). Putative Acr and anti-CRISPR-associated (Acra) prophages were detected through previously described Acr systems6 (blastp, e ≤ 0.001 and score ≥ 30). Putative Acr and Acra proteins were identified by searching for HTH-domain-containing proteins identified based on HTH domains in the SMART database (see above) in inovirus sequences displaying a match to a CRISPR spacer extracted from the same host genome.

Microscopy and PCR investigation of a predicted provirus in *M. profundi* MobM. *M. profundi* strain MobM cells were grown in anaerobic DSMZ medium 479 at 37 °C with 5 mM methanol added as a methanogenic substitute instead of trimethylamine6. After 37 h of growth, anaerobic mitomycin C was added to the culture at a final concentration of 1.0 μg·ml−1 to induce the provirus. Samples were collected before and 4 h after induction and were filtered with 0.22-μm pore size polyethersulfone filters (Millipore, Fisher Scientific) to obtain a ‘cellular’ ((≥0.22 μm) and a ‘viral’ (<0.22 μm) fraction.

The four types of samples (with or without induction, cellular and viral fractions) were prepared and imaged at the Molecular and Cellular Imaging Center, Ohio State University, Wooster, OH, USA. An equal volume of 2× fixation (6% glutaraldehyde and 2% paraformaldehyde in 0.1 M potassium phosphate buffer pH 7.2) was added directly to the culture post-induction. Of the medium, 30 μl was applied to a formvar and carbon-coated copper grid for 5 min, blotted and then stained with 2% uranyl acetate for 1 min. Samples were examined with a Hitachi H7500 electron microscope and imaged with the SIA-L12C (16 megapixels) digital camera.

PCRs were initially run for induced and non-induced samples on both size fractions with three pairs of primers: one internal to the predicted provirus (B primers), one spanning the insertion site (P primers) and one spanning the junction of the predicted excised circular genome (C primers). The reactions were conducted for 35 cycles with denaturation, annealing and extension cycles of 0.5, 0.5 and 1.0 min at 95°C, 52.0 and 72.0°C, respectively. For C primers, numerous nonspecific amplification products were obtained with these conditions, and another set of PCRs was conducted with higher annealing temperatures of 56.5 °C and 57.5 °C, both in triplicates. The PCR product was then cleaned to remove polymerase, free dNTPs and primers (Zymo Research) and subsequently used as templates for Sanger sequencing. The resulting chromatograms were analysed using the R2 packages sangerseqR, sangeranalyseR and readR.

The extracted primary sequences were aligned to the MobM genome using blastn and MUSCLE, and the alignment was visualized with Jalview.

Experimental characterization of hypothetical proteins from self-targeted *Pseudomonas* inoviruses. Hypothetical proteins predicted on inovirus prophages, which were (1) found in *Pseudomonas* genomes, (2) predicted to be targeted by at least one CRISPR spacer from the same genome, and (3) for which no acr locus could be identified anywhere else in the same genome, were selected for further functional characterization. The ten candidate genes were first codon optimized for expression in *Pseudomonas* using an empirically derived codon usage table. Codon optimization and vendor defined synthesis constraints removal were performed using BOOST5. Synthetic DNA were obtained from Thermo Fisher Scientific and cloned in between the Sacl and Pstl sites of an *Escherichia coli* plasmid. Colony PCR was conducted with primers specific to the putative gene to verify the presence of the insertion before testing.

**P. aeruginosa** strains (PA01:p-lac I-C CRISPR–Cas, PA14 and 4386) were cultured on LB agar or liquid media at 37 °C. The pHERD30T plasmids were electroporated into *P. aeruginosa* strains, and LB was supplemented with 50 μg·ml−1 gentamicin to maintain the pHERD30T plasmid. Phages DMS3macrIF1 were amplified on PAO1, and phage JBD44a was amplified on PA14 lysogens were electroporated with 100 ng plasmid DNA, allowed to recover for 1 h in LB at 37 °C and plated on LB agar plates supplemented with 50 μg·ml−1 gentamicin and 0.1% arabinose. Colonies were enumerated after growth for 14 h at 37 °C. Transformation efficiency (TE) was calculated as colonies per microgram DNA, and the percentage TE was calculated by normalizing the TE of the CRISPR RNA-expressing plasmids to the TE of an empty vector.

To evaluate the effect of an acr locus from a co-infecting prophage on self-targeted inoviruses, strain PA14 ACRISPRI1/ACRISPRI2 (P1I) was lysogenized with phage DMS3macrIF1 by streaking out cells from solid media, inoculating and screening for colonies resistant to superinfection by DMS3macrIF1. Lysogeny was confirmed by plaque induction. The same plasmid transformation approach was then used to assess the effect of inovirus self-targeting on host cell viability.

