

Chromosomal integrons are genetically and functionally isolated units of genomes

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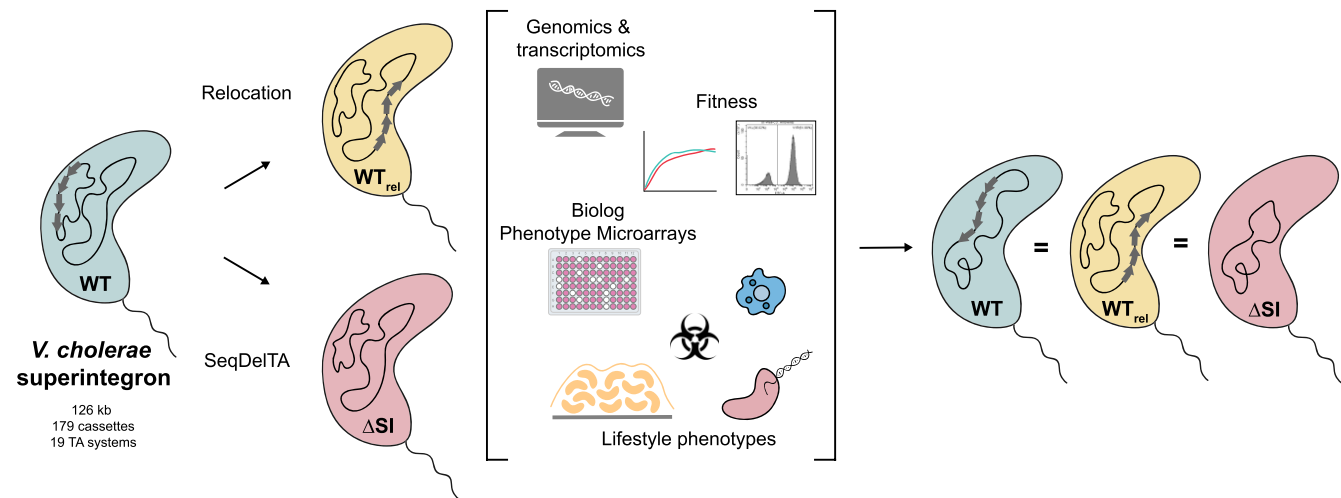
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Abstract

Integrons are genetic elements that increase the evolvability of bacteria by capturing new genes and stockpiling them in arrays. Sedentary chromosomal integrons (SCIs) can be massive and highly stabilized structures encoding hundreds of genes, whose function remains generally unknown. SCIs have co-evolved with the host for aeons and are highly intertwined with their physiology from a mechanistic point of view. But, paradoxically, other aspects, like their variable content and location within the genome, suggest a high genetic and functional independence. In this work, we have explored the connection of SCIs to their host genome using as a model the Superintegron (SI), a 179-cassette long SCI in the genome of *Vibrio cholerae* N16961. We have relocated and deleted the SI using SeqDelTA, a novel method that allows to counteract the strong stabilization conferred by toxin–antitoxin systems within the array. We have characterized in depth the impact in *V. cholerae*'s physiology, measuring fitness, chromosome replication dynamics, persistence, transcriptomics, phenomics, natural competence, virulence and resistance against protist grazing. The deletion of the SI did not produce detectable effects in any condition, proving that—despite millions of years of co-evolution—SCIs are genetically and functionally isolated units of genomes.

Graphical abstract



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Introduction

Integrations are genetic elements that allow bacteria to adapt swiftly to changing environments (1–3). They recruit new genes encoded in small mobile genetic elements called integron cassettes (IC) (or gene cassettes), stocking them in an array to form a genetic memory of adaptive functions. According to the needs of their hosts, integrons can also modulate cassette expression by reshuffling their order within the array (4,5). The integron platform comprises the gene encoding the integrase (*intI*)—the recombinase that governs all reactions—the integration site where cassettes are incorporated (*attI* site), and two promoters in opposite orientations that drive the expression of the integrase (P_{int}) and that of cassettes in the array (P_c), which are generally promoterless (Figure 1A). Integrations are well known for their role in the rise of multidrug resistance and the spread of resistance genes among clinically relevant Gram-negative species (6–8). Encoded on plasmids, these Mobile Integrations (MIs) act as vehicles for more than 170 resistance genes against the most relevant families of antibiotics (9,10). Yet, MIs are only a small subset of all integrations that have been mobilized from the chromosomes of environmental bacteria by transposons. The vast majority of integrations in nature are sedentary chromosomal integrations (SCIs), which are found in a variety of phyla and have an ancient origin (11–13). These SCIs are a highly variable region of the genome, representing a hotspot for genetic diversity (14). On the antipodes of what is observed for MIs, the functions encoded in SCIs are generally not related to antimicrobial resistance and are mostly unknown (15). The colossal repertoire of cassettes in the environment holds a high potential for biotechnology that is yet to be explored (16).

The paradigm of SCI is the Superintegration (SI) (17), a massive structure encoded in the secondary chromosome of *Vibrio cholerae*, the causative agent of cholera disease. In strain N16961, where it was first described, the SI is 126 kb long, embodying 3% of the genome of this bacterium and containing 179 ICs. Although the model of integron predicts that the SI should be mostly silent, a recent RNA-seq analysis shows that many cassettes are expressed at biologically relevant levels (18). The functions of cassettes are mostly cryptic, except for the notable exception of 19 cassettes that encode toxin–antitoxin (TA) systems (19,20) and a chloramphenicol resistance gene (21) (Figure 1B). TA cassettes are scattered along the SI and serve to stabilize the array through a post-segregational killing mechanism: if the TA system is lost, the labile antitoxin is degraded before the more stable toxin, leading to cell death (22,23).

Despite being the best-studied SCI, the knowledge on the SI is limited for two reasons. First, the long cassette array interferes with the genetic tools commonly used in the field to deliver recombination experiments that would allow the characterization of the system. Second, its integrase cannot be studied in heterologous genetic backgrounds because it seems to require additional host factors (24–26).

SCIs have co-evolved with the genome of the host for millions of years (27,28). Indeed, large SCIs like the SI are ubiquitous in *Vibrio* species, and the phylogenetic signal of integrases mimics that of the species (1). This suggests that their acquisition predates the radiation of the genus, an event that occurred >300 million years ago (29). The timescale of this co-evolution process has allowed for the intertwining between host physiology and integron activity at many different levels.

A blatant example is that integrase expression is under the control of the host's SOS response. Yet, many other aspects are rather more subtle and represent better the depth of such connections (30). Examples of this are (i) the balanced interplay between the integrase and single-strand binding (SSB) proteins of the host on *attC* sites, that impede *attC* site folding to avoid its formation unless needed for recombination, i.e. when the integrase is expressed (31,32); or (ii) the fact that SCIs have a specific orientation related to the origin of replication, to maximize their carrying capacity (23,33). A striking observation is that some aspects of SCIs remain extremely plastic, such as their content and their location—despite their size, SCIs can be found in either chromosome depending on the species of *Vibrio*. Hence, it remains unclear whether the intertwining between host and integron is restricted to the functioning of the platform and the mechanistic of recombination (P_{int} and P_c promoters and *attC* sites), or if it is extensive to the array of functions encoded in the variable part. In other words, we ignore if cassettes are streamlined to act as plug-and-play add-ons to the pool of functions encoded in the genome without interacting with it, or whether they can modify or interfere with the host physiology in any other way.

In this work, we have addressed the question of whether chromosomal integrations are genetically and functionally isolated units of the genome by relocating and deleting the SI from *V. cholerae*. To overcome the high stabilization of the structure by the TAs, we have designed an approach called SeqDelTA (Sequential Deletion of Toxin–Antitoxin Systems) that exploits natural competence and homologous recombination to deliver a set of 16 consecutive allelic replacements. At each step, a toxin is inactivated without altering the cognate antitoxin, deleting a piece of the SI. After the last allelic replacement, we produce the scarless deletion of the SI through a counter-selectable suicide vector. The resulting strain has been sequenced, and unintended mutations located elsewhere in the genome have been corrected. Incidentally, this strain represents a milestone in the study of chromosomal integrations, since it provides, for the first time, the possibility of delivering experimental studies using common tools, without interference from the SI. To understand how the deletion of the SI affects *V. cholerae*, we have characterized in depth this strain from a variety of perspectives, including chromosome replication dynamics, fitness, natural competence, antibiotic persistence, transcriptomics, phenomics, virulence and protist grazing assays. We observe no significant variations in the physiology of the Δ SI strain compared to the wild-type. We conclude that SCIs are functionally isolated units of the genome, despite eons of co-evolution with their hosts. This has strong implications in the type of functions one can expect to find in ICs.

Materials and methods

SI relocation from Chr2 to Chr1

All strains, plasmids and primers used for the SI relocation are listed in Supplementary Tables S1, S4 and S5. To relocate the whole SI, we used a genetic tool based on the recombination of two bacteriophage attachment sites (34). Prophage excision from the host chromosome relies on site-specific recombination between two sequences flanking the phage, termed attachment sites *attL* and *attR*. This recombination is carried out by phage-specific recombinases 'Int' and their cognate RDF

