

# Microarray-based Detection of Antibiotic Resistance Genes in *Salmonella*

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**Abstract** In the presented study, 143 *Salmonella* isolates belonging to 26 different serovars were screened for the presence of antibiotic resistance genes by microarray analysis. The microarray contained a total of 223 oligonucleotides representing genes encoding for resistance to the following antibiotic classes: aminoglycoside,  $\beta$ -lactam, chloramphenicol, MLS, sulfonamide, tetracycline, trimethoprim, and vancomycin. To a large extent, the microarray data were consistent to the general findings concerning antibiotic resistance in *Salmonella*. Most of the analyzed isolates, harbored three or more resistance genes with the highest numbers found in isolates belonging to the *Salmonella* serovars Typhimurium, Paratyphi B var. Java, Bredeney, Saint Paul and Heidelberg and the only Give isolate investigated.

**Keywords** Antibiotic Resistance · Microarray · Oligonucleotides · *Salmonella* · Serovars · Typhimurium

## Introduction

The intensive use of antimicrobial agents in both animal husbandry and public health has resulted in the drastic increase in antibiotic-resistant bacteria (WHO 1997). Drug resistance is becoming a worldwide problem, which is also demonstrated by the emergence of methicillin-resistant *Staphylococcus aureus* (MRSA), multidrug resistance, and extended spectrum beta-lactamase (ESBL) producing bacteria (de Neeling et al. 1998; WHO 2005; Stürenburg and

Mack 2003). To combat this problem, it will be necessary to increase knowledge on the mechanisms, diversity, and distribution underlying the observed antibiotic resistance (AR) characteristics (Aarts et al. 2006). Molecular tools rather than phenotypic methods can provide this kind of information. Furthermore, molecular research will give a more precise insight on reservoirs of antibiotic resistance (AR) genes and will support the assessment of the dissemination risk of these determinants by horizontal gene transfer. As a result of the ongoing research on resistance, the number of described AR genes and AR-related mutations has increased enormously. Consequently, multi-detection and screening methods will be necessary, rather than single-plex assays, to identify the gene(s) or mutation (s) responsible for the resistance phenotype. A promising tool is the microarray, which allows the detection of a large set of DNA targets simultaneously. This multitarget technique has already proven to be a powerful and rapid method to screen for the presence of multiple antibiotic resistance genes (Aarts et al. 2006 and Holzman 2003). Initially, microarrays were restricted to detect particular classes of AR genes, for example tetracycline or erythromycin resistance determinants (Call et al. 2003 and Volokhov et al. 2003). However, the increase in multidrug-resistant bacteria, the fact that over 500 AR genes have been described, excluding mutation-based resistance, and new resistance determinant are still being identified, require microarrays for broad screening purposes (Frye et al. 2006 and Perreten et al. 2005). Consequently, the previously described thematic AR genes microarray (van Hoek et al. 2005) was expanded with a considerable number of oligonucleotides and now contains 223 oligonucleotides representing over 430 AR genes. These genes confer resistance determinants belonging to the following antibiotic classes: aminoglycoside,  $\beta$ -lactam, chloramphenicol, macro-

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lide lincosamide and streptogramin (MLS), sulfonamide, tetracycline, trimethoprim, and vancomycin. They are found in both gram-positive and gram-negative bacteria. This expanded microarray was used for the screening of a large set of *Salmonella* strains belonging to 26 different serovars including *Salmonella* Typhimurium, represented by various different phage types such as DT104.

## Materials and Methods

### *Salmonella* Isolates

In total, 143 *Salmonella* isolates were included in this study. The serovars and number of isolates were *S. Agona* (11), *S. Anatum* (4), *S. Bareilly* (1), *S. Blockley* (4), *S. Brandenburg* (1), *S. Bredeney* (3), *S. Derby* (4), *S. Dublin* (1), *S. Enteritidis* (8), *S. Give* (1), *S. Hadar* (10), *S. Heidelberg* (7), *S. enterica* subsp. *houtenae* 48:g.z51:- (1), *S. Infantis* (5), *S. Kentucky* (1), *S. Livingstone* (2), *S. London* (3), *S. Mandaka* (4), *S. Newport* (2), *S. Paratyphi B* var. *Java* (10), *S. Rough* (1), *S. Saint Paul* (3), *S. Senftenberg* (2), *S. Tennessee* (1), *S. Typhimurium* (44), and *S. Virchow* (9). Isolates were obtained from various sources in Europe, i.e., BFR, Germany; CIDC-Lelystad, The Netherlands; FRIKI, The Netherlands; INRA, France; Institut Pasteur, France; ISS, Italy; National Institute of Public Health and the Environment (RIVM), The Netherlands. Part of the strains investigated belonged to the RIKILT collection. All *Salmonella* isolates were grown overnight in Brain Heart Infusion broth (Merck, Haarlem, The Netherlands) at 37° C.

### DNA Isolation and Labeling

DNA was isolated from pure cultures using the Wizard® genomic DNA purification kit (Promega Benelux b.v., Leiden, The Netherlands) according to the manufacturer's manual. A total of 4 µg of isolated DNA was fluorescently labeled according to van Hoek et al. (2005).

### Oligonucleotide Design and Microarray Fabrication

Oligonucleotide probes were designed according to van Hoek et al. (2005) representing genes belonging to the following antibiotic resistance classes: aminoglycosides, β-lactams, chloramphenicol, MLS, sulfonamides, tetracyclines, trimetoprim, and vancomycin. Furthermore, oligonucleotides were included, identifying genes from the *mar* operon, integron-specific integrases, the left border of the first characterized *Salmonella* Genomic Island (SGI1) and *Salmonella* species-specific sequences. Several polymerase chain reaction (PCR) primers were designed to verify the hybridization results obtained by microanalysis.

**Fig. 1** Thematic AR genes oligonucleotide microarray: (a) the microarray layout, (b) examples of hybridization results. On the surface of one slide, two microarrays were spotted. The microarray consists of two grids, each of which contains three subgrids (i.e., the oligonucleotides are spotted in triplicate except the *Salmonella* specific spots which are six times present on a microarray). The gray marked oligonucleotide names in the subgrids are the *Salmonella*-specific spots. These spots are encircled in the hybridization examples (b). The black marked names were not included in the analysis of the *Salmonella* strains. They represent oligonucleotides specific for other bacterial species or which do not work properly (nonspecific, high-background signals)

From the GenBank public DNA database, genes of interest and homologous sequences were retrieved. Related sequences were aligned using the ClustalX Software. Potentially gene-specific primers and oligonucleotides were designed based on the regions within the open reading frame with the highest level of homology using the Generunner software package (Hasting Software, Inc. Hastings-on-Hudson, NY). The main criteria designing the oligonucleotides were the absence of secondary structures at the hybridization temperature that was used, very long GC stretches (>10) and repeats. To check for potential interference with other sequences, Blast searches were performed with the selected primers and oligonucleotides against the public DNA databases.

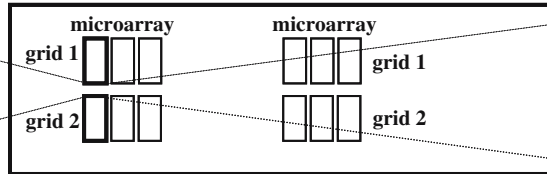
All oligonucleotides were modified with a 5' C6-amine linker to enhance binding to the microarray glass slides. The oligonucleotides were manufactured by Biolegio (Nijmegen, The Netherlands) and spotted at a concentration of 50 µM in 5×SSC on SCHOTT Nexterion E slides (Isogen Life Science, IJsselstein, The Netherlands) using a MicroGrid II Microarrayer (BioRobotics Ltd., Cambridge, UK). After spotting, the microarrays were washed and blocked according to the manufacturer's instructions (SCHOTT JENA<sup>er</sup> GLAS GmbH, Jena, Germany). The microarrays were prehybridized in hybridization buffer (5×SSC, 0.2% sodium dodecyl sulfate (SDS), 5×Denhardt's solution, 50% (v/v) formamide, 0.2 mg/ml denatured herring sperm DNA) in a humid hybridization chamber at 42° C for at least 4 h. Subsequently, the slides were rinsed as described by van Hoek et al. (2005).

### Microarray Hybridization and Analysis

Hybridization conditions of the microarray with fluorescently labeled total DNA, scanning, and analysis of the resulting images were performed according to Franssen-van Hal et al. (2002) and van Hoek et al. (2005) All hybridization signals (determined as arbitrary units by the ArrayVision™ Software) were first corrected for spot area and background signal surrounding each individual spot.

