Insertion sequence AS5 (ISAS5) is involved in the genomic plasticity of Aeromonas salmonicida

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The genome of the fish pathogen Aeromonas salmonicida subsp salmonicida harbors a large number of insertion sequences (ISs), many of which are located on plasmids. In the present study, we analyzed the small plasmid profile of A. salmonicida strains to identify evidences of plasmid alterations. Ten out of 78 strains analyzed displayed an unconventional plasmid profile. However the HER1104 strain was unique, having a positive PCR signal for pAsa1 plasmid despite not carrying this plasmid. Instead, HER1104 was bearing a plasmid at higher molecular weight than pAsa1. We characterized this new larger plasmid, which we called pAsa1B since it is a derivative of pAsa1 containing one more complete IS (ISAS5) than the parental plasmid. An additional 96 bp relic of ISAS5 was also present in pAsa1B. These results prove that ISAS5 is another active mobile genetic element in A. salmonicida subsp salmonicida and provided further proof of the genomic plasticity of this bacterium.

Introduction

Aeromonas salmonicida subsp salmonicida (A. salmonicida) is the Gram-negative bacterium responsible for furunculosis, a disease that mainly affects fish in the salmonid family.1,2

A. salmonicida strains usually contain many plasmids. Three small high-copy-number plasmids [pAsa1 (5.4 kb), pAsa2 (5.2 kb) and pAsa3 (5.6 kb)] have been found in the vast majority of the A. salmonicida strains that have been analyzed.3,4 These cryptic plasmids only bear genes for their own replication, mobilization and stability. pAsa1, another high-copy-number plasmid, is also commonly found in A. salmonicida strains.4,5 While this 6.4 kb plasmid is very similar to pAsa3, it also possesses the aopP gene that encodes a type three secretion system (TTSS) effector.6 The aopP gene is specific to pAsa1. This plasmid also possesses one copy of ISAS11, an insertion sequence (IS).7 These four plasmids represent the conventional set of plasmids seen in A. salmonicida strains and can easily be analyzed by EcoRI digestion followed by electrophoresis.4

A 155 kb low-copy-number plasmid (pAsa5) is also present in most A. salmonicida strains. It carries the vast majority of TTSS genes.8,9 Other plasmids such as pASOT, pASOT2, pASOT3, pRAS1, PRAS3.1, pRAS3.2, pAR-32, pAsa4 and pAsa6, which range in size from 11 to 167 kb, have also been identified in sub-groups of A. salmonicida strains.8,10 Most of these occasional plasmids carry antibiotic resistance genes.

The genomic plasticity of A. salmonicida is likely IS dependent given the involvement of ISs in gene inactivation and plasmid rearrangements7 and the fact that the A. salmonicida genome contains a large number of ISs.7 The tapA gene, which encodes the A protein involved in the A-layer at the surface of the bacterium, can be disrupted by the spontaneous transposition of ISASI and ISASI2.16,17 ISASI can also be inserted in abcA, a gene involved in O-polysaccharide synthesis and transport.18 Recently, it has been shown that three ISASI1 of pAsa5 trigger the rearrangement of this plasmid, resulting in the excision of the TTSS locus (loss profile 1) or the excision of this locus with an additional region of the plasmid (loss profile 2).7,8 These rearrangements of pAsa5 provoke the loss of virulence by A. salmonicida.8 More complex rearrangement profiles of pAsa5 has been also observed for some A. salmonicida strains, suggesting that ISs other than ISASI1 may be active in provoking multiple, sequential rearrangements of pAsa5.5,15

In the present study, we wanted to bring additional evidences supporting the hypothesis that the high content of mobile genetic elements in the genetic material of A. salmonicida might contribute to frequent rearrangements and rapid evolution of the bacterium’s genome. Therefore, we analyzed the small plasmids in A. salmonicida strains using EcoRI digestion and sequence analysis.

