

## RESEARCH ARTICLE

# Halopseudomonas species: Cultivation and molecular genetic tools

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**Abstract**

The *Halopseudomonas* species, formerly classified as *Pseudomonas pertucinogena* lineage, form a unique phylogenetic branch within the Pseudomonads. Most strains have recently been isolated from challenging habitats including oil- or metal-polluted sites, deep sea, and intertidal zones, suggesting innate resilience to physical and chemical stresses. Despite their comparably small genomes, these bacteria synthesise several biomolecules with biotechnological potential and a role in the degradation of anthropogenic pollutants has been suggested for some Halopseudomonads. Until now, these bacteria are not readily amenable to existing cultivation and cloning methods. We addressed these limitations by selecting four *Halopseudomonas* strains of particular interest, namely *H. aestusnigri*, *H. bauzanensis*, *H. litoralis*, and *H. oceani* to establish microbiological and molecular genetic methods. We found that C<sub>4</sub>-C<sub>10</sub> dicarboxylic acids serve as viable carbon sources in both complex and mineral salt cultivation media. We also developed plasmid DNA transfer protocols and assessed vectors with different origins of replication and promoters inducible with isopropyl-β-D-thiogalactopyranoside, L-arabinose, and salicylate. Furthermore, we have demonstrated the simultaneous genomic integration of expression cassettes into one and two *attTn7* integration sites. Our results provide a valuable toolbox for constructing robust *chassis* strains and highlight the biotechnological potential of *Halopseudomonas* strains.

**INTRODUCTION**

Microbes play an important role as production platforms in various sectors of the biotechnological and chemical industries. Their use under industrial process conditions requires tolerance to different stressors, such as toxic reaction intermediates or products, osmotic stress, or pH shifts (Bitzenhofer et al., 2021). Several bacteria are well established as biotechnological workhorses, including *Escherichia coli*, *Pseudomonas putida*, *Pseudomonas taiwanensis*, *Corynebacterium glutamicum*, and *Bacillus subtilis* (Blombach et al., 2022; Gießelmann et al., 2019; Loeschcke & Thies, 2015;

Reva et al., 2006; Rodrigues et al., 2014; Tabor et al., 2011; van Dijn & Hecker, 2013). These bacteria were optimised towards stress tolerance, e.g. by engineering metabolic fluxes towards bioproduction, reducing the genome, additional stress-resistance genes such as those enabling compatible solute production, or applying adaptive laboratory evolution (Cárdenas Espinosa et al., 2023; Gießelmann et al., 2019; Kuepper et al., 2020). A different strategy harnesses billion years of natural evolution by establishing microorganisms that have naturally adapted to specific challenges (Blombach et al., 2022; Czajka et al., 2017; Fatma et al., 2020; Riley & Guss, 2021). Here, their natural

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habitat already serves as an important indicator of the type of natural stress tolerance to be expected (Czajka et al., 2017; Volmer et al., 2015).

Among such potentially stress-tolerant bacteria, many members of the *Halopseudomonas* genus were isolated from habitats with toxic contaminations, elevated or constantly changing temperatures, or high osmotic pressure. Most of the currently identified 25 species have been described for the first time in the last 20 years (Bollinger, Thies, Katzke, & Jaeger, 2020). Previously, they were placed as the *Pseudomonas pertucinogena* lineage within the genus *Pseudomonas*, but considerable molecular and metabolic differences from other *Pseudomonas* spp. called for a recent reclassification as *Halopseudomonas* (syn. *Neopseudomonas*) within the *Pseudomonadaceae* (Rudra & Gupta, 2021).

Halopseudomonads appear to have in common a small genome with a size of approximately 4 Mb, which is uncommon among *Pseudomonaceae*. This implies a limited metabolic flexibility of these bacteria, which, in contrast to the versatile group of *Pseudomonas* spp., may reflect a specific adaptation to ecological niches. Consequently, Halopseudomonads are described to metabolise hardly any sugars but grow on organic acids like acetic acid, lactic acid, and succinic acid (Pascual et al., 2012; Sánchez et al., 2014; Wang & Sun, 2016; Zhang et al., 2011). Further, they metabolise Tween20 and sebacic acid. Genome analyses revealed genes encoding interesting biocatalysts such as esterases, halohydrin dehalogenases, and  $\omega$ -transaminases or the biosynthesis potential to produce ectoine and polyhydroxyalkanoates (Bollinger, Thies, Katzke, & Jaeger, 2020). Furthermore, a role of Halopseudomonads in the degradation of anthropogenic pollutants such as oil or plastic waste was suggested (Bollinger, Thies, Katzke, & Jaeger, 2020; Gomila et al., 2017; Villela et al., 2023). Until now, *Halopseudomonas* spp. have not been studied in detail, presumably associated with a lack of reference genomes, metabolic models, and robust molecular genetic tools. We have investigated the growth of four selected strains under typical laboratory conditions. Furthermore, we characterised different plasmid-based expression systems and evaluated the genomic integration of expression modules into two *attTn7*-sites via Tn7-transposition using *H. litoralis* as one of the studied model strains.

