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Occurrence, integrity and functionality of *Aca*ML1—like viruses infecting extreme acidophiles of the *Acidithiobacillus* species complex



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ABSTRACT

General knowledge on the diversity and biology of microbial viruses infecting bacterial hosts from extreme acidic environments lags behind most other econiches. In this study, we analyse the *Aca*ML1 virus occurrence in the taxon, its genetic composition and infective behaviour under standard acidic and SOS-inducing conditions to assess its integrity and functionality. Occurrence analysis in sequenced acidithiobacilli showed that *Aca*ML1-like proviruses are confined to the mesothermophiles *Acidithiobacillus caldus* and *Thermithiobacillus tepidarius*. Among *A. caldus* strains and isolates this provirus had a modest prevalence (30%). Comparative genomic analysis revealed a significant conservation with the *T. tepidarius Aca*ML1-like provirus, excepting the tail genes, and a high conservation of the virus across strains of the *A. caldus* species. Such conservation extends from the modules architecture to the gene level, suggesting that organization and composition of these viruses are preserved for functional reasons. Accordingly, the *Aca*ML1 proviruses were demonstrated to excise from their host genomes under DNA-damaging conditions triggering the SOS-response and to produce DNA-containing VLPs. Despite this fact, under the conditions evaluated (acidic) the VLPs obtained from *A. caldus* ATCC 51756 could not produce productive infections of a candidate sensitive strain (#6) nor trigger it lysis.

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1. Introduction

General knowledge of the diversity and biology of microbial viruses from extreme environments lags behind most other econiches. Since the early 90s viruses infecting acidophilic archaea (*Sulfolobales* and *Thermoplasmales*) have been recovered from enrichment cultures of samples originating in acidic hot springs

Abbreviations: VLPs, viral-like particles; MGEs, mobile genetic elements; ICEs, integrative conjugative elements; RISC, reduced inorganic sulfur compound; PFGE, pulse field gel electrophoresis.

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[1,2] and moderately hot acidic mine drainages [3]. Although the number of known acidophilic archaeal viruses is limited, their morphological diversity is exceptional, comprising spindles, filaments, bottles and turreted icosahedral morphotypes never observed among bacterial viruses [2,4]. The genes they encode are underrepresented in current databases and their proteins products carry rather unusual folds [1,5]. Also, they have utterly unique life cycles, with mechanistic innovations in virion egress [6]. Like their hosts, these viruses show adaptations to the extreme acidity (and high temperature) of the econiches they inhabit, being stable at pH values lower than 3 (and temperatures above 80 °C) [7]. Their uniqueness has raised many relevant mechanistic and evolutionary questions [4] and their properties, as naturally occurring nanoparticles, have promising biotechnologically applications [8].

In contrast, to their archaeal counterparts, very few viruses infecting bacterial acidophiles have been detected or described in

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the literature. These include the ØAcl bacteriophage from *Acidiphilium* sp. ATCC 55305 [9], the ThGEI-J prophage from *Thiomonas* sp. 3As [10], the *Aca*ML1 provirus from *Acidithiobacillus caldus* ATCC 51756 [11] and a putative virus (AMDV1), reconstructed from metagenomic data from an acid mine drainage biofilm, presumably infecting group II and III *Leptospirillum* spp. [12]. Even if virions have been found to occur in cognate samples [9,10,13] these bacterial viruses, featuring temperate life cycles, remain poorly characterized.

All four aforementioned bacterial species have representatives in the biotechnological process of bioleaching. In particular, the acidithiobacilli are ubiquitous in industrial acidic biotopes, with several characterized species that play key roles in metal recovery [14]. Representative members of this group have recently been sequenced [15] and a wide variety of mobile genetic elements [16] have been recognized to populate the genomes of the different species through comparative genomic analyses. Despite this fact no temperate virus other than the one infecting the A. caldus type strain (AcaML1) has been detected so far in this complex taxon [17], and little is known about the influence of this biological agent on the biology of its host and its performance in bioleaching. On the other hand, viruses are useful tools in genetic engineering of cognate hosts and, as such, their description and characterization holds the promise of developing reproducible genetic tools for these biotechnologically important acidophilic model microorganisms.

Herein, we explore the *Aca*ML1 virus occurrence, composition and behaviour under standard and SOS-inducing conditions, to evaluate its integrity and functionality. Given the absence of reproducible genetic modification tools for the acidithiobacilli, identification of a lysogenic bacteriophage capable of infecting and transducing genetic information into members of this genus could represent an important contribution to the field.

2. Materials and methods

2.1. Nucleotide sequence data analysis

Completely sequenced genomes of Acidithiobacillus spp. and Thermithiobacillus tepidarius (Table 1: Supplementary Table 1) were obtained from the NCBI website (http://www.ncbi.nlm.nih.gov). The genome of strain DSM 16786 was obtained form the site: http:// biominingdb.cmm.uchile.cl/genomes/At_ferrooxidans_Wenelen. AcaML1-like proviruses from A. caldus strains were sequenced from PCR amplicons spanning the different AcaML1 modules, using primers detailed in Supplementary Table 2. ORFs were identified using Meta-GeneMark [18] and the predicted protein products derived were functionally annotated using well-established pipelines [11]. AcaML1 (JX507079) CDSs were used as queries to search across the predicted proviral ORFs from A. caldus strains using BlastP [19]. Queries producing no matches were searched by TBlastN [19] against the reconstructed proviral genomes. An E-value of 10(-5) was used as cut-off in both of the alignments. Orthologs in predicted proteomes of each strain were cross-compared using bidirectional BlastP, a reciprocal 95% identity [E,10(-10)] cut-off and in-house BioPerl scripts. Global similarity between queries and ORFs with significant hits was calculated using the Smith-Waterman algorithm implemented in the SSEARCH [20]. Sequence nucleotide identity between proviruses was performed with the Python3 module pyani (https://github.com/widdowquinn/pyani). Gene clusters synteny analysis was performed using genoPlotR package [21].