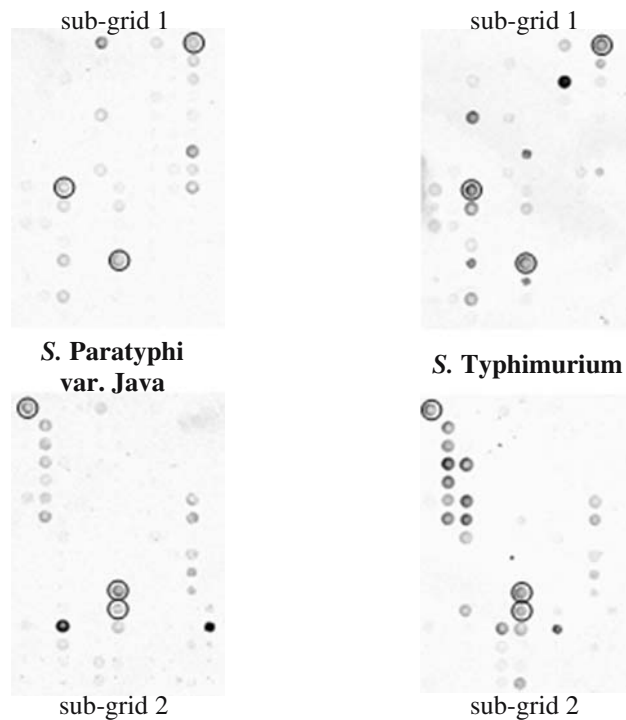
**a**

blaACC 50mer_I	blaFOX 50mer	blaPER 50mer_II	dfrA1 60mer	dfrA17 50mer	tetA 60mer	tetQ 50mer	Sspp 60mer
blaACC 50mer_II	blaIMP 50mer	blaPSE 50mer	dfrA2 60mer	dfrA19 60mer	tetB 60mer	tetS 50mer	Sspp 50mer
blaACC 50mer_III	blaKPC 50mer	blaPSE 60mer	dfrA3 50mer_I	dfrA 50mer_I	tetC 50mer	tetT 50mer	<i>Streptococcus</i>
blaACT 48mer	blaMIR 47mer	blaROB 50mer_I	dfrA3 50mer_II	dfrB 50mer_I	tetD 60mer	tetU 50mer	sul1 60mer_I
blaCARB 50mer_I	blaMOR 50mer_I	blaTEM-60mer	dfrA5 50mer	dfrB 50mer_II	tetE 60mer	tetV 60mer	sul1 50mer
blaCARB 50mer_II	blaOXA 50mer_I	blaVIM 50mer_I	dfrA6 50mer	dfrD 50mer	tetG 60mer	tetW 50mer	sul1 60mer_II
blaCARB 50mer_III	blaOXA 50mer_II	blaVIM 50mer_II	dfrA7 60mer	dfrA12 60mer_I	tetH 60mer	tetX 50mer	sul2 50mer
blaCARB 50mer_IV	blaOXA 50mer_III	blaVIM 50mer_III	dfrA8 50mer	<i>E. coli</i>	tetI 50mer	tetY 50mer	
blaCMY 50mer_I	blaOXA 50mer_IV	sipB/C 60-mer	dfrA9 50mer	<i>E. coli</i>	tetJ 50mer	tetZ 50mer	sulA 50mer
blaCMY 50mer_II	blaOXA 50mer_V	marA 50mer	dfrA10 60mer	<i>Lactobacillus</i>	tetL 50mer	tet30 60mer	<i>Enterococcus</i>
blaCTX-M 50mer_I	blaOXA 50mer_VI	marB 50mer	dfrA11 60mer	<i>Bifidobacterium</i>	tetK 50mer	tet31 60mer	<i>Staphylococcus</i>
blaCTX-M 50mer_II	blaOXA 50mer_VII	<i>E. coli</i>	dfrA12 60mer	<i>Lactococcus lactis</i>	tetM 50mer	tet32 60mer	tet36 60mer
blaCTX-M 50mer_III	blaOXA 50mer_VIII	marC 50mer	dfrA13 60mer	invA 60mer_II	tetO 50mer_I	tet33 60mer	tet37 60mer
blaDHA 50mer_I	blaOXA 50mer_IX	<i>E. coli</i>	dfrA14 60mer	Stmm 50mer	tetO 50mer_II	tet34 60mer	otrA 50mer
blaDHA 50mer_II	blaPER 50mer_I	<i>E. coli</i>	dfrA15 50mer		tetA(P) 50mer	tet34 50mer	otrB 60mer
			dfrA16 60mer		tetB(P) 50mer	tet35 60mer	otrB 50mer
							otrB 60mer
							tetW 60mer



sipB/C 60-mer	aacC 50mer_VII	aphA 50mer_IV	catB 50mer_I	vanA 60mer_I	ereA 50mer_I	msr 50mer_II	mefE 60-mer
aac6-aph2 50mer	aadA1 50mer	aphA 50mer_V	catB 50mer_II	vanB 50mer	ereA 50mer_II	vat 50mer_I	vat 60mer_I
aacA 50mer_I	aadA1 60mer_I	aphA 50mer_VI	catB 50mer_III	vanC1 50mer	ereB 50mer	vat 50mer_II	
aacA 50mer_II	aadA2 60mer_I	strA 50mer	catB 50mer_IV	vanC2/C3 50mer	ermA 50mer	vat 50mer_III	
aacA 50mer_III	aadA2 60mer_II	strB 50mer	catB 50mer_V	vanD 50mer		vat 50mer_IV	
aacA 50mer_IV	aadA 60mer_I	strA 60mer	cat 50mer_I	vanE 50mer	erm 50mer_I	vat 50mer_V	<i>Bifidobacterium</i>
aacA 50mer_V	aadA1 60mer_II	strB 60mer	cat 50mer_II	dfrA12 60mer_II	erm 50mer_II	vga 50mer_I	<i>Bifidobacterium</i>
aacA 50mer_VI	aadB 50mer	strA-strB 60mer	cat 50mer_III	vanC2/C3 60mer	erm 50mer_III	ygb 50mer_I	<i>Lactococcus lactis</i>
aacA 50mer_VII	aadD 50mer	aphA-3 60mer	cat 50mer_IV	vanA 60merII	erm 50mer_IV	ygb 50mer_II	<i>Lactobacillus</i>
aacA 50mer_VIII	aadE 50mer	aadE 60merI	cat 50mer_V	vanB 60mer	erm 50mer_V	ermA 60mer_II	<i>Bifidobacterium</i>
aacC 50mer_I	aph(2') 50mer_I	aadE 60merII	cat 50mer_VI	invA 60mer_I	erm 50mer_VI	ermB 60mer_II	<i>Streptococcus</i>
aacC 50mer_II	aph(2') 50mer_II	int11 59mer	cat 50mer_VII	Sspp 60mer		ermC 60mer_II	<i>Lactococcus lactis</i>
aacC 50mer_III	aph(2') 50mer_III	int12 60mer	cmIA 50-mer	Sspp 50mer	mefE 50mer	ermA 60mer_I	<i>Bifidobacterium</i>
aacC 50mer_IV	aphA 50mer_I	int13 60mer	cmIB 50-mer		mph 50mer_I	ermB 60mer_I	<i>Streptococcus</i>
aacC 50mer_V	aphA 50mer_II	int14 60mer	flor 50mer	Seis 60mer	mph 50mer_II	ermC 60mer_I	<i>Enterococcus</i>
aacC 50mer_VI	aphA 50mer_III	SGII LB 50mer	flor 60mer	Stmm 60mer	msr 50mer_I	mefA 59mer	<i>Staphylococcus</i>
							aphE 60mer
							sat 60mer_I
							sat 60mer_II
							sat 60mer_III

**b**



Only hybridization signals with a signal-to-noise ratio higher than three ( $S/N > 3$ ) were taken into account for further analysis in Excel. With the Excel software the fluorescent signals were corrected for labeling and hybridization conditions. The average hybridization signals obtained with the *Salmonella* specific oligonucleotides (invA 60mer\_I, invA 60mer\_II, sipB/C 60mer, and Ssp 60mer) were used for this correction factor. As a rule, corrected hybridization signals with a ratio of 0.5 or more indicated the presence of a particular antibiotic resistance gene.

## PCR

The PCR tests to verify the microarray hybridization results were performed in a total volume of 50  $\mu$ l containing approximately 40 ng of bacterial DNA, 10 pmol of each primer, 1 $\times$ PCR buffer (Invitrogen BV, Breda, The Netherlands), 3 mM MgCl<sub>2</sub>, 0.2 mM of each deoxyribonucleoside triphosphate (dNTP), 1 U *Taq* DNA polymerase recombinant (Invitrogen BV, Breda, The Netherlands). The following PCR program was used: 95° C for 3 min; 35 cycles of 95° C for 30 s, 55° C for 30 s, 72° C for 30 s; 72° C for 10 min. The PCR products were analyzed by electrophoresis on a 2% agarose gel.

## Results

In a previous study, the successful application of microarray screening for the presence of antibiotic resistance genes in *Salmonella* was described (van Hoek et al. 2005). In the present paper, both the number of oligonucleotides on the microarray and the amount of *Salmonella* strains investigated were increased. The oligonucleotides have been designed in such a way that they are gene-specific and not species-specific. Consequently, the microarray is able to detect and identify AR genes in various species. The microarray now contains 223 oligonucleotide probes identifying over 430 AR genes encoding for resistance against aminoglycosides (#43),  $\beta$ -lactams (#39), chloramphenicol (#16), MLS (#32), sulfonamides (#5), tetracyclines (#42), trimethoprim (#23), and vancomycin (#9). Furthermore, oligonucleotides representing the multiple antibiotic resistance (mar) locus, different integron classes, and the *Salmonella* genomic island 1 (SGI1) were also added to the microarray (Fig. 1). The oligonucleotide names, lengths, sequences, GC %, and represented gene(s) are given in Table 1. Oligonucleotides already described in Mättö et al. (2007) and van Hoek et al. (2005) are indicated. Phenotypic and genotypic well-defined control strains were used to check whether the designed oligonucleotides could detect the corresponding gene(s) (indicated with an asterisk

in Table 1). Furthermore, for confirmation purposes part of the microarray data was verified by PCR using gene-specific assays (Table 2).

A total of 143 *Salmonella* isolates encompassing 26 different serovars were screened for the presence of AR genes using the oligonucleotide microarray. For serovar Typhimurium, a total number of 44 isolates were investigated, which belonged to 14 different phage types, including the multiresistant phage type DT104. An overview of the microarray screening results is shown in Table 3. The data are presented per serovar and if applicable (*S. Typhimurium*) by phage type. The amount of strains investigated is indicated and the number of isolates showing a hybridization signal with a certain oligonucleotide. Only results from oligonucleotides that gave a hybridization response with at least one of the *Salmonella* isolates are included in Table 3. As a result of the lack of control strains for all oligonucleotides present on the microarray, some results might be less reliable and this could lead to an overestimation of the number of AR genes present in an isolate. In contrast, the number of detected AR genes could also be underestimated by not knowing whether the oligonucleotides are actually able to detect the corresponding AR gene(s). However, various experiments with DNA from control strains available resulted in hybridization signals with oligonucleotides representing the AR gene(s) present in these bacteria (data not shown), indicating that the chosen criteria for the design of the oligonucleotides were appropriate.

Some antibiotic resistance genes were represented by two or three oligonucleotides that differed in length (see Table 1). In the case of *aadA1*, *tet(A)*, and *sul1* the 50-mer oligonucleotide resulted in lower numbers of positive strains in comparison to the corresponding 60-mer. However, for other genes, such as *dfrA14* and *floR*, the 50- and 60-mer oligonucleotides gave similar results. As some of the AR genes were only represented by a 50-mer oligonucleotide, this could have influenced the number of positive strains.

A total of 51.4% of the microarray data was checked by PCR (approximately 2,500 PCR tests were performed). More than 95% of the PCR results were in agreement with the microarray data (not shown). Similar percentages were found by Malorny et al. (2007) with an oligonucleotide multiprobe microarray developed for the molecular characterization of *Salmonella*. In those cases of discordant results, the observed hybridization values were often very low and close to the cutoff values between positive and negative hybridization signals.