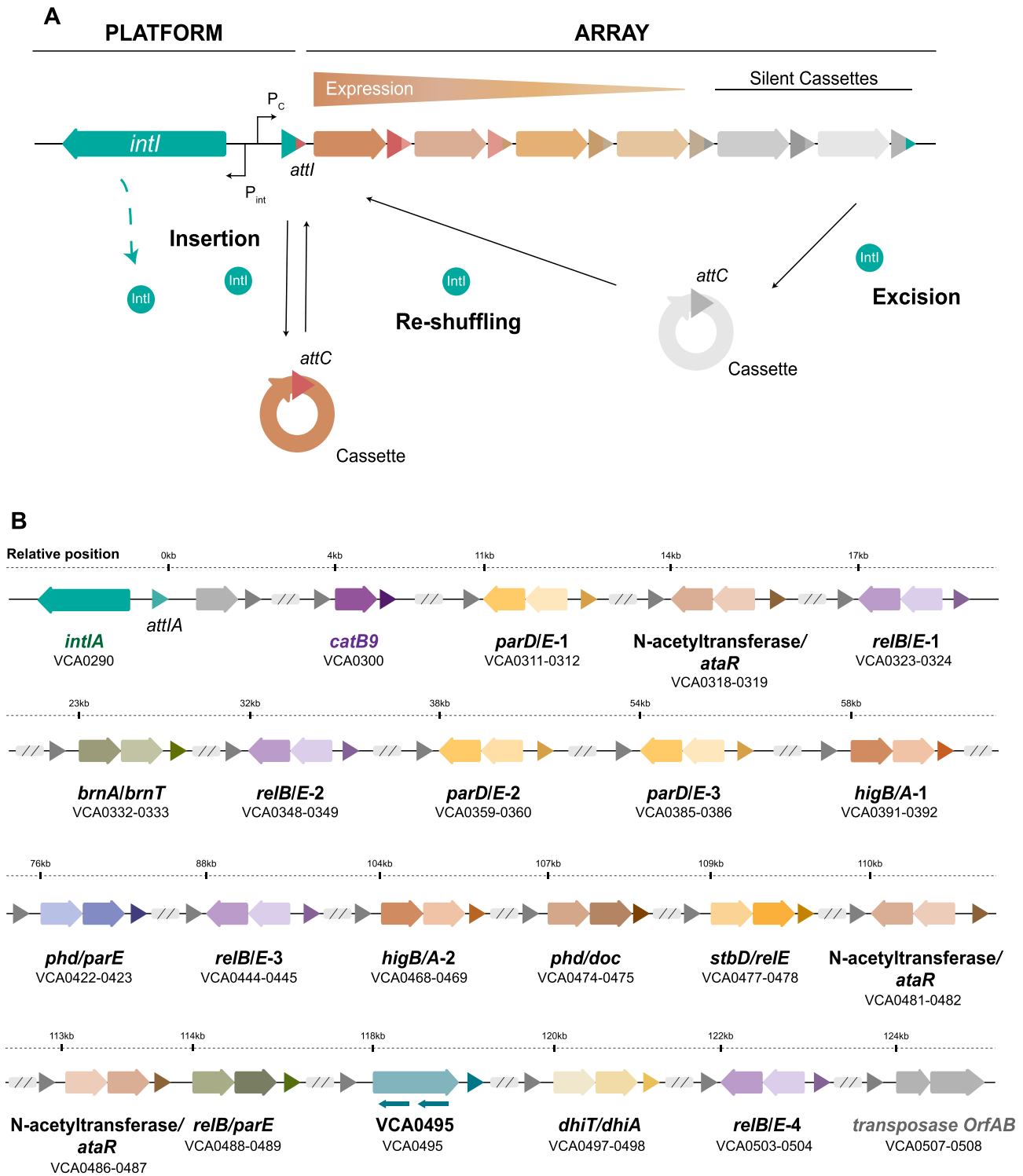


Figure 1. (A) Schematic representation of an integron. On the left side, the functional platform includes the integrase gene (*int1*), the cassette promoter (P_c), the integrase promoters (P_{int}) and the integron insertion site (*attI*). On the right side, the variable array of ICs is shown. Hybrid *attI* and *attC* sites are marked with corresponding colors. The arrows within the cassettes indicate the orientation of the open reading frame, with the color intensity of each arrow representing the expression level. The diagram also illustrates the primary reactions mediated by the *IntI* recombinase: cassette insertion ($attI \times attC$), excision ($attC \times attC$) and reshuffling. (B) Diagram showing the organization and relative position of the relevant genes within the SI cassette array. Arrows represent individual genes contained within cassettes, while gray triangles denote the *attC* sites. TA systems are illustrated with colored arrows: darker arrows indicate toxins and lighter arrows indicate antitoxins. The orientation and position of each arrow reflect the transcriptional direction of the genes. Additionally, smaller arrows mark regions corresponding to non-coding RNAs.

(recombination directionality factor), or excisionase, Xis. This tool uses the attachment sites from bacteriophages HK and λ (Supplementary Figure S1). The *attL/attR* pairs of each site were associated with parts of a genetic marker that is reconstituted when the sites recombine: HK sites were associated with parts of a *bla* gene (ampicillin resistance) and the λ sites with parts of *lacZ*. Co-expression of the HK and λ integrases/excisionases triggers the recombination of the partner *attL/attR* sites, which allows for the reconstitution of fully functional *bla* and *lacZ* markers and the selection of clones in which the SCI relocation has occurred. Blue colonies growing in LB supplemented with carbenicillin and X-gal (5-bromo-4-chloro-3-indolyl- β -D-galactopyranoside) were polymerase chain reaction (PCR) verified and used for growth curves. The SI was relocated between genes VC0018 and VC0019 in both orientations (Figure 2A).

SeqDelTA

All strains, plasmids and primers for the deletion of the SI are listed in Supplementary Tables S2, S3, S4 and S5. To generate the Δ SI mutant of *V. cholerae* N16961, we developed SeqDelTA, as depicted in Figure 3. After identifying the 19 TA systems contained in the *V. cholerae* N16961 SI, we prepared successive constructs designed with homology regions (HR) and a selective marker. A left homology region (LHR) was designed and maintained throughout the procedure. This region corresponded to the 5' region of the *catB9* gene (VCA0300), and was amplified with primers VCA 299 F and LRHI VCA 300 R. For the successive deletions, different right homology regions (RHD) were designed to maintain antitoxin gene but to eliminate part of the corresponding toxin gene, ensuring that the event was not lethal to the bacteria (Figure 3 and Supplementary Table S2). As selective markers, three different antibiotic resistance genes (Zeo^R, Cm^R and Carb^R) were used and amplified with the same primers, facilitating the consequent design and assembly of the HRs. These three fragments (LHR, resistance marker and RHR) were then assembled by splicing by overlap extension PCR (SOE-PCR), placing the resistance marker between the LHR and the RHR. Each successive construct was introduced by natural transformation (described below) into the *V. cholerae* strain obtained from the previous TA system deletion. This strategy allowed the removal of the resistance marker introduced in the previous step while advancing the deletion of TA systems by changes in the RHR. When necessary, due to gene orientation and/or gene order in the TA system, alternative LHRs were designed, and two resistance markers were maintained in consecutive allelic replacements before the new step allowed deletion of both (Supplementary Table S2).

The last deletion was performed using the R6K origin of replication-plasmid pMP7 as described in (25,35). pMP7 replication is dependent of the cell π protein and encodes the *ccdB* toxin under the control of the inducible promoter P_{BAD}. Five hundred bp of each side of the SI were amplified with primers RHI link pMP7 F and LRHI + D R for the LHR, and RHD F and RHD link pMP7 R for the RHR. The two fragments were then joined by SOE-PCR and digested with enzymes NaeI y HindIII. Finally, the insert was ligated with the empty pMP7 backbone, giving rise to pMP7_ Δ *intIA attIA* zeo^R. We started the cloning steps by transforming the pMP7_ Δ *intIA attIA* zeo^R plasmid into *ccdB*-resistant *Es-*

cherichia coli π 3813 (*thy::pyr* +) competent cells. Transformants were selected in LB agar plates containing Cm (25 μ g/ml), thymidine (dT; 0.3 mM) and glucose (1%). For conjugal transfer of pMP7_ Δ *intIA attIA* zeo^R into *V. cholerae*, the plasmid was first transformed into the donor strain *E. coli* β 3914 (*dap::pir* +). Transformants were selected in LB agar plates supplemented with Cm (25 μ g/ml), 2,6-diaminopimelic acid (DAP; 0,3 mM) and glucose (1%). Conjugation assay was performed by growing both donor (*E. coli* β 3914) and recipient (*V. cholerae* A066) strains to a OD₆₀₀ of 0.2. Then, cells were mixed in a ratio of 1/10 (donor/recipient) and transferred to a mating filter placed on a LB agar plate supplemented with DAP (0.3 mM) and glucose (1%) and conjugation was performed overnight at 37°C. Integration of the pMP7_ Δ *intIA attIA* zeo^R into the *Vibrio* genome was selected by washing the filter in 5 ml of LB and plating serial dilutions on Cm (2.5 μ g/ml) and glucose (1%) plates, but lacking DAP. Afterward, Cm-resistant colonies were grown in liquid LB medium and plated in LB agar plates supplemented with L-arabinose (0.2%) to express *ccdB* and select for the second crossover that implies the excision of the pMP7 backbone. After this process, we Sanger-sequenced the SI region and selected a *V. cholerae* Δ SI strain (A101).

Natural transformation assays

Transformation assays were performed as previously described in (36) with some modifications. Briefly, strains were inoculated from frozen stocks into LB broth and grown rolling overnight at 30°C. Overnight cultures were diluted 1:100 and grown to an OD₆₀₀ of 1.0. Cells were then washed and resuspended in DASW (defined artificial seawater) (Instant Ocean[®]) 0.5 \times and inoculated in a final volume of 1 ml DASW with 150 μ l of chitin (Apollo Scientific[®]) slurry. Cells were incubated in static conditions at 30°C for \sim 18 h. Then, 550 μ l of supernatant was removed from the reactions, and the purified transforming DNA (tDNA) was added. DNA and cells were incubated for \sim 18 h in static conditions at 30°C. When needed, reactions were outgrown by adding 1 ml LB broth and shaking at 30°C for 2 h. Cells were plated in LB agar with the corresponding antibiotic.

To measure natural competence frequencies, cells were incubated with 2 μ g of a PCR product carrying a kanamycin resistance cassette flanked by 1 kb long homology regions targeting the *lacZ* gene. Primers and strains used for this assay are listed in Supplementary Tables S3 and S5. In these assays, cells were incubated for 8 h with tDNA before plating in selective medium (LB plates supplemented with kanamycin 75 μ g/ml). Frequencies were calculated as the number of colony-forming units (CFUs) growing in kanamycin over the total CFUs.

DNA isolation, Whole Genome Sequencing (WGS) and data analysis

Genomic DNA of the obtained *V. cholerae* Δ SI strain (A101) was extracted using the DNeasy[®] Blood and Tissue Kit (QIAGEN) following the manufacturer's protocol. The DNA quantity was determined using a BioSpectrometer (Eppendorf). WGS sequencing was performed at Institut Pasteur using Illumina paired-end reads, at MiGS (Pittsburgh, USA), now SeqCoast Genomics (Portsmouth, USA) and inhouse at MBA (Madrid) using MinIon. Data analysis was accomplished with Geneious Prime software and the genetic variants were

identified by mapping the generated reads to the *V. cholerae* N16961 reference genome (Accession number: chromosome 1 CP028827.1; chromosome 2 CP028828.1).

Marker Frequency Analysis (MFA)

Marker frequency analysis (MFA) was performed as described in (37). Cells were grown in LB at 37°C with shaking and genomic DNA was isolated using the DNeasy® Tissue Kit (Qiagen) from exponential and stationary phase cultures. Genomic DNA was sequenced with Illumina technology. Short reads were aligned to the reference genome using Bowtie2 (38) and R2R. The number of reads starting per bp (*N*) were calculated for 1 and 10 kbp windows. The exponential phase data were normalized with a factor calculated as the deviation of observed *N* from calculated *N* for each window of the stationary phase culture data. Any window including repeated sequences was excluded and *N* was plotted as a function of the absolute position, centered on the replication origins.

