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PERSPECTIVE

Using the class 1 integron-integrase gene as a proxy for anthropogenic pollution

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Around all human activity, there are zones of pollution with pesticides, heavy metals, pharmaceuticals, personal care products and the microorganisms associated with human waste streams and agriculture. This diversity of pollutants, whose concentration varies spatially and temporally, is a major challenge for monitoring. Here, we suggest that the relative abundance of the clinical class 1 integron-integrase gene, *intl1*, is a good proxy for pollution because: (1) *intl1* is linked to genes conferring resistance to antibiotics, disinfectants and heavy metals; (2) it is found in a wide variety of pathogenic and nonpathogenic bacteria; (3) its abundance can change rapidly because its host cells can have rapid generation times and it can move between bacteria by horizontal gene transfer; and (4) a single DNA sequence variant of *intl1* is now found on a wide diversity of xenogenetic elements, these being complex mosaic DNA elements fixed through the agency of human selection. Here we review the literature examining the relationship between anthropogenic impacts and the abundance of *intl1*, and outline an approach by which *intl1* could serve as a proxy for anthropogenic pollution.

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Introduction

Humans produce and use a diverse array of compounds in domestic, industrial and agricultural settings. These compounds can contaminate ecosystems, elevating local concentrations of pollutants such as heavy metals, synthetic organic compounds and radioactive isotopes. Together with microbiological contaminants, they create a zone of impact emanating from human activities. Managing impacts requires monitoring to assess the efficacy of preventative or remedial measures, by measuring the quantities and distribution of individual pollutants. However, because some 80000 different compounds are now traded in the marketplace, testing for all pollutants is not feasible (Rockstrom et al., 2009). Focussing on just one class of pollutant is also problematic, because the composition of pollutants varies both geographically and temporally. Furthermore, diverse classes of pollutants, such as antibiotics and endocrine disrupting compounds, have significant biological effects at extremely low concentrations (Diamanti-Kandarakis *et al.*, 2009; Gillings, 2013).

An alternative to direct detection is to use a proxy that exhibits rapid responses to diverse environmental pressures and could thus be a generic marker for anthropogenic pollutants. We propose that the class 1 integron-integrase gene, *intI1*, could serve as such a marker, because:

- it is commonly linked to genes conferring resistance to antibiotics, disinfectants and heavy metals (Liebert et al., 1999; Partridge et al., 2001);
- 2. it has penetrated into diverse pathogenic and commensal bacteria of humans and their domestic animals (Goldstein *et al.*, 2001; Stokes and Gillings, 2011);
- 3. the abundance of *intI1* can rapidly change in response to environmental pressures, because the class 1 integron resides in diverse bacterial species that themselves have rapid generation times, and it is often located on mobile genetic elements that can readily transfer between bacteria; and

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4. the common 'clinical' forms of *intI1* are xenogenetic, that is, recently assembled under selection pressures imposed by human activities (Gillings *et al.*, 2008a).

Independent studies have already begun to note remarkable correlations between *intI1* and associated genetic elements with various measures of human impact (Gaze *et al.*, 2011; Pruden *et al.*, 2012; Jechalke *et al.*, 2013b). Here we review the recent evolutionary origins of the clinical class 1 integron, examine a series of case studies using *intI1* as an environmental marker of human pollution, and suggest methods for using this gene as a proxy for human impact.

The evolutionary history of the class 1 integron

Integrons are an ancient and common feature of bacterial genomes, where they usually reside on chromosomes (Gillings, 2014a). They have three core features: an integron-integrase gene (intI), a recombination site (attl) and a promoter (P_C). These features allow capture and expression of exogenous genes as part of gene cassettes that are recombined into the attI site using the integrase activity encoded by intI (Boucher et al., 2007; Cambray et al., 2010) and subsequently expressed from P_C (Collis and Hall, 1995) (Figure 1). This allows genes to be acquired and expressed with minimal disturbance to the existing genome. Integrons sample cassettes from an extraordinarily diverse pool that encodes functions of potential adaptive significance. Consequently, they are a hot spot of genomic diversity in a range of genera (Gillings et al., 2005; Boucher et al., 2011; Hall, 2012; Wu et al., 2013).