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Table 1. Small plasmid profiles for the A. salmonicida strains displaying an unconventional plasmid profile following EcoRI digestion

| Name       | Origin* | Source and/or reference† | Plasmid profile‡ | aopP by PCR¶ | Tetracycline resistance¶ |
|------------|---------|--------------------------|-------------------|--------------|--------------------------|
| 01-B526    | Brook trout (QC, Canada) | FMVUM19 | Cryptic + pAsal1 | P            | S                        |
| HER1110    | *(Japan) | FhRC20 | pAsa3 only | N            | S                        |
| HER1104    | *(France) | FhRC | Cryptic + 8.5 kb | P            | S                        |
| HER1085    | *       | FhRC | Cryptic + pAsal1 + 14 kb | P            | S                        |
| HER1084    | *(France) | FhRC20 | Cryptic + 7.6 kb | N            | S                        |
| A449       | Brown trout (France) | 21 | Cryptic | N            | R                        |
| 2009–157K5 | Brook trout (NB, Canada) | FOC | Cryptic + pAsal1 + 11 kb | P            | R                        |
| 2010–47K18 | Brook trout (NB, Canada) | FOC | Cryptic + pAsal1 + 11 kb | P            | R                        |
| 2009–195K29| Brook trout (NB, Canada) | FOC | Cryptic + pAsal1 + 11 kb | P            | R                        |
| 2009–144K3 | Brook trout (NB, Canada) | FOC | Cryptic + pAsal1 + 11 kb | P            | R                        |
| 2004–208   | *       | FOC | pAsa1 + pAsa2 + pAsal1 +3.0 kb + 7.6 kb | P            | S                        |

The 01-B526 strain is shown as an example of strain displaying the conventional profile. *Further information not known or not traceable. †FMVUM: Laboratoire de bactériologie clinique, Faculté de médecine vétérinaire, Université de Montréal (Montreal, QC, Canada); FhRC: Félix d’Hérelle Reference Center, Département de biochimie, de microbiologie et de bio-informatique, Université Laval (Quebec City, QC, Canada); MAPAQ: Ministère de l’agriculture, des Pêcheries et de l’alimentation du Québec; FOC: Aquatic Animal Health Department, Fisheries and Oceans Canada (NB, Canada); NB: New Brunswick; NS: Nova Scotia; QC: Quebec. Cryptic means that the bands for the pAsa1, pAsa2 and pAsal1 plasmids were visible on the gel. The sizes of bands other than pAsa1, pAsa2, pAsa3, or pAsal1 are indicated in kb. P: positive PCR; N: negative PCR. R: resistant; S: sensitive.

salmonicida strains in order to identify unusual plasmid profiles that might provide evidence for the genomic plasticity of this bacterial species. This work led to the characterization of a new plasmid showing that ISAS5 is another active mobile genetic element in A. salmonicida.

Results

The small plasmid profile of 78 A. salmonicida strains was analyzed in order to identify those displaying unconventional plasmid profiles. After EcoRI digestion, the electrophoretic patterns of plasmids under 50 kb in size revealed that 68 of the 78 strains had bands corresponding to pAsa1, pAsa2, pAsa3 and pAsal1 (data not shown). This is considered the common profile for small plasmids in A. salmonicida. Moreover, all the strains with the common set of small plasmids also tested positive by PCR for the aopP gene, which is on the pAsal1 plasmid (data not shown).

On the other hand, 10 strains displayed unusual plasmid profiles (i.e., additional bands other than those of the common profile were present while others were absent) (Table 1). Five strains (HER1085, 2010–47K18, 2009–157K5, 2009–195K29 and 2009–144K3) displayed the bands for the complete set of common small plasmids but also had an additional band either at 14 or 11 kb. Two strains (HER1110 and A449) exhibited a smaller set of plasmids than expected. In the case of HER1110, only the pAsa3 plasmid was present, while A449 was missing the pAsal1 plasmid as previously reported. The band corresponding to pAsal1 was also missing in HER1084 but this strain had an additional band at 7.6 kb. The 2004–208 strain did not harbor the pAsa3 plasmid, but its plasmid profile included two additional bands at 3.0 and 7.6 kb. Lastly, HER1104 contained pAsa1, pAsa2, and pAsa3 but not pAsal1. However, HER1104 had a fourth band corresponding to a plasmid about 8.5 kb (Fig. 1).