Quantification and statistical analysis. Sequence similarity searches were conducted with thresholds of E-value ≤ 0.001 and bit score ≥ 30 or 50, the former being used mainly for short proteins. The different classifiers (random forest, conditional random forest and generalized linear model) used to identify inovirus sequences were evaluated using a tenfold cross-validation approach. For all boxplots, the lower and upper hinges correspond to the first and third quartiles, respectively, and the whiskers extend no further than 1.5 times the interquartile range.

**Reporting Summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**

The following files are available at https://genome igeni.dee.gov/portal/PhylS/Tag/PhylTag_home.html: Gb_files_inoviruses.zip; GenBank files of all representative genomes for each inovirus species; Ref_PCs_inoviruses.zip: PCVs from the references (raw fasta, alignment fasta and hmm profile); iPFs_inoviruses.zip: protein families from the extended inovirus data set (raw fasta, alignment fasta and hmm profile); MobM_C_primer_amplicon.fasta: multiple sequence alignment of the C primer products with the *Methanobulus* MobM genome (NZ_FOU01000007), confirming that C primer products span the junction of the excised genome. Accession numbers of all inovirus sequences used as reference are listed in Supplementary Table 1. Accession numbers of all genomes and metagenomes mined, including detailed information for each (meta)genome in which some inovirus sequences were detected are available in Supplementary Table 2. Finally, the list of all inovirus genome accession numbers, along with taxonomic and environmental distribution information, is provided in Supplementary Table 3.

**Code availability**

The set of scripts and models used to detect inovirus sequences is available at https://bitbucket.org/royouzi/invirius/src/master/Invirius_detector/.

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**References**


**Acknowledgements**

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**Author contributions**


**Competing interests**

The authors declare no competing interests.

**Additional information**

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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

Data were collected using a custom set of scripts specifically designed to identify inovirus genomes. These are available at https://github.com/simroux/Inovirus/tree/master/Inovirus_detector

Data analysis

For specific reference inoviruses, genes were predicted de novo using Glimmer v3. Sequence similarity searches were conducted using blast+ v2.7.1, hmmer 3.1b2, hhpred [online at https://toolkit.tuebingen.mpg.de/#/tools/hhpred] and hhsearch v2.0.15. Sequences were clustered using InfoMap 0.18.25, mummer 3.0, SDT 1.0, and cd-hit 4.7. Viral sequences (non-inoviruses) were automatically detected using VirSorter v1.0.5. Signal peptide and transmembrane domains were predicted using SignalP 4.1 and TMHMM 2.0c. Trees were built using FastTree2 and IQ-Tree 1.5.5, based on alignments computed with muscle 3.8 or MAFFT v7.294b and automatically trimmed with trimAL v1.4 or BMGE v1.12. Alignments were manually inspected using Jalview v10.0.2. Statistical analyses, sanger sequenced reads interpretation, and automatic classifier design were conducted in R v3.4.3. Secondary structure of putative inovirus major capsid proteins were predicted using Phyre v2.0. Figures were generated with R 3.4.1 using the ggplot2 package, Cytoscape v3.6.1, iTOL v4.4.1, and python v3.6.2 using matplotlib v2.0.2.

Constraints in sequences to be synthesized were automatically identified and adjusted using BOOST v1.3.3.

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All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
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- A list of figures that have associated raw data
- A description of any restrictions on data availability

The following data are available at https://genome.jgi.doe.gov/portal/Inovirus/Inovirus.home.html:
Gb_files_inoviruses.zip: GenBank files of all representative genomes for each inovirus species.
Ref_PCs_inoviruses.zip: Protein clusters from the references (raw fasta, alignment fasta, hmm profile).
iPFs_inoviruses.zip: Protein families from extended inovirus dataset (raw fasta, alignment fasta, hmm profile).
MobM_C_primer_amplicon.fasta: Multiple sequence alignment of the C primer products with Methanologus MobM genome (NZ_FOUJ01000007) confirming that C primer products span the junction of the excised genome.

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size
No sample size calculation was performed, as the largest collection of publicly available data possible was mined.

Data exclusions
No data were excluded.

Replication
None of the findings was found to be impossible to replicate. This includes PCR amplification of the putative archaeal inovirus provirus, which was repeated either two of three times with similar results (see Supplementary Fig. 11), and the superinfection experiments which were conducted twice and produced similar results.

Randomization
None of the analyses involved allocation of samples to different groups.

Blinding
None of the analyses required blind investigation since the study does not involve a treatment vs control trial (with the exception of "obvious" negative controls such as "no template" PCR).

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