Correction of *V. cholerae* ΔSI mutations

All the strains, plasmids and primers are listed in Supplementary Tables S3, S4 and S5. Three mutations were found in *V. cholerae* ΔSI strain (A101): *rocS* (VC0653), *cry2* (VC01392) and *rpoS* (VC0534). While *rocS* and *rpoS* mutations led to frameshifts, *cry2* contained a non-synonymous mutation. To revert the mutations to the WT variant, allelic exchanges were performed using the pMP7, as previously described. To provide homology for the allelic replacement, 1000 bp-fragments of the N16961 strain genes *rocS*, *rpoS* and *cry2* were amplified with primers *rocS*_pMP7 F/R, *rpoS*_pMP7 F/R and *cry2*_pMP7 F/R, respectively, which contain 20 bp-homology fragments with the pMP7 cloning site; pMP7 backbone was amplified with primers pMP7_bb_Gibson F/R. Cloning of the amplified fragments was performed using Gibson assembly (39), giving rise to pMP7_*rocS*, pMP7_*rpoS* and pMP7_*cry2*. We performed the mating and allelic exchanges as previously mentioned, starting with pMP7_*rocS*, then pMP7_*rpoS*, and finally, pMP7_*cry2*. After each step, we Sanger-sequenced the targeted gene and selected a *V. cholerae* strain corrected for *rocS*+ (A677), *rocS*+, *rpoS*+ (A684), and finally *rocS*+, *rpoS*+, *cry2*+ (B522). We verified by Illumina whole-genome sequencing that *V. cholerae* ΔSI (B522) did not contain any other unintended mutations.

RNA extraction and preparation for RNA-seq

Total RNA was extracted from *V. cholerae* WT and ΔSI cultures grown in LB at 37°C in both exponential (OD₆₀₀ 0.8) and stationary (OD₆₀₀ 2.8) growth phases using the RNeasy® Mini Kit (QIAGEN), following the manufacturer's protocol. To eliminate any residual DNA, RNA was treated using the TURBO DNA-free™ Kit (Invitrogen). RNA concentration was measured using a BioSpectrometer (Eppendorf), and RNA integrity was determined using a Qubit™ 4 fluorometer (Invitrogen) with the RNA IQ Assay Kit (Invitrogen). Ribodepletion RNA library and sequencing was performed at the Oxford Genomics Centre using a NovaSeq6000 sequencing system (Illumina). Three biological samples per condition were sequenced.

RNA-seq data analysis

Raw Illumina reads were trimmed and adapter-removed using Trim Galore v0.6.6 (<https://github.com/FelixKrueger/TrimGalore>) with a quality threshold of 20 and removing reads <50 bp. Trimmed paired reads were mapped to the *V. cholerae* N16961 reference genome (accession number: chromosome 1 CP028827.1; chromosome 2 CP028828.1) using BWA-MEM v0.7.17 (40). featureCounts from the Rsubread v2.10.2 package (41) was used to obtain read counts per annotated feature, including coding sequences (CDS), non-coding RNA, transfer-messenger RNA, ribonuclease P, tRNA, antisense RNA and signal recognition particle RNA. Differential expression analysis was performed from raw counts using DESeq2 v1.36.0 (42), by comparing the expression data of the deletion mutant against the WT strain.

Differentially expressed genes (DEGs) of CDSs were annotated with Gene Ontology (GO) terms retrieved from UniProt on 7 July 2023 (43). Gene set enrichment analysis (GSEA) was performed to identify sets of biological processes, molecular functions or cellular components with different expression profiles between the SI deletion mutant and WT strains, in both exponential and stationary phases. For this, a pre-ranked list of DEGs, ordered by log₂ fold changes, was provided to the GSEA function of the clusterProfiler v4.8.1 (44). Genes belonging to the SI (positions 31 2057–43 8942 bp of chromosome 2) were not included to prevent biased enrichment results caused by the acute downregulation of the superintegron genes. GSEA was run with default parameters, correcting for multiple tests with the Benjamini–Hochberg procedure.

Reverse transcription-quantitative PCR (RT-qPCR)

Independent samples of total RNA from *V. cholerae* WT and ΔSI cultures grown in LB at 37°C in both exponential (OD₆₀₀ 0.8) and stationary (OD₆₀₀ 2.8) were isolated as described previously. To eliminate any residual DNA, RNA was treated using the TURBO DNA-free™ Kit (Invitrogen) and RNA concentration was measured using a BioSpectrometer (Eppendorf). Before cDNA synthesis, a previous DNA wipe-out step was performed using the QuantiTect® Reverse Transcription Kit (QIAGEN) following the manufacturer's instructions; cDNA synthesis was carried out afterward using the same kit using the temperature steps: 42°C for 15 min; 95°C for 3 min.

Reverse transcription quantitative PCR (RT-qPCR) was performed in an Applied Biosystems QuantStudio 3 using the Fast SYBR™ Green Master Mix (Thermo Fischer Scientific). Primers used are listed in Supplementary Table S5, and *gyrA* was used as housekeeping gene. Relative changes in gene expression for the SI deletion mutant with respect to the WT were determined according to the threshold cycle method ($2^{-\Delta\Delta CT}$). Mean values were obtained from two independent biological replicates with three technical replicates each.

Note on the controls used for the search for phenotypes

The original *V. cholerae* N16961 isolate contains mutations in the *hapR* gene and is not naturally competent. The deletion of the SI has been performed in a well-known derivative strain in which natural competence is restored by the insertion of a transposon carrying a functional *hapR* gene (strain A001 in the Supplementary Table S3). It is known that *V. cholerae* cultures have a tendency toward the *hapR*⁻ genotype. Through genome sequencing we observed this phe-

nomenon in our stock of the parental strain, which seemed to rise through homologous recombination between both *hapR* alleles. Since *hapR*⁻ strains have certain phenotypes, such as distinct growth curves or a decreased virulence, we performed most phenotypic experiments using as a control a strain of *V. cholerae* N16961, in which mutations in *hapR* have been corrected without the addition of a second allele (A096 in our strain list). In RNA-seq and MFA experiments we could use as a control the parental strain (A001), by confirming through sequence analysis that all replicates contained a functional *hapR* allele.

Growth curves

Overnight cultures were prepared by inoculating single colonies from LB agar plates in LB liquid medium and incubating for 16–20 h at 37°C. Cultures were then diluted 1:1000 in fresh medium and 200 µl were transferred into 96-well plates (Nunc, Thermo Scientific). OD₆₀₀ was measured every 15 min for 16 h (36 h for 20°C) using a Biotek Synergy HTX plate reader (Agilent) at different temperatures (20, 30, 37 and 42°C). V_{max} and area under the curve (AUC) were determined using Gen5 software and MATLAB, version R2022a (Mathworks), respectively. At least 10 independent replicates were included in the experiment and significant differences were assessed by performing an unpaired *t*-test.

Growth curves in the presence of subinhibitory concentrations of norfloxacin, ceftriaxone and tannic acid were performed by diluting the compounds at the desired concentrations in MH medium. Overnight cultures of *V. cholerae* WT and ΔSI were diluted 1:1000 in the antimicrobial-containing MH media in 96-well plates (Nunc, Thermo Scientific). OD₆₀₀ was measured every 20 min for 24 h using a Biotek Synergy HTX plate reader (Agilent). At least three biological replicates were included in the experiment.

Competition assays

Competition assays were performed by flow cytometry to measure the fitness values of *V. cholerae* WT, the ΔSI mutant and the corrected intermediate deletion strains, relative to an *E. coli* DH5α strain as a common competitor carrying a pSU38 plasmid with the *gfp* under the control of the constitutive promoter P_CS (strain A370, [Supplementary Table S3](#)). The competition procedure was performed as described in (45) with some modifications. Briefly, pre-cultures were prepared by inoculating single colonies from a LB agar plate in 200 µl of liquid LB in a 96-well plate (Nunc, Thermo Scientific). After 22 h of growth at 37°C with 250 rpm shaking, cultures were mixed at 1:1 proportion and diluted 1:400 in fresh medium. In order to confirm the initial proportions of *E. coli* carrying *gfp* and the non-fluorescent *V. cholerae* strains, cells were diluted 1:400 in NaCl (0.9%) and 30,000 events per sample were recorded using a CytoFLEX S flow cytometer (Beckman Coulter). Bacterial mixtures were competed for 22 h at 37°C with 250 rpm shaking, and final proportions were determined as stated above. The fitness values of both *V. cholerae* WT and ΔSI, relative to *E. coli* P_CS::*gfp* were calculated using the formula: $w = \ln(N_{\text{final,gfp-}}/N_{\text{initial,gfp-}})/\ln(N_{\text{final,gfp+}}/N_{\text{initial,gfp+}})$, where *w* is the relative fitness of the non-GFP-tagged *V. cholerae* strains, *N*_{initial,gfp-} and *N*_{final,gfp-} are the numbers of non-GFP-tagged *V. cholerae* at the beginning and end of the competition, and *N*_{initial,gfp+} and *N*_{final,gfp+} are the numbers of *E. coli* P_CS::*gfp* cells at the beginning and end of the competition,

respectively. Twelve/six biological replicates were performed for each competition experiment and significant differences were assessed by performing unpaired *t*-test.

Phenotypic microarray assay

Phenotype Microarrays (Biolog) PM1-PM20 were tested as previously described in (46) with some modifications. Briefly, single colonies of *V. cholerae* WT and ΔSI were inoculated in liquid LB medium and grown overnight at 37°C. For PM1 and PM2 plates inoculation, 120 µl of Redox Dye mix D (100×), 1 ml of sterile water and 880 µl of a 85% T-cell suspension in NaCl (0.9%) were added to 10 ml of IF-0 (1.2×), resulting in a final volume of 12 ml per plate. For PM3-PM8 plates inoculation, which requires an appropriate carbon source, 120 µl of sodium pyruvate (2 M) was added as an additive and mixed with 120 µl of Redox Dye mix D (100×), 880 µl of sterile water and 880 µl of a 85% T-cell suspension in NaCl (0.9%). Then, this solution was added to 10 ml of IF-0 (1.2×), resulting in a final volume of 12 ml per plate. For PM9-PM20 plates inoculation, 120 µl of Redox Dye mix D (100×), 1 ml of sterile water and 880 µl of 1:200 dilution of the 85% T-cell suspension in NaCl (0.9%) were added to 10 ml of IF-10 (1.2×), resulting in a final volume of 12 ml per plate. Each plate was inoculated with 100 µl per well of the respective prepared solution and then sealed using Breathe-Easy® sealing membranes (Sigma-Aldrich). Plates were then transferred to a Biostack Microplate Stacker (Agilent) to process several plates per day and incubated at 37°C using a Memmert IF750 (Mettler). OD₅₉₀, which indicates the reduction of the tetrazolium dye, was measured every 20 min during 24 h using a Biotek Synergy HTX plate reader (Agilent). Experiments were conducted in duplicate and growth characteristics were determined by calculating the AUC using MATLAB, version R2022a (Mathworks).

Antibiotic persistence assay

Three colonies of *V. cholerae* WT and ΔSI were inoculated in LB liquid medium and grown overnight at 37°C. Overnight cultures were diluted 1:100 in 1 ml of fresh LB medium in 24-well plates and grown at 37°C on 150 rpm shaker for 3 h. After 3 h incubation, cultures were challenged individually with 10-fold minimum inhibitory concentration (MIC) of ciprofloxacin (MIC = 0008 µg/ml) or ampicillin (MIC = 16 µg/ml) and incubated for 6 h at 37°C on 150 rpm shaker. The CFUs/ml were enumerated at times 0 h (before adding the antibiotics), 2, 4 and 6 h by serially diluting the cells and plating on LB agar.