In general, multiple AR genes were detected per isolate, especially in those belonging to the serovars Typhimurium, Paratyphi B var. Java, Bredeney, Saint Paul, and Heidelberg, and the only Give isolate investigated. Common AR genes found in the different serovars were *strA* conferring

**Table 1** Microarray oligonucleotides

Oligonucleotide name <sup>a</sup>	Sequence (5'–3')	GC %	Gene(s) represented <sup>d</sup>
<b>Aminoglycosides</b>			
aac6-aph2 50mer <sup>b</sup>	gattgttattaatggaatatagatatgatgataatgccacaatgttaa	24.0	<i>aac(6')-aph(2'')</i> *, <i>aacA/aphD</i>
aacA 50mer_I	ttgtcagaccagattatcaaaataaaggattggcaagatcctgcttaag	36.0	<i>aacA1</i>
aacA 50mer_II	cacgccgacactgcygacgtacaggaacagactgtgccaagcgtttta	52.0	<i>aacA4</i> , <i>aac(6')-Ib</i>
aacA 50mer_III	tgctgtagcaccgacggagaagcactagggttgcccagctttcgatcc	58.0	<i>aacA5</i> , <i>aacA7</i>
aacA 50mer_IV <sup>b</sup>	ccggccgcacatcggtgagtggtgggtggYgacgaagagcgaccgactc	66.0	<i>aac(6')-II</i>
aacA 50mer_V	atcggttgcttgacggaaactccatcgcgttcgcacagctgtacgtg	54.0	<i>aac(6')-IIb</i>
aacA 50mer_VI	agttaaacaagggtgggtacaaagctcgtacgctcgtcgtggaactc	50.0	<i>aac(6')-IIc</i>
aacA 50mer_VII	gcatcatttattcgatggcagacgggtggcgttgcttgcggatgc	56.0	<i>aac(6')-Iy</i>
aacA 50mer_VIII	tatgcttgggaatatgctggtatgataaactgattggMtgaccgatta	40.0	<i>aac(6')-Ib</i> , <i>aac(6')-II</i> , <i>aac(6')-Iq</i>
aacC 50mer_I	caaagttaggtggctcaagatggcgcacatcgcacatgtaggctcggc	52.0	<i>aac(3)-Ia</i> , <i>aacC1</i>
aacC 50mer_II	catcgaaggctaggatgcacactccctgattagccactgaagcgtgt	56.0	<i>aac(3)-Ib</i>
aacC 50mer_III <sup>c</sup>	agctgaaacgctgacggagcctcacgaactcggcagccttggggaaag	60.0	<i>aac(3)-IIa</i> , <i>aacC2</i>
aacC 50mer_IV	ctggtggcaatagaaggatacgtgctgatgctggcgcggcgtggatac	58.0	<i>aac(3)-IIIa</i> , <i>aacC3</i>
aacC 50mer_V	gccgttcgcgacactacagccacgcaatggcgcgatgatcggaggttcg	62.0	<i>aac(3)-IIIb</i>
aacC 50mer_VI	cgctgtgggagagcggggaaccctgatggtgactgcggctggaacgacg	72.0	<i>aac(3)-VII</i>
aacC 50mer_VII	ggacctcagtgaggcggactacaataatggctgctcctccagaagcgtgc	64.0	<i>aacC9</i>
aadA 60mer_I	agccatacagtgatattgattgctgttactgtgctgcacggctcgtgatgactgtcc		<i>aadA6</i> , <i>aadA10</i> , <i>aadA11</i> , <i>aadA13</i>
aadA1 50mer <sup>c</sup>	ggcctgaagccacacagtgatattgattgctgttaccggtgaccgtaag	50.0	<i>aadA1*</i>
aadA1 60mer_I <sup>c</sup>	ggcctgaagccacacagtgatattgattgctgttaccggtgaccgtaag	48.3	
aadA1 60mer_II	cccgtcactactgaagctagacagccttactgtgacaagaagaagatcgtggcctc	49.2	
aadA2 60mer_I <sup>c</sup>	cttgaccgggtcctgaacaggatctattcagggcgtgagggaaccttgaagctatg	51.7	<i>aadA2*</i>
aadA2 60mer_II	ctaagcaagcttactgggacaaaagaagatcacttggcctcacgagatcacttgg	48.3	
aadB 50mer	gtccgtgtaacagctgggagcgcgatcatctgggattactttactatgc	53.3	<i>aadB*</i>
aadD 50mer	aggcaaatggcgtaatattcgtgtgcaaggaccgacaacatttctacat	49.2	<i>aadD</i>
aadE 50mer <sup>b</sup>	cgtttatactaatgatgactggcttaataatftgggaatataataatg	28.0	<i>aadE*</i>
aadE 60mer_I <sup>b</sup>	caaggagatgatgattgctgcaatgattttggaatgtaaaccttattgttaaaag	33.3	
aadE 60mer_II <sup>b</sup>	taaagtgtagcaagaactataaagtattgaaaggtatataatccgaggatttggggag	35.0	
aph(2') 50mer_I	ggatgcccttgcataatgatgaagcagctttttgaaagagtacattcca	42.0	<i>aph(2')-Ib</i>
aph(2') 50mer_II	gaaggcttaaggcgaaggatcaggactgatttctgaagggttagagct	50.0	<i>aph(2')-Ic</i>
aph(2') 50mer_III	ccggaggtggtttttacaggaatgccaacagaaactgacaaatgtcttt	44.0	<i>aph(2')-Id</i>
aphA 50mer_I	tctatcgtattggtggaagcccaatgcccagaggtgtttctgtaaacat	44.0	<i>aphA1*</i> , <i>aphA1-IAB</i>
aphA 50mer_II	caggatcctctgcatctcactctgctcctgcccagaaagatccatcat	50.0	<i>aph(3')-IIa</i> , <i>aphA2</i> , <i>nptII*</i>
aphA 50mer_III <sup>b</sup>	gatctgcccgatggtgattgcaaaaactgggaagaagacactccatttaa	46.0	<i>aphA3*</i>
aphA 50mer_IV	attgcttctctataaaggagcactcaatctgttaaatcaattgcta	34.0	<i>aphA6</i>
aphA 50mer_V	catgagtgagttaaaggggaaacacatagattgctttattgatccaa	38.0	<i>aphA7</i>
aphA 50mer_VI	ctgatttcttgcggcctgcatgagatcccaacgattgaatgcccttc	52.0	<i>aph(3')-Id</i>
aphA 60mer_I <sup>b</sup>	tttctcggaaagatgaagatgaacaaagccctgaaaagattatcgagctgatcgg	43.3	<i>aphA3*</i>
aphE 60mer <sup>b</sup>	acacggctgctgccacgggtgatctctgctcctcccaatcgtcctccatccggagacc	63.3	<i>aphE</i>
sat 60mer_I	gtgaaggttcgatggtgcacatcaccgaccaaggcttgaactatcaccagaagtgtga	46.7	<i>sat2<sup>e</sup></i>
sat 60mer_II <sup>b</sup>	aagcgatgccgactgttctcagcttgcggctgcttcttctcaggtcacagctga	55.0	<i>sat3</i>
sat 60mer_III <sup>b</sup>	cccagcgaaccattgaggtgataggttaagattataccgaggtatgaaaacgagaattgg	43.3	<i>sat4</i>
strA 50mer <sup>c</sup>	acggcggcttggatggtgtcccgcaatggcctgcaatcccgactcttacc	58.0	<i>strA*</i>
strB 50mer	cgggtgctcggctgtgagaacaactcgtatggtgctcgaatatgccgggga	54.0	<i>strB*</i>
<b>β-lactams</b>			
blaACC 50mer_I	gcctacagctattatgccggaagatattaaaaataccacacagctgatg	40.0	<i>bla<sub>ACC-1</sub>*</i>
blaACC 50mer_II	tgagcaaacctcctctctattagcatgaatcaaacctcactgaaagg	44.0	<i>bla<sub>ACC-2</sub></i>
blaACC 50mer_III	atatcggttactcaagtacggcaaaaactcactcaggatctgatgtggaa	42.0	<i>bla<sub>ACC-3</sub></i>
blaACT 48mer	gaagccggactcctcagataattcactcagaaaggccttaccct	52.1	<i>bla<sub>ACT-1</sub></i> , <i>bla<sub>ACT-2</sub></i>
blaCARB 50mer_I	tctcccgaatagaaaagcaagtaggacaagaataacgctcgtgatgac	44.0	<i>bla<sub>CARB-4</sub></i>
blaCARB 50mer_II	aaattggtgagcaaatagcgaagacagtaattatggagaatagccgtaac	38.0	<i>bla<sub>CARB-5</sub></i> , <i>bla<sub>CARB-8</sub></i>
blaCARB 50mer_III	caactcctaaggcaatagccagcagcttaaatcaattatttgggttcc	38.0	<i>bla<sub>CARB-6</sub></i> , <i>bla<sub>CARB-7</sub></i> , <i>bla<sub>CARB-9</sub></i>
blaCARB 50mer_IV	tctagatcgtgctgagcctgagctcaatgaaggtaaactcgggtatttga	46.0	<i>bla<sub>CARB-7</sub></i> , <i>bla<sub>CARB-9</sub></i>
blaCMY 50mer_I	atcaagaccagctcggcggatctgctgcttggtaagccaacatcgg	56.0	<i>bla<sub>CMY-1</sub></i> , <i>bla<sub>CMY-10</sub></i> , <i>bla<sub>CMY-11</sub></i>
blaCMY 50mer_II	ggcgagcagcctgaagcagcatttggcccagttgatggagcagaccctgc	62.0	<i>bla<sub>CMY-8</sub></i> , <i>bla<sub>CMY-9</sub></i> , <i>bla<sub>CMY-19</sub></i> , <i>bla<sub>MOX-1</sub></i>
blaCTX-M 50mer_I	ggcttaccagcgtcgtggactgcaggtgataagaccggcagcggcgacta	62.0	<i>bla<sub>CTX-M-9</sub> group<sup>f</sup></i>

Table 1 (continued)