2.2. Bacterial strains, growth conditions and collection

Bacterial strains used in this study are described in Table 2. *A. caldus* strains were grown in modified 9 K [22] supplemented with 5 gr/L elemental sulfur at 40 °C and pH 3.5 or in mineral salts

Table 1Occurrence of *Aca*ML1 in genomes of *Acidithiobacillus* spp. and *T. tepidarius*.

Species	Strains	Genome ID	G+C%	Size (Gb)	Viral Genes	AcaML1
T. tepidarius	DSM3134	AUIS01	66.8	2,96	34	+/-
A. caldus	ATCC51756	NZ_CP005986	61.4	2,99	72	+
	DX	LZYE01	61.0	3,12	7	_
	ZBY	LZYF01	61.0	3,16	6	_
	ZJ	LZYG01	61.0	3,14	6	_
	MTH-04	LXQG01	61.3	2,87	7	_
	SM-1	NC_015850	60.9	0,32	8	_
	S1	LZYH01	60.9	2,79	9	_
A. ferrooxidans	ATCC23270	NC_011761	58.8	0,30	9	_
	ATCC53993	NC_011206	58.9	2,89	9	_
	DSM 16786	AFWEN	58.4	3,7	11	_
	GGI-221	AEFB01	58.6	3,17	6	_
	Hel18	LQRJ01	58.6	3,11	8	_
	YQH-1	LJBT01	58.6	3,11	2	_
	DLC-5	JNNH01	57.6	4,18	10	_
A. ferriphilus	BY0502	LVXZ01	56.8	2,98	6	_
A. ferrivorans	SS3	NC_015942	56.6	3,21	7	_
	PQ33	LVZL01	56.6	3,30	9	_
	YL15	MASQ01	56.6	3,00	7	_
	CF27	CCCS02	56.4	3,43	12	_
A. thiooxidans	ATCC 19377	AFOH01	53.2	3,02	4	_
	ZBY	LZYI01	53.2	3,79	10	_
	A01	AZMO01	53.1	3,82	11	_
	DMC	LWSB01	53.1	3,85	10	_
	BY-02	LWRZ01	53.1	3,81	10	_
	JYC-17	LWSD01	53.1	3,8	10	_
	A02	LWSA01	53.0	3,72	10	_
	DXS-W	LWRY01	52.9	3,95	12	_
	GD1-3	LWSC01	52.9	3,95	12	_
	Licanantay	JMEB01	52.8	3,9	9	_
	CLST	LGYM01	52.4	3,97	11	_
A. albertensis	DSM14366	MOAD01	52.6	3,50	10	_

Table 2A caldus strains and isolates used in this study.

Strain	Origin	DOI
ATCC 51756	Kingsbury coal spoil U.K.	10.1111/j.1574-6968.1983.tb00426.x
BC13	Birch Coppice Colliery, U.K.	10.1111/j.1574-6968.1983.tb00426.x
MNG	Arsenopyrite pilot plant, South Africa	10.1128/AEM.00864-08
F	Nickel pilot plant, South Africa	10.1128/AEM.00864-08
#6	Fairview mine, South Africa	10.1128/AEM.00864-08
CSH12	Continuous bioreactor, Australia	10.1128/AEM.00864-08
MEL1	Copper bioleaching heap, Chile	10.1016/j.resmic.2014.07.014

medium (MSM) with trace elements [23] supplemented with 5 mM potassium tetrathionate at 40 °C and pH 2.5, under aerobic conditions (200 rpm). Cultures were scaled up to a final volume of 500 ml and brought to late exponential growth phase to obtain sufficient biomass for downstream procedures. Cell and viral concentrates were obtained by tangential flow filtration using Minimate filtering cassettes (Pall) with 1000 kDa or 100 KDa pore-size. The subcellular fraction ('viral fraction') was processed as in Covarrubias et al. [13].

2.3. General DNA techniques

A. caldus stationary phase cultures to be used for nucleic acid purification, and/or for PFGE, were centrifuged at 10,000 rpm for 15 min at 4 $^{\circ}$ C. The harvested cells were stored as cell pellets at -80 $^{\circ}$ C for DNA isolation or were resuspended in 1X MSM for PFGE applications. DNA isolation and routine manipulations were carried out following standard protocols [24].

Polymerase chain reaction (PCR) products were amplified with Dreamtaq (Thermo Fisher Scientific) or the high-fidelity Herculase II fusion DNA polymerase (Agilent Technologies) depending on the experiment. Oligonucleotide primers used in each case are listed in Supplementary Table 2. Each PCR reaction contained 10 ng of template DNA, 0.5 mM of required primers and 0.2 mM of each deoxyribonucleotide in a volume of 25 ml of 1X PCR buffer containing 1.5 mM MgCl2. PCR conditions were as follows: initial denaturing step at 95 °C for 5 min followed by 28–30 amplification cycles (denaturation at 95 °C for 20 s, annealing at the appropriate temperature depending on the specific primer pairs for 20 s and elongation at 72 °C) and a final elongation step at 72 °C for 10 min. Amplicons to be used in nested PCR reactions or as Southern blot probes were purified from the agarose gels with the Mini Elute Gel Extraction Kit (Qiagen) and the Purelink Purification Kit (Thermo Fisher Scientific), respectively. DNA sequencing was carried out at Macrogen Inc.