Resistance to protozoan predation

Tetrahymena ellioti strain 4EA was routinely cultured at 28°C in 2% proteose peptone supplemented with 10 µM FeCl₃ and 250 µg/ml of both streptomycin sulfate and penicillin G. Log-phase *V. cholerae* bacteria (OD₆₀₀ = 0.6) and ciliates (3 × 10⁵ cells/ml) were washed and suspended in 0.01 M Tris-HCl pH 7.5 buffer. Co-cultures were established by combining ciliates and bacteria (1:200 ratio) in a 96-well plate (Costar) for up to 48 h at 28°C. Control bacteria were incubated in 0.01 M Tris-HCl pH 7.5 buffer in the absence of ciliates for the same period. Every 24 h, intracellular bacteria were released by lysing protozoan cells with 1% Triton X-100. A number of viable bacteria were quantified by CFU counts on LB agar plates.

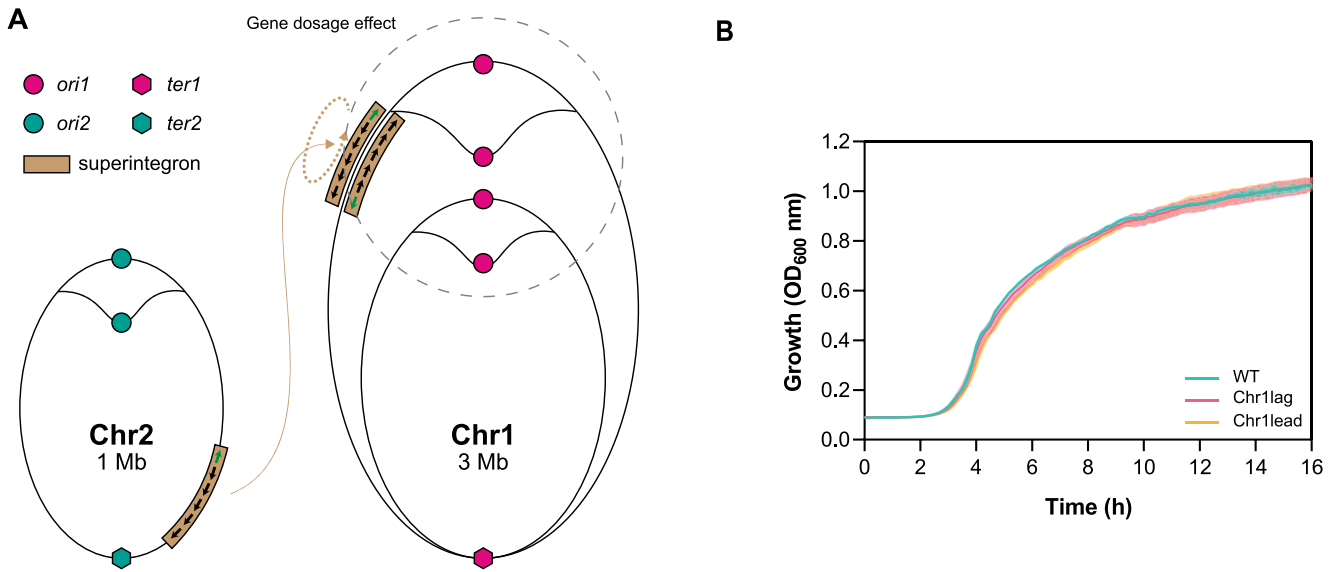


Figure 2. (A) Scheme of the relocation of the SI from Chr2 to Chr1, where it is inserted in both orientations (modified from (76)). (B) Growth curves of WT and relocated-SI strains (Chr1lag and Chr1lead). No significant growth differences are observed.

Acanthamoeba castellanii strain Neff was routinely propagated at 30°C in peptone-yeast-glucose (PYG) medium (Pronadisa) supplemented with 250 µg/ml of both streptomycin sulfate and penicillin G. For grazing resistance experiments, 24 h old amoebae were washed in AC buffer (4 mM MgSO₄·7H₂O, 0.4 mM CaCl₂, 3.4 mM sodium citrate dihydrate, 0.05 mM Fe(NH₄)₂(SO₄)₂·6H₂O, 0.05 mM Na₂HPO₄·7H₂O, 2.5 mM KH₂PO₄, 0.05 mM NH₄Cl, pH 6.5) and seeded at a concentration of 1 × 10⁵ cell/well in 96-well tissue culture plate (Costar). Amoebae were allowed to adhere to the bottom of the wells for 1 h before addition of bacteria. Log-phase *V. cholerae* cells (OD₆₀₀ = 0.6) were washed in AC buffer and added to the amoebae at a multiplicity of infection (MOI) of 500. Plates were centrifuged at 500 × g for 10 min to synchronize bacterial uptake. Co-cultures were incubated statically at 30°C for up to 48 h. Control bacteria were incubated in AC buffer in the absence of amoebae for the same period. Every 24 h, intracellular bacteria were released by lysing protozoan cells with 1% Triton X-100. The number of viable bacteria were quantified by CFU counts on LB agar plates.

Caenorhabditis elegans virulence assay

Caenorhabditis elegans N2 Bristol individuals were routinely maintained at 20°C on potato dextrose agar (PDA; Sigma-Aldrich) plates seeded with *E. coli* OP50. Overnight LB cultures of the *V. cholerae* strains and *E. coli* OP50 were diluted 1:100 and grown at 37°C until an OD₆₀₀ of 0.8 was reached. Then, 50 µl from each strain were spread on 6-cm-diameter plates containing PDA medium. Plates were incubated for 16–20 h at 37°C to form a bacterial lawn and 50 L4-stage hermaphrodite individuals were then placed on the PDA plates and incubated at 20°C. Live *C. elegans* were scored every 48 h for 8 days. An individual was considered dead when it no longer responded to touch. *E. coli* OP50 was used as a negative control of virulence. At least, three independent experiments were performed.

Biofilm formation assay

Overnight cultures were diluted 1:100 in fresh LB medium. Hundred microliters were transferred into 96-well Sero-cluster™ plates (Costar) and incubated at 37°C for 20 h without shaking. After incubation, all the wells were washed with sterile distilled water for three times and dried. Biofilm mass was stained with 125 µl crystal violet (1%) per well and incubated for 15 min at room temperature. Crystal violet was removed by performing three washes with sterile distilled water. The stained biofilm mass was then detached from the wells using acetic acid (30%) and transferred to a new 96-well plate (Nunc, Thermo Scientific). OD₅₇₀ was measured using a Biotek Synergy HTX plate reader (Agilent). At least 20 independent experiments were performed per strain. One-way analysis of variance (ANOVA) was used to determine significant differences among the three groups.

Swarming motility assay

Swarming motility assays were performed as described in (47). Briefly, 3 µl of overnight cultures were soaked on motility agar plates containing 10 g/l tryptone, 5 g/l NaCl and 0.25% bacto-agar. The swarming motility halos were measured and compared after growth at 37°C for 6 h. At least three independent experiments were performed per strain. One-way ANOVA was used to determine significant differences among the three groups.

Results

Genomic position of the SI does not alter *V. cholerae* growth

SCIs are present in all *Vibrio* species, suggesting their acquisition before the genus radiation (1). Despite their size, the genomic location of SCIs in *Vibrio* species is remarkably plastic: they can be found in the same position as the *V. cholerae* SI but also in other unrelated locations, e.g. in the chromosome 1 (48). However, the orientation is usually conserved: the bot-

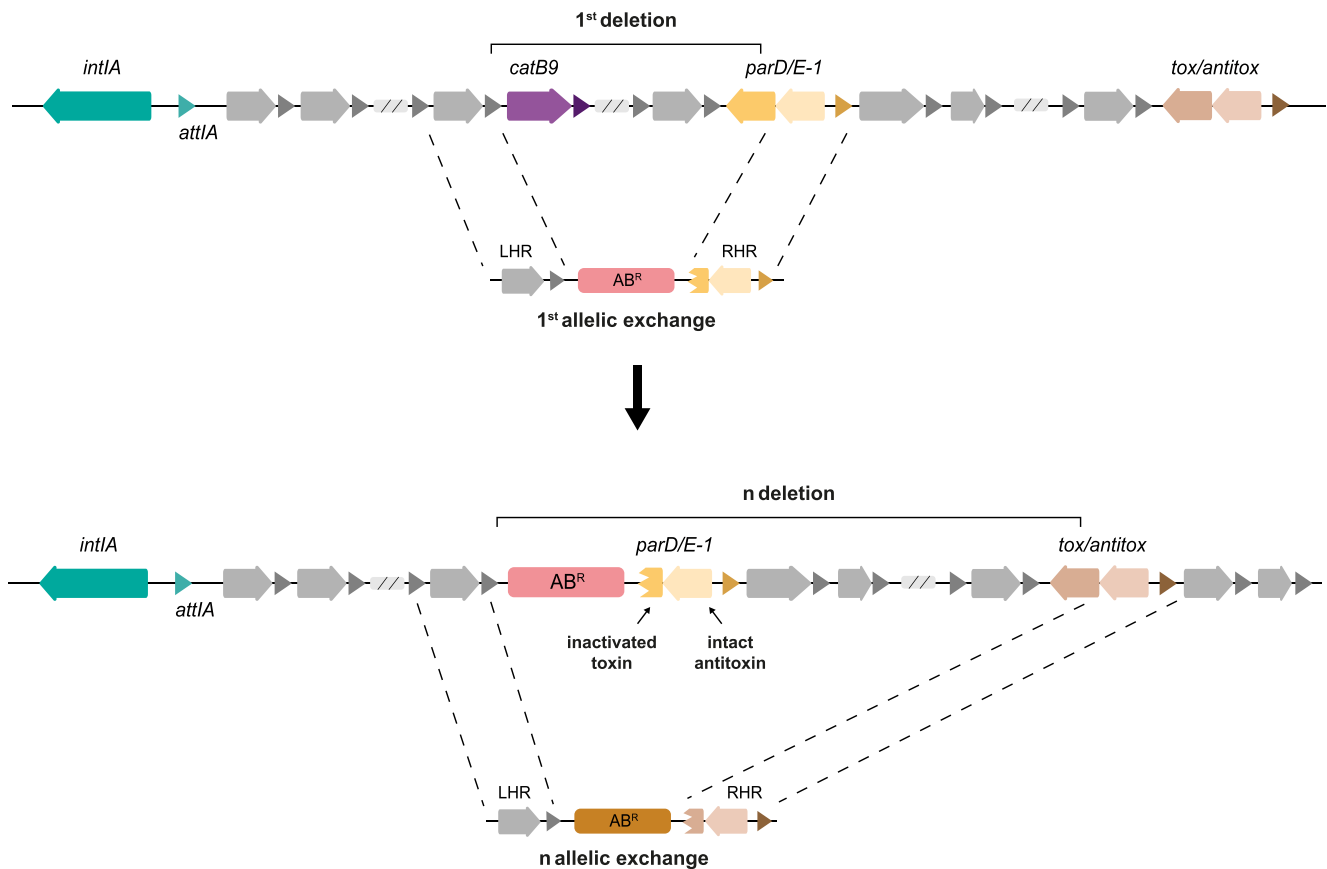


Figure 3. Schematic representation of two successive allelic replacements using SeqDelTA. The LHR is conserved during the deletion process, allowing the recycling of the resistance markers at each step. The RHR is redesigned at each step to inactivate the following TA module and delete the cassette cargo in between. The inactivation of the TA system is performed by knocking the toxin in but leaving the antitoxin intact. A total of 18 sequential replacements were performed. The last deletion step was carried out using a suicide plasmid.

tom strands of *attC* sites are usually carried by the leading strand template (33) to avoid the loss of cassettes (23).

