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2 **A broad-spectrum *mer* operon in a multi-drug resistant strain of the fish pathogen,**  
3 ***Aeromonas salmonicida*.**

4 **Erin Parry**  
5 **Colby College Biology Department**  
6 **Honors Thesis**  
7 **Submitted May 17, 2006**

8 **Abstract**

10 *Aeromonas salmonicida* AS03, a potential fish pathogen, was isolated from  
12 Atlantic salmon, *Salmo salar*, in 2003. This strain was found to be resistant to  $\geq 1000$  mM  
14 HgCl<sub>2</sub> and  $\geq 32$  mM phenylmercuric acetate as well as multiple antimicrobials. Mercury  
16 (Hg) and antibiotic resistance genes are often located on the same mobile genetic  
18 elements, so the genetic determinants of both resistances and the possibility of horizontal  
20 gene transfer were examined. Specific PCR primers were used to amplify and sequence  
22 distinctive regions of the *mer* operon. *A. salmonicida* AS03 was found to have a  
24 pDU1358-like broad-spectrum *mer* operon, containing *merB* as well as *merA*, *merD*,  
26 *merP*, *merR* and *merT*, most similar to *Klebsiella pneumoniae* plasmid pRMH760. To our  
28 knowledge, the *mer* operon has never before been documented in *Aeromonas* spp. PCR  
and gene sequencing were used to identify class 1 integron associated antibiotic  
resistance determinants and the Tet A tetracycline resistance gene. The transposase and  
resolvase genes of Tn1696 were identified through PCR and sequencing with Tn21  
specific PCR primers. We provide phenotypic and genotypic evidence that the *mer*  
operon, the aforementioned antibiotic resistances, and the Tn1696 transposition module  
are located on a single plasmid or conjugative transposon that can be transferred to *E. coli*  
DH5 $\alpha$  through conjugation in the presence of low level Hg and absence of any antibiotic  
selective pressure. Additionally, the presence of low-level Hg or chloramphenicol in the

mating media was found to stimulate conjugation, significantly increasing the transfer  
30 frequency of conjugation above the transfer frequency measured with mating media  
lacking both antibiotics and Hg. This research demonstrates that mercury indirectly  
32 selects for the dissemination of the antibiotic resistance genes of *A. salmonicida* AS03.

### **Introduction**

34 *Aeromonas salmonicida*, a common fish pathogen, is the bacterium responsible  
for furunculosis in temperate and coldwater salmonid fish (1, 36). Outbreaks of this  
36 pathogen can be controlled by vaccination or, more commonly, by antibiotic  
administration. However, excessive usage of antibiotics in fish feed has caused the  
38 emergence of multiply antibiotic resistant fish pathogens (36). As a result of these multi-  
drug resistant fish pathogens, countries such as Denmark, Norway, Sweden, and the  
40 United Kingdom have begun to monitor antibiotic usage in fish farms (36). The  
acquisition of antibiotic resistance genes by *A. salmonicida* confounds disease control  
42 and is thus economically costly to the aquaculture industry(35). Mobile genetic elements  
such as Tn1721 and class 1 integrons isolated from *A. salmonicida* strains are commonly  
44 located on resistance plasmids (36). Aeromonad resistance plasmids are often IncU  
plasmids or genetically similar to IncU plasmids, suggesting that they share a common  
46 origin and have since diverged due to different selective pressures(9, 37). IncU plasmids  
belong to the plasmid incompatibility group U, which consists of similar plasmids that  
48 cannot co-exist in any bacterial strain and are thus said to be incompatible (37) Therefore,  
the resistance determinants associated with fish farm pathogens are of further  
50 significance because they are very similar to those found in pathogenic enterobacteria in  
humans and hospital effluents (37). Antibiotic resistances associated with resistance

52 plasmids, such as oxytetracycline resistance, have been transferred from *Aeromonas sp.*  
to other bacterial species, including *E. coli* (11).

54 Mercury (Hg) resistance is prevalent among environmental bacterial isolates due  
to the selective pressure of elevated mercury contamination (3). The most common  
56 mechanism for bacterial mercury resistance involves the reduction of toxic mercury(II) to  
volatile Hg(0) by the mercuric reductase enzyme (3, 4). The gene encoding mercuric  
58 reductase, *merA*, is usually located in the *mer* operon, where it is associated with other  
genes coordinating mercury resistance. In addition to *merA*, the typical *mer* operon  
60 contains two genes responsible for mercury transport into the cell (*merT* and *merP*) and  
two regulatory genes (*merD* and *merR*). Some *mer* operons also contain the accessory  
62 genes *merB*, *merC* and *merF* (25). Since mercury resistance determinants are often  
located on mobile genetic elements such as plasmids and transposons, the *mer* operon can  
64 be transferred between species (3, 43). Several studies have demonstrated the transfer of  
mercury resistance via conjugation between bacterial species(3, 43). Although excessive  
66 antibiotic usage is largely responsible for the growing prevalence of antibiotic resistance,  
non-antibiotic selective agents, including mercury and other heavy metals, have been  
68 implicated as potential vehicles for the indirect selection of antibiotic resistance (34).  
Antibiotic and mercury resistance genes are frequently adjacent to each other on mobile  
70 genetic elements and furthermore located the same integrons (5, 28, 38, 43). This  
correlation between antibiotic and mercury resistance has been documented in the fecal  
72 and intestinal bacteria of primates with mercury amalgam fillings, where bacteria  
containing a *mer* operon are more likely to exhibit multiple antibiotic resistance(38).  
74 Furthermore, McArthur and Tuckfield observed increased antibiotic resistance in

bacterial strains exhibiting mercury resistance isolated from the sediment of mercury-  
76 polluted streams (28). Although many plasmid-borne antibiotic resistant genes have been  
characterized in *Aeromonas sp.*, no aeromonad to our knowledge has been documented to  
78 contain the *mer* operon conferring mercury resistance (1, 36).