2.4. Pulsed field gel electrophoresis assays

Cellular suspensions of 10(8) cells/ml were used in the preparation of gel plugs in a 1:1 (v/v) proportion with 1% Pulse Field Certified Agarose (BioRad). Plugs were incubated for 1 h at 50 °C in cell lysis buffer (1X TE; 50 mM Tris, 50 mM EDTA, pH 8.0) with 1% Sarcosyl and 20 mg/ml proteinase K under constant agitation (200 rpm). Plugs were washed 4 times in preheated (50 °C) distilled water and 3 times with preheated 1X TE buffer. The plugs were stored in 1X TE at 4 °C until use. Samples were digested with the 10U/ul XbaI (Thermo Fisher Scientific) in the recommended reaction buffer at 37 °C for 4 h. DNA fragments were separated in a 1% Pulse Field Certified Agarose with 0.5X Tris-Borate EDTA (TBE) running buffer at 14 $^{\circ}$ C. The pulsed field gel electrophoresis was performed in a CHEF-DR III apparatus (BioRad). Orthogonal-field alternation was achieved by varying the pulse time between 1 and 30 s for 13 h, with a rotation angle of 120° and voltage of 6 V/cm. The 5 kb ladder CHEF DNA size standard (BioRad) was used to estimate the molecular weight of the resolved DNA samples. Gels were stained with SYBR Safe DNA gel StainTM (Thermo Fisher Scientific) for 30 min under constant agitation and washed for 10 min in distilled water before being photographed. Images were stored as TIFF files.

2.5. DNA probe labelling and southern blot

An internal fragment of the gp39 gene (0.3 kb in length) was labelled with Biotin-14-dATP using the Thermo Fisher Scientific Nick Translation System according to manufacturer's recommendations. PFGE-resolved DNA was transferred onto positively charged nylon membranes Hybond-XL (General Electric) by capillary neutral transfer using the ULTRAhyb hybridization buffer at ambient temperature for 2 h. Transferred DNA was immobilized by UV cross-linking in a UV oven (BioLink) at 1134 J/m2. The membranes were placed into glass hybridization bottles and prehybridized in ULTRAhyb buffer (Thermo Fisher Scientific) at 42 °C for 30 min in a hybridization oven. The denaturated (90 °C for 10 min). labelled DNA probe was added to the prehybridization solution and incubated for 16 h at 42 °C under slow-speed rotation. The membranes were washed at 42 °C, twice with the high-stringency buffer from the NorthernMax Kit (Thermo Fisher Scientific) for 5 min and twice with low-stringency buffer from the same Kit for 15 min. The washed membranes were incubated with Streptavidine (Biolegend) diluted 1:5000 (v/v) in high-stringency buffer at 42 °C for 1 h under constant shaking. To remove Streptavidine excess, a final wash with this buffer was carried on at room temperature for 5 min. The membrane was revealed using the Pierce ECL western blotting substrate (Thermo Fisher Scientific) and conventional autoradiographic methods.

2.6. DNA damaging treatments to assess proviral excision

To produce DNA damage, and induce proviral excision, 10(10) total cells concentrated 100 fold were UV irradiated (200 J/m2, 254 nm for 3 min) or incubated with iron (III) sulfate (200 mM) for 45 min or mitomycin C (1 µg/ml) for 8 h). Induced cells were recovered for one or two generation times (8-16 h, respectively) in fresh 1X MSM media (with potassium tetrathionate) at 40 °C and 200 rpm. All experiments were performed in triplicate. After recovery, cells were harvested for excision analysis using either Nested PCR or PFGE combined with Southern blot as described above.

2.7. Infection assays to assess host lysis and virus like particles production

A. caldus ATCC 51756 cultures were processed as in 2.6. to induce the AcaML1 provirus lytic cycle. Cells were centrifuged to recuperate the spent supernatants and the supernatants processed as in section 2.2 to obtain the virus-enriched concentrates. Equivalently treated strain #6 cultures (virus negative) and untreated A. caldus ATCC 51756 supernatant concentrates served as controls. Each

supernatant concentrate (100X) was added to an exponentially growing culture of *A. caldus* strain #6 (phage sensitive) 24 h after inoculation [10(5) cell/ml]. Cell counts were recovered every 20 h and plotted against time. The Live/Dead BacLight kit (Thermo Fisher Scientific) was used to assess the relative numbers of bacteria with intact plasma membranes from those with compromised membranes.

Virus enriched supernatants were filtered through a 0.022 μ m Anopore Inorganic Membrane (Anodisc) and observed by epifluorescence microscopy after staining with SYBR Green I (Thermo Fisher Scientific), according to established procedures [24]. In selected samples the viral like particles (VLPs) were precipitated by the addition of NaCl (1 M) and polyethylene glycol 8000 (10%) [25], and analysed by electron microscopy using a Tecnai Spirit operated at 120 kV and a 2 K \times 2 K CCD camera (Magnification 48,500 \times -4,95 Å/pixel). Specimens were prepared for EM using the conventional negative staining procedure with 2% uranylacetate.