We ignore whether SCI relocations in different species have been innocuous for the cell or have entailed complex changes that required subsequent adaptation. To test this, here we used the Φ HK recombination machinery to relocate the *V. cholerae* SI. We moved it from the right arm of chromosome 2—close to the replication terminus—to chromosome 1, near the origin of replication (between VC0018 and VC10019), where we placed it in both orientations (Figure 2A and Supplementary Figure S1). This relocation could have a potential impact on cell growth because of the higher gene dosage effect near the Ori of chromosome 1. Additionally, it could also generate conflicts between replication and transcription machineries depending on the orientation of the array. We measured growth of both mutants and could not observe differences with the WT (Figure 2B), suggesting that SCIs are genetically independent units of the genome.

SeqDelTA, a tool to erase chromosomal integrons

To further assess the level of genetic and functional independence, we sought to delete the SI. The deletion of the SI in *V. cholerae* is a longstanding milestone in the field that should allow to deliver the kind of experiments that have advanced the field in the last years using the Class 1 integron as a model. Previous efforts to delete the SI were unfruitful because a one-

step deletion approach was used in the presence of—at that time—unknown TA systems. Post-segregational killing due to these uncharacterized TAs made the deletion impossible. The discovery and characterization of these TA systems (19,20), has enabled the development of SeqDelTA, a multi-step approach that allows the sequential inactivation of TA systems while deleting, at each step, the cassette cargo between two TAs (Figure 3).

The SeqDelTA approach exploits the natural competence of *V. cholerae* to generate a series of sequential allelic replacements. To do so, we introduce linear DNA fragments containing a resistance marker flanked by left and right homology regions (LHR and RHR). At each step, these homology regions correspond to specific sequences, allowing the deletion of SI fragments. The SI spans from gene VCA0291 (the integrase) to VCA0506 (nucleotides 309,750—435,034). As LHR, we generally targeted the 5' region of *catB9* chloramphenicol resistance cassette at the ninth position of the array (VCA0300). This choice was initially made to avoid interference with expression from the P_C promoter, since the strain is susceptible to chloramphenicol, although we have recently shown that this gene is transcriptionally active (18). In contrast, RHRs vary at each step of the deletion process, allowing the deletion of an SI fragment spanning from the LHR until the next TA system in the array. They are specifically designed to inactivate the toxin in the TA while keeping the antitoxin gene intact, ensuring the viability of the replacement. Sequen-

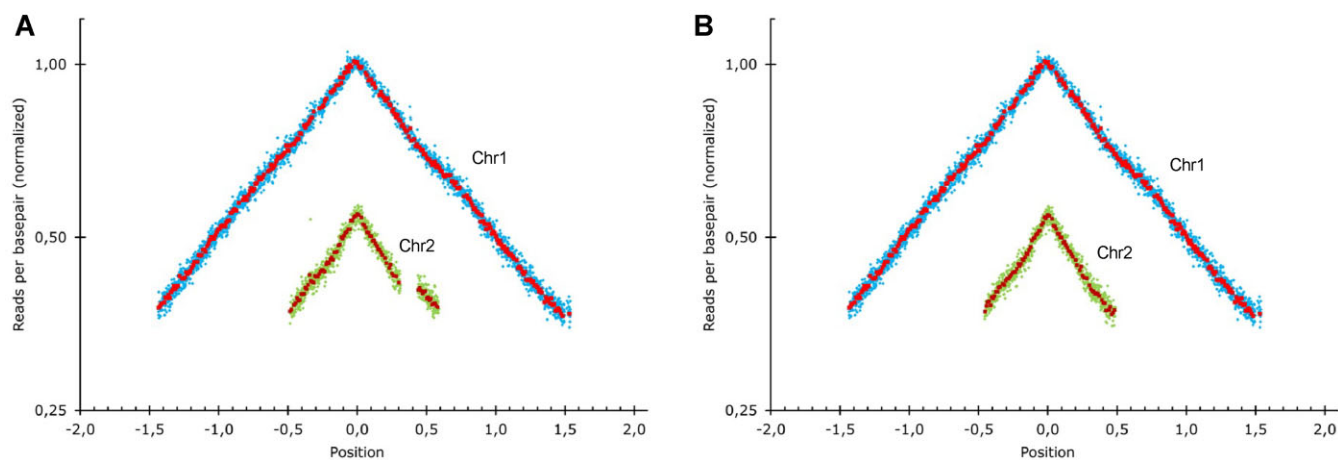


Figure 4. Marker frequency analysis of the two chromosomes of *V. cholerae* Δ SI mapped against *V. cholerae* WT genome (A) and against *V. cholerae* Δ SI genome (B). The gap on chromosome 2 in panel (A) reflects the SIs deletion. The genome position is represented relative to the *oriC* (set to 0). Blue and green rhombus represent the average of 1 kbp windows, and red and green squares represent the average of 10 kbp windows in Chr1 and Chr 2, respectively.

tial deletions facilitate the removal of the resistance marker from the previous step, allowing for easy alternation of resistance markers. This scheme for SeqDelTA was generally kept constant. In the cases in which the orientation and/or the order of the genes in the TA system were not favorable (e.g. the *phd/parE* TA system where the toxin is positioned after the anti-toxin [see Figure 1B]), we used alternative LHRs and maintained two resistance markers in consecutive allelic replacements. Then, a third replacement allowed the removal of both markers. After each allelic exchange, several colonies growing on the appropriate antibiotic were verified through PCR and checked phenotypically for the loss of the previous marker where applicable. The growth curve of PCR-verified clones was then analyzed to avoid the hitchhiking of mutations with important deleterious effects. A clone without growth defects was chosen to continue the deletion process. After the last allelic replacement, we had erased all cassettes from VCA0300 (*catB9*) to VCA0503 (the last toxin). We then used a counter-selectable integrative vector (pMP7 (35)) to deliver the scarless deletion of the remainder of the SI, including the integrase, the first 8 and the last 2 cassettes from the array, as well as a transposase inserted immediately downstream the last cassette (VCA0291 to VCA0508, see Figure 1B). A PCR from the borders of the integron confirmed the deletion of the SI.

Whole genome sequencing (WGS) and marker frequency analysis (MFA)

We performed WGS on the obtained *V. cholerae* Δ SI using Illumina short reads and confirmed the scarless deletion of the SI. We also used this approach to perform MFA, a technique that allows to assess the dynamics of chromosome replication. MFA compares the abundance of reads across the genome between exponential and stationary growth conditions. In fast growing bacteria as *V. cholerae*, this comparison reveals the gene dosage gradient from the origin to the terminus of replication as an inverted 'V'. MFA allows to observe the relative timing of replication of both chromosomes, and the speed of replication across the genome (37). As shown in Figure 4, no major alterations in chromosome replication dynamics could be observed, compared to the WT strain.

MFA can reveal the presence of large chromosomal rearrangements (such as inversions), through inconsistencies in the slope of the inverted V shape, but lacks resolution for smaller rearrangements like IS movements. We therefore sequenced *V. cholerae* Δ SI using MinION long reads to detect the presence of any rearrangement that could go undetected by the assembly of short reads. The combination of small and long reads allowed us to obtain a high-quality genome sequence in which we could rule out genomic rearrangements while detecting the presence of unintended single-nucleotide polymorphisms (SNPs). The *V. cholerae* Δ SI strain bore three unintended mutations in *rocS* (VC0653), *cry2* (VC01392) and *rpoS* (VC0534) genes. RocS is a diguanylate cyclase involved in the smooth-to-rugose switch in our strain. The Δ SI strain had a single base deletion that altered the reading frame of *rocS*. We could observe the loss of motility and increase in biofilm formation described in *rocS*⁻ mutants (49,50). We were able to map the emergence of this mutation to the third deletion step of SeqDelTA. Interestingly, 12 clones from two independent experiments had been kept from this deletion step, of which 11 had mutations in *rocS*. We found eight different types of mutations among them, pointing to an unusually high mutation rate. Indeed, the sequence of *rocS* is rich in homopolymeric tracts and is likely to act as a contingency locus, allowing for a frequent smooth-to-rugose switch (51). *cry2* presented a non-synonymous (Pro129Lys) mutation that appeared in the 12th step. This gene encodes a protein annotated as a photolyase, for which no experimental evidence of its function could be found. RpoS is an alternative sigma factor related to stress response and entry in stationary phase. It is a major regulator of cell physiology and influences many relevant phenotypes, such as oxidative stress, natural competence, motility or colonization (52–54). In the last deletion step 16, *V. cholerae* Δ SI acquired a single-base insertion in *rpoS* giving rise to a frameshift.

To understand the impact of the deletion of the SI in the bacterium's physiology we need to avoid the interference of the mutations in these three genes. To do so, we restored the WT sequence of the three alleles using the counter-selectable suicide vector pMP7. We then verified the restoration of the three genes, as well as the absence of other mutations arising during this process again through WGS using short and long reads. As a result, we could confirm that the Δ SI strain