Class 1 integrons, which are often found in close proximity to *mer* operons, are  
80 found in many gram-negative clinical isolates and have been shown to carry a high  
diversity of antibiotic resistance cassettes. Class 1 integrons have been documented to  
82 associate with transposons encoding both mercury and tetracycline resistance, thus  
mobilizing the integron and the transposon's resistance determinants together (27, 30).  
84 The genes encoding class 1 integrases associated with transposons carrying mercury  
resistance are identical to those encoding integrases not associated with transposons,  
86 suggesting that the incorporation of integrons into transposons occurred recently and is an  
ongoing process (15). Integron-borne antimicrobial resistances could therefore be easily  
88 spread to pathogenic bacteria via transposons, especially with the indirect selective  
pressure exerted by mercury (30).

90 The bacterial strain characterized in this study, *A. salmonicida* AS03, was isolated  
from hatchery-grown Atlantic salmon, *Salmo salar* in 2003. *A. salmonicida* AS03 was  
92 screened for resistance to HgCl<sub>2</sub>, phenylmercuric acetate (PMA), and a total of 26  
antimicrobial agents. Distinctive regions of the *mer* operon, class 1 integron genes, and  
94 *Tet A*, a tetracycline resistance determinant typical of many multiple-drug resistant  
aeromonads isolated from aquaculture facilities, were partially characterized  
96 genotypically. We provide evidence that the *mer* operon and multiple antibiotic  
resistance genes in *A. salmonicida* AS03 can be concomitantly transferred to *Escherichia*

98 *coli* DH5 $\alpha$  by conjugation and are, therefore, associated with at least one mobile genetic  
element. Furthermore, low levels of either mercury or the antibiotic chloramphenicol  
100 stimulated higher transfer frequencies of conjugation.

## **Materials and Methods**

102 **Bacterial isolates and cultivation conditions.** An isolate of *Aeromonas salmonicida*,  
designated strain AS03, was found to be resistant to both mercury and antimicrobials.  
104 Strain AS03 was isolated from 7 of 60 Atlantic salmon (*Salmo salar* L.) during a routine  
health inspection in 2003 of hatcheries in Northeastern North America. *Escherichia coli*  
106 DH5 $\alpha$  (Invitrogen Life Technologies, Carlsbad, CA) was used as the recipient in the  
conjugation experiments. *A. salmonicida* AS03 and *E. coli* DH5 $\alpha$  were grown on Brain  
108 Heart Infusion (BHI) agar (Difco, Detroit, MI) at 22°C and at 37°C, respectively. Both  
strains were stored in BHI broth with 50% (vol/vol) glycerol at -80°C.

110 **Genomic DNA extraction and confirmation of strain identity.** 16S rRNA gene  
sequencing was done to confirm the identity of the *A. salmonicida* AS03 strain. Total  
112 genomic DNA was extracted (E.N.Z.A. Bacterial DNA Kit, Omega Bio-tek, Doraville,  
GA) according to the manufacturer's protocol. A hypervariable, 174-200 bp region of the  
114 16S rDNA gene was PCR amplified using primers and cycling conditions described by  
Muzyer *et al*(29) and sequenced on an ABI Prism 310 Genetic Analyzer (Applied  
116 Biosystems, Foster City, CA). Isolate identification was based on 16S rDNA sequence  
similarities (Blastn against GenBank at <http://www.ncbi.nlm.nih.gov/BLAST>).

118 **Mercury minimum inhibitory concentrations (MICs).** Hg MIC values for both HgCl<sub>2</sub>  
and phenylmercuric acetate (PMA; Sigma-Aldrich, Inc., St.Louis, MO) were determined  
120 for *A. salmonicida* AS03 and *E. coli* DH5 $\alpha$  by the method of Wang *et al.*(42) using BHI

plating media solidified with Noble agar (Difco, Sparks, MD). Bacterial suspensions ( 4  
122 .0 µl; 2.0 McFarland) of fresh (24-48 hr) solid plating media cultures in sterile,  
demineralized water were inoculated on BHI plating media containing 0, 50, 100, 250,  
124 500, 750, and 1000 µM HgCl<sub>2</sub> , and BHI plating media containing 0, 0.5, 1.0, 2.0, 4.0,  
8.0, 16.0 and 32.0 µM PMA and incubated for two days at 22°C. Hg MICs were recorded  
126 as the lowest concentration of Hg that inhibited bacterial growth. Two control strains,  
mercury-susceptible *Bacillus subtilis* 168 and mercury-resistant *E. coli* SK1592, were run  
128 in parallel with the experimental strains(18)