3. Results and discussion

3.1. Search for AcaML1-like viruses in sequenced acidithiobacilli

To investigate whether AcaML1-like lysogenic proviruses were present in other strains of the species, their occurrence was evaluated in all publically available genome sequences of the taxon (Table 1; Supplementary Table 1). Thermithiobacillus tepidarius was also included in the analysis as the closest ancestor sequenced. BLASTp and HMM searches failed to reveal the presence of viral proteins related to AcaML1 or other known bacteriophages in the genomes of the Acidithiobacillus spp. strains analysed. A few viral proteins distantly related to the AcaML1 prophage, encoding integrases, excisionases and phage-type regulators were however widespread in all genomes investigated (Supplementary Table 1). These proteins typically conform viral gene modules of larger mobile genetic elements of the ICE type (Integrative Conjugative Elements), where they play a role in the control and regulation of integration and excision [26,27]. A DNA segment potentially encoding a full provirus was predicted in the genome of T. tepidarius. At the protein level, this provirus presents similarities with the A. caldus AcaML1 provirus, with equivalent gene products related to the infectious cycle of the virus (excisionase, recombinases), lysis of the host (lysozyme), and a structural gene (capsid protein), but it lacks orthologous genes related to the

morphogenesis of the base plate and tail (Supplementary Table 1). According to available genomic evidence, AcaML1-like proviruses are confined to the mesothermophilic acidithiobacilli, T. tepidarius and A. caldus.

3.2. Prevalence of AcaML1-like proviruses in A. caldus strains and industrial isolates

To investigate whether lysogenic proviruses like AcaML1 were present in other strains of A. caldus, occurrence of AcaML1 signatures was evaluated in a collection of seven A. caldus isolates (Table 2) using pulsed-field gel electrophoresis (PFGE) combined with Southern blot. Hybridization of a biotinilated probe, directed to the viral gene gp39 (encoding the viral capsid), to the PFGE fingerprints of all seven isolates (Fig. 1A) revealed the presence of lysogenic proviruses similar to AcaML1 in four of the strains tested, namely the type strain (ATCC 51756), strain F, BC-3 and CSH12 (Fig. 1B). Under the conditions evaluated a single band was identified in the four positive lanes, implying that only the integrated form of the virus could be detected in the type strain and the other AcaML1 positive isolates. PFGE-based size estimations indicate that all these proviruses are similar in size to AcaML1 (aprox. 60 kb). This result was confirmed by PCR amplification of selected gene markers encompassing the different gene modules of the AcaML1 provirus genome. All 19 gene markers tested could be amplified from total DNA preparations obtained from strains BC13, F and CSH-12 indicating, as well, global conservation of the gene content of the AcaML1-like proviruses present in A. caldus strains (Fig. 2).

3.3. Comparative genomics of AcaML1-like proviruses from A. caldus strains

AcaML1 proviruses present in A. caldus strains F, BC-13 and CSH-12 were PCR amplified using high-fidelity and high-processivity Herculase II fusion DNA polymerase and sequenced at Macrogen on a module-to-module basis using primers indicated in Supplementary Table 2. Sequences obtained (MH142219, MH142220) were aligned against the AcaML1 genome (JX507079) and cross-compared to each other. The putative provirus identified in the whole genome sequence of the neutrophilic relative of A. caldus, T. tepidarius, was included in this analysis for comparative reasons. Results of the whole genome multiple alignments are shown in Fig. 3.

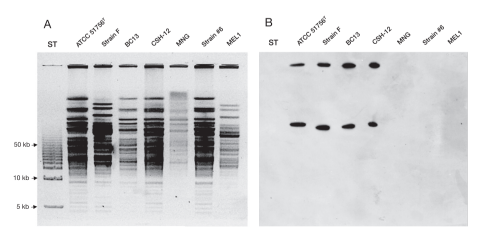


Fig. 1. Prevalence of AcaML1-like prophages in A. caldus strains. **A.** PFGE profiles of Xbal digested genomic DNA of A. caldus strains. Staining was done with SYBR Green I. The molecular weight standard used is the 5 kb ladder DNA size standard (BioRad). **B.** Southern blot of the gel in panel A using a 300 bp long probe specific for the gene gp39 encoding the viral capsid.

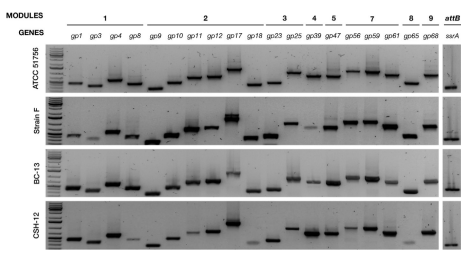


Fig. 2. Occurrence of *Aca*ML1 genes in *A. caldus* strains. Different predicted viral genes spanning gene modules 1 to 9 were amplified by end point PCR using primers indicated in Supplementary Table 1. The housekeeping gene *ssrA*, predicted integration site for the provirus (*attP* site), was used as PCR positive control. The molecular weight standard used is the Thermo Scientific GeneRuler 1 kb DNA Ladder.

The genomic divergence between AcaML1-like proviruses and the predicted provirus integrated in the genome of *T. tepidarius* (26.1%) is slightly lower than the genomic divergence existing between the two species (31.1%, value calculated for both type strains), being thus coherent with the phylogenetic distance between hosts. Average nucleotide identity between the sequenced A. caldus proviruses is higher than 99.9%, highlighting the conservation of the provirus at DNA sequence level between acidophilic strains. Such conservation is puzzling, considering the geographical origin of the strains (Table 2). Recent evidence from studies on the aerosolization of sand-dust and organic aggregates in sea sprays, showing high fluxes and high deposition rates for viruses across the free atmosphere [28], could provide an explanation for this observation. However, in the absence of additional information on the genetic background of the host strains and/or genomic data of the A. caldus populations present at each site it is unfeasible to draw further conclusions on this respect.