Four strains (2010–47K18, 2009–157K5, 2009–195K29 and 2009–144K3) from New Brunswick (Canada) displayed the same profile, with an additional band at about 11 kb detected following the EcoRI digestion (Table 1). These four strains were resistant to tetracycline (Table 1). The pRAS3 plasmid (11.8 kb) is known to bear a TetC gene that confers resistance to tetracycline. We confirmed by PCR analyses that 2010–47K18, 2009–157K5, 2009–195K29 and 2009–144K3 bear this plasmid (data not shown), which is likely the one detected at 11 kb after electrophoresis. The A449 strain also displayed a resistance to tetracycline, but in this case this is due to the presence of a tetracycline resistance gene found on its pAsa4 plasmid.

Strains without pAsal1 were negative by PCR for the presence of the aopP gene, except for HER1104 (Table 1). In theory, it was expected that the aopP gene was present only in strains displaying pAsal1. This discrepancy between the plasmid profile and the PCR result for HER1104 set this strain apart from all others we analyzed, and suggested that elements from pAsal1 are still present in the genome of HER1104. Since pAsal1 contains ISAS11, which is known to be active in genome rearrangements, we suspected that the additional 8.5 kb band in HER1104 may be the result of modifications to pAsal1. The 8.5 kb band was thus isolated and characterized.
DNA restriction fragments encompassing the entire 8.5 kb plasmid were cloned and sequenced. The sequence analysis revealed a high level of identity with the sequence of pAsa1 and ISAS5. This unexpected result suggested that the additional plasmid in HER1104 might be a combination of pAsa1 and ISAS5. This 2614 bp IS belongs to the IS21 family.4,22 A449 contains 12 complete and three partial versions of ISAS5 in its genome.7

To obtain the complete sequence of the additional plasmid in HER1104, primers were designed from the sequences of the clones. The sequences obtained with these primers made it possible to identify the insertion site of ISAS5 in the plasmid. A second set of primers was designed to fill in the gaps and to get a complete circular sequence. Due to its high degree of similarity with pAsa1, the new plasmid (8989 bp) was named pAsa1B. The nucleotide sequence of pAsa1B was deposited in DDBJ/EMBL/GenBank under accession number KC686700.

The linear maps of pAsa1 and pAsa1B are shown in Figure 2. While the two plasmids were nearly identical, pAsa1B contained two ISAS5 elements. One was complete and was identical to other ISAS5 in the genome of A. salmonicida. It was inserted in mobA, resulting in the interruption of this gene. A fragment of ISAS5 was also found in the non-coding region between the repA and the mobA genes in pAsa1B (Fig. 2). This 96 bp fragment corresponded to part of the non-coding region upstream from the istA gene in ISAS5.22 This ISAS5 fragment was associated with the absence of a 121 bp sequence ranging from position 603 through 723 of pAsa1, which was replaced by the ISAS5 portion.

It has already been reported that pAsa5 in HER1104 is altered.8 Under stressful conditions, two typical pAsa5 alterations are most likely produced, that is, loss profile 1 and loss profile 2.2,4 As already mentioned, loss profile 1 results from the deletion of the TTSS locus, while loss profile 2 results from an additional 40 kb deletion upstream from this locus.7 The alteration of pAsa5 in HER1104 fits the loss profile 2 category (Fig. 3A). Loss profile 2 was observed less frequently than loss profile 1. However, until now, all loss profile 2 plasmids have been explained by homologous recombination of flanking ISAS11 (A and C).7

To confirm that pAsa5 loss profile 2 in HER1104 was due to the recombination of ISAS11A with ISAS11C, 11AF-11CR PCR was performed to amplify the recombined ISAS11A/C as well as the flanking sequences on either side of the reclosed plasmid.

Discussion

The goal of the present study was to discover additional evidence of the genomic plasticity of A. salmonicida. We thus analyzed the small plasmid profiles of 78 strains from different parts of the world, but mainly from Canada. The vast majority of the strains harbored only the cryptic plasmids pAsa1, pAsa2 and pAsa3 in addition to pAsa1. Strains with an unusual profile can be classified into two categories: (1) strains grown in stressful conditions and (2) strains possessing antibiotic resistance related to the presence of a small plasmid bearing an antibiotic resistance gene.