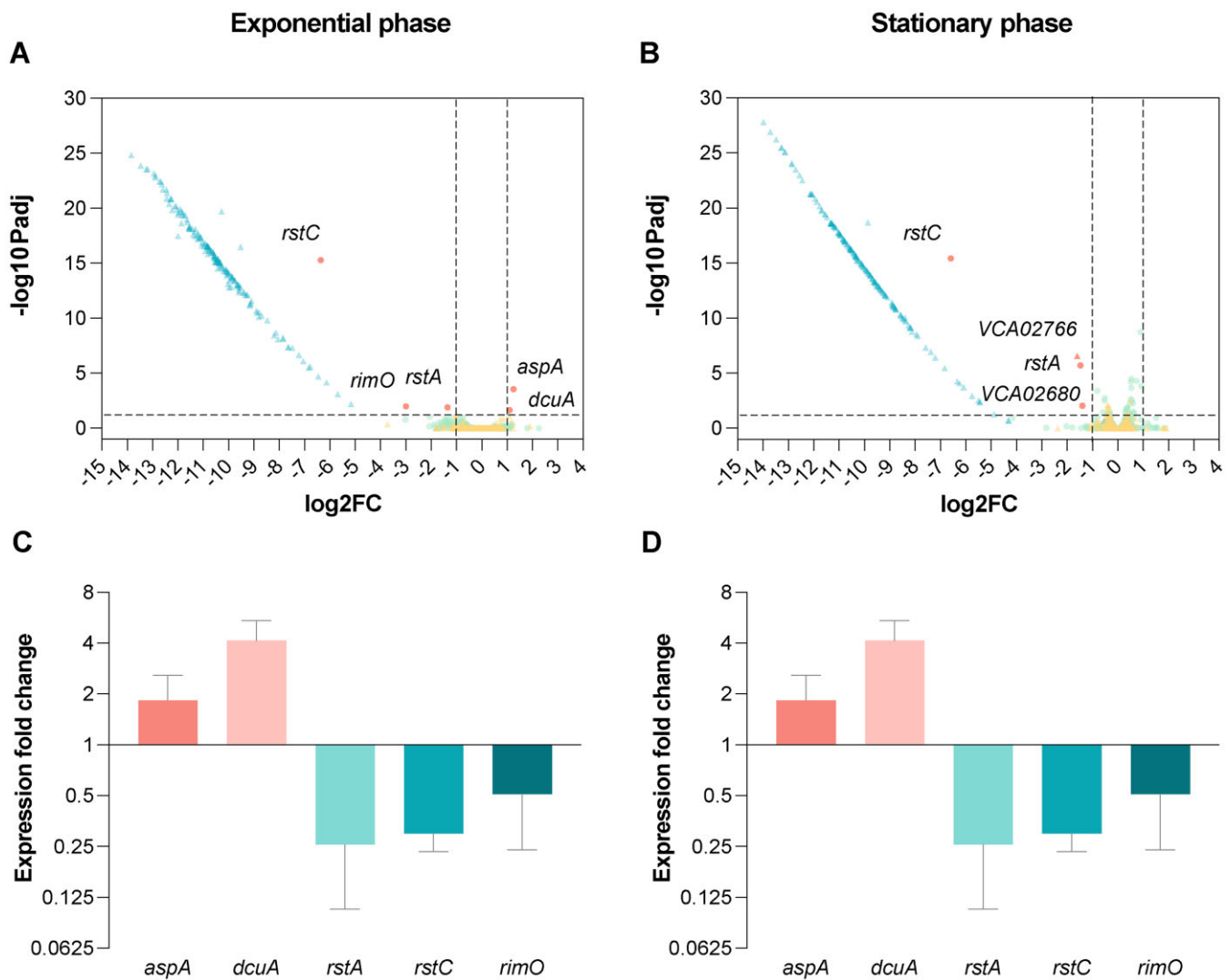


Figure 5. (A) Volcano plots showing gene expression measured by RNA-seq in *V. cholerae* Δ SI in comparison to the WT strain in exponential and (B) stationary growth phase. Green dots represent chromosome 1 genes; yellow triangles represent chromosome 2 genes, while blue triangles represent those genes from the SI; red dots or triangles represent downregulated or upregulated genes from chromosome 1 or 2, respectively. The vertical dashed lines indicate the log₂ fold change (FC) cutoffs, and the horizontal dashed line indicates the threshold of the *P*_{adj} value (<0.05). (C) Validation of the DEGs by RT-qPCR during exponential and (D) stationary growth phase. Differential expression values represent the fold change in gene expression compared to the WT strain. Error bars indicate standard deviation of two biological replicates with three technical replicates each.

is genetically identical to the WT except for the absence of the SI and the loss of one of the two copies of satellite phage RS1.

Transcriptomics

Regulatory networks are generally finely tuned and often affected by minor genetic changes. To address the level of functional isolation of the SI, we sought to assess the changes in expression patterns of the genome in the absence of the SI. Indeed, the deletion of the SI could have pleiotropic effects through direct and indirect links to regulatory networks within the cell. To unveil such effects, we have performed transcriptomic analysis of the Δ SI and the parental strain in exponential and stationary phase (Figure 5A and B). As expected, all genes contained within the SI gave a clear signal of decreased expression. Unexpectedly, the transcriptome of the rest of the genome remained fundamentally unaltered, with

few exceptions that were confirmed through RT-qPCR using independent RNA extractions.

The loss of one of the tandem copies of Φ RS1 in the Δ SI strain (either due to homologous recombination between both copies or to the excision of one copy of the prophage) led to the downregulation in both exponential and stationary phase of *rstA* (VC01411) and *rstC* (VC01409) that was confirmed through RT-qPCR on independent samples. In the WT strain, *rstA* is found three times as part of the Φ CTX (VC01397) and as part of the two adjacent copies of the satellite phage Φ RS1 (VC01407, VC01411), while *rstC* is exclusively found in the latter two copies (VC01409, VC01413) (Supplementary Figure S2). This downregulation through the loss of a Φ RS1 copy, a common variable region in *V. cholerae* strains (55,56), serves as an additional validation of transcriptomic results.

Besides prophage-related genes, in exponential growth phase we also found a 2-fold upregulation of *aspA*

(VC00205; aspartate ammonia-lyase) and *dcuA* (VC00204; C4-dicarboxylase transporter) and a 6-fold downregulation of *rimO* (VC00523; ribosomal protein S12 methylthiotransferase). *aspA* and *dcuA* colocalize in *V. cholerae* chromosome 1. While *aspA* catalyzes the reversible conversion of L-aspartate to fumarate (57), *dcuA* is involved in the L-aspartate/fumarate antiport (58). Additionally, RNA-seq data showed genes VC02680 and VCA02766 to be downregulated 2-fold approximately in stationary phase, but RT-qPCR failed to confirm such downregulation (Figure 5C and D).

A GSEA can reveal the collective up- or downregulation of related genes, even if individually they show no significant changes in expression patterns. GSEA was performed here on the group of DEGs, revealing that the Δ SI strain showed an enrichment of the iron ion transmembrane transport process (GO:0034755; NES = 2,126432; *P*adj = 0.00714409) and the arginine biosynthetic process (GO:0006526; NES = 1 959 273; *P*adj = 0.02539212) in exponential phase and of isoleucine biosynthetic process in stationary phase (GO:0009097; NES = 2 020 381; *P*adj = 0.0129091). However, neither *aspA*, *dcuA*, nor *rimO* are annotated with GO terms belonging to these biological processes. Collectively, these data indicate that deleting the SI does not have important consequences at the transcriptome level in *V. cholerae*.

Phenotype search

The working model of integrons suggested that most of the array of the SI should be silent, and only the cassettes closest to the P_c are expressed (see Figure 1A). Hence, it would be unlikely to find many phenotypic changes after the deletion of the SI. Yet, this seems not to be true, with independent reports showing both the presence of transcription start sites scattered along the SI and a large proportion of cassettes expressed at biologically relevant levels (18), in accordance with our RNA-seq results (59). To address the potential emergence of phenotypic changes and, therefore, the discovery of cassette functions, we implemented a broad approach to characterize a maximum of phenotypes.

Growth and Fitness

The deletion of *V. cholerae* SI implies the loss of 3% of the bacterial genome. Thus, it is expected that elimination of this genomic load would result in some growth and/or fitness consequences. To explore this possibility, we performed growth curves of 24 independent *V. cholerae* Δ SI colonies and compared with the WT in rich LB medium. As shown in Figure 6A, growth curves of both strains present similar shapes after 16 h of growth. We did not observe significant growth differences between strains measured as V_{max} . We then measured the AUC, which combines more information about the growth curve, such as lag phase, V_{max} , and the carrying capacity. The results showed similar AUC values for both strains. To confirm that the deletion of the SI does not influence bacterial fitness, we increased the sensitivity by performing competition assays using flow cytometry. Using a fluorescent *E. coli* strain as a competitor, we found no significant differences between the fitness values given by *V. cholerae* WT and Δ SI (Figure 6A). We hypothesized that this lack of cost could be the serendipitous net result of deleting costly and beneficial cassettes. To test this, we performed the same competitions using intermediate deletion strains (previously corrected for unintended point muta-

tions). Again, we saw no significant differences between the fitness of the parental and these intermediate deletion strains (Supplementary Figure S3) supporting the lack of cost of the SI.

V. cholerae is known to thrive in a variety of environments different from the human host. Its natural setting is brackish or marine water, where temperature is lower (12–30°C versus 37°C). To test the impact of deleting the SI in these conditions, we performed growth curves at 20 and 30°C. We also included 42°C as the heat-shock response could be induced, and the SI could pose an adaptive advantage in such conditions. We show that the growth of the WT and Δ SI strains is similar at all temperatures (Figure 6B). Altogether, these results indicate that deletion of the *V. cholerae* SI does not significantly affect bacterial growth.

Biolog Phenotype Microarrays

To address if we can observe phenotypic differences in the absence of the SI, we have used Biolog Phenotype Microarrays® (PMs), a phenomics technology that allows the testing of bacterial respiration and growth in nearly 2000 conditions (46). PMs test carbon utilization, nitrogen sources, phosphorus and sulfur compounds, biosynthetic pathways, osmotic, ionic and pH effects, as well as a broad set of chemicals and antibiotic compounds at four different concentrations. By using tetrazolium violet as a redox dye in the mixture, we monitor bacterial respiration for 24 h and compare the metabolism of two independent replicates of the WT and Δ SI strain in all the available conditions (Supplementary Figure S4).

To search for differences between the two strains, the mean AUCs given by both replicates of WT and Δ SI in a specific compound were graphically compared using Matlab software (R2022a). In Figure 7, we show the correlation of AUC values for WT and Δ SI strains in each condition. Distance to the origin of the axis represents metabolic activity, and deviations from the diagonal represent differences between strains. Generally, dots fall within the diagonal, showing that both strains behave similarly in the presence of the different compounds, which also supports the similarity in growth rates across a broad variety of conditions. It is of note, that the Δ SI shows a potential decrease in respiration when using dipeptides and tripeptides as nitrogen sources (PM6 and PM8) (see ‘Discussion’ section). Within antimicrobial-containing plates (PM11–20), we generally did not find differences between the WT and Δ SI. However, we have highlighted some isolated cases in Figure 7 where the AUC of one of the strains is at least the double of the other and the standard deviation does not cross the bisector. These compounds are ceftriaxone (G4, PM11), cadmium chloride (D4, PM14), methyl viologen (E12, PM15), norfloxacin (B3, PM16), tannic acid (F10, PM17), sodium metasilicate (E4, PM18), iodonitro tetrazolium violet (D8, PM19), thioridazine (C1, PM20) and benserazide (A10, PM20). Despite the lack of consistent results with other molecules of the same family, we believed these results deserved further verification. We selected three compounds (ceftriaxone, norfloxacin and tannic acid) and measured the MIC but found identical values for both strains: norfloxacin 50 ng/ml; ceftriaxone 25 ng/ml; tannic acid 250 ng/ml. We also performed growth curves using subinhibitory concentrations of the three compounds in MH but could not observe the same phenotype previously obtained using the phenotype microarrays (Supplementary Figure S5).