**Antibiotic MICs.** The *in vitro* activity of antimicrobial agents against *A. salmonicida*  
130 AS03 and *E. coli* DH5α isolates was assessed by the MIC method following the National  
Committee for Clinical Laboratory Standards (NCCLS) guidelines. MICs were  
132 determined using the Sensititre<sup>®</sup> (MJ Panel; Trek Diagnostic Systems, Westlake, OH)  
dried susceptibility panels. Manufacturer's instructions were followed for *E. coli* DH5α,  
134 however, modifications for *A. salmonicida* AS03 included incubation at 22°C for 48h.  
Antibiotic MIC assays were done in triplicate for each strain, and MIC was determined as  
136 the lowest concentration of the antimicrobial agent able to inhibit growth. As  
recommended by NCCLS and Sensititre<sup>®</sup> guidelines, the following reference strains were  
138 included as internal standards: *Escherichia coli* (ATCC 25922), *Pseudomonas*  
*aeruginosa* (ATCC 27853), and *Enterococcus faecalis* (ATCC 29212). For specific  
140 antimicrobials that were not included in the Sensititre MJ panel, additional susceptibility  
testing was performed on BHI plating media with antimicrobial discs containing  
142 streptomycin (10 µg), tobramycin (10 µg), kanamycin (30 µg) and chloramphenicol (30  
µg) (Sensi-Disc; Becton, Dickinson and Co., Sparks, MD). *A. salmonicida* cultures were

144 incubated at 22°C while *E. coli* cultures were incubated at 37°C before zones of  
inhibition were measured.

146 **PCR amplification and sequencing of *mer* genes.** Primers (Table 1) were used to probe  
for and amplify regions of the *mer* operon using genomic DNA as a template (25). The  
148 short (288 bp) *merA* PCR product was cloned according to the manufacturer's protocol  
(TA Cloning Kit ; Invitrogen Life Technologies, Carlsbad, Calif.) and sequenced. All  
150 other PCR products were cleaned (E.N.Z.A. CyclepureKit or E.Z.N.A Gel Extraction Kit;  
Omega Bio-tek, Doraville, GA) according to manufacturer's protocol and sequenced. All  
152 *mer* sequences were collected and aligned in Sequencher 4.6 (Gene Codes Corporation,  
Ann Arbor, MI). *mer* gene sequences were entered in the blastn database to confirm gene  
154 identity and identify similar sequences.

**PCR amplification and sequencing of *Tn21* genes and antibiotic resistance genes.**

156 Primers (Table 1) were used to amplify *Tn21*, class 1 integron genes, and the tetracycline  
resistance gene Tet A. These reactions were carried out on a gradient thermocycler  
158 (iCycler, BioRad Hercules, CA) for 35 cycles of 94 °C for 1 min, 53-57°C for 1 min and  
72°C for 2.5 min. The 1 kb PCR product of *in-F* and *in-B*, the 1 kb Tet A amplicon, a 411  
160 bp *tnpA* amplicon, and a 240 bp *tnpR* amplicon were sequenced

**Conjugation experiments and plasmid profiling.** The ability of *A. salmonicida* AS03  
162 to transfer mercury- and antibiotic resistance genes to *E.coli* DH5α (Invitrogen Life  
Technologies; Carlsbad, CA) was examined by conjugation experiments using the solid  
164 plating medium method (21). Growing colonies of the *A. salmonicida* AS03 were mixed  
with approximately equal amounts of growing recipient strain *E. coli* DH5α on BHI

166 plates, with mercury and antibiotic amendments as specified below, and incubated at  
22°C from 1 to 2 days

168 The frequency of transfer was determined according to the method of Sorum, et  
al. (37) with the following modifications. Transfer frequency was measured by dividing  
170 the number of transconjugants per ml by the number of recipients per ml in the mating  
mixture. The total number of recipients was estimated after overnight incubation on BHI  
172 solid plating media without selection agents at 37°C, at which the growth of the donor  
strain was suppressed, while selection of transconjugants was done on BHI agar plates  
174 with either 8 µg ml<sup>-1</sup> of chloramphenicol or 250 µM HgCl<sub>2</sub> following incubation at 37°C.  
These concentrations of chloramphenicol and HgCl<sub>2</sub> were found to inhibit growth of the  
176 *E. coli* DH5α recipient. All *E. coli* DH5α transconjugants were subjected to mercury and  
antibiotic MIC assays at a temperature of 37°C to determine their phenotypic resistance  
178 characteristics. The transfer frequencies were calculated on the basis of the average  
number of colonies counted from duplicate plates after appropriate serial dilution (2.0  
180 McFarland) of the mating mixture.

Several conjugation experiments were performed to determine if the transfer of  
182 resistance genes is stimulated by exposure of the donor *A. salmonicida* AS03 to mercury  
and chloramphenicol. Three experiments were conducted under each condition of either  
184 chloramphenicol or mercury exposure of the donor strain and mating mixture. To assess  
the stability of the resistance determinants in the *E. coli* DH5α transconjugants, both  
186 antibiotic and mercury MICs were done on the transconjugants after 10 subculture  
passages on BHI plating media at 37°C.

188           **(i) Conjugation in the presence of chloramphenicol.** *A. salmonicida* AS03 was  
grown overnight on BHI solid plating media amended with 8  $\mu\text{g ml}^{-1}$  chloramphenicol.  
190 Donor and recipient strains were mixed as described above on BHI agar plates containing  
1  $\mu\text{g ml}^{-1}$  chloramphenicol, a concentration determined to be sub-lethal to recipient *E.*  
192 *coli* DH5 $\alpha$ . The mating mixtures were incubated and the frequencies of transfer were  
determined as described above.