Figure 1. HER1104 displays an uncommon small plasmid profile. Plasmids under 50 kb in size were extracted from A. salmonicida A449, 01-B526, and HER1104 and were digested with EcoRI. The digested DNA samples were separated by pulsed field gel electrophoresis. The three strains tested contained pAsa1, pAsa2, and pAsa3. It was already known that A449 does not possess pAsa1 while this plasmid is present in 01-B526.6,7 EcoRI-digested HER1104 DNA does not contain a band corresponding to pAsa1 but does contain a band corresponding to an unknown plasmid with a higher molecular weight, which was named pAsa1B after further analyses.

Figure 2. Maps of the pAsa1 and pAsa1B plasmids. Genes are illustrated by arrows while IS elements are illustrated by rectangles. The numbers between the two plasmids indicate the identity (in %) for the plasmid regions indicated. As shown by the maps, pAsa1B contains all the elements found in pAsa1 except that a 121 bp fragment was replaced by the insertion of the 96 bp ISAS5 fragment in the first third of the plasmid. In addition, pAsa1B bears a complete ISAS5 inserted in the mobA gene.

Figure 3. PCR analyses of A449 and A449-R2 strains served as controls. The A449-R2 strain, which was derived from A449 and displayed a pAsa5 loss profile 2 due to the recombination of ISAS11A with ISAS11C,7 was used as a positive control. HER1104 generated the same PCR products as A449-R2, confirming that the loss profile 2 seen in HER1104 is also caused by an ISAS11 recombination (Fig. 3B).
The present study is the first to describe the mobility of IS from the IS21 family (ISAS5 in this case) in *A. salmonicida*. Moreover, the presence of a partial ISAS5 on pAsal1B compared with pAsal1 provides support for the idea that ISs in *A. salmonicida* can generate incomplete and complex integrations and rearrangements as already suggested.7,15 If the two ISAS5-related sequences are not taken into consideration, the sequence identity between pAsal1 and pAsal1B is higher than 99.9%, indicating that pAsal1B is a direct derivative of pAsal1 in which two ISAS5 (one complete and one partial) have been integrated.

A cluster of four different ISs located downstream from the TTSS locus in pAsa5 (Fig. 3A) may be the source of the ISAS5 observed in pAsal1B. In this cluster, ISAS11C is directly followed by an ISAS5 (ISAS5A). It is thus possible that, due to its relative proximity, the ISAS11 recombination detected in pAsa5 promoted the mobility of ISAS5A and its activation and transposition in pAsal1B. However, since the 11CR primer hybridizes in ISAS5A, we cannot infer that this IS copy is the source of the transposition event into pAsal1 that generated pAsal1B. If ISAS5A is the source of the ISAS5 in pAsal1B, this event occurred by replicative transposition, a mechanism already observed in the IS21 family.22 In fact, the presence of mobile genetic elements of the IS21 family such as ISAS5 has been reported in numerous bacterial genomes. These ISs have been shown to be involved in different types of genetic rearrangements, including recombinations, the creation of new composite transposons, replicon fusion, interplasmid recombinations and deletions by intramolecular transposition.22,25-27

The transposition of ISAS5 in the pAsal1 replicon reported here is another illustration of the genetic activity of these IS and their impact on the genome of their host. IS21-related genetic events can occasionally leave remnants of the IS in the "donor" DNA when the transposition mechanism is defective. These sequence relics have been reported in every genome bearing these transposons, including *A. salmonicida*.9,22,24,28,29 The 96 bp portion of the ISAS5 sequence detected in pAsal1B is an example of an aborted transposition. The fact that two transposition events by ISAS5 occurred in the pAsal1 replicon confirms the involvement
of this IS in *A. salmonicida* genome plasticity and reinforces the idea that the high content of mobile genetic elements in the genetic material of this species might indicate that *A. salmonicida* is subject to frequent rearrangements and rapid evolution.