Oligonucleotide name <sup>a</sup>	Sequence (5'–3')	GC %	Gene(s) represented <sup>d</sup>
blaCTX-M 50mer_II	gctaaatcagcgcgttgaatcaagaagcgcactggttaactacaatc	44.0	<i>bla</i> <sub>CTX-M-2</sub> group <sup>g</sup>
baCTX-M 50mer_III	gctgatggcagcgcaaccgtcacgctgtttaggaagtgtccgctgt	60.0	<i>bla</i> <sub>CTX-M-1</sub> group <sup>h</sup>
blaDHA 50mer_I	agggtccggatgctgtaaaaaaccgtgctggatctgtgaattctatcag	52.0	<i>bla</i> <sub>DHA-1</sub> , <i>bla</i> <sub>DHA-2</sub> , <i>bla</i> <sub>MOR</sub>
blaDHA 50mer_II	atgatcattaaccgctgacccaacgaggtgcactgcagccgaccgggt	60.0	
blaFOX 50mer	atagtctggccagccattgagMaactgatgagccagaccctgctgccc	57.0	<i>bla</i> <sub>FOX</sub>
blaIMP 50mer <sup>c</sup>	ctctcatttcatagYgacagcacRggBggaaatagagtgttaattctc	46.3	<i>bla</i> <sub>IMP</sub> <sup>*</sup>
blaKPC 50mer	gcgccgctgacggaaagcttcaaaaaactgacactgggctctgactgg	58.0	<i>bla</i> <sub>KPC</sub>
blaMIR 47mer	tgccaaaaccgtcgtcggaggcagtgataacaagggtgctggtggcac	59.6	<i>bla</i> <sub>MIR</sub> , <i>bla</i> <sub>ZEG-1</sub>
blaMOR 50mer_I	ggaaggggatcacactgctggatctggtacttacaccgagggcggtg	64.0	<i>bla</i> <sub>MOR</sub> , <i>bla</i> <sub>DHA-1</sub> , <i>bla</i> <sub>DHA-2</sub>
blaOXA 50mer_I	taccaatgacttagctgctcatcaaaaggaatatttccagcatcaacat	40.0	<i>bla</i> <sub>OXA-10</sub> group <sup>i</sup>
blaOXA 50mer_II	tcaacattcaaaattcctaagctctaatagctctgaaaccggcgccat	40.0	<i>bla</i> <sub>OXA-5</sub>
blaOXA 50mer_III	ccattaaagggtaccctattcaagaggtagagtttttcccaattagc	40.0	<i>bla</i> <sub>OXA-23</sub> , <i>bla</i> <sub>OXA-27</sub> , <i>bla</i> <sub>OXA-49</sub> , <i>bla</i> <sub>OXA-73</sub>
blaOXA 50mer_IV	cgatgacctgacataaccgattacctttaaattagaactcaagaag	36.0	<i>bla</i> <sub>OXA-24</sub> , <i>bla</i> <sub>OXA-25</sub> , <i>bla</i> <sub>OXA-26</sub> , <i>bla</i> <sub>OXA-33</sub> , <i>bla</i> <sub>OXA-40</sub> , <i>bla</i> <sub>OXA-72</sub>
blaOXA 50mer_V	ttcgtgatgagttccagattttgatgggacggcgttaacaggggcttt	48.0	<i>bla</i> <sub>OXA-02</sub> , <i>bla</i> <sub>OXA-15</sub> , <i>bla</i> <sub>OXA-32</sub> , <i>bla</i> <sub>OXA-34</sub> , <i>bla</i> <sub>OXA-102</sub>
blaOXA 50mer_VI	gagctatttgcaaaagagatcggtgaagacaaggctcgacgctattgaa	44.0	<i>bla</i> <sub>OXA-03</sub> , <i>bla</i> <sub>OXA-21</sub>
blaOXA 50mer_VII	tgataatccgattctagcactgcttttctcagctgttgactgtctc	40.0	<i>bla</i> <sub>OXA-20</sub> , <i>bla</i> <sub>OXA-37</sub>
blaOXA 50mer_VIII	ggaacagcaatcatacacaagaacgctggatgcaatttctgtgtttgg	42.0	<i>bla</i> <sub>OXA-01</sub> , <i>bla</i> <sub>OXA-04</sub> , <i>bla</i> <sub>OXA-30</sub> <sup>*</sup> , <i>bla</i> <sub>OXA-31</sub> , <i>bla</i> <sub>OXA-47</sub>
blaOXA 50mer_IX	acacaacaattaggtatgactcatttaagaattatgtgatcattca	30.0	<i>bla</i> <sub>OXA-29</sub>
blaOXA 50mer_X	gcccgttccacctcaagctggcgtgcccgtatggcttcgaccacg	68.0	<i>bla</i> <sub>OXA-22</sub>
blaPER 50mer_I	ttataaaagctgtagtactgctcctcagcctactgatgtattctttagt	38.0	<i>bla</i> <sub>PER-1</sub> , <i>bla</i> <sub>PER-3</sub>
blaPER 50mer_II	tctgttactgttaatcgtgctcagctattcaaaaactgctgccaat	40.0	<i>bla</i> <sub>PER-2</sub>
blaPSE 50mer <sup>c</sup>	gtcactgtaatttactacgttcagctattccggcgggatggaacattgc	48.0	<i>bla</i> <sub>PSE-1</sub> (= <i>bla</i> <sub>CARB-2</sub> ) <sup>*</sup>
blaPSE 60mer <sup>c</sup>	cactgtaatttactacgttcagctattccggcgggatggaacattgctcagg	50.0	
blaROB 50mer_I	atgacattgcccattatgtgaagcagccgctgctgttagcgcacaacg	68.0	<i>bla</i> <sub>ROB-1</sub>
blaTEM 60mer <sup>c</sup>	ctcaccagtcacagaaaagcatcttaccggtgatgacagtaagagaattatgcagtcg	45.0	<i>bla</i> <sub>TEM</sub> <sup>*</sup>
blaVIM 49mer_I	atgttaaaagtattagtagttattgtctacatgaccgctctgtca	34.7	<i>bla</i> <sub>VIM-1</sub> , <i>bla</i> <sub>VIM-4</sub> , <i>bla</i> <sub>VIM-5</sub>
blaVIM 50mer_II	ttttgagtaagtattggtctattgaccgctctatcatggtctattgcg	40.0	<i>bla</i> <sub>VIM-2</sub> , <i>bla</i> <sub>VIM-3</sub> , <i>bla</i> <sub>VIM-6</sub> , <i>bla</i> <sub>VIM-8</sub> , <i>bla</i> <sub>VIM-9</sub> , <i>bla</i> <sub>VIM-10</sub> , <i>bla</i> <sub>VIM-11</sub> , <i>bla</i> <sub>VIM-14</sub>
blaVIM 50mer_III	aattcgacgctttctggtgtatcagctcattcgtcatggcctgactg	48.0	<i>bla</i> <sub>VIM-7</sub>
<b>Chloramphenicol</b>			
cat 50mer_I	cttttagaactggtfacaatagcagcgagagtgattggttgggataag	40.0	<i>cat</i> ( <i>pC194</i> ), <i>cat-TC</i> <sup>*</sup> , <i>cat</i>
cat 50mer_II	cacctgaatatatcatcattaccctgggtgagttttgacggatttaacct	40.0	<i>catII</i>
cat 50mer_III	aacaccagaaaatcattaaatattcagcattaccctgggttaattttg	28.0	<i>catIII</i> , <i>catA3</i>
cat 50mer_IV	tatttaacaatgtgaaatgtactacagctatgactccaatatagaat	26.0	<i>catB</i>
cat 50mer_V	cagccttggactgagtgtaagtctgactttaaatcatttttagcagatt	36.0	<i>catD</i> , <i>catP</i>
cat 50mer_VI	aattacctgaggatattagaactatagcgacgcttttgaatttcatgcc	34.0	<i>catQ</i>
cat 50mer_VII	gcaagatgtggcgtgttacgggtgaaacctggcctatttccctaaagggt	50.0	<i>catA1</i> , <i>cat</i>
catB 50mer_I	gcaaaatcagtcgatcattccagcgggctgcccagacaggtataggaag	52.0	<i>catB2</i>
catB 50mer_II	gttgataagttgatcatcggtagtttctctctatcgggagtgggcctc	48.0	<i>catB3</i>
catB 50mer_III	caaaattggagacggtgcccgtgatgtagtgcctggttgacaaaag	50.0	<i>catB8</i> <sup>*</sup>
catB 50mer_IV	catgcaagaagaccagctttttcaagttcaacggacgcttcaaaaagg	46.0	<i>catB6</i>
catB 50mer_V	caaggcctcagcagtgattgataagtacattccatttttctatcagga	42.0	<i>catB9</i>
cmlA 50mer	aatggctcagctgctactccccgtaagtgcctgaactctgtgtgac	52.0	<i>cmlA</i> , <i>cmlA1</i> <sup>*</sup> , <i>cmlA4</i> , <i>cmlA5</i> , <i>cmlA6</i> , <i>cmlA7</i>
cmlB 50mer	tcactacggcttctggtctctatgcttctgcttccggcgatagcg	54.0	<i>cmlB</i>
floR 50mer <sup>c</sup>	gctgtgctgtttgcgggagcgtctgttggggatcggcgaactttacgg	60.0	<i>floR</i> <sup>*</sup>
floR 60mer	gctgtggatgctgctgttggcggagcgtctgttggggatcggcgaactttacgg	61.7	
<b>marRAB locus</b>			
marA 50mer	tccaaatggcacctgcaacggatgttaaaaaagaRaccggctcattcatt	50.0	<i>marA</i> <i>Salmonella</i> , <i>E. coli</i> , <i>Shigella</i>
marB 50mer	gggtctgttacttaccctccggatagcattgcagaacaaactttgt	50.0	<i>marB</i> <i>Salmonella</i>
marC 50mer	attattttctgcccgtggcgtgatcctgtggggatgcttaccagcttc	50.0	<i>marC</i> <i>Salmonella</i>
marR 50mer	ggacggcggcaatttggagcaatgcatcaacgaccagggcaagacc	58.0	<i>marR</i> <i>Salmonella</i>

Table 1 (continued)