194           **(ii) Conjugation in the presence of mercury.** The donor strain, *A. salmonicida*  
AS03, was grown overnight on BHI solid plating media with added 25  $\mu\text{M HgCl}_2$ .  
196 Donor and recipient strains were mixed as described above on BHI agar plates containing  
25  $\mu\text{M HgCl}_2$ , and the mating mixtures were incubated as described above. Transfer  
198 frequencies were done as described above except that selection for *E. coli* DH5 $\alpha$   
transconjugants was done on BHI plating media with 250  $\mu\text{M HgCl}_2$ .

200           Plasmid DNA was isolated from *A. salmonicida* AS03, *E. coli* DH5 $\alpha$  recipient,  
and the *E. coli* DH5 $\alpha$  transconjugant using the E.N.Z.A. Plasmid miniprep kit (Omega  
202 Bio-tek, Doraville, GA). Plasmids were visualized using pulsed field gel electrophoresis  
(PFGE) in a 0.8% agarose gel at 175 V at 4°C. Genomic DNA was extracted from *A.*  
204 *salmonicida* AS03, *E. coli* DH5 $\alpha$  recipient, and the *E. coli* DH5 $\alpha$  transconjugant as  
described above in order to serve as template DNA in the following PCR amplifications.  
206 PCR primer sets A1-s.F and A5-n.R, *In-F* and *In-B*, *tnpR-F* and *tnpR-B*, Tet A FW and  
Tet A RV, and p5-II and p2-II (Table 1) were used to probe the *E. coli* DH5 $\alpha$  recipient  
208 and transconjugant for the presence of the *merA*, *aadA7*, *tnpR*, Tet A, and *tnpA* genes that  
were present in *A. salmonicida* AS03.

210 **Nucleotide sequence accession numbers.** The sequence of the 16S rRNA gene has been  
 deposited in the GenBank database under the following accession number: strain AS03  
 212 XXXXXXXXXX. Partial sequences of the *merA*, *merD*, *merR-T*, *merB*, Tet A, and *aadA7*  
 genes of strain AS03 have been deposited in the GenBank database under accession  
 214 numbers XXXXXX through XXXXXXXX

216

**Table 1:** PCR primers used in this study

Primer	Gene	Sequence	Source or Reference
A1-s.F	<i>merA</i>	TCCGCAAGTNGCVACBGTNGG	(41)
A5-n.R	<i>merA</i>	ACCATCGTCAGRTARGGRAAVA	(41)
R1-F	<i>merR-merT</i>	GCGGATTTGCCTCCACGTTGA	(25)
T1-R	<i>merR-merT</i>	CCAGGCAGCAGGTTCGATGCAAG	(25)
P1-F	<i>merP-merA</i>	GGCTATCCGTCCAGCGTCAA	(25)
A0-R	<i>merP-merA</i>	GTCGCAGGTCATGCCGGTGATTTT	(25)
B1-F	<i>merB</i>	TCGCCCATATATTTTAGAAC	(25)
B2-R	<i>merB</i>	GTCGGGACAGATGCAAAGAAA	(25)
D2-F	<i>merD</i>	CCAGGCGGCTACGGCTTGTT	(25)
D3-R	<i>merD</i>	GGTGGCCAACTGCACTTCCAG	(25)
A6-F	<i>merA-merD</i>	GCCGACCAGTTGTTCCCCTACCTGACG	(25)
D1-R	<i>merA-merD</i>	CGCACGATATGCACGCTCACCC	(25)
<i>In-F</i>	Integron	GGCATCCAAGCAGCAAGC	(40)
<i>In-B</i>	Integron	AAGCAGACTTGACCTGAT	(40)
p5-II	Tn21 <i>tnpA</i>	TACTGCCGCGCATCAAGATC	(10)
p2-II	Tn21 <i>tnpA</i>	AGAAAGTTTCGTCCTGGGCTG	(10)
tnpR-F	Tn21 <i>tnpR</i>	GGCGACACCGTGGTGGTGCATAGC	(6)
tnpR-B	Tn21 <i>tnpR</i>	CGGTAAGCCCCGCGTTGCTTGGC	(6)
qacF	<i>qacEA1-sul1</i>	ATCGCAATAGTTGGCGAAGT	(35)
Sul R	<i>qacEA1-sul1</i>	GCAAGGCGGAAACCCGCGCC	(35)
tetA FW	Tet A	GTAATTCTGAGCACTGTTCGC	(17)
tetA RV	Tet A	CTGCCTGGACAACATTGCTT	(17)

218

## Results

220 **16S rDNA gene sequencing**

The 16S rDNA sequence of *A. salmonicida* AS03 was identical (100%) to that of *A.*

222 *salmonicida* (AY297782.1), *A. salmonicida* subspecies subspecies *Achromogenes*

(AY910844.1) and *A. salmonicida* isolate 3-St 2-6 (DQ133187.1).

224

226

228

230 **Table 2.** Antibiotic minimum inhibitory concentrations ( $\mu\text{g/ml}$ ) as determined by Sensititre panels for *A. salmonicida* AS03, *E. coli* DH5 $\alpha$ , and the transconjugant.