Hundreds of integron classes have been described, defined on the basis of the relative homology of *intI* (Cambray *et al.*, 2010; Boucher *et al.*, 2011). Of these, the class 1 integrons, so named because they were first to be discovered, had properties that meant that they were well equipped to move by lateral DNA

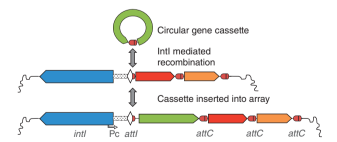


Figure 1 Integron structure and function. Integrons consist of a gene for an integron-integrase (intl) that catalyses recombination between the attC site of circular gene cassettes and the attendant integron recombination site, attl. This activity results in the sequential insertion of multiple, different cassettes to form a tandem cassette array that, in some cases, might contain hundreds of different genes. Inserted genes are expressed by an integron-encoded promoter, Pc.

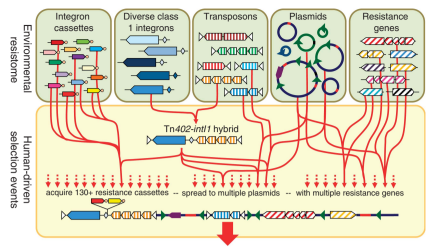
transfer into a wide range of commensal and pathogenic bacteria, and to accumulate diverse antibiotic resistance genes once humans tried to control bacteria with antimicrobial compounds. These fortuitous properties included: location on the chromosomes of Betaproteobacteria whose habitats intersect the human food chain; ability to move between chromosomal locations and between species (Gillings et al., 2008a); carriage by 0.002% of cells in an unaffected soil (Gaze et al., 2011) compared with as many as 5% of cells in affected soil, fresh water and biofilms (Gaze et al., 2005; Hardwick et al., 2008); ability to acquire a wide range of gene cassettes (Biskri et al., 2005); and frequent association with qac genes that encode versatile efflux pumps (Gaze et al., 2005; Gillings et al., 2009a).

When metagenomic DNA is examined from environmental sources, diverse genes belonging to *intI1* can be detected. In contrast, all examples of *intI1* recovered from clinical contexts have essentially identical DNA sequences, showing that there was a single common ancestor for the 'clinical' class 1 integron that has spread antibiotic resistance among Gram-negative pathogens (Gillings *et al.*, 2008b). Consequently, the class 1 integrons now circulating freely within human-dominated ecosystems have a conserved DNA sequence that, in the main, distinguishes them from the diverse class 1 integrons present in the more general environment.

The best explanation for the origin of the clinical class 1 integron is that a chromosomal class 1 integron from an environmental betaproteobacterium was captured by a transposon of the Tn402 family (Figure 2). This integron carried a gene cassette encoding resistance to disinfectants (qacE), and subsequently captured a gene for sulphonamide resistance (sul1), deleting the terminus of the qacE cassette (Kholodii et al., 1995; Gillings et al., 2008a; Gillings, 2014a).

The Tn402 transposon has the unusual property of targeting the res sites of plasmids (Minakhina et al., 1999) and, consequently, the Tn402-class 1 integron hybrid was able to transpose into a wide variety of plasmids (Figure 2) that then enabled lateral transfer into an equally wide variety of bacterial species. One of the most successful of these insertion events associated the Tn402-integron with a mercury resistance operon (mer) to spawn the Tn21 element that itself went on to generate a series of complex derivatives (Liebert et al., 1999; Partridge et al., 2001). The Tn402-integron has also subsequently generated extensive internal variation by deletion of parts of qacE, sul1 and/or the Tn402 transposition machinery (Hall et al., 1994; Brown et al., 1996; Partridge et al., 2001). Variation in the cassette array has been generated by the collective acquisition of over 130 different antibiotic resistance gene cassettes (Figure 2) (Partridge et al., 2009), conferring resistance to the majority of antibiotics used to control Gram-negative pathogens (Mazel,