Oligonucleotide name <sup>a</sup>	Sequence (5'–3')	GC %	Gene(s) represented <sup>d</sup>
<b>MLS</b>			
ereA 50mer_I	catgaaaccgcacgttgatattgactcactgttggcgtccattgatg	46.0	<i>ere(A)*</i>
ereA 50mer_II	gttcccatgggacagcatctcgcagagagggagggggattaccgtgc	62.0	<i>ere(A2)</i>
ereB 50mer	tgatataccagaaatggaggttcatactaccacaaataggagatagtc	38.0	<i>ere(B)*</i>
ermA 50mer <sup>b</sup>	gtgactaaagagcggtaaacccctctgagaatataaaagtgaattcaaac	38.0	<i>erm(A)*</i>
ermA 60mer_I <sup>b</sup>	gttctttcactaaaaaccaattccgacagcgttgaagcatgcaaatgctactaatatt	35.0	
ermA 60mer_II <sup>b</sup>	caacgagcttRgggttRctRttaatggtggaRatggatataaaaatKctYaaaaaagta	30.0	
ermB 60mer_I <sup>b</sup>	attcgtgctactttaattcacaagaatattctacagttaattcccaacaacagagg	35.0	<i>erm(B)*</i>
ermB 60mer_II <sup>b</sup>	ggattctacaagcgtacctggatattcaccgaactagggtgctctgacactcaa	46.7	
ermC 60mer_I <sup>b</sup>	ggcagaagttgataattctatattaagtagtggctcaagagaatatttcatcctaaacc	31.7	<i>erm(C)*</i>
ermC 60mer_II <sup>b</sup>	acgcaaaattgttttgatagtagctaatgagattttaactcgtggaatacgggtt	30.0	
erm 50mer_I	ccttcaatagaaactcacaaaaagtattttcagggaagcttcaaatc	34.0	<i>erm(F), erm(FS), erm(FU)</i>
erm 50mer_II	caaatagtaaatgatgataactgaaatttaccatttccagccacaatcc	30.0	<i>erm(G)</i>
erm 50mer_III	taagtcccaacaacaagcatataaaatctacggtaatatatacttatt	28.0	<i>erm(GM)</i>
erm 50mer_IV	gctaaaagggtgctaatacaaatcgttctactagcactattttaaagac*	32.0	<i>erm(GT)*, erm(T)</i>
erm 50mer_V <sup>c</sup>	aacattacattgctgttctggtgctccgaatatgcgtaaaaggagccg	48.0	<i>erm(D), erm(J), erm(K)</i>
erm 50mer_VI	agattattaagaaatattattagaagagtaaatcacaactgatgatg	22.0	<i>erm(Q)</i>
mefA 59mer <sup>b</sup>	tccttgcatctggaatgtgataggggctattattagggttattgggaattacca	39.0	<i>mef(A)*</i>
mefE 50mer <sup>c</sup>	aaggagagatgaaagaggtgtggttctgacagacaaaagcagattgt	40.0	<i>mef(E)*</i>
mefE 60mer <sup>b</sup>	gctagcaggagccttattattaggaagattaggggctcgaagcagattactaatt	40.0	
mph 50mer_I	cgcagccacaatagatccagaatacaaaatattgatggaattgaac	36.0	<i>mph(BM), mph(C)</i>
mph 50mer_II	gctgactgtcaatgagcttggctcactatagatcgtgacgccaccg	56.0	<i>mph(A), mph(K)</i>
msr 50mer_I	agtgaactccatatactatgcatacaaccgacagatgagtggtggtga	44.0	<i>msr(A), msr(SA)</i>
msr 50mer_II	taggtgcaaatggtgtagtgaagacaactttacttgaagctatttaccac	38.0	<i>msr(A), msr(SA), msr(B)</i>
vat 50mer_I	tattgaaattgggactacacctattatgatgaccagtaaatcccaccg	42.0	<i>sat(G), vat(E), vat(E-3)–vat(E-8)</i>
vat 50mer_II	ttttgaaaaattagaanaattgttgaggttggagaatactcatattatgat	24.0	<i>sat(A), vat(D)</i>
vat 50mer_III	gaatgaattcaataacaacttatcttttaataataatgggaaatggttgg	28.0	<i>vat(B)</i>
vat 50mer_IV	attgtcgtggtgaaatccctaaaaattataagaaaaaggttttctgatgg	32.0	<i>vat(A), vat</i>
vat 50mer_V	ggtgggacctagatagagacgataaatgaaaatattgattgcatcctg	40.0	<i>vat(C)</i>
vat 60mer_I	cgtgccaaccacgtaattgaaaggtatctgacttatcatttaataatttttaggtggcga	40.0	<i>sat(G), vat(E), vat(E-3)–vat(E-8)</i>
vga 50mer_I	gatgaaccaacaactttctgtatgggRctatagagggcttggatc	30.0	<i>vga(A), vga(A)<sub>LC</sub>, vga</i>
vgb 50mer_I	aaatatgataaagttgcatcaattgatgaaaaattacagatgccacc	46.0	<i>vgb</i>
vgb 50mer_II	cagggaattagaanaatctctaccaacaatgcagcggctccagtg	42.0	<i>vgb(B)</i>
<b>Sulfonamide</b>			
sul1 50mer <sup>c</sup>	gccccgcaccggaacatcgtgcacgtgctgcaaccttcaaaaagctg	60.0	<i>sul1*</i>
sul1 60mer_I <sup>c</sup>	tttctgagccccgcaccggaacatcgtgcacgtgctgcaaccttcaaaaagctgaa	53.3	
sul1 60mer_II	gatttttctgagccccgcaccggaacatcgtgcacgtgctgcaaccttcaaaaagc	53.3	
sul2 50mer <sup>c</sup>	gcgctcaagcagatgcaattcccgtctgctgacagttatcaMccccg	61.0	<i>sul2*</i>
sulA 50mer	caccctagctgctcatMttYcctcattttggtttgWcaagcttttac	44.0	<i>sulA</i>
<b>Tetracycline</b>			
otrA 50mer <sup>b</sup>	tcgtctggacgatctcaaggtcaacctcatcgacccccgggcccactcc	62.0	<i>otr(A)*</i>
otrB 50mer <sup>b</sup>	ggtcaactcaccatcggcgtcggcatctcggcagcgtcaccacctgc	64.0	<i>otr(B)</i>
otrB 60mer <sup>b</sup>	cgcaagcccatgacctgatctccatcgtggttctatcggcggctcgtgctgctg	63.3	
tet30 60mer	ccctgtcaatggcctcaactgttctcgcgtgttttctgcccgaagccgaaagg	56.7	<i>tet(30)</i>
tet31 60mer	gctcttatcatggtcattatctctctccctaaagagcaatcaccccaaaagaaatcgag	43.3	<i>tet(31)</i>
tet32 60mer	gttatttttagctgatgatacttgaaactgaacgacatctgggaaatgaaaaactcctg	35.0	<i>tet(32)</i>
tet33 60mer	ggtgactgttctggtccatctcgccacatctgccgttttctgctgctctctctca	56.7	<i>tet(33)</i>
tet34 50mer	atcatgatcaccagcgtgatgaccgtgctaaaagcggcagaaggtgat	48.0	<i>tet(34)</i>
tet34 60mer	cagctgctgaaaaacagatgccagctgcaacagtggaaggtatttggcgggtgagccgt	51.7	
tet35 60mer	tatcgacgcagctacctatgcatcattatcagtgccgtttgtgtcatgtattgctttatca	41.7	<i>tet(35)</i>
tet36 60mer	attaacatctagctgtacaatacactacatatacaaaaagacagagaacaattttaga	28.3	<i>tet(36)</i>
tet37 60mer	tgaagaactcaggtattcacattgattatctgttaacacatggctctgtatggc	38.3	<i>tet(37)*</i>
tetA 60mer <sup>c</sup>	ccaggcaggtgatgaggaaactcagggcagctgcaaggctcactgcgcgctcacc	66.7	<i>tet(A)*</i>
tetA(P) 50mer <sup>b</sup>	ttgtagcacagattgtatgggattagggtctactttatcagtgctcg	44.0	<i>tetA(P)</i>
tetB 60mer <sup>c</sup>	tatcgcttaatgaggttattcttctccttggccttgaaaaatgtctgaccgattggt	41.7	<i>tet(B)*</i>

Table 1 (continued)