The genomes of the *A. caldus* lysogenic proviruses had an average length of 59.4 kb, including the redundant ends and encoded 72 ± 2 protein coding ORFs. The *T. tepidarius* provirus was predicted to be significantly larger (84.6 kb) with 94 predicted protein coding genes. In both cases a ~10-bp direct repeat flanking the proviruses was identified (DR_{Aca}: CCACCATTTA; DR_{Ttp}:

CCATCGACACG). The GC content of the AcaML1-like proviruses averaged 65.5%, being higher than those of the host genomes (61.72%; [26]). A lower G + C content of phage genomes in comparison to those of their hosts is a widespread phenomenon, for example found for $Escherichia\ coli\$ phages T4 and JS98 [29]. In contrast, increased G + C content has seldom been reported (e.g. [30]).

Predicted orfs cover 96.7% of the sequence and are located on different strands. One group of genes points leftward (32 genes) while the rest of the prophage (40 genes) is oriented rigthward with respect to the predicted origin of replication of the host genome. The "lefthand" side of the genome groups candidate early genes, while the ORFs on the "righthand" side are predicted to be intermediate and late transcription genes. Results from both BlastP searches and curation using phage-dedicated resources (see Materials and Methods) indicated that 59.4%, of the predicted proteins in *Aca*ML1 have significant sequence similarity to proteins of known function or unknown function but of viral origin, including nucleotide metabolism, DNA replication/recombination, components of viral particles and others. The characteristics of these ORFs and their corresponding predicted proteins are described in Supplementary Table 1. Genes with functional assignments were named accordingly. Gene function assignment and

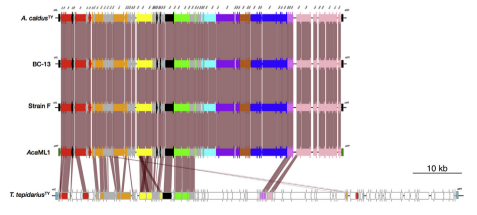


Fig. 3. Whole genome multiple alignment of *Aca*ML1-like proviruses present in the *Acidithiobacilaceae*. Reconstructed proviral genomes were aligned against the *Aca*ML1 genome (JX507079) and the *T. tepidarius* predicted provirus (conting AUIS01000006, coordenates 2854 to 86496). Similarity between predicted proteins was calculated using BlastP (Cut-off values: E–05; >30% identity) and displayed graphically with genoPlotR.

architecture analysis allowed a tentative subdivision of the *Aca*ML1 provirus genome into modules as follows: replication/regulation, packaging, structure/morphogenesis, lysis and lysogeny (Fig. 3). This modular organization is typical for temperate phages from *Myoviridae* family [31].

3.4. Gene products from the leftward region

Divergently oriented gene clusters 1 and 2 encode signature proteins implicated in lysogeny establishment and regulation and control of the lysogeny-lytic switch. Gene module/cluster 1 contains 8 ORFs (gp1-gp8) and includes genes coding for a putative transcriptional regulator (Gp1), the small (Gp2) and large (Gp3) subunits of a site-specific recombinase (pfam00239), similar to the integrases of P2 temperate phages [32] that catalyze unidirectional integration of the prophage into the host genome and an excisionase (pfam06806), necessary to enable excisive recombination by the phage integrase [33]. Genes gp4-gp7 have no predicted function. Divergently oriented to gp8 is another ORF (gp9) which encodes a putative transcriptional regulator, baring the HTH motif of MerR family regulators. This gene is the best candidate for the cllike phage repressor and forms part of the 2nd cluster/module of AcaML1 together with other 15 genes (gp9 - gp26) with ascribed functions to phage replication and regulation. The deduced amino acid sequence of ORF gp11 bares significant similarity to a phage antirepressor protein (COG3617) with a conserved N-terminal BRO domain (pfam02498). Presumably, Gp11 is implicated in the maintenance of lysogeny by modulating the activity of the repressor [34], although other roles in replication and/or transcription have also been proposed for proteins carrying BRO domains [35]. Open reading frame gp17 encodes for a phage primase (P4 family), which is predicted to synthesize the RNA primer at the origin of replication required for replication initiation and gp18 an RNAse H endonuclease predicted to degrade the RNA primer during replication. Gp23 contains a CRO domain (pfam09048); members of this family of proteins typically act as early gene expression repressors and are required to reach the lytic growth stage [36]. Two orfs gp24 and gp25, encoding two highly similar methyltransferases, are probably related to DNA modification associated to phage replication.

3.5. Gene products from the rightward region

The right part of the genome represents the "late region" and is made up of gene products playing a role in viral particle formation and assembly, DNA packaging and host lysis for viral particles release. The first gene module in this region of the genome (gp34 – gp42) encodes several functions directly related with the formation of the procapsid shell and the genome packaging of complex double-stranded DNA viruses into the viral head. Proteins Gp34 and Gp35 were identified as the small (GpNu) and the large subunits (GpA) of a terminase (pfam05876), respectively. The amino acid sequence which exhibits weak similarity to terminases of other phages, contains two partially conserved domains: one in the Nterminal half for a homing endonuclease and one in the C-terminal part for a terminase [37]. The terminase hetero-multimeric complex is responsible for packaging DNA in lamboid like phages such as P2, through recognition, cleavage and translocation of DNA concatemers into de viral head. Immediately adjacent to the terminase is gp36 which encodes for the phage protein GpW (pfam02831), proven necessary for the stabilization of DNA within the phage head and for attachment of tails onto the head during morphogenesis [38]. As in other known prophages the gene encoding the portal protein (Gp37, pfam05136), the serine protease responsible for procapsid maturation (Gp38, pfam01343, [39]), the head decoration protein D (Gp39, pfam02924, [40]) and the major capsid protein E (Gp40, pfam03864) follow the terminase. Although portal proteins are not globally conserved, Gp37 is highly similar to the portal protein of Comamonas aquatica (91% similarity), an aquatic Burkholderiales bacterium. In other phages the portal protein serves as the entry and exit door for phage DNA during genome packaging and ejection, as a heads-tail connector and possibly as a nucleator for capsid assembly [40]. The major capsid protein belongs to the E superfamily, typically found in lamboid like phages like P2 [41]. P2 forms a 60 nm icosahedral capsid, with T = 7 symmetry, from 415 copies of capsid protein derived from gene product N (gpN) to package its 33.5 kb DNA [42]. A number of experimental studies have suggested that virion sizes are a function of genome sizes and that an allometric relationship between the genome length and the volumen of the virion has been recognized [43]. Since AcaML1s genome is 26.1 kb longer than P2. AcaML1 virion size is predicted to be somewhat larger.