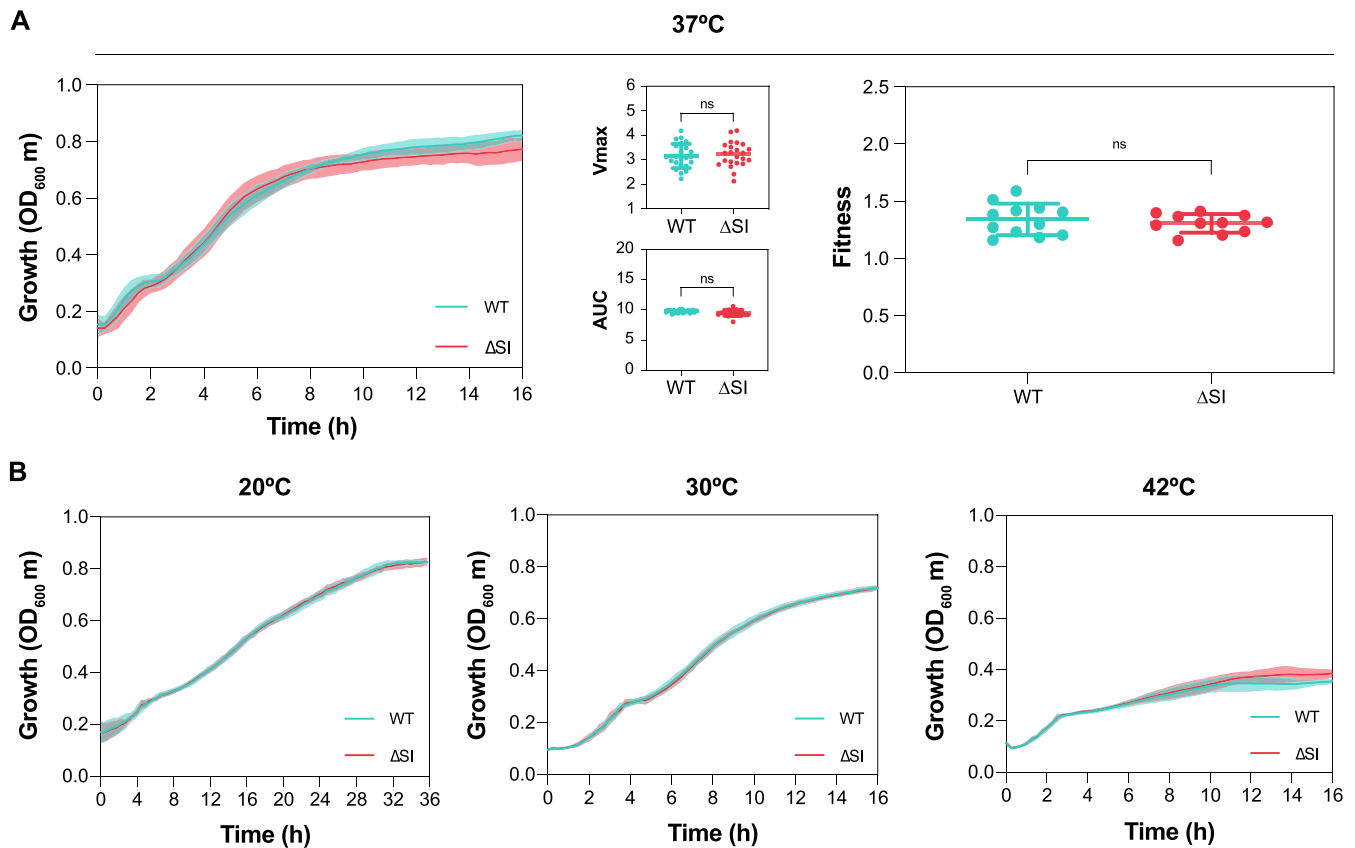


Figure 6. (A) Growth curves of *V. cholerae* WT (blue) and Δ SI (red) strains in LB at 37°C. Growth parameters including V_{\max} and the AUC were extracted from growth curves in LB by measuring the optical density at 600 nm (OD₆₀₀). Values correspond to the measurement of 24 independent colonies. Relative fitness of *V. cholerae* WT and Δ SI strains compared with *E. coli* DH5 α Pcs::gfp was performed in LB by inoculating cells at a ratio of 1:1. Fitness values were determined from 12 independent experiments by flow cytometry. The *P*-values were calculated by comparing each measure with that of the WT strain using an unpaired *t*-test. Ns: not significant. **(B)** Growth curves of *V. cholerae* WT (blue) and Δ SI (red) strains in LB at 20, 30 and 42°C, respectively. Values correspond to the measurement of 10 independent colonies.

Antibiotic persistence, grazing resistance, virulence, biofilm formation, natural competence and swarming motility

Antibiotic persistence is the ability of a bacterial population (or a part of it) to survive exposure to bactericidal drugs (60), without becoming resistant to the antibiotic. TA systems have been described to play a role in triggering persistence in bacteria by reversibly intoxicating the bacterium and producing a transient arrest of metabolic activity that renders antibiotics ineffective. This role is controversial, at least for some TAs, since several studies were later shown to be biased by the activity of prophages (61). Given the deletion of a variety of TA systems in the Δ SI mutant we decided to test antibiotic persistence upon exposure to high concentrations of antibiotics. We challenged growing cultures of *V. cholerae* WT and Δ SI to 10-fold the MIC of ampicillin and ciprofloxacin, and viability was measured along a 6 h-period (Figure 8A). Similar decreases in the number of viable cells were observed for both strains, of varying intensities depending on the antibiotic molecule, suggesting that TA systems encoded in the SI do not play a role in antibiotic persistence in *V. cholerae*.

V. cholerae interacts with various free-living protozoa in aquatic environments, as the model organisms *Tetrahymena pyriformis* and *Acanthamoeba castellanii* (62). These interactions can be complex and may impact the survival and infectivity of the pathogen. For instance, it has been reported that *T. pyriformis* and other ciliates can enhance *V. cholerae*'s en-

vironmental persistence by expelling hyper-infectious forms (63). Other studies indicate that predation by protozoa, particularly ciliates, can eliminate *V. cholerae* from the environment (64). Additionally, *V. cholerae* can survive intracellularly within *A. castellanii*, potentially providing a protected niche (65). Given the relevance of these interactions, we aimed to investigate whether the *V. cholerae* SI contains factors that may contribute to grazing resistance by protozoa. To assess this, survival assays were conducted by co-incubating *V. cholerae* WT and Δ SI with the protozoan predators *T. ellioti* and *A. castellanii*. Although we observed a different degree of grazing between the ciliates and the amoeba, there were no differences between *V. cholerae* WT and Δ SI in terms of intracellular survival within both protozoa under the tested conditions (Figure 8B).

V. cholerae expresses well-known virulence factors to colonize and cause infection in the mammalian host, as the cholera toxin or the toxin-coregulated pili (66,67). The nematode *C. elegans* has been widely used as an invertebrate infection model to screen and identify virulence factors of several human pathogens, including *V. cholerae* (68,69). TA systems in the SI are reported to be associated with virulence. For instance, the RelBE TA systems of the SI play a role in intestine colonization in a mouse model (70). However, evidence on the implication of other elements in the SI in the virulence of *V. cholerae* is lacking. To address this question, we performed

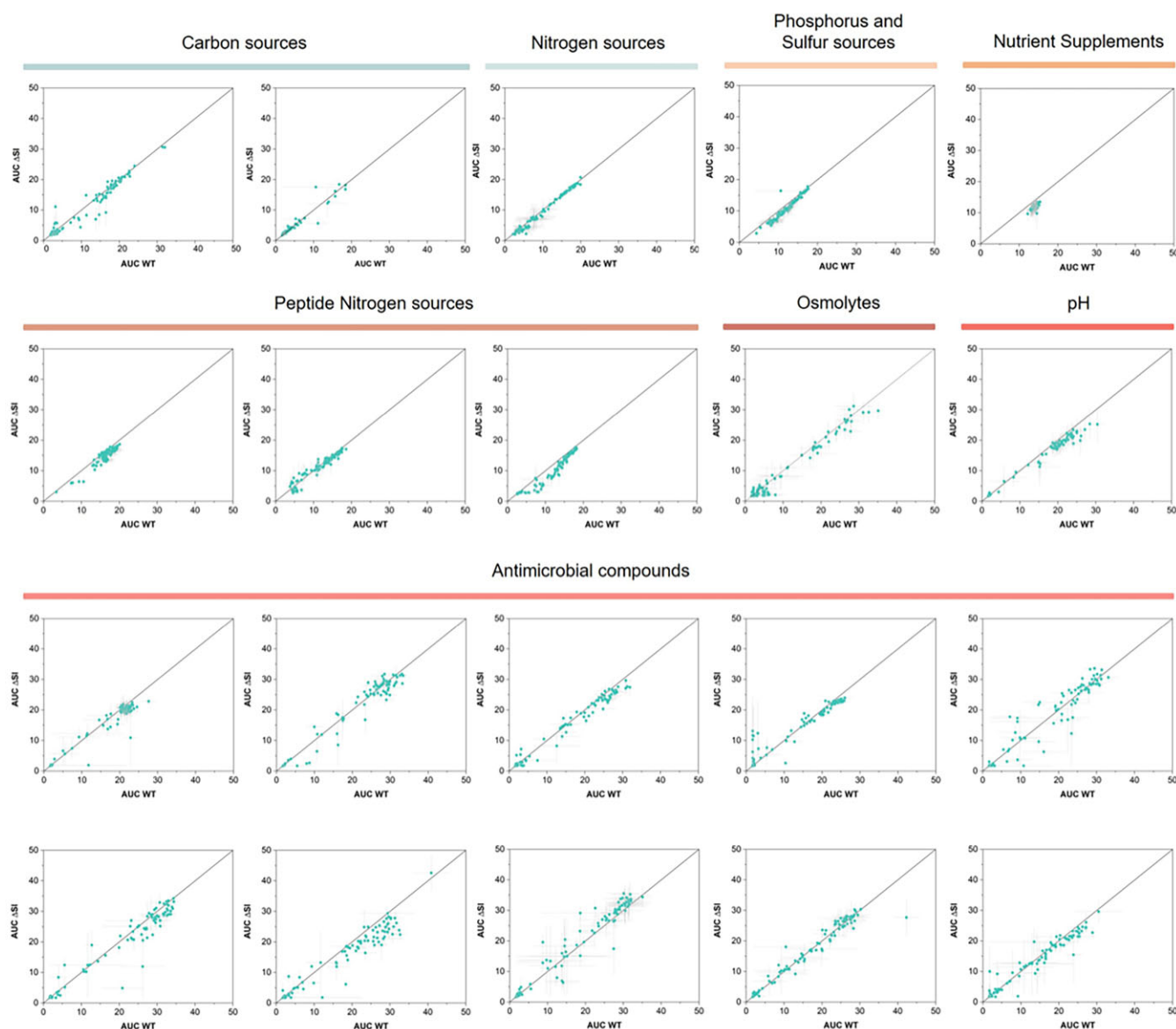


Figure 7. Comparison of AUC values of *V. cholerae* WT and Δ SI in different substrates given by Biolog Phenotype Microarrays (PM1–PM20). The AUC values are the mean of two biological replicates for each strain. Error bars indicate the standard error. The points that are above the bisector indicate those conditions where the absence of the SI is beneficial for the bacteria, while those points that lay under the bisector indicate the conditions where the presence of the SI is beneficial for the bacteria.

a *C. elegans* killing assay. Hermaphrodites L4 worms raised on *E. coli* OP50 were transferred onto bacterial lawns of WT and Δ SI strains. The number of *C. elegans* individuals were counted every 48 h for 8 days. *E. coli* OP50 was used a control. As shown in Figure 8C, we did not observe any differences in lethality between *V. cholerae* WT and the Δ SI mutant.