Oligonucleotide name <sup>a</sup>	Sequence (5'–3')	GC %	Gene(s) represented <sup>d</sup>
tetB(P) 50mer <sup>b</sup>	agaaatacaagaaaagctttcattatgcaagagaaggaagctatatac	30.0	<i>tetB(P)</i>
tetC 50mer <sup>b</sup>	tgcggtattcggaaactctgcacgcccctgcctcaagccttcgctcactgctc	58.0	<i>tet(C)*</i>
tetD 60mer <sup>b</sup>	agcgcaggtatcagctttatcacactgctaaacctctggcgctgtgtgtgtttt	46.7	<i>tet(D)*</i>
tetE 60mer <sup>b</sup>	tgggtatggataattggctgctggattatgtatgttcattgattatactgaggttttc	36.7	<i>tet(E)*</i>
tetG 60mer <sup>c</sup>	cgcttttcgcaagttttcattatcaactgatcgcccaagtgccctgcagccctatggg	58.3	<i>tet(G)*</i>
tetH 60mer <sup>b</sup>	gggcgaaaaaacaccattatgatcagatgtctattgatgatgggct	40.0	<i>tet(H)</i>
tetJ 60mer <sup>b</sup>	ttttcattactttgtttccaagaaactcaaacacaaaaatttcgactga	30.0	<i>tet(J)</i>
tetK 50mer <sup>b</sup>	atgtttatattgttatggcgctgattatctttactaaaacagtatac	26.0	<i>tet(K)*</i>
tetK 60mer <sup>b</sup>	gtagtagacaaggagtaggactgctgctcctcactgattatggtgtgtgtagctag	45.0	
tetL 50mer <sup>b</sup>	gtaatggtgtgattgctgctgctatattccaagaaaataggggtaaagc	42.0	<i>tet(L)*</i>
tetL 60mer <sup>b</sup>	cggctacattggtgggatactgttgatagaagaggtcctttatagctgttaaacatcgg	43.3	
tetM 50mer <sup>b</sup>	aaagctggacaagaattgtagagccatattcttattttaaaRttatgct	30.0	<i>tet(M)*</i>
tetM 60mer <sup>b</sup>	gaagtKattacKaataaattttatcatcaacacatcgaggctMgtctgaactttcgga	36.7	
tetO 50mer <sup>I<sup>b</sup></sup>	tcatcaacgctgaaggtcaactgcaactatgcccggcaggttttaagat	44.0	<i>tet(O)*</i>
tetO 50mer <sup>II<sup>b</sup></sup>	ctttctgggctctgctgctggtgtccatagaccgctccctattggaagc	56.0	
tetQ 50mer <sup>b</sup>	acattgtgattgaagaccgctttgtcctttccataaaactcatatag	38.0	<i>tet(Q)*</i>
tetS 50mer <sup>b</sup>	gacatcataaataagcagactgtgaatctaaattgaaaccttattgta	28.0	<i>tet(S)*</i>
tetS 60mer <sup>b</sup>	tatgtagatacagtaactcacgaaattgtgctatcttttttaggtgaggtcctaaatggag	36.7	
tetT 50mer <sup>b</sup>	aattgacaaatgtaaggatgataaagtaattcaagtaaaataatagag	26.0	<i>tet(T)*</i>
tetU 50mer <sup>b</sup>	attggtcagataattgctagacatacaaaataataatcggaattgtctg	30.0	<i>tet(U)</i>
etV 60mer <sup>b</sup>	gccgaccggatcaaccagcgcaccatcatcattgccgtcagagtggtcaactcgtcacg	60.0	<i>tet(V)</i>
tetW 50mer <sup>b</sup>	ggataagctctccgccgatattatcatcaagcagacggtgctgctgtccc	54.0	<i>tet(W)*</i>
tetW 60mer <sup>b</sup>	cattcaagcggcagctcctcctcagtgccacagatgaaagtYaacattgtggatac	45.0	
tetX 50mer <sup>b</sup>	aaaagcgggattgttcaaaactattatgacttagccttaccatgggtg	54.0	<i>tet(X)*</i>
tetY 50mer <sup>b</sup>	gccagctttttgctgtgtttttctatgagctgattggcgagcgcc	50.0	<i>tet(Y)</i>
tetZ 50mer <sup>b</sup>	atcgactacctgctgctgcactgacggacacgctgtgggtcttttacct	56.0	<i>tet(Z)</i>
<b>Trimethoprim</b>			
dfrA 50mer <sup>I</sup>	acaactgaccactgggaatacactgtgaatggcacggaaaacttttaatt	38.0	<i>dfrA</i> , <i>dfrC</i>
dfrA 50mer <sup>II</sup>	aggctcaccgagacaaaagtggtgctgtatggccgcaagacatttgatc	54.0	<i>dfrA13</i> , <i>dfrA21*</i> , <i>dfrA22</i> , <i>dfrA23</i>
dfrA1 60mer <sup>c</sup>	gcggtcgtaacacggttcaagttttacatctgacaatggaacgtaKtgatcttccatca	40.8	<i>dfrA1*</i>
dfrA2 60mer	atcgctRcgcaagaatactcgYgcccgttggcagggtcaagtYgtcggKtggtatgtgca	57.5	<i>dfrA2</i>
dfrA3 50mer <sup>I</sup>	gaagtggattgtctgtggaaggagatgcaattttccccgcaatagaccg	48.0	<i>dfrA3</i> , chromosomal
dfrA3 50mer <sup>II</sup>	cattgacgctcagttgaacgggtataccattttccccgattacctatcgc	50.0	<i>dfrA3</i> , plasmid
dfrA5 50mer	cctggacggcccataatgacaacgtaataatgattcccgtcagcgaagag	50.0	<i>dfrA5</i>
dfrA6 50mer	atgaaaatattcttattggcagctgtttccgagaatggagtaattggctc	40.0	<i>dfrA6</i>
dfrA7 60mer	gtgttctccaaatcgaataatgacagtagtgcaggaaaggaattcaagctcaaatg	38.3	<i>dfrA7</i>
dfrA8 50mer	cttgcctcgaatgagaagctgcacactgcatgattgacgccaag	54.0	<i>dfrA8</i>
dfrA9 50mer	gtgggaagagtgctgacgagaactagctgctcactggtgacaactctac	52.0	<i>dfrA9</i>
dfrA10 60mer <sup>c</sup>	tcattgtggtgtgtgtttattatctgaagcgatagaactgctagcactgtttacatga	38.3	<i>dfrA10*</i>
dfrA12 60mer <sup>I</sup>	tcgcagactcactgagggaaaagctgttgcctatggggcgaagacctttgagctatcgg	51.7	<i>dfrA12*</i>
dfrA12 60mer <sup>II</sup>	cgtagttgtttcaacgctgtcgcacgctatcctttggcatcgaactcggcaatgaact	51.7	
dfrA14 50mer <sup>c</sup>	aatgatgacaatgtatgttattcagtcacatcgaagagccatggacag	40.0	<i>dfrA14*</i>
dfrA14 60mer <sup>c</sup>	ttggacatcaaatgatgacaatgtatgttattcagtcacatcgaagagccatggacag	40.0	
dfrA15 50mer	gccgttgaactcgttcaagctcacttccagtgatgagaatgtattgtg	50.0	<i>dfrA15*</i>
dfrA16 60mer	gagatggagacatagttttcctgaaatcccagatacattcaagttggtatttgagcaag	38.3	<i>dfrA16*</i>
dfrA17 50mer	aaaacgtcctagttttcctcaatagaaaatgctttgaaagagctatca	32.0	<i>dfrA17</i>
dfrA19 60mer	atctcgtgctgtgatcaacctgtatgagaataaccgatcaacgattgctgctattggtg	45.0	<i>dfrA19</i>
dfrB 50mer <sup>I</sup>	aacgctcccgtgcaagggcagtttgcctcctccctgagtgccacctttgg	62.0	<i>dfrB2</i>
dfrB 50mer <sup>II</sup>	caacacaacaatggagtcagctactctgctgctgcccagtttgcgctccc	52.0	<i>dfrB3</i>
dfrD 50mer	gagtaatcggcaaggataacgacattccatggagaatttctagtattgg	42.0	<i>dfrD</i>
<b>Vancomycin</b>			
vanA 60mer <sup>I<sup>b</sup></sup>	gcggaatgggaaaacgacaattgctattcagctgtactctcggcgataaaaaatgcac	45.0	<i>vanA*</i>
vanA 60mer <sup>II<sup>b</sup></sup>	cggtgtSgatatttttaacaagataacggccgctgactgaacgaagtaatacYctg	41.7	
vanB 50mer <sup>b</sup>	taccctgtctttgtgaagccggcagcggctcaggttcttcttggcgtaac	56.0	<i>vanB*</i>
vanB 60mer <sup>b</sup>	cgcactacatcgaatcaaaaaacggYgtatggaagctatgcaagaagccatgtacgg	45.0	
vanC1 50mer <sup>b</sup>	tggttgcctatgctgctcctccgcaattatgatgaacaatggctcttgc	50.0	<i>vanC-1</i>



**Table 1** (continued)

Oligonucleotide name <sup>a</sup>	Sequence (5'–3')	GC %	Gene(s) represented <sup>d</sup>
vanC2/C3 50mer <sup>b</sup>	ttgactgtcggctgtgtgacgccattcattagtagacggcttttcg	46.0	<i>vanC-2/3</i>
vanC2/C3 60mer <sup>b</sup>	actctttgactgtcggctgtgtgacgccattcattagtagacggcttttcgatttg	43.3	
vanD 50mer	gagattgccgcaaacatagatacaaaaaatcatcagccttattatattgg	32.0	<i>vanD</i>
vanE 50mer	aagggacaagacacctacaaaaagtcgatgcgtttgcgaaaatacatggatt	40.0	<i>vanE</i>
<b>Integron related</b>			
intI1 59mer <sup>c</sup>	cgagcagctgtcgcgtgcacgggcatgtggctgaaggaccagccgagggccgcagcg	72.9	<i>intI1*</i>
intI2 60mer <sup>c</sup>	atgaatgcttgcgtttgcgggttaaagattttgataatggctgcatcactgtgc	40.0	<i>intI2*</i>
intI3 60mer	accactgtctcaagcagggcacagacatccgaacgggtcgaagagtttggggcattcg	56.7	<i>intI3</i>
intI4 60mer	cgccgcatcatatgaacgaacagtactacaaaaagcgggtgagaagatcggctcaagaa	46.7	<i>intI4</i>
SGI1 LB 50mer	ttctgtattgggaagtaaactcctaataaataaaaaacgaagtaaaa	24.0	SGI1 left border*
<b>Salmonella specific</b>			
invA 60mer_I	taagcgaacgtgtttccgtgcgtaatatgaaataattatggaagcgcctcattgtggg	40.0	<i>invA Salmonella*</i>
invA 60mer_II	tgcttcttactaataacagctgcgtttacgaccYgaattMctgafYctggtactaatgg	40.0	
Seis 60mer <sup>c</sup>	gggagccaatataatgaccaagcaaaactcgaattgacggcctgcaggttggcgaggt	48.3	<i>sefA S. Enteritidis*</i>
sipB/C 60mer <sup>c</sup>	agcgctaaagatattcgaatagattgtattagcagcagtaaaagtcagtgacctgggg	41.7	<i>sipB/C Salmonella*</i>
Sspp 50mer <sup>c</sup>	tgaaggaaattacgctgcatttattgatcagaatacggccctgctgg	44.0	putative DNA/RNA endonuclease <i>Salmonella*</i>
Sspp 60mer <sup>c</sup>	cgtaaaaaagtgaaagaaattacgctgcatttattgatcagaatacggccctgctgg	43.3	
Stmm 50mer <sup>c</sup>	actgaggatgtgaaaaatgtacaagttgcaaatgctgatttgacagaggc	40.0	<i>flhC S. Typhimurium*</i>
Stmm 60mer <sup>c</sup>	actgaggatgtgaaaaatgtacaagttgcaaatgctgatttgacagaggctaaagccgca	41.7	

<sup>a</sup> The antibiotic class is indicated in bold and italics.

<sup>b</sup> Described by Mättö et al. (2007).

<sup>c</sup> Described by van Hoek et al. (2005).

<sup>d</sup> For genes with an asterisk, control strains were available.

<sup>e</sup> Also called *sat1* (Partridge and Hall 2005).

<sup>f</sup> The *bla*<sub>CTX-M-9</sub> group includes *bla*<sub>CTX-M-9\*</sub>, *bla*<sub>CTX-M-13</sub>, *bla*<sub>CTX-M-14</sub>, *bla*<sub>CTX-M-16</sub>, *bla*<sub>CTX-M-17</sub>, *bla*<sub>CTX-M-19</sub>, *bla*<sub>CTX-M-21</sub>, *bla*<sub>CTX-M-24</sub>, *bla*<sub>CTX-M-27\*</sub>, *bla*<sub>CTX-M-38</sub>, *bla*<sub>CTX-M-51</sub>, *bla*<sub>TOHO-2</sub>, *bla*<sub>UOE-2</sub> (Bonnet 2004).

<sup>g</sup> The *bla*<sub>CTX-M-2</sub> group includes *bla*<sub>CTX-M-2\*</sub>, *bla*<sub>CTX-M-5</sub>, *bla*<sub>CTX-M-20</sub>, *bla*<sub>CTX-M-31</sub>, *bla*<sub>CTX-M-35</sub>, *bla*<sub>CTX-M-43</sub>, *bla*<sub>CTX-M-56</sub>, *bla*<sub>CTX-M-59</sub>, *bla*<sub>KLUA</sub>, *bla*<sub>TOHO-1</sub> (Bonnet 2004).

<sup>h</sup> The *bla*<sub>CTX-M-1</sub> group includes *bla*<sub>CTX-M-1</sub>, *bla*<sub>CTX-M-3</sub>, *bla*<sub>CTX-M-10</sub>, *bla*<sub>CTX-M-11</sub>, *bla*<sub>CTX-M-12</sub>, *bla*<sub>CTX-M-15\*</sub>, *bla*<sub>CTX-M-22</sub>, *bla*<sub>CTX-M-28</sub>, *bla*<sub>CTX-M-32</sub>, *bla*<sub>CTX-M-33</sub>, *bla*<sub>CTX-M-34</sub>, *bla*<sub>CTX-M-36</sub>, *bla*<sub>CTX-M-37</sub>, *bla*<sub>CTX-M-38</sub>, *bla*<sub>CTX-M-42</sub>, *bla*<sub>CTX-M-52</sub>, *bla*<sub>CTX-M-53</sub>, *bla*<sub>CTX-M-54</sub>, *bla*<sub>CTX-M-55</sub>, *bla*<sub>CTX-M-57</sub>, *bla*<sub>CTX-M-58</sub>, *bla*<sub>CTX-M-60</sub>, *bla*<sub>CTX-M-61</sub>, *bla*<sub>CTX-M-64</sub>, *bla*<sub>CTX-M-66</sub>, *bla*<sub>UOE-1</sub> (Bonnet 2004).