Blast analysis of the predicted phage products revealed several proteins as being involved in phage tail morphogenesis, recognition of the host and genome delivery upon infection. Four gene products of AcaML1 could be associated to the baseplate formation including orthologs for the genes encoding the spikes of the tails end (Gp43: protein V pfam04717, [44]), the wedge protein (Gp45: protein W, pfam04965, [45]), the receptor-binding protein (Gp46: protein J, pfam04865, [46]) and assembly protein I (Gp47: protein I, pfam09684) of P2-type phages. Gp45 has a conserved lysozyme domain, similar to that of T4-like phages protein GP5, predicted to aid in penetration of the peptidoglycan layer during the initial infection process [47]. ORFs gp48-gp53 encode predicted tail fibers sharing some degree of similarity with P2 phages important for adsorption of phage to the outer membrane of the bacterial cell [48]. Gp49 is a phage-associated protein with predicted endosialidase activity. This enzymatic activity has been found in the tail spikes of certain phages where it catalyzes the degradation of the bacterial capsule [49]. Gp53 shares distant similarity in its N-terminal domain with proteins of the lectin C superfamily. Lectin-like domains have been found associated to receptor binding proteins in bacteriophages where they confer binding variability [50]. These enzymes may be tail components involved in adsorption of the phage or/and DNA injection. Next to the tail fibers encoding genes are two genes, gp56 and gp57, encoding the major tail sheath FI (pfam04984) and the tail tube protein FII (pfam04985) of Myoviridae bacteriophages [51], respectively. Gp58 encodes a small tail protein (protein E, pfam06528) that stabilizes the tail in P2 bacteriophages [52]. Distal to this cluster is gene gp59, encoding the protein that determines tail size by working as a template for measuring tail length during assembly [53] better known as tape measure protein (pfam10145). It consists of 1026 residues and is one of the longest open reading frames of the AcaML1 genome. The protein has a well-conserved core region that is often found in phage tail tape measure proteins. Assuming 1.5 Å per amino acid residue [53], the calculated length of the AcaML1 tail would be 154 nm. The function of the distal genes in this cluster *gp60-gp62* is less clear. Gp60 corresponds to protein U (pfam06995) of P2-like phages and is likely to play a role in tail assembly [51]. Gp61 is similar to protein X of P2 (pfam05489) and bares a LysM domain (pfam01476) with predicted peptidoglycan degrading activity [54]. Gp62 shows similarity with protein D (pfam05954), another phage associated protein with unclear function.

The contiguous cluster of four genes encodes proteins involved in programmed lysis. As other dsDNA phages, *Aca*ML1 seems to make use a dual lysis system consisting of a holin and an endolysin [55,56]. Protein Gp63 encodes a low complexity, highly conserved, hydrophilic 94 amino acids long protein with a single transmembrane region (amino acid position 66–85) that places the N-

teminus in the cytoplasm and the C-terminal end of the protein in the periplasm, resembling other known class III holins [57]. Holins facilitate the release of phage particles from infected bacteria during viral-induced cell lysis [58]. A second small ORF (gp64), downstream of the putative holin gene, encodes a 158 aa long protein of unknown function. Orthologs of this gene frequently concur with gp63 orthologs and the predicted protein products are extremely well conserved (e.g. in *T. tepidarius*). Predicted protein Gp65 is 160 amino acids long and contains the signature of known phage endolysins of the muramidase type (pfam00959) and lack predicted secretory signals. Muramidases cleave the glycosidic beta 1,4-bonds between the N-acetylmuramic acid and the N-acetylglucosamine of the peptidoglycan from outside the cytoplasm, thereby enabling progeny virions to be liberated [59]. Because of their unique ability to cleave peptidoglycan in a generally speciesspecific manner (shared homology is not found within the catalytic domain), endolysins provide a means of selective and rapid killing of bacteria with no effect on the rest of the native microflora. Access to peptidoglycan depends on the small hydrophobic holins, which enable endolysin molecules to cross the inner membrane. These properties confer a biotechnological potential to both types of proteins.

The distal gene module in *Aca*ML1 contains three genes (*gp66* – *gp69*) encoding a DNA restriction-modification system and two insertion sequences. Gp66 (pfam00145) is a predicted site-specific

DNA methylase that modifies cytosine to 5-methylcytosine. In turn, Gp67 and Gp68 encode the two subunits of a predicted McrBC (pfam10117) restriction enzyme cutting DNA in multiple positions within two distant locations (between 40 pb and 3 Kb apart) baring the 5-methylcytosine modification preceded by a purine (RmC[N] RmC) [60]. Presence of RM-systems in phages and other mobile genetic elements is frequent, playing a relevant role in the stabilization of the mobile element within the genome [61] or being target of frequent horizontal transfer between bacteria [62]. Similar to other cases appointed in the literature (e.g. [63]), the RM-system of AcaML1 is flanked by transposases of the IS5 and IS21 families (gp70-74) (pfam05598 and pfam13751). These genes could facilitate transfer of the RM gene cassettes between the host chromosome and horizontal transfer vector such as phages [62].