The discovery and characterization of pAsal1B revealed for the first time the mobile nature of ISA55 in *A. salmonicida*. It is the fourth IS, after ISA51, ISA52 and ISA511 for which experimental evidences indicate the involvement of an IS in gene inactivation or plasmid rearrangements through its transposition in this bacterium. The present study also provided additional proofs demonstrating that the genomic plasticity of *A. salmonicida* is complex. As already mentioned, stressful conditions lead to the activation of plasmid rearrangements in *A. salmonicida*. In the future, it would be interesting to determine which conditions specifically provoke the mobilization of ISA55.

### Material and Methods

**Bacterial strains and growth conditions.** The *A. salmonicida* strains used in our study are listed in Table S1. In addition to these strains, A449-R2 was also used.7 The strains were grown on furunculosis agar medium (10 g of bacto-tryptone, 5 g of yeast extract, 1 g of t-tyrosine, 2.5 g of NaCl and 15 g of agar in one liter of water) at 18°C for 3 d. *Escherichia coli* MC1061 was used for cloning.

**Plasmid DNA isolation and restriction enzyme profiles.** Plasmid miniprep kits (Feldan, 9K-006-0010) were used as recommended by the manufacturer to extract < 50 kb plasmids from the strains listed in Table 1. The extracts (25 μl) were digested with EcoRI [New England Biolabs (NEB), R3101S]. The digested DNA samples were separated either by regular gel electrophoresis or by pulsed field gel electrophoresis with the following parameters: a pulse time ranging from 0.2 to 13.0 sec, a run time of 16 h at 6 V/cm and a 120° angle. The DNA bands under UV illumination. agarose gels were stained with ethidium bromide to visualize the DNA.

**Tetracycline resistance analyses.** Strains listed in Table 1 were analyzed for their resistance to tetracycline. A homogeneous bacterial lawn of 10^8 bacteria was made on Mueller-Hinton agar and were analyzed for their resistance to tetracycline. A homogeneous bacterial lawn of 10^8 bacteria was made on Mueller-Hinton agar. The plates were incubated at 18°C for 24 h on the petri dish. Afterwards, the plates were incubated at 18°C. The results were taken an incubation of 24 h and were interpreted depending on the presence or absence of a growth inhibition zone.

**PCR analyses.** PCR analyses were performed to test the presence of the aopP gene in all the *A. salmonicida* strains analyzed, to confirm the presence of pRAS5 in some strains, and to verify the rearrangement status of pAsal1B in HER1104 strain. The PCR primers used are listed in Table 2. DNA templates were prepared by lysing one bacterial colony of every *A. salmonicida* strain in 25 μl of SWL buffer (50 mM KCl, 10 mM Tris, pH 8.3, 2.5 mM MgCl₂, 0.45% NP-40 and 0.45% Tween 20).31 The lysates were heated at 95°C for 5 min. The PCR mixture contained 4 μl of 5 × GoTaq buffer (Promega, M791A), 1.6 μl of 2 mM dNTP, 1.3 μl of forward and reverse primers (100 ng/μl each), 0.1 μl of GoTaq [5 U, (Promega, M3005)], 10.7 μl of H₂O and 1 μl of DNA template. The PCR program was as follows: 2 min 30 sec at 95°C, 30 cycles of 30 sec at 95°C, 30 sec at 60°C and 2 min at 68°C, followed by a final extension for 5 min at 68°C. The samples were separated on 1% agarose gels, which were stained with 0.5 μg/ml ethidium bromide. The PCR reactions were performed at least twice.

**Cloning and sequencing of pAsal1B.** EcoRI-digested plasmid DNA from HER1104 was deposited in 18 wells of a 0.4% agarose gel prestained with ethidium bromide and was migrated for 5 h at 70 V. Bands in the 8 kb region were excised under UV illumination.