<sup>i</sup> The *bla*<sub>OXA-10</sub> group includes *bla*<sub>OXA-7</sub>, *bla*<sub>OXA-10</sub> (= *bla*<sub>PSE-2</sub>), *bla*<sub>OXA-13</sub>, *bla*<sub>OXA-14</sub>, *bla*<sub>OXA-16</sub>, *bla*<sub>OXA-17</sub>, *bla*<sub>OXA-19</sub>, *bla*<sub>OXA-28</sub>, *bla*<sub>OXA-35</sub>, *bla*<sub>OXA-56</sub>, *bla*<sub>OXA-74</sub>, *bla*<sub>OXA-101</sub>.

resistance to the aminoglycoside antibiotic streptomycin, an extended-spectrum- $\beta$ -lactamase *bla*<sub>TEM</sub> gene, the genes *marA*, *marC*, and *marR* belonging to the multiple antibiotic resistance (*mar*) locus and the tetracycline resistance gene *tet(A)*. Also general, but less frequently present, were the aminoglycoside resistance genes *aadA1* and *aadA2*, the sulfonamide resistance gene *sul1*, and the trimethoprim resistance gene *dfrA1*. MLS and vancomycin resistance genes were nearly absent in the *Salmonella* investigated, with one exception, i.e., *mph(A)* or *mph(K)* was demonstrated in one Heidelberg and three Blockley isolates.

In contrast to the high frequency of *strA*, the related *strB* gene was not detected by microarray analysis, although an increased incidence in streptomycin-resistant *Salmonella* has been reported because of the presence of both *strA* (*aph(3'')-Ib*) and *strB* (*aph(6'')-Id*) (Sundin and Bender 1996 and Pezzella et al. 2004). The absence of a hybridization signal

is probably the result of the bad performance of the designed strB 50-mer oligonucleotide. This was confirmed by the design of three new oligonucleotides, i.e., strA 60mer (5'-gccatggtgatccctgcatgccgaacttcattggtggaccctaaactcttcaatgcacgg-3') strA-strB 60mer (5'-tgccgattgaccctctgacttggggKtgatgttcatgccgctgttttctgctcattg-3', specific for the 3' end of *strA* and 5' end of *strB*) and strB 60mer (5'-gatgagcaatgctcctggaactgcgtgggctacatggcgatctgcatcatgaaacatcat-3'). All three oligonucleotides gave a hybridization signal when tested with a control strain containing both *strA* and *strB*.

As listed in Table 1, the microarray contained a considerable number of oligonucleotides representing the four classes of ESBLs, e.g., class A: *bla*<sub>CTX-M</sub> type genes; class B: *bla*<sub>VIM</sub> type genes; class C: *bla*<sub>CMY</sub> type genes; class D: *bla*<sub>OXA</sub> type genes. Forty-nine out of the 143 examined *Salmonella* harbored a *bla*<sub>TEM</sub> gene, whereas *bla*<sub>PSE-01</sub> was

**Table 2** PCR primers

Antibiotic class or integrase	Primer name	Sequence (5'–3')	Primer name	Sequence (5'–3')	Gene(s)	Length PCR product	
Aminoglycoside	aacA4_F	tggggcgggagaagaagc	aacA4-R	tgcttcgccaagtaact	<i>aacA4</i>	184 bp	
	aacC2_F	tgggtgcccgcctaacc	aacC2-R	caaagcaatcgagaatg	<i>aacC2</i>	195 bp	
	aadA1_F <sup>a</sup>	atgaggggaagtgtgatcgc	aadA1_R <sup>a</sup>	ttccaaaaggctgtgatcaaa	<i>aadA1</i>	216 bp	
	aadA2_F <sup>a</sup>	gcagcgaatgacattcttg	aadA2_R <sup>a</sup>	catccttcggcgcgattttg	<i>aadA2</i>	284 bp	
	aphA1_F	gatttatatgggtatag	aphA1_R	cgggaagaggcataaatg	<i>aphA1</i>	223 bp	
	sat2_F	aagactctgctgctatggc	sat2_R	tctgtgctcccagagaac	<i>sat2</i>	346 bp	
	strA_F	cgaacgagagctaccgg	strA_R	ttccgagcccaccaagg	<i>strA</i>	140 bp	
	strB_F	tgctgatgaactgcgcg	strB_R	ggagaagggcagaaggc	<i>strB</i>	227 bp	
β-lactam	blaACC-1_F	tgaagctgtattccctg	blaACC-1_R	tgttttgcccgtacc	<i>bla<sub>ACC-1</sub></i>	202 bp	
	blaCTX-M-g1_F	gtacagcaaaaactgccc	blaCTX-M-g1_R	ctttcacttttctcagc	<i>bla<sub>CTX-M-1</sub></i> , <i>bla<sub>CTX-M-3</sub></i> , <i>bla<sub>CTX-M-10</sub></i> , <i>bla<sub>CTX-M-11</sub></i> , <i>bla<sub>CTX-M-12</sub></i> , <i>bla<sub>CTX-M-15</sub></i> <sup>*</sup> , <i>bla<sub>CTX-M-22</sub></i> , <i>bla<sub>CTX-M-28</sub></i> , <i>bla<sub>CTX-M-32</sub></i> , <i>bla<sub>CTX-M-33</sub></i> , <i>bla<sub>CTX-M-34</sub></i> , <i>bla<sub>CTX-M-36</sub></i> , <i>bla<sub>CTX-M-37</sub></i> , <i>bla<sub>CTX-M-38</sub></i> , <i>bla<sub>CTX-M-42</sub></i> , <i>bla<sub>CTX-M-52</sub></i> , <i>bla<sub>CTX-M-53</sub></i> , <i>bla<sub>CTX-M-54</sub></i> , <i>bla<sub>CTX-M-55</sub></i> , <i>bla<sub>CTX-M-57</sub></i> , <i>bla<sub>CTX-M-58</sub></i> , <i>bla<sub>CTX-M-60</sub></i> , <i>bla<sub>CTX-M-61</sub></i> , <i>bla<sub>CTX-M-64</sub></i> , <i>bla<sub>CTX-M-66</sub></i> , <i>bla<sub>UOE-1</sub></i>	170 bp	
	blaCTX-M-g2_F	cgctgcatgcccagggc	blaCTX-M-g2_R	gcaaaaagttcatcgccagc	<i>bla<sub>CTX-M-2</sub></i> , <i>bla<sub>CTX-M-5</sub></i> , <i>bla<sub>CTX-M-20</sub></i> , <i>bla<sub>CTX-M-31</sub></i> , <i>bla<sub>CTX-M-35</sub></i> , <i>bla<sub>CTX-M-43</sub></i> , <i>bla<sub>CTX-M-56</sub></i> , <i>bla<sub>CTX-M-59</sub></i> , <i>bla<sub>KLUA</sub></i> , <i>bla<sub>TOHO-1</sub></i>	136 bp	
	blaCTX-M-g5_F	gagcttggcgcagcg	blaCTX-M-g5_R	cgctcactttatcgggc	<i>bla<sub>CTX-M-5</sub></i> , <i>bla<sub>KLUA</sub></i>	87 bp	
	blaCTX-M-g9_F	ggcaatacagccgccc	blaCTX-M-g9_R	cagcggcgcacagccctg	<i>bla<sub>CTX-M-9</sub></i> , <i>bla<sub>CTX-M-13</sub></i> , <i>bla<sub>CTX-M-14</sub></i> , <i>bla<sub>CTX-M-16</sub></i> , <i>bla<sub>CTX-M-17</sub></i> , <i>bla<sub>CTX-M-19</sub></i> , <i>bla<sub>CTX-M-21</sub></i> , <i>bla<sub>CTX-M-24</sub></i> , <i>bla<sub>CTX-M-27</sub></i> <sup>*</sup> , <i>bla<sub>CTX-M-38</sub></i> , <i>bla<sub>CTX-M-51</sub></i> , <i>bla<sub>TOHO-2</sub></i> , <i>bla<sub>UOE-2</sub></i>	135 bp	
	blaOXA-g1_F	tatggcatttggatgccc	blaOXA-g1_R	gttttctatggctgag	<i>bla<sub>OXA-01</sub></i> , <i>bla<sub>OXA-04</sub></i> , <i>bla<sub>OXA-30</sub></i> , <i>bla<sub>OXA-31</sub></i> , <i>bla<sub>OXA-47</sub></i>	352 bp	
	blaPSE-1_F <sup>a</sup>	cgctatctgaaatgaaccag	blaPSE-1_R <sup>a</sup>	tttcgctctgccattgaagc	<i>bla<sub>PSE-1</sub></i>	229 bp	
	blaTEM_F <sup>a</sup>	tgggtgcacagtggggttac	blaTEM_R <sup>a</sup>	gtagctccttcgctctcc	<i>bla<sub>TEM</sub></i>	328 bp	
	Chloramphenicol	catB3_F	gtagtttctgctctatc	catB3_R	cttcttagcgggattgct	<i>catB3</i>	302 bp
		cmlA_F	cgctcgtcgacatgtggc	cmlA_R	gccaaagctgagacacacc	<i>cmlA</i> , <i>cmlA1</i> , <i>cmlA4</i> , <i>cmlA5</i> , <i>cmlA6</i> , <i>cmlA7</i>	272 bp
Sulfonamide	floR_F <sup>a</sup>	ccttctcgtcttctctcg	floR_R <sup>a</sup>	ggtaggatgaaggtgaggaa	<i>floR</i>	255 bp	
	sul1_F <sup>a</sup>	gccttcgacggagcggggt	sul1_R <sup>a</sup>	aggcatgatcaaccctcgg	<i>sul1</i>	363 bp	
	sul2_F	caaggcagatggcattccc	sul2_R	gtcgcacggcgggtgcctc	<i>sul2</i>	211 bp	
Tetracycline	tetA_F <sup>a</sup>	gccggcgcctcaagcaattt	tetA_R <sup>a</sup>	ccacgtttgataagaagcc	<i>tet(A)</i>	148 bp	
	tetB_F <sup>a</sup>	acgtgaatttattgctcgg	tetB_R <sup>a</sup>	atacagatccaaagcgac	<i>tet(B)</i>	205 bp	
	tetC_F	gcgggatatcgccattccg	tetC_R	cgtagaggatccacaggacg	<i>tet(C)</i>	206 bp	
	tetD_F	aaaccggcgggtacagacaga	tetD_R	aaaccgaccgccgctgtc	<i>tet(D)</i>	200 bp	
	tetG_F <sup>a</sup>	gactggcttcgttctctgg	tetG_R <sup>a</sup>	ttcgaatggtctcgtagt	<i>tet(G)</i>	308 bp	
Trimethoprim	dfrA1_F <sup>a</sup>	ccaaaggtgaacagctcctg	dfrA1_R <sup>a</sup>	atatgtatgtctactctg	<i>dfrA1</i>	271 bp	
	dfrA2_F	gttgcWgggcagtttgcgct	dfrA2_R	gcagccacaggataaat	<i>dfrA2</i>	185 bp	
	dfrA10_F	ttgagagcttcttagaa	dfrA10_R	accggtacatacacatcagc	<i>dfrA10</i>	158 bp	
	dfrA12_F	cctcgtttgacgcgctc	dfrA12_R	attggcggcgggaagaacg	<i>dfrA12</i>	204 bp	
	dfrA14_F <sup>a</sup>	tctgtgggtgcgaagacg	dfrA14_R <sup>a</sup>	atgggtaattgttctcgg	<i>dfrA14</i>	204 bp	
	dfrA16_F	tacaaaagcttgattcc	dfrA16_R	aatagttaattgttagact	<i>dfrA16</i>	142 bp	
Integrase	intI1_F <sup>a</sup>	atcgggccttgatgttac	intI1_R <sup>a</sup>	gcgcgctgaaaggtctgg	<i>intI1</i>	256 bp	
	intI2_F <sup>a</sup>	gcaggttatgatactcg	intI2_R <sup>a</sup>	gctgtttctgctttccc	<i>intI2</i>	157 bp	

<sup>a</sup> Described by van Hoek et al. (2005)



almost exclusively found in *S. Typhimurium* DT104. The plasmid-encoded CTX-M type genes were occasionally detected, the class C  $\beta$ -lactamase gene *bla*<sub>ACC-1</sub> was detected twice and a class D ESBL *bla*<sub>OXA</sub> was found once.