Antimicrobials by class (a)	<i>Aeromonas salmonicida</i> AS03 Donor	<i>Escherichia coli</i> DH5 $\alpha$ recipient	<i>Escherichia coli</i> DH5 $\alpha$ transconjugant
Tetracyclines			
Tetracycline (0.25-8)	>8	1	>8
Penicillins			
Ampicillin (0.5-16)	>16	2	>16
Ampicillin/Sulbactam (8/4-16/8)	>16/8	0	>16/8
Mezlocillin (4-64)	0	0	16
Ticarcillin/Clauvulanic Acid (4/2-64/2)	8/2	0	64/2
Amoxicillin/Clauvulanic Acid (0.5/0.25-16/8)	>16/8	4/2	>16/8
Piperacillin (4-64)	0	0	16
Cephems			
Cefazolin (1-16)	>16	2	>16
Cephalothin (8-16)	>16	0	>16
Cefoxitin (2-16)	>16	0	>16
Ceftazidime (1-16)	16	0	>16
Ceftriaxone (4-32)	8	0	16
Cefuroxime (2-16)	8	0	>16
Aminoglycosides			
Amikacin (4-16)	0	0	0
Gentamicin (0.25-8)	0.5	0	0
Fluoroquinolones			
Ciprofloxacin (0.06-2)	0.25	0	0
Norfloxacin (4-8)	0	0	0
Ofloxacin (0.25-4)	0.5	0	0
Lomefloxacin (0.5-2)	2	0	0
Folate Pathway Inhibitors			
Trimethoprim/Sulfamethoxazole (0.5/9.5-2/38)	>2/38	0	0
Sulfisoxazole (256)	>256	0	>256

a Ranges tested in MIC in  $\mu\text{g/mL}$

232

**Hg and Antibiotic minimum inhibitory concentrations of *A. salmonicida* AS03.**

234 *Aeromonas salmonicida* AS03 demonstrated resistance to  $\geq 1000 \mu\text{M}$   $\text{HgCl}_2$  and  $\geq 32 \mu\text{M}$   
PMA as well as maximal resistance to 11 antimicrobials. In addition, some low-level  
236 resistance to ceftriaxone, ciprofloxacin, ofloxacin, and trimethoprim/sulfamethoxazole  
was observed (Table 2). Antimicrobial susceptibility disc assays showed that *A.*  
238 *salmonicida* AS03 was also resistant to 10  $\mu\text{g}$  streptomycin and 30  $\mu\text{g}$  chloramphenicol,  
and susceptible to 10  $\mu\text{g}$  tobramycin and 30  $\mu\text{g}$  kanamycin

240 **Genes encoding the *mer* operon.** Regions of the *merA*, *merB*, *merD*, *merR*, *merP*, and  
*merT* genes were amplified and sequenced in *A. salmonicida* AS03. All *mer* gene PCR  
242 amplifications of *A. salmonicida* AS03 produced single PCR bands of the following  
expected sizes: 288 bp for *merA*, 134 bp for *merA-P*, 225 bp for *merR-T*, 502 bp for  
244 *merB*, 219 bp for *merD*, and 181 bp for *merA-D*. Blastn searches confirmed the identities  
of the amplicons. All genes demonstrated the most sequence similarity to *Klebsiella*  
246 *pneumoniae* pRMH760 (>95% sequence similarity), which contains a pDU1358-like  
*mer* locus (accession no. AY123253). PCR mapping of the *merD-merA* region verified  
248 the presence of *merB* between *merA* and *merD* (3, 24, 25).

**Antibiotic resistance genes.** PCR amplification with class 1 integron primers *in-F* and  
250 *in-B* yielded several amplicons in *A. salmonicida* AS03. A 1 kb amplicon from this  
amplification was successfully sequenced and identified through blastn analysis as  
252 *aadA7*, which encodes an aminoglycoside 3' adenylyltransferase enzyme conferring  
resistance to streptomycin and spectinomycin. This gene exhibited the most similarity to  
254 *aadA7* in *Escherichia coli* (AF234167.1), *Vibrio cholerae* (DQ196322.1) and *Salmonella*  
*enterica* (AY463737) and explains the streptomycin resistant phenotype of *A.*

256 *salmonicida* AS03 determined by antimicrobial susceptibility disks. PCR primers *qacF*  
and *sulR* yielded an approximately 800 bp PCR product which was identified by blastn  
258 analysis on the reverse sequence as *sull*. This *sull* gene resembled *sull* genes of *E. coli*  
plasmid PQR-1 (AY655485) and *K. pneumoniae* pRMH760 (AY123253). The forward  
260 sequence confirmed the presence of *qacEΔI* and was most similar to *qacEΔI* in *E. coli*  
pSa (L06822.4) and pRMH760 (AY123253) in the blastn database. PCR and gene  
262 sequencing with primers *tetA FW* and *tetA RV* amplified an approximately 1 kb amplicon,  
which demonstrated the most sequence similarity with *tetA* from multiple genera,  
264 including *E. coli* pAPEC-OR-R (AY214164), *S. S. typhimurium* pU302L (AY333434),  
and *A. salmonicida* pRAS1 (AJ131405).