Association of a single DNA sequence variant of *intl1* with diverse xenogenetic elements, each composed of a mosaic of plasmids, transposons and resistance determinants

Figure 2 The recent evolutionary origin of the clinical class 1 integron and its incorporation into diverse xenogenetic elements. The raw materials for the assembly of the complex mosaic DNA elements that now carry the clinical *intI1* were all present in the environmental resistome. A single sequence variant from the diverse pool of class 1 integrons in natural environments was captured by a Tn402 transposon, thus forming a Tn402-intI1 hybrid, and giving the integron greater mobility. This hybrid integron, in total, has captured at least 130 different gene cassettes encoding resistance to diverse antibiotics. At the same time, the Tn402 portion of the hybrid element targeted the *res* sites of plasmids, transposing the whole hybrid molecule into a diverse collection of plasmids. This, in turn, promoted movement of clinical *intI1* between different bacterial species by conjugation of those plasmids. Human selection events have also independently fixed the acquisition of diverse resistance genes onto the collection of plasmids invaded by the hybrid integron. These independent acquisitions resulted in the accumulation of genes for resistance to metals, antibiotics, disinfectants and other compounds, along with other genetic elements such as insertion sequences and transposons. As a result, a single molecular species (the clinical *intI1* sequence variant) has become associated with an ever expanding and diverse set of plasmids, transposons and resistance genes. These mosaic elements can be thought of as xenogenetic, in the sense that they owe their current structures and abundance to human activity.

2006; Cambray *et al.*, 2010; Stokes and Gillings, 2011).

Consequently, the 'clinical' intI1 variant is now found on a range of different mobile elements that are freely transmissible between diverse commensal and pathogenic bacteria associated with humans and domestic animals (Nandi et al., 2004; Bailey et al., 2010; Djordjevic et al., 2013; Liu et al., 2013). This 'clinical' *intI1* is also closely linked to various genes that confer phenotypes of environmental significance, such as antibiotic, disinfectant and heavy metal resistances (Figure 2) (Liebert et al., 1999; Norman et al., 2009; Gillings et al., 2009b; Moura et al., 2010; Heuer et al., 2012; Domingues et al., 2013). Finally, 'clinical' intI1 comprises a single molecular species with essentially identical DNA sequences, regardless of the diverse genetic and cellular landscapes they now inhabit (Figure 2).

Antibiotics and antibiotic resistance genes as pollutants

Between 30% and 90% of ingested antibiotic is excreted unchanged by both animals and humans (Sarmah *et al.*, 2006). Antibiotics are only partly removed by wastewater treatment (Giger *et al.*, 2003; Watkinson *et al.*, 2007) and, depending on the antibiotic, can still be found at levels between 10 and 1000 ng l⁻¹ in secondary effluent (Le-Minh *et al.*, 2010). Antibiotics can enter soils via animal manure used for fertilization (Chee-Sanford *et al.*,

2009), whereas other antibiotics are excreted preferentially in urine (Subbiah *et al.*, 2012). As much as 80% of the antibiotics used in aquaculture flow into the environment (Cabello *et al.*, 2013). Consequently, there is a zone around human activities that is enriched with antibiotics.

The use of antibiotics has vastly increased the abundance of 'clinical' class 1 integrons, such that they are now present in up to 80% of enterobacteria in humans and farm animals (Tenaillon et al., 2010; Marchant et al., 2012; Liu et al., 2013). Consequently, large numbers of bacteria containing integrons are released into the environment, with one estimate suggesting that disposal of sewage sludge in the United Kingdom adds 1019 integroncontaining bacteria to waste streams each year (Gaze et al., 2011). Wastewater treatment is not designed to remove DNAs, and the abundance of *intI1* often increases during the water treatment process (LaPara et al., 2011; Ma et al., 2011b, 2013; Chen and Zhang, 2013; Cheng et al., 2013; Du et al., 2014). This might be a consequence of selection driven by the antibiotics, disinfectants and heavy metals that are also inefficiently removed during water treatment (Baker-Austin et al., 2006; Selin, 2009; Hegstad et al., 2010; Rosewarne et al., 2010). As a result, any bacteria that carry class 1 integrons associated with resistance determinants, or that are able to acquire them by lateral gene transfer, would increase in abundance during various stages of water treatment.