3.6. Induction of replication intermediaries of AcaML1-like proviruses by DNA damage

Based on the above prevalence and genomic analyses, attempts to establish the capacity of the proviruses to produce functional viral progeny were made in all four AcaML1 positive (A. caldus ATCC 51756, BC13, CSH12, F). As initial strategy mitomycin C, a DNA damaging agent that produces DNA-adducts and lesions that trigger the SOS response, was used to induce proviral excision. Standard mitomycin C concentrations ($1 \mu g/ml$) for this type of

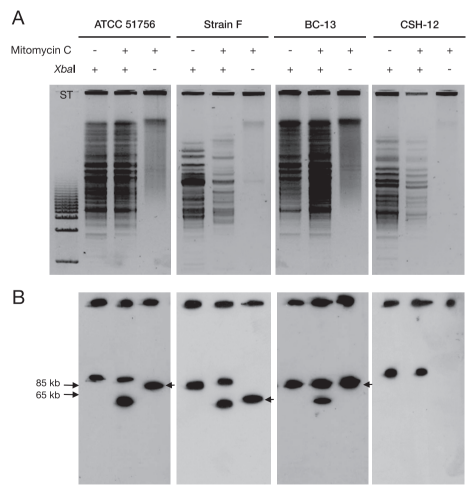


Fig. 4. Excision of *Aca*ML1-like proviruses from the chromosome of *A. caldus* strains upon mitomycin C treatment. **A.** Pulsed-field gel electrophoresis profiles of *Xba*l digested genomic DNA of *A. caldus* strains. Staining was done with SYBR Green I. Molecular weight standard used is the 5 kb ladder DNA size standard (BioRad). **B.** Southern blot of the gel in the left panel using a 300 bp long probe specific for the gene *gp39*, encoding the viral capsid. Excised forms of *Aca*ML1 are indicated with arrows in each panel.

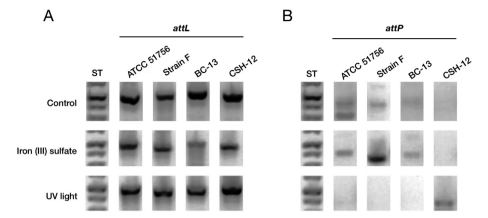


Fig. 5. Iron (III) sulfate and UV exposure as elicitors of excision of AcaML1-like proviruses from the chromosome of A. caldus strains. The chromosomal junction formed to the left of the integrated provirus (attL) and the viral attachment site (attP) were amplified by nested PCR using primers indicated in Supplementary Table 2. DNA damage was induced with iron (III) sulfate (200 mM for 45 min) or UV light exposure (200 J/m², 254 nm for 3 min). The molecular weight standard used is the Thermo Scientific GeneRuler 1 kb DNA Ladder.

assays were used [64]. Excision was evaluated through PFGE and Southern blot (as in 3.2) using both *Xba*I digested and undigested DNA (Fig. 4). The results obtained indicate that the proviruses present in the chromosome of *A. caldus* ATCC 51756, strain F and BC13 respond to the induction trigger and produce an episomal form of the virus ranging in size between 65 and 85 Kb. No episomal form of *Aca*ML1 could be identified in strain CSH-12, despite of the fact that the integrated form was confirmed in both mitomycin C treated and untreated cells.

Other DNA damaging agents, iron (III) sulfate and UV exposure, were also tested as inducers of provirus excision. Capacity of AcaML1 to excise out of the respective host genomes upon treatment with these DNA damage elicitors was assayed by nested PCR. Targeted regions (shown in Supplementary Figure 1) included the chromosomal junctions formed at both sides of the integrated provirus (attL and attR) and the viral and bacteria chromosomal attachment sites (attP and attB sites, respectively). Representative results are shown in Fig. 5 and full results are summarized in Table 3. Positive amplicons of the attL and attP products were obtained in 200 mM ferric iron treated cell of the type strain, strain F and BC-13. In turn, UV light at the exposure dose tested (200 J/m2, 254 nm for 3 min) only elicited excision in the CSH-12 strain (Fig. 5B; Table 3). These results demonstrate that all four A. caldus strains evaluated carry proviruses capable of producing the episomal replication intermediary upon excisive induction.

3.7. Infection assays using AcaML1-like proviruses from A. caldus strains

To evaluate if the induced viruses prompted host lysis and produced viral particles, infections assays were set up. No lawn-based plate assay is available for *A. caldus*, thus lytic capacity of *AcaML1* could not be tested using conventional plaque formation assays. To assess host killing, the effect of Viral-Like Particles (VLPs) on growth decline (with cell lysis) and/or loss of viability (without cell lysis) was evaluated using conventional liquid cultures and modified 9 K media supplemented with elemental sulfur as energy source. *A. caldus* ATCC 51756 was used as source of *AcaML1* VLPs. In order to avoid the possibility of superinfection related immunity issues, *A. caldus* strain #6 (which proved negative to the presence of lysogenic *AcaML1* in 3.2) was chosen as virus sensitive strain for the infection assays. Production of VLPs was triggered by exposing *A. caldus* ATCC 51756 mid log cultures to ferric iron for 8 h (as in 2.6). VLPs were recovered from

TFF-concentrated culture supernatants, treated with NaCl (1 M) and polyethylene glycol 8000 (10%) to precipitate the virus, and visualized using epifluorescence microscopy after SYBR-Green staining (Fig. 6B). Mid log cultures of strain #6, grown for 24 hs (I) in fresh 9 K media (F9K) were challenged with the *Aca*ML1 VLPs concentrates or with spent media concentrates of control cultures (ferric iron-treated *A. caldus* strain #6). To uncover variations in growth and/or viability with respect to untreated and control samples, lysis of the sensitive strain was assessed by direct cell counts and phase contrast microscopy, and by Live-Dead staining and epifluorescence microscopy.