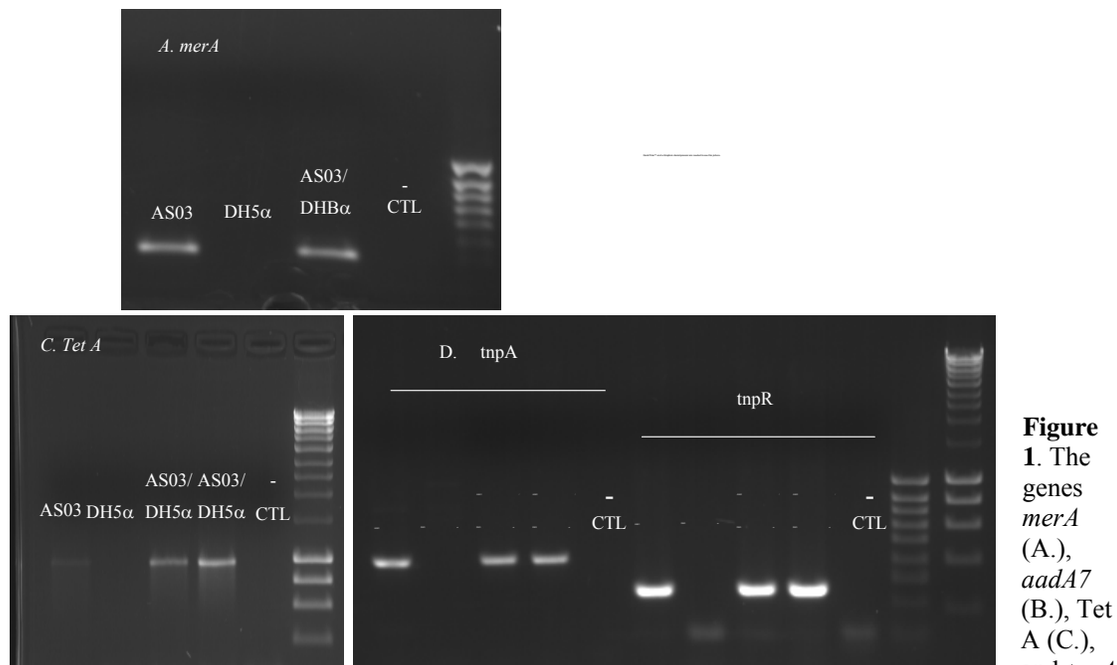
266 **Transposon genes.** Because this strain exhibited simultaneous antibiotic and Hg  
resistance phenotypes, the presence of Tn21, which carries both resistance determinants,  
268 was assayed using primers described by Dahlberg et al. and Guerra et al.(6, 10). A 411 bp  
fragment was amplified and sequenced using primers *p5-II* and *p2-II*, which was more  
270 similar to the *tnpA* from Tn1696 than that of *tnpA* of Tn21, and was almost identical to  
the Tn1696 *tnpA* of *P. arguinosus* R1033 (U12338.2) and *K. pneumoniae* pRMH760  
272 (AY123253). Similarly, the 240 bp amplicon obtained from PCR with primers *tnpR-R*  
and *tnpR-F* was identified as *tnpR* from Tn1696 and was near identical to the Tn1696  
274 *tnpR* of *P. arguinosus* p1033 and *K. pneumoniae* pRMH760.

**Conjugation.** *A. salmonicida* AS03 was able to transfer both mercury and antibiotic  
276 resistance to *E. coli* DH5α in conjugation experiments. Initial conjugation experiments  
showed that, regardless of the selective agent, either 8 μg ml<sup>-1</sup> chloramphenicol or 250  
278 μM HgCl<sub>2</sub>, the *E. coli* DH5α transconjugant had multiple antibiotic resistances with MIC

levels either identical to or exceeding that of the donor *A. salmonicida* AS03 strain,  
280 especially among the tetracycline, penicillin, and cephem classes of antibiotics (Table 2).  
The folate pathway inhibitor sulfisoxazole was also expressed maximally in the  
282 transconjugant. Notably, antibiotics of the fluoroquinolone class were not transferred,  
even though the donor, *A. salmonicida* AS03, exhibited low levels of resistance to some  
284 of the antimicrobials in both the fluoroquinolone classes.

Similarly, mercury MICs in the transconjugant were identical under conditions of  
286 either chloramphenicol or HgCl<sub>2</sub> selection. The donor *A. salmonicida* AS03 exhibited  
high resistance to mercury with MICs of  $\geq 1$  mM HgCl<sub>2</sub> and  $\geq 32$   $\mu$ M PMA, whereas the  
288 *E. coli* DH5 $\alpha$  recipient showed low levels of mercury resistance with MICs of 100  $\mu$ M  
HgCl<sub>2</sub> and 16  $\mu$ M PMA. After mating with *A. salmonicida* AS03, the *E. coli* DH5 $\alpha$   
290 transconjugant had mercury MICs of  $\geq 1$  mM HgCl<sub>2</sub> and  $\geq 32$   $\mu$ M PMA. No plasmids  
were isolated from the *E. coli* DH5 $\alpha$  recipient nor the transconjugant, while at least  
292 three plasmids, estimated to be between 4-10 kB using the BAC-Tracker Supercoiled  
DNA Ladder (Epicentre, Madison, WI), were visualized in the *A. salmonicida* donor.  
294 PCR amplification from the genomic DNA of the *E. coli* DH5 $\alpha$  transconjugant showed  
the presence of the *merA*, *aadA7*, *tnpR*, *Tet A*, and *tnpA* genes that were previously  
296 sequenced in the *A. salmonicida* AS03 donor. These genes were not amplified in the *E.*  
*coli* DH5 $\alpha$  recipient and, therefore, must have been acquired by the *E. coli* DH5 $\alpha$   
298 transconjugant via conjugative transfer from the *A. salmonicida* AS03 donor (Figure 1).