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Resistance genes and DNA vectors are increasingly being recognized as environmental contaminants (Pruden et al., 2006; Stalder et al., 2014), and their abundance in natural environments and wild animals has been increasing since the first human use of antibiotics (Knapp et al., 2009; Gillings, 2013). The complex DNA molecules that now bear class 1 integrons often also carry genes for resistance to diverse antibiotics, disinfectants and other environmental contaminants, all embedded in a mosaic of mobile elements. These individual components often have a separate phylogenetic origin, each having been acquired in a separate event, and then fixed by human selection. Because human activities have had a direct role in the selection of sequential gene acquisitions, these complex mosaics of resistance elements can be thought of as xenogenetic. Such xenogenetic molecules have properties of both pollutants and invasive species, as they are pollutants that can replicate (Storteboom et al., 2010; Gillings and Stokes, 2012; Pruden et al., 2012). Methods to control pollution by antibiotics and their respective resistance genes have been suggested, including limiting the use of antibiotics in agriculture, and improving treatment of urban, industrial and hospital waste water (Pruden et al., 2013; Berglund et al., 2014).

Intl1 as a potential marker of anthropogenic pollution

The 'clinical' *intI1* gene has key advantages as a generic marker of anthropogenic influence. These include: universal presence and high abundance in the commensal bacteria of humans and domestic animals, a consequently high representation in waste streams, low abundance in less affected environments and a uniform and highly conserved DNA sequence. Based on these properties, a number of research groups have used quantitative analysis of *intI1* to track human influence (Table 1).

Examining the relationship between pollutants, antibiotic resistance and class 1 integrons reveals a number of general trends (Table 1). *IntI1* is poorly removed during water treatment, and its abundance often increases downstream from water treatment plants and human habitation. The intI1-carrying bacteria are abundant in manure, in digestates from biogas plants and in pesticide biopurification systems (Dunon et al., 2013; Jechalke et al., 2013a). Mesocosms designed to test land application of wastewater solids show that intI1 has a slow decay rate (Burch et al., 2014). In this regard, intI1-carrying bacterial populations are similar to other persistent pollutants, such as metals, antibiotics disinfectants.

The co-occurrence of integrons, resistance genes and pollutants is probably causal, as co-selection of antibiotic resistance genes and integrons occurs in environments polluted with heavy metals and disinfectants (Baker-Austin *et al.*, 2006; Hegstad *et al.*, 2010; Seiler and Berendonk, 2012). This coselection is most likely caused by the physical location of class 1 integrons on a range of transposons and plasmids that also carry genes for resistance to antibiotics, heavy metals and disinfectants (Table 1), and consequently, class 1 integrons can be selected via simple linkage. Similarly, *intI1* abundance has been associated with pesticide pollution, via the co-occurrence of integrons and genes for degradative pathways on IncP-1 plasmids (Dealtry *et al.*, 2014b).

Although the class 1 integron integrase gene does not directly confer resistance to any particular pollutant, its linkage to a diverse suite of antibiotic, metal and disinfectant resistance genes means that it is an excellent de facto measure of the general level of resistance determinants. For instance, there is a strong correlation between the abundance of *intI1* in reclaimed water and the abundance of antibiotic resistance genes such as *sul1* and *tetG* (Wang *et al.*, 2014). Similarly, at the scale of whole watersheds, there is a strong correlation between *sul1*, which is commonly linked to intI1, and the upstream capacities of wastewater treatment and animal feeding operations (Pruden et al., 2012). Because resistance determinants confer selective advantages on those bacterial cells that carry them, intI1 abundance should then reflect the general response of the bacterial community to selection imposed by anthropogenic pollution. Consequently, intl1 abundance should be a good measure of general selective pressure. In contrast, targeting specific resistance determinants such as tet or sul is not a generic measure, as abundance of these genes is dependent on both their presence in a waste source and the presence of specific antibiotics to which they confer resistance.