The microarray also included oligonucleotides representing sequences of class 1 and class 2 integrons (integrase genes) and sequences of SGII. As summarized in Table 3, most of the DT104 isolates gave hybridization signals with oligonucleotide intI1 59mer representing the integrase of a class 1 integron. Class 2 integrons were almost exclusively detected in Paratyphi B var. Java, but, also two Agona, the investigated *houtenae* strain, and one of the DT104 isolates harbored *intI2*.

A total of 44 *Typhimurium* strains have been investigated by microarray analysis, including 19 DT104 or Pt506 isolates. The DT nomenclature is according to the English and the Pt based on the Dutch phage-typing system. DT104 corresponds with Pt506 and they are grouped as DT104 in Table 3.

The microarray hybridization data of the *Typhimurium* isolates revealed a high frequency of the commonly SGII localized AR determinants: the *aadA2* gene coding for an adenylyltransferase mediating resistance to the aminoglycoside streptomycin and the aminocyclitol spectinomycin, the ESBL *bla*<sub>PSE-1</sub> gene, the sulfonamide resistance gene *sulI*, the chloramphenicol/florfenicol resistance gene *floR* and the tetracycline resistance gene *tet(G)*. Most of the DT104 strains investigated here also harbored these AR genes; however, some of them only showed hybridization signals with a subset of these determinants, which suggests the presence of variant SGII. From four of the investigated isolates (S/921495, S/960275, S/954435, and S/960081), it was already known that they contain variable SGII (Boyd et al. 2002). These strains had already been analyzed on a smaller oligonucleotide microarray (see van Hoek et al. 2005).

In *S. Paratyphi* B var. Java, the streptomycin resistance gene *aadA1* was found in all strains, although, for one of them one of the 3 *aadA1* oligonucleotides (i.e., *aadA1* 60mer\_II) did not give a hybridization signal. The related resistance genes *aadA2* and *strA* were also detected in this serovar, but less frequent. In addition, all investigated Paratyphi B var. Java isolates contained *dfpA1* and *sat2*.

Among the 10 investigated Hadar isolates, seven contained *bla*<sub>TEM</sub> and *strA*; furthermore, in eight *tet(A)* was identified. Less frequently *aphA1* and *sulI* were demonstrated. The genes *strA*, *bla*<sub>TEM</sub>, and *tetA* were found in similar incidences in isolates belonging to the serovars Heidelberg and London. Within *S. Enteritidis*, responsible for the greatest part of the human cases of salmonellosis, most likely related to the consumption of raw shell eggs, the number of AR genes per isolate was relatively low on average.

## Discussion

Taking into account the number of antibiotic resistance genes described in the literature so far, it would take quite an effort to investigate bacterial isolates by PCR. Microarray analysis allows the screening of a large number of targets simultaneously and as such circumvents the shortcomings of PCR. The potential of miniaturization in addition to multiplexing offers a considerable advantage of microarray analysis over other molecular methods for clinical or epidemiological applications. As summarized by Garaizar et al. (2006), microarrays have been shown to be helpful for quick detection of antibiotic resistance, determination of virulence and pathogenicity, species determination, genome comparison, and molecular epidemiological typing of strains.

In this paper, the successful application of microarray analysis was demonstrated again using a microarray containing 223 oligonucleotides representing more than 430 AR genes for the screening of a large set of *Salmonella* isolates. To a large extent, the obtained hybridization results confirmed the general findings concerning antibiotic resistance in *Salmonella*. Although many antibiotic resistance genes are represented by the microarray, the used setup is, however, not suited for the detection of mutation-mediated resistance, like for instance mutations within the gyrase gene leading to nalidixic acid resistance.

The level and extent of resistance among *Salmonella* varies in different geographical locations. Nevertheless, there is a correlation between the length of time an antimicrobial agent has been used and its corresponding resistance (McDermott 2006). A general increase is noticed with respect to *Salmonella* strains with a multidrug resistance phenotype including resistance to ampicillin, chloramphenicol, sulfonamides, streptomycin, and tetracycline (Brisabois et al. 1997). The microarray data presented here support those findings because in the majority of the isolates investigated multiple AR genes were detected. The most frequently detected genes in this study belong to the inducible *mar* operon, which controls the intrinsic levels of susceptibility to structurally different antibiotics. The occurrence of these genes within *Salmonella enterica* have been described by, for instance, Sulavik et al. (1997) and Kunonga et al. (2000), who demonstrated the presence of the *marR*, *marA*, and *marB* genes in *S. Typhimurium* and 30 different serovars, respectively, and by Randall and Woodward (2001) describing 44 serovars of *Salmonella* with a conserved *marA*. The results reported here concerning the widespread occurrence of *mar* genes in the genus of *Salmonella* are in agreement with the data mentioned above, although the *mar*-related oligonucleotides were not tested with well-defined control strains, nor were the results confirmed by PCR. Probably as a consequence *marB* was not detected, whereas *marR* and

*marA*, also part of the transcriptional unit TU2 called *marRAB* and *marC*, a gene belonging to transcriptional unit TU1, were found.

The second most frequently detected gene was the ESBL *bla*<sub>TEM</sub>, although *strA*, *sul1*, and *tet(A)* were also common among the *Salmonella* strains investigated. It is a well-known fact that like other Enterobacteriaceae, also ESBL-producing *Salmonella* strains have been emerging worldwide during the last decade (Bonnet 2004 and Hasman et al. 2005). Integrons (in particular class 1) are frequently found in *Salmonella* and are often associated with AR genes cassettes. These elements have the ability to integrate and excise genes and play a role in the dissemination of AR among different bacteria (Miko et al. 2003; Rowe-Magnus and Mazel 2002; Carattoli 2001 and Martinez-Freijo et al. 1998). Within the Typhimurium DT104 isolates, recognized as an emergent and multiresistant pathogen, different combinations of AR genes were found in this study that are most likely associated with the class 1 integron elements located on the SGII. Strains belonging to phage type DT104 are very commonly characterized by the clustering of five resistance genes, i.e., *bla*<sub>PSE-1</sub>, *floR aadA2*, *sul1*, and *tet(G)*, on the so-called Salmonella Genomic Island 1, resulting in the penta resistance type ACSSuT (Boyd et al. 2001). However, the phenomenon of variant SGII elements has been identified by Boyd et al. (2002) and the described strains were also analyzed here. The microarray results were in full accordance with the published data (see also van Hoek et al. 2005).

Class 2 integron sequences were found in the Paratyphi B var. Java isolates investigated, all harbored *aadA1*, *sat2* and *dfrA1*, which are known to be part of the *Tn7* transposon a known class 2 integron (Partridge and Hall 2005). The microarray results are in good agreement with Miko et al. (2003). These authors investigated 85 multi-resistant D-tartrate positive *Salmonella enterica* subsp. *enterica* serovar Paratyphi B (also known as *S. Paratyphi B* var. Java) isolates for the presence of integrons and detected in all of them the chromosomally located *Tn7*-like class 2 integron harboring the same *dfrA1-sat2-aadA1* gene cassette.

*S. Hadar* is often found as a cause of human infections in many European countries, and high frequencies of resistance are reported for ampicillin, streptomycin, and tetracycline (Threlfall et al. 2000). These phenotypic data were supported by the obtained microarray results, as *bla*<sub>TEM</sub>, *strA*, and *tet(A)*, respectively, were identified in nearly all *S. Hadar* investigated. These multiple AR genes, however, could not be linked to the presence of a class 1 or 2 integron; this phenomenon has been reported before in this serovar (Randall et al. 2004).

According to the Dutch monitoring program of antimicrobial resistance and antibiotic usage in animals and other

reports (for review, see McDermott) the highest level of resistance is found in *Salmonella* isolates belonging to serovar Typhimurium (McDermott 2006; Mevius et al. 2004 and Mevius and van Pelt 2005). The most common phenotypes in *S. Typhimurium* display resistance to amoxicillin, tetracycline, sulphamethoxazole, trimethoprim, chloramphenicol, and florfenicol with the highest levels of resistance in strains with a DT104 phage type. This is also reflected by the microarray screening presented here as AR genes related to the above-mentioned phenotypes were found in high frequencies within serovar Typhimurium and phage type DT104. Besides *S. Typhimurium*, also within *S. Paratyphi* var. Java and *S. Derby*, the percentage of resistance to amoxicillin, tetracycline, sulphamethoxazole, trimethoprim, and chloramphenicol/florfenicol is relatively high (Martinez-Freijo et al. 1998; Mevius et al. 2004 and Mevius and van Pelt 2005) compared to the other serovars. As a consequence a high number of AR genes within isolates of these serovars were identified with the microarray screening.

Part of the oligonucleotides presented in this study have already been described in previous studies (van Hoek et al. 2005 and Mättö et al. 2007). The latter publication demonstrated the applicability of the designed oligonucleotide microarray not only for pathogens, but also for the screening of gram-positive *Bifidobacterium* spp. strains that are generally associated with good intestinal health. Broad screening of the presence of AR genes both within the pathogenic and in the environmental or food-associated (harmless) bacterial population will help to identify reservoirs for AR determinants and to critically evaluate the use of antibiotics both in agriculture and for the treatment of infections in humans.

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