Spent culture supernatant from ferric iron-treated A. caldus ATCC 51756 cultures had very little effect on the cell counts of fresh cultures of strain #6 with respect to the controls at all times tested (Fig. 6A). Similarly, only minor effects were observed on the viability (V) of strain #6 (Fig. 6C; Supplementary Figure 2), with 38% of cell death at 48 h after treatment with virus-positive spent media concentrates (SC9KTY) and 30% death in similarly treated the cells with virus-negative spent media concentrates (SC9K6). Supporting statistics for these tests are provided in Supplementary Table 3. These results suggest that AcaML1 preparations had no or very little lytic effect of the chosen sensitive strains. Similar results were obtained with other virus-negative A. caldus strains (MNG, MEL1, data not shown). Other forms of resistance to AcaML1, other than superinfection, cannot presently be discarded. Seeking for an explanation to these results, electron microscopy was used to inspect the AcaML1 concentrates. In all preparations evaluated icosahedral tail-less virus-like particles were observed (Fig. 6C). These results explain the neglectable levels of lysis observed in the

Table 3PCR-based detection of the episomal *Aca*ML1 form in *A. caldus* strains upon treatment with DNA damage elicitors.

attL-attR/attB-attP Strain	Control	Mitomycin C	Iron(III) sulfate	UV light
ATCC 51756	+/*	+/*	+/+	+/-
F	+/*	+/*	+/+	+/-
BC-13	+/-	-/+	+/+	+/-
CHS-12	+/-	+/*	++/-	+/+

a. Mitomycin C (1 μ g/ml; 8 h).

b. Iron (III) sulfate (200 mM; 45 min).

c. UV (200 j/m²; 3 min).

⁽⁺⁾ attR and attL present; (-) attR and attL absent; (*) one att site of the pair absent.

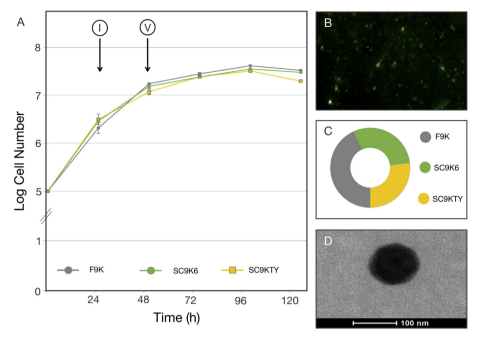


Fig. 6. AcaML1 infection assays using A. caldus sensitive strain #6. **A.** Growth curve of strain #6 cultures challenged with spent media concentrates of A. caldus ATCC 51756 (SC9KTY) and #6 (SC9K6) ferric iron-treated cultures or with spent media concentrates of strain #6 untreated cultures (F9K). Infection (I) occurred 24 h after set up of the experiment. **B.** Fluorescence microscopy of SYBR Green I stained A. caldus ATCC 51756 spent media concentrates after iron (III) sulfate induction showing fluorescent viral-like particles (VLPs). **C.** Viability counts obtained 48 h after onset of the assay (V) using Live-Dead staining of treated (SC9KTY), untreated (F9K) and control (SC9K6) samples. **D.** Electron microscopy of VLP recovered form AcaML1-enriched concentrates.

infection assays and cast doubt on the capacity of the *Aca*ML1 provirus to assemble and maintain functionally stable viral particles under the acidic conditions tested. A definite answer to this question will require additional research in the near future.

4. Conclusions

Mesothermophilic Acidithiobacilliaceae family members, A. caldus and T. tepidarius, host in their genomes lysogenic proviruses with redundant ends, ranging in size between 59 and 85 kb. Comparative genomic analyses performed herein revealed partial conservation of the gene content and gene modules architecture between A. caldus and T. tepidarius proviruses, suggesting that their organization and composition is preserved for functional reasons. Among A. caldus strains the integrated proviruses had only modest prevalence (30%), but were highly conserved at both the amino acidic and DNA sequence levels. In all four AcaML1-provirus positive strains (ATCC 51756, BC13, F and CSH12) the proviral genome is organized in a leftward region that encodes signature proteins implicated in lysogeny establishment and regulation and control of the lysogeny-lytic switch, and a rightward region encoding genes predicted to play a role in viral particle formation and assembly, DNA packaging and host lysis for viral particles release.

Induction assays, performed under DNA-damaging conditions triggering the SOS-response, demonstrated that the *Aca*ML1-positive strains carry proviruses capable of producing the episomal replication intermediary upon excision, and that DNA-containing VLPs are produced. Despite this fact, under the conditions evaluated (acidic) the VLPs obtained from *A. caldus* ATCC 51756 could not produce productive infections of a candidate sensitive strain (#6) nor trigger it lysis. The neglectable levels of lysis observed in the infection assays correlate to the presence of icosahedral tail-less virus-like particles in all preparations evaluated by transmission electron microscopy. This aspect casts doubt on the capacity of the *Aca*ML1 provirus to assemble and maintain

functionally stable viral particles under the conditions tested, which emulated the acidity of their usual habitats. Further studies are required to define if the lack of tails is an experimental artefact of the current experimental design or an adaptive biological response.

Conflicts of interest

None declared.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.resmic.2018.07.005.

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