Towards practical application of *intl1* as a marker

Resistance genes and their vectors originate from environmental sources, where they form part of the resistome (D'Costa et al., 2006; Wright, 2010). This is also the case for *intI1*, which occurs naturally in environmental samples (Figure 2). The use of generic intl1 PCR primer pairs (Stokes et al., 2006) effectively amplifies both clinical and environmental variants of intI1, potentially contributing noise to quantification of intI1 shed from human sources. In environmental samples, intI1 exhibits considerable sequence diversity (Gillings et al., 2008b), whereas the clinical intI1 has a uniform, conserved sequence. For example, the Fungene database (http://fungene.cme.msu.edu/index.spr) (Fish et al., 2013) has over 500 sequences with >99% identity to intI1. These are mostly from clinical isolates, although a few are from environmental strains.



 Table 1
 Environmental and laboratory studies examining the relationship between diverse pollutants, antibiotic resistance genes and class 1 integrons

System	Location	Sample, method	Comments	Reference
Hospital effluent Medical center effluent Sewage treatment Sewage treatment Sewage treatment Sewage mesocosms	France France USA China China USA	E. coli isolation, PCR Aerobic digester, qPCR	IntI1 abundance increases because of effluent IntI1 abundance increases because of effluent IntI1 has longest half-life of genes tested IntI1 abundance increases IntI1 abundance increases in effluent Efficiency of intI1 removal dependent on	
Wastewater treatment	China	Water, qPCR	treatment system Efficiency of <i>intI1</i> removal dependent on	Du et al. (2014)
Wastewater treatment	China	Water, qPCR	treatment system Efficiency of <i>intI1</i> removal dependent on	Chen and Zhang (2013)
Wastewater treatment	China	Water, sediment, qPCR	treatment system intI1 increases in abundance downstream	Zhang et al. (2009)
Wastewater treatment	USA	Water, sediment, qPCR	from city Intl1 abundance significantly increases in effluent	LaPara et al. (2011)
Wastewater treatment	UK	Bacterial isolation, PCR	Shows co-selection of intI1 and disinfectant	Gaze et al. (2005)
Freshwater microcosm	USA	Bacterial isolation	resistance Shows co-selection of antibiotic and metal resistance	Stepanauskas et al. (2006)
Waste streams	UK	Sludge, manure, qPCR	Shows selection of <i>intI1</i> by waste antibiotics/disinfectants	Gaze et al. (2011)
River catchment	Cuba	Sediment, water, qPCR	Ab resistance correlates with degree of pollution	Graham <i>et al.</i> (2011)
River catchment	Pakistan	Water, qPCR	IntI1 and other gene abundance increases with human impact	Khan et al. (2013)
River catchment	USA	Sediment, water, qPCR	IntI1 abundance increases with industrial pollution	McArthur et al. (2011)
Stream catchment	Australia	Sediment, qPCR	IntI1 abundance increases with human impact	Hardwick et al. (2008)
Freshwater habitats	Canada	Water, floc, microarray	IntI1 cassette abundance increases with human impact	Drudge et al. (2012)
Estuary, catchment	France	E. coli isolation, qPCR	IntI1 and Ab resistance correlates with degree of pollution	Laroche et al. (2009)
Estuary	USA	Sediment, water, qPCR	IntI2 abundance increases with human impact	Uyaguari et al. (2013)
Estuary	Canada	Sludge, PCR	IntI1 and diverse cassettes associated with industrial waste	Koenig et al. (2009)
Various	Worldwide	PCR, cloning	IntI2 abundance increases with human impact	Rodríguez-Minguela et al. (2009)
Environ. gradient	USA	Sediment, qPCR	IntI1 abundance increases with metal/anti- biotic pollution	Wright et al. (2008)
Environ. gradient	China	Sediment, sequencing	Integron and plasmid abundance increases with impact	Chen et al. (2013)
Environ. gradient	Argentina	Bacterial isolation, PCR	Trend for <i>intI1</i> to increase in abundance with urbanization	Nardelli <i>et al.</i> (2012)
Environ. gradient	Australia	Sediment, qPCR	IntI1 abundance increases with heavy metal pollution	Rosewarne et al. (2010)
Environ. gradient	Worldwide	Soil, sediment, PCR	IncP plasmid abundance increases with pesticide impact	Dealtry et al. (2014a)
Swine production Slaughterhouse water	Not stated Portugal	Soil and water, qPCR Bacterial isolation, PCR	IntI1 and other genes increase in abundance IntI1 increased in abundance during treatment	Hong <i>et al.</i> (2013) Moura <i>et al.</i> (2007)
Farm manuring Farm manuring Farm manuring Manure, wastewater Manure treatment	Germany UK Germany China China	Soil, rhizosphere, qPCR Soil, qPCR Soil, manure, PCR Water, manure, qPCR Manure, qPCR array	IntI1 and other genes increase in abundance IntI1 increased in abundance IntI1 and other genes increase in abundance IntI1 and other genes increase in abundance Transposons and resistance genes increase in	Cheng et al. (2013)
Animal microbiota	Various	E. coli isolation, PCR	abundance IntI1 increases in frequency with increased	Skurnik et al. (2006)
Archived soils	Scotland	Soil, qPCR	human contact Correlation of resistance genes with copper	Knapp et al. (2011)
Diverse	Various	Review	pollution Shows co-selection of antibiotic and heavy	Baker-Austin et al. (2006)
Diverse	Various	Review	metal resistance Shows co-selection of antibiotic and heavy	Seiler and Berendonk (2012)
	Various	Review	metal resistance Shows co-selection of antibiotic and disin-	Hegstad et al. (2010)