300



**Figure 1.** The genes *merA* (A.), *aadA7* (B.), Tet A (C.), and *tnpA*

326 and *tnpR* (D.) are present in the *A. salmonicida* AS03 donor and the *E. coli* DH5α transconjugant but absent  
 328 in the *E. coli* DH5α recipient, showing that these genes were transferred to the *E. coli* DH5α transconjugant  
 330 from *A. salmonicida* AS03 via conjugation. The notation AS03/DH5α is used to denote the transconjugant  
 on the gel pictures.

332 Conjugative transfer of resistance determinants to the *E. coli* DH5α recipient  
 varied under different conditions of the donor and mating mixture. The highest transfer  
 334 frequencies were observed when the donor was grown on BHI plating media amended  
 with either 8 μg ml<sup>-1</sup> chloramphenicol or 25 μM HgCl<sub>2</sub> prior to conjugation and the  
 336 mating mixtures contained either chloramphenicol or mercury at levels sub-lethal to the  
*E. coli* DH5α recipient (1 μg ml<sup>-1</sup> chloramphenicol or 25 μM HgCl<sub>2</sub>). Statistically, there  
 338 was no significant difference between the transfer frequencies of these two exposure  
 conditions. However, exposure to either chloramphenicol or mercury stimulated  
 340 significantly greater transfer frequencies than observed in the control, which contained  
 neither mercury nor chloramphenicol (Table 3). Regardless of the exposure condition,  
 342 transconjugants maintained the same mercury and antibiotic resistance phenotypes they  
 had demonstrated immediately after conjugation following ten sequential transfers on

344 BHI plating media, indicating that all of the transferred resistance determinants were  
stably inherited.

346

348 **Table 3.** Transfer frequencies of a conjugation between an *E. coli* DH5 $\alpha$  recipient and an *A. salmonicida*  
AS03 donor on solid BHI media under different conditions

Conjugation Condition	Transfer Frequency <sup>a</sup> $\pm$ SE
BHI agar	$(2.20 \pm .48) \times 10^{-2}$
25 $\mu$ M Hg in BHI agar	$(4.85 \pm .94) \times 10^{-2}$
8 $\mu$ M Chloramphenicol in BHI agar	$(6.60 \pm 1.0) \times 10^{-2}$

350 <sup>a</sup> Transfer frequency is the number of CFU on selective BHI media containing 8  $\mu$ g ml<sup>-1</sup> chloramphenicol  
divided by the number of CFU on non-selective BHI media

352

### Discussion

354 Hg resistance in *A. salmonicida* AS03 was found to be encoded by a broad-  
spectrum pDU1358-like *mer* operon. Plasmid pDU1358 is a multiple-antibiotic resistance  
356 IncC incompatibility group plasmid, specifying broad-spectrum mercurial resistance, that  
was originally isolated from a hospital strain of *Serratia marcescens* (16) *A. salmonicida*  
358 AS03 exhibited a similar amplicon pattern to that of pDU1358 locus 8 and a dissimilar  
pattern to that of Tn21 based upon the results reported by Liebert et al.(25). Additionally,  
360 the *mer* sequence in blastn with the highest degree of similarity to all *A. salmonicida mer*  
sequences, pRMH760, is a pDU1358-like *mer* locus. pRMH760 is a conjugative plasmid  
362 that carries resistances to chloramphenicol, ampicillin, gentamicin, kanamycin,  
neomycin, streptomycin, spectinomycin, sulfamethoxazole, tobramycin, and  
364 trimethoprim as well as a Tn1696 transposition module and a broad-spectrum *mer*  
locus(32). The mobile genetic element in *A. salmonicida* AS03 was also shown to carry  
366 ampicillin, streptomycin, spectinomycin, sulfamethoxazole, and trimethopim resistances

in addition to the broad-spectrum *mer* locus and Tn1696 transposition module. Hence,  
368 similar integron-containing composite transposons could be present in *A. salmonicida*  
AS03 and pRMH760. Several gram-negative isolates have been reported to possess more  
370 than one type of *mer* locus (25, 33). Data from this study does not support the existence  
of more than one type of *mer* locus in *A. salmonicida* AS03. Whereas Liebert et al. (20)  
372 found multiple *mer* loci in their bacterial isolates, as determined by the presence of  
multiple PCR bands, using the same PCR primer pairs (B1 and B2, D2 and D3, P1 and  
374 AO and R1 and A5), we observed only a single amplicon for the same PCR  
amplifications in *A. salmonicida* AS03.

376 Class I integrons are prevalent in antibiotic-resistant clinical isolates of gram-  
negative bacteria (32). Each class 1 integron may contain several gene cassettes  
378 encoding antibiotic resistance, and the global reservoir for these cassettes is large (22,  
27). Our data support the presence of at least one class 1 integron in *A. salmonicida*  
380 AS03. The class 1 integron genes *sul1*, *qacEΔ1* and *aadA7* were sequenced from this  
strain. The aminoglycoside resistance gene that encodes for adenylyltransferase,  
382 *aadA7*, was first found in an *E. coli* strain in which the *mer* operon was located on the  
mobile genetic element Tn21(27). *aadA7* has never been reported in *A. salmonicida* (2,  
384 12, 23, 27, 31). The tetracycline resistance gene, Tet A, has previously been found in *A.*  
*salmonicida*, purportedly as a result of the widespread use of tetracycline in veterinary  
386 medicine (22). This tetracycline resistance determinant was probed for in this study as  
this gene has been found in many *A. salmonicida* strains and, along with Tet E, is the  
388 most common tetracycline resistance determinant isolated from aquatic bacteria(21, 22)

The transposase genes *tnpA* and *tnpR* amplified and sequenced in this study show  
390 a higher degree of similarity to these genes found on Tn1696 than those carried on Tn21.  
This transposition module has not previously been reported in *A. salmonicida*. However,  
392 the integron In4, which is typically associated with Tn1696, has been detected in *A.*  
*salmonicida* (36). It is possible that the *mer* operon and associated antibiotic resistance  
394 genes in *A. salmonicida* AS03 are under the control of the Tn1696 transposition module  
as part of a composite transposon.