### Table 2. Primers used in this study

| Primer name | Sequence (5′-3′) | Ref. |
|-------------|-----------------|-----|
| MT5-R1      | AAGTCTCGATCTGTCCTGATA | This study |
| MT7-R1      | TTTGGAAGCGTGTTCCTGATGCC | This study |
| MT9-R1      | AGGACCGCAAACAGAACATTTGCG | This study |
| MT11-F1     | TTCACTTACTGGCAATGAGCCTT | This study |
| MT13-R1     | GTTTGATATCCAAAGTGGAGGG | This study |
| MT15-F1     | TGGCCCTATGGCATATCCAACAA | This study |
| MT18-R2     | TGGTTGATATGGGAGTGGCCAGT | This study |
| MT19-R1     | CTCTATGCTGCATGGCATAA | This study |
| MT21-F1     | CTCAAGTCCTCGTCTTCAACT | This study |
| MT22-R1     | GTCACTGAAAGGAGAATCGATGAG | This study |
| MT23-R1     | CGGTCAGTCTCAGTTGCTAGG | This study |
| MT24-R1     | GTCTTTCAAGCGCAAGGACAAG | This study |
| MT25-F1     | TGGGTTAGATGGTCTGAGGAAA | This study |
| MT26-R1     | GACCAATGGTAGCTGATTCC | This study |
| MT27-F1     | CTGTCCTGATCCATCCCATATC | This study |
| MT28-R1     | GATAAGCAGGCTTCAAGAGATG | This study |
| MT29-F1     | TTCTTATGCTGCTTCTGGATG | This study |
| MT30-R1     | CGGATAGAAGTGTGATGTCC | This study |
| MT31-F1     | TTGTGTTGATGAGTTGGTAGG | This study |
| MT32-R1     | TTAGAGAATGACTGGCCCTAA | This study |
| MT33-F1     | TTCACCAGCTTGCCAGTTAT | This study |
| MT33-F2     | TAAGGTGATCTCTGCTGGTTTG | This study |
| MT34-F1     | CTATGCTGCCAATCCAGAAAC | This study |
| MT34-F2     | ATCGATGCCAACATCGAACAA | This study |
| MT35-R1     | CATGGATGATGCTGAGTGGAATA | This study |
| MT35-R2     | GGTCAACTGGCAAGAGTAGG | This study |
| MT36-R1     | CAACACCAGCTTCCCATCAGTA | This study |
| MT36-R2     | CATATCCAGCTCATGCCCAA | This study |
The DNA was sent to the genome analysis platform (IBIS, Université Laval, Quebec City, QC, Canada) for sequencing. Nucleotide BLAST analyses of the sequences were performed using tools available on the National Center for Biotechnology Information website (www.ncbi.nlm.nih.gov). To complete the assembly of pAsa1B, the DNA was sequenced by primer walking. The primers used are listed in Table 2. Primers were designed using primer design software available online on the Integrated DNA Technologies website (www.idtdna.com). The sequences of pAsa1 and pAsa5 were used to design the required primers (GenBank accession numbers: CP0000646 and AJ508382).

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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Supplemental Materials
Supplemental materials may be found here: www.landesbioscience.com/journals/mge/article/25640

Table 2. Primers used in this study (continued)

| Primer name | Sequence (5′-3′) | Ref. |
|-------------|-----------------|------|
| pAsa1 F | TAAATGGGTTAGTACAGGA | 4 |
| pAsa1 R | TGCAATTTTGTAAAAGATTGTT | 4 |
| pRAS3deb-F | CATGACATTCGGCATCAGCTTCAA | This study |
| pRAS3deb-R | TCCTCAGCGGGAATCCTCATACT | This study |
| pRAS3mil-F | GCGGTCTGAAACAAACCTCGTTT | This study |
| pRAS3mil-R | ACTGTGGGCGTGGCTCAAACATC | This study |
| tapA F | ACATGGAAGAAGATAACGGC | 30 |
| tapA R | AGAGCTCATTGTTAGCAG | 30 |
| traC F | TGCACTATCCCGACCCTACCC | 8 |
| traC R | TCGTAATCGGCGTCTTGTC | 8 |
| exsD F | AGAAGTGTACCTGACCAAAAGGCAA | 8 |
| exsD R | TGGCAACAGCTGTGCCAGAAGACC | 8 |
| resD F | TCAGAAACTTGGGACATGCTCAC | 8 |
| resD R | TGATGTCAGACCTTCCCTGAGACA | 8 |
| 11AF | AATAGGGTCCGAAGCTGGTGTTA | 7 |
| 11CR | AACTGCAAGAGATAGAGCTCTGAG | 7 |
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