 $Abbreviations:\ Ab,\ antibiotic;\ \textit{intI2},\ class\ 2\ integron-integrase\ gene;\ qPCR,\ quantitative\ PCR.$

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Sequences for environmental variants of *intI1* are still in the minority in databases, and the region of intI1 for which most data are available is that amplified by primers HS464/HS463a (Gillings et al., 2008b). Examination of the sequence data (Supplementary Table S1) reveals a number of nucleotide positions where the clinical *intI1* can be distinguished from most reported environmental variants. A primer pair targeting intI1 nucleotide positions 165-184 and 456-476 (intI1F165 5'-CGA ACGAGTGGCGGAGGGTG-3' and intI1R476 5'-TAC CCGAGAGCTTGGCACCCA-3') is one possibility for specifically amplifying the clinical version of *intI1*. Because these primers target the clinical *intI1* sequence, but not the diverse intI1 variants known to be present in environmental bacteria, they should allow a more precise quantitative analysis. As more complete sequences from environmental variants of intI1 become available, better regions for discrimination could be identified.

Sample collection and processing

Environmental monitoring of human impact and the efficacy of remediation could be conducted using quantitative analysis of intI1 abundance. Careful consideration should be given to sampling strategies and data generation. Samples of sediment, soil or water should be taken in a uniform, reproducible manner. The likelihood of temporal variation should be taken into account. For instance, sewage treatment water can vary considerably over a 24-h period, and composite or flow proportionate samples should be considered. Ideally, each sampling time or point should be represented by at least triplicate samples, to be treated as triplicates in all subsequent steps such as DNA extraction and quantitative PCR (qPCR). Each sampling point for soil or sediment can be laid out in a grid to capture microvariation. At the minimum, samples should be identified by date, GPS coordinates and land use. The GSC (Genomic Standards Consortium) provides a guide to collection of environmental data under its MIMARKS environmental packages (Yilmaz et al., 2011), conveniently implemented by RDP according to habitat type with prepopulated googlesheets (http:// rdp.cme.msu.edu/wiki/index.php/RDP_MIMARKS_ GoogleSheets). For a detailed description of one multipurpose soil sampling procedure, see the BASE website (http://www.bioplatforms.com.au/ special-initiatives/environment/soil-biodiversity/ sample-collection-procedure). Soil can be stored at 4 °C, or snap-frozen immediately upon collection, and maintained frozen during transport to minimize changes to microbial populations.