396 The conjugation data strongly indicate that the antibiotic resistance determinants  
in *A. salmonicida* AS03, with the exception of those encoding resistance to  
398 fluoroquinolones, the *mer* operon and Tn1696 transposition module were located together  
on a mobile genetic element capable of conjugative gene transfer. Supporting this  
400 observation is the fact that, with the exception of fluoroquinolone resistance, the  
antibiotic resistance determinants previously reported in *A. salmonicida* are encoded on  
402 plasmids, in which resistance genes are often closely linked on mobile genetic  
elements(22, 24, 27, 30, 43). The plasmid pRMH760 carries many of the same antibiotic  
404 resistance determinants in addition to the *mer* operon and Tn1696 transposition module  
and these genes were similarly transferred to *coli* DH5 $\alpha$  (32). Multiple conjugation  
406 experiments were conducted under different mating mixture conditions of  
chloramphenicol and mercury exposure. However, all transconjugants exhibited identical  
408 antibiotic and mercury resistance phenotypes (Table 2). These data strongly suggest that  
these genes are located on a single mobile genetic element as opposed to multiple genetic  
410 elements.

Because these resistance genes are transferable via conjugation, it is likely that the  
412 mobile genetic element containing the *mer* operon, the Tn1696 transposition module, and  
the genes encoding the transferred antibiotic resistances is either a conjugative R plasmid  
414 or conjugative transposon. Plasmids were not detected in either the *E. coli* DH5 $\alpha$   
recipient or the transconjugant that had phenotypically similar mercury and antibiotic  
416 resistance capabilities as the donor *A. salmonicida* AS03. The resistance genes could  
either be present on a low-copy number plasmid that was not detectable by standard  
418 experimental protocols or that they were located on a conjugative transposon that, after  
conjugative transfer, integrated in the recipient's chromosome. Previous studies  
420 involving conjugative transposons in various conjugation systems have reported the  
inability to detect plasmids in the transconjugants (19) Due to the high levels of  
422 antimicrobial and mercury resistance in the transconjugant, it would be expected that  
plasmids containing these resistance determinants to be in high copy number. Therefore,  
424 our data suggest that the mercury and antibiotic resistance genes, as well as the  
transposition module of Tn1696, were encoded on a conjugative transposon. Conjugative  
426 transposons are known to carry both mercury and antibiotic resistance genes. The  
conjugative transposon SXT, characterized in *Vibrio cholerae*, encodes resistance to  
428 sulfamethoxazole, trimethoprim, chloramphenicol, and streptomycin (20). In this study,  
all of antibiotic genes encoded on SXT were observed to be transferable in our  
430 conjugation system (Table 2).

Furthermore, it was demonstrated that low-level Hg and chloramphenicol can  
432 stimulate conjugation as evidenced by increased the transfer frequencies (Table 3). Due  
to atmospheric deposition, mercury levels in the environment are rising and its presence

434 in the landscape is a threat to wildlife as well as human health in the Northeastern United  
States and Atlantic Canada (4, 14, 26, 39). High mercury levels have been documented in  
436 freshwater and marine fish in Atlantic Canada and the United States and fish  
consumption advisories have been issued in all five Canadian Provinces as well as thirty-  
438 nine states in the U.S (7, 8, 13, 14). Our study provides evidence that environmental  
mercury might have provided the direct selection pressure necessary to facilitate the  
440 acquisition of broad-spectrum Hg resistance, as well as the indirect selection pressure that  
facilitated multiple-drug resistance by *A. salmonicida* AS03. Due to Hg pollution, bodies  
442 of water contain low-levels of Hg. Not only can this mercury continually select for the  
presence of plasmids or transposons with mercury resistance genes but this mercury can  
444 stimulate the transfer of these resistance genes between bacterial species. In addition,  
antibiotic resistance genes were transferred to *E. coli* DH5 $\alpha$  along with mercury  
446 resistance genes in the complete absence of antibiotics and in the presence of low-level  
mercury in the mating mixtures. This demonstrates that mercury can in fact directly select  
448 for the transfer of mercury resistance genes and indirectly select for the horizontal  
transfer of antibiotic resistance genes in a system in which the antibiotic and mercury  
450 resistance genes are both located on mobile genetic elements. This potentially pathogenic  
aeromonad is significant not only because it is resistant to many antibiotics but because  
452 its antibiotic and mercury resistances are located on mobile genetic elements, which may  
include conjugative R-plasmids or conjugative transposons, and can disseminate these  
454 genes to other organisms by horizontal gene transfer. This genetic element can easily be  
transferred to other bacterial strains, such as salmonid commensal bacteria or other  
456 pathogenic hatchery bacteria when low-level Hg acts as a selection agent.

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