Biofilm formation, natural competence, and motility are key behaviors in the lifestyle of *V. cholerae*, with strong implications for its survival in the environment, pathogenesis, dispersal and genetic variability. Motility is involved in regulating biofilm formation by participating in important processes, such as surface attachment or biofilm dispersal (71). Increased biofilm formation also entails a downregulation of natural competence through quorum sensing control (72,73). We investigated the role of the SI in these processes by measuring natural competence, biofilm formation, and swarming motility. As shown in Figure 8D–F, none of the processes was

significantly affected in the Δ SI strain. As a control for biofilm formation, we included a *V. cholerae* strain with a frameshift mutation in *hapR* (a regulator of quorum sensing), leading to an increase in biofilm production. For motility, we tested a strain with an amino acid change mutation in *rocS* as a non-motile control.

Discussion

The working model of integrons is based on semiconservative recombination and the strong expression of cassettes upon integration, highlighting that cassettes must be adaptive to be maintained in the array. This, together with the intimate intertwining between integrons and their hosts (through the SOS response controlling the expression of the integrase (74), or through host factors (26,30)) put integrons at the service of the host to provide adaptation on demand (2,26,75). A clear

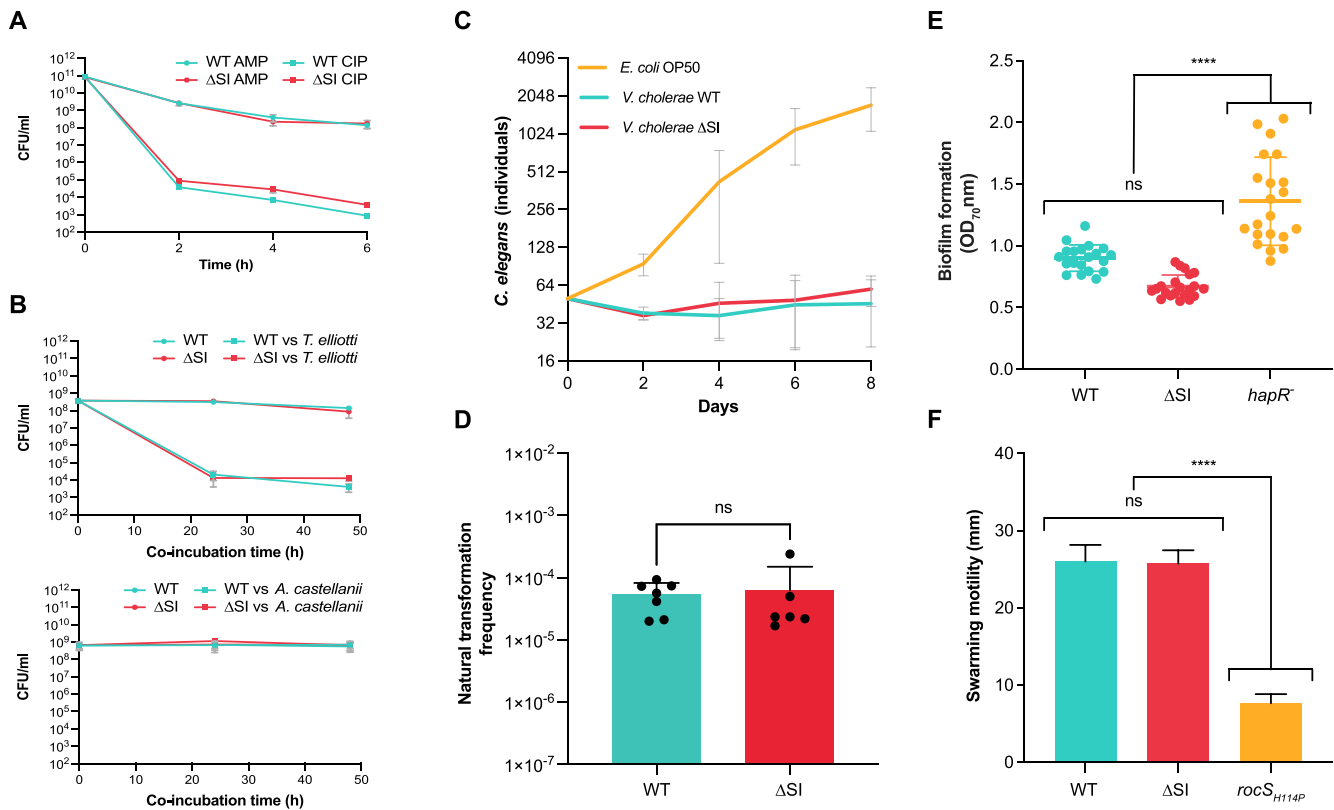


Figure 8. (A) Antibiotic persistence assay of *V. cholerae* WT and ΔSI mutant challenged with 10-fold the MIC of ampicillin (AMP) and ciprofloxacin (CIP), individually. CFUs/ml are plotted against time. Error bars indicate standard deviations from three independent biological replicates. (B) *V. cholerae* WT and ΔSI survival under predation by *T. ellipti* (MOI 1:10000) or *A. castellanii* (MOI 1:500). The number of surviving bacteria, incubated either in the absence (circles) or presence (squares) of ciliates/amoeba, was quantified by CFU counting after 0–48 h of co-incubation. Error bars indicate standard deviations from at least three independent biological replicates. (C) Virulence assay of *V. cholerae* WT and ΔSI mutant in a *C. elegans* model. *E. coli* OP50 was used as a negative control. Error bars indicate standard deviations from three independent biological replicates. (D) Natural transformation assay of *V. cholerae* WT and ΔSI mutant. Transformation frequencies are given on the Y-axis and were calculated as the number of CFUs growing in kanamycin over the total CFUs. Bar charts show the mean ± s.d. from biological replicates (individual dots) from at least three independent experiments. (E) Biofilm formation of *V. cholerae* WT and ΔSI mutant. A *V. cholerae hapR* (frameshift) mutant was included as a positive control of biofilm formation. Data from at least 20 independent colonies is represented. (F) Swarming motility assay of *V. cholerae* WT and ΔSI mutant. A *V. cholerae rocS* (H114P) mutant was included as a non-motile control strain. Error bars indicate standard deviation of at least three biological replicates. *P*-values (****) were calculated by pairwise comparison using the one-way ANOVA test ($P < 0.0001$). Ns: not significant.

example of this is their role in the rise and spread of multidrug resistance during the last decades. In *Vibrionaceae*, integrons provide genetic diversity upon which natural selection can act (48). It is therefore assumed that SCIs provide great adaptability too. Yet, a better understanding of how they do so remains out of reach because the functions encoded in the cassettes of chromosomal integrons are mostly unknown (15). Apart from antibiotic resistance genes and TA systems, only a handful of cassettes have been characterized (for a review, see (76) and (77)).

Chromosomal integrons have co-evolved with their hosts during aeons. It is therefore plausible that hosts have co-evolved with integrons and have adapted to the presence of such massive structures. Also, the functions encoded in cassettes could be intimately linked to the host's physiology, somehow interfering or modulating housekeeping functions. Indeed, *in silico* analysis of cassettes in *Vibrionaceae* revealed potential functions like information storage, cellular processes and metabolism (15). This is certainly at odds with the variable nature of the array of cassettes, which would imply that cassettes provide exclusively 'self-contained' or independent functions that do not interfere with those encoded in the rest

of the genome. Here, we have tested the hypothesis that SCIs are genetically and functionally independent of the host by relocating and deleting the SI of *V. cholerae* and testing a broad variety of phenotypes of importance in the lifestyle of this bacterium. The result is surprisingly clear: the ΔSI mutant behaves almost identically to the WT strain in most conditions tested, and differences (like those observed in the use of di-peptides) are, at best, very mild. This tilts the balance towards the 'plug-and-play' model of cassette function. Indeed, our data suggest that cassettes that are expressed have been selected to be as neutral as possible. In line with our results, a recent CRISPRi study in *V. cholerae* shows that guides against the SI are not deleterious, except when they silence the anti-toxin genes (78). In this sense, it is possible that SCIs are biased to encode functions that are exclusively activated under specific cues and with little transcriptional and translational demand. A good example of this is the case of TAs. These systems often repress their own expression under normal conditions but can be triggered by infecting phage, delivering their function (abortive infection) without the need for additional RNA or protein synthesis (79). It is of note that coding such low-cost triggerable functions is not *conditio sine qua non*,

since the shuffling ability of integrons can also serve to switch on and off the expression of cassettes. This has been shown for the *catB9* cassette in the SI, which can move from ninth to second position, providing antibiotic resistance (80).

Focusing on the subtle differences that we have observed, we note that the deletion of the SI has a limited impact on gene expression, with only three genes (*aspA*, *dcuA* and *rimO*) being mildly affected. *aspA* and *dcuA* are involved in nitrogen metabolism. AspA carries out the reversible conversion of L-aspartate to fumarate, releasing ammonia for nitrogen assimilation (81). This enzyme has a specific interaction with DcuA, which functions as an L-aspartate/fumarate antiporter under both aerobic and anaerobic conditions (58). This interaction has been proposed to form a metabolon, where DcuA facilitates the uptake of aspartate, while AspA converts it to fumarate (82). The GSEA analysis revealed an enrichment in arginine and isoleucine biosynthetic processes in the Δ SI strain. Arginine synthesis is derived from glutamate, which, in turn, originates from L-aspartate. Additionally, the precursor molecule for isoleucine biosynthesis is pyruvate, a product of glycolysis, and an indirect byproduct of the aspartate metabolism (83). These findings, coupled with the observation that the Δ SI mutant exhibits impaired growth when utilizing certain dipeptides and tripeptides as nitrogen sources, may suggest a potential connection between these processes. However, it is important to note that these pathways are not directly linked, and further experimental data would be needed to ascertain whether a change in metabolic balance has occurred within the cell. Given the subtle signal from our data, we remain cautious about the biological significance of these observations.

The obtention of a Δ SI mutant of *V. cholerae* is a long-awaited milestone in the field, unblocking the study of SCIs. Getting rid of the interference of the cassettes within the SI allows to use this strain as a chassis to easily perform many of the experiments that have been delivered in *E. coli* with the class 1 integron, allowing to enrich the field with data from other integron models. It is also the perfect chassis to unveil cassette functions in a classical *in trans* approach, indeed one of the key questions that remains unanswered in the field and one with an important biotechnological potential.

Data availability

The RNA-seq data included in this publication have been deposited in NCBI's Gene Expression Omnibus and are accessible through GEO Series accession number GSE247496.

Supplementary data

Supplementary Data are available at NAR Online.

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Conflict of interest statement

Universidad Complutense de Madrid and Institut Pasteur have published a patent covering SeqDelTA and the *V. cholerae* Δ SI strain B522. Patent reference number: ES 2 970 040 A1. Inventors: Escudero, J.A.; Trigo da Roza, F.; Blanco, P.; Mazel, D. and Lopéz-Igual, R.

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