IntI1 monitoring could be used for analysis of water samples, such as wastewater effluents, feedlot runoff and affected streams, rivers, lakes and oceans. Water samples can be collected by bulk grab techniques, using methods described for coliform

monitoring. Water samples contain a particulate fraction, and many microbes, including microbes carrying intI1, attach to particulate matter suspended in the water. Most methods employ filtration, with 0.22 µm cutoff capturing the majority of bacteria and other particulates. The filter is then directly subject to DNA extraction. However, extracellular DNA may also be of interest, and this will pass through filters under some conditions. Recent studies have introduced techniques for analysing extracellular forms of antibiotic resistance genes (Mao et al., 2013). Further assessment and standardization of filter membrane composition and pore size employed for analysis of *intI1* in water samples for different purposes (extracellular versus intracellular) would be of interest.

Sufficient sample should be taken for multiple analyses, and for archival storage. To ensure representative subsampling, the cone and quarter method can be used (Ferrari *et al.*, 2008). The DNA extraction method employed should be suitable for diverse cell types and for removal of inhibitors present in soil, sediment, manure, sludge and other intractable substrates (Yeates and Gillings, 1998; Gillings, 2014b). The integrity of extracted DNA should be assessed using agarose electrophoresis and the concentration estimated photometrically. Alternatively, double-stranded DNA could be quantified using fluorometric methods (Singer *et al.*, 1997).

Concentrations of *intI1* can then be determined using real-time qPCR, correcting by the total bacterial abundance as measured by 16S rRNA gene PCR performed on the same sample. Ideally, three independent environmental samples should be processed in parallel to control for variation introduced during processing. PCR inhibition caused by co-extracted compounds can be overcome using bovine serum albumin (Gaze et al., 2011), an environmental master mix or template dilution. Primers for amplification of 16S rRNA genes should be specific for bacteria (Nadkarni et al., 2002). Primer sets need to be optimized across a range of concentrations and annealing temperatures. Standard curves for each target gene need to be determined, and positive control standards of known copy number prepared by PCR (Hardwick et al., 2008; McKinney and Pruden, 2012). The qPCR results could also be normalized by the total DNA in a sample, which would generate an idea of the relative abundance of *intI1* in relation to the entire metagenome. If a housekeeping or other gene is used to normalize intI1 abundance, it should be established that it has a similar amplification efficiency to that of *intI1*.

Rapidly advancing molecular technologies will add new capabilities for understanding human impact. Highly parallel qPCR equipment from several vendors allows analysis of multiple primer sets and samples, such that hundreds of antibiotic resistance genes, mobile elements and their variants



can be analysed (Looft *et al.*, 2012; Zhu *et al.*, 2013), allowing more comprehensive spatial and temporal studies. Furthermore, amplicons can be sequenced, providing diagnostic-level insight into the probable origin of these genes. A recent example of the latter shows clusters of *intI1* and antibiotic resistance gene identities at a country scale and at an intercontinental scale for *intI1* (Johnson *et al.*, 2014).

As DNA sequencing becomes more efficient and cheaper, direct sequencing of metagenomic samples may replace qPCR approaches. In such an analysis, clinical *intI1* sequences could be extracted from the sequence data and normalized to a single copy housekeeping gene. Already, such approaches are being used, based on high-throughput next-generation sequencing methods.

Conclusion

The clinical version of the intI1 gene has some unique advantages as a universal marker of selective pressures imposed by anthropogenic pollution. Its recent emergence into human-dominated ecosystems means that it has a homogenous and conserved DNA sequence, simplifying detection. It has seen a rapid increase in abundance and geographic distribution, fuelled by the extensive use of antibiotics and its insertion into diverse mobile elements, coupled with its penetration into a wide range of bacterial species associated with human-dominated ecosystems. During this expansion, it has become closely linked with genes that confer resistance to disinfectants and heavy metals, as well as the wide range of antibiotic resistance determinants for which it is well known. Consequently, versions of the clinical intI1 gene are capable of conferring diverse advantages to those cells that carry them, and these advantageous phenotypes correspond with the selective agents that are most likely to be present in human waste streams.

Conflict of Interest

The authors declare no conflict of interest.

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Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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