## EXPERIMENTAL PROCEDURES

### Bacterial strains and plasmids

*Escherichia coli* DH5 $\alpha$   $\lambda$ pir (Penfold & Pemberton, 1992) was used for cloning; the strain S17-1  $\lambda$ pir (Simon et al., 1983) was used for the conjugational transfer of expression plasmids. The *E. coli* strains were either

cultivated on LB agar plates or in liquid LB medium (Luria/Miller, Carl Roth<sup>®</sup>) at 37°C, supplemented with 10  $\mu$ g/mL gentamycin (Gm), 100  $\mu$ g/mL ampicillin (Amp) or 50  $\mu$ g/mL kanamycin (Km) when needed. *H. aestus-nigri* VGXO14 (Sánchez et al., 2014), *H. bauzanensis* BZ93 (Zhang et al., 2011), *H. litoralis* 2SM5 (Pascual et al., 2012), and *H. oceani* KX20 (Wang & Sun, 2016) were grown on LB-agar plates supplemented with a total of 3% (w/v) NaCl at 30°C for 48 h. The increased salt concentration prevented the cells from drying out and enabled storage of the plates for up to one month at 4°C. For the cultivation of the R-strains (VGXO14R, BZ93R, 2SM5R, and KX20R), 25  $\mu$ g/mL rifampicin (Rif) was supplemented. When other antibiotics were needed, 25  $\mu$ g/mL Km or 25  $\mu$ g/mL Gm were added. To identify and distinguish the four different species from one another, hydrolysis of synthetic plastic material was determined by halo formation on Impranil DLN agar plates. Therefore, 4 mL Impranil<sup>®</sup> DLN-SD (Covestro AG) were added to 1 L of LB-agar (containing 3% (w/v) NaCl and antibiotics when needed) and mixed thoroughly (Molitor et al., 2020). The liquid cultivation, unless stated otherwise, was performed in LB medium enriched with 45 mM succinate (Merck KGaA, Cas: 6106-21-4) (LB<sub>suc</sub>) as a complex medium. The used mineral salt medium (MSM medium) was prepared according to (Hartmans et al., 1989), supplemented with 18 mM sebacic acid (Merck KGaA, Cas: 111-20-6) as the sole carbon source. The carbon source was supplemented in C-equimolar concentrations derived from comparable studies (C-equimolar to 30 mM adipic acid [Ackermann et al., 2021]). Sebacic acid was prepared as a sterile-filtered 10-fold concentrated stock solution in H<sub>2</sub>O while adjusting to pH 10 with sodium hydroxide and supplemented to the medium before cultivation. The cultivation was either performed in 100 mL Erlenmeyer flasks (10% filling volume) at 130 rpm or in a 48-well Round Well Plate<sup>®</sup> (1 mL filling volume) sealed with “Breathable rayon film seals for biological cultures” (VWR<sup>®</sup>, Radnor) at 1000 rpm in the microbioreactor BioLector I (Beckman Coulter GmbH) for at least 24 h at 30°C unless otherwise stated. Two to three colonies were added to the cultivation medium for pre-culture inoculation. Bacterial strains harbouring a vector-based antibiotic resistance gene were cultivated under the required selection pressure (25  $\mu$ g/mL Gm or 25  $\mu$ g/mL Km). The growth rate at the logarithmic growth phase was calculated using the following equation (Christian et al., 1982):

$$r(t) = \frac{\ln(N_2) - \ln(N_1)}{\Delta t}$$

$r(t)$ , growth rate [ $\text{h}^{-1}$ ];  $N_1$ , mean of a biological triplicate of the biomass at early logarithmic state;  $N_2$ , mean of a biological triplicate of the biomass at late logarithmic state;  $t$ , time [h].

All bacterial strains and plasmids used in this study are listed in [Table S1](#), along with their construction and genetic properties.

## Cultivation of selected *Halopseudomonas* spp. for target gene expression

Main cultures of *H. aestusnigri* VGXO14, *H. bauzanensis* BZ93, *H. litoralis* 2SM5, and *H. oceani* KX20 were inoculated to an optical density  $OD_{580\text{ nm}}$  of 0.1 (determined with a Photometer, cuvette: 1 cm path length) and grown in Round Well Plates<sup>®</sup> using the microbioreactor BioLector I (Beckman Coulter GmbH). The cell density was monitored online every 20 min via scattered light intensity at 620 nm, and GFP fluorescence intensity was measured using an  $Ex_{508\text{ nm}}/Em_{532\text{ nm}}$  filter, whereas mCherry fluorescence was detected with an  $Ex_{580\text{ nm}}/Em_{610\text{ nm}}$  filter. The induction of heterologous gene expression was achieved during early logarithmic growth phase (approximately 4.5 h after inoculation) by adding the inducer molecule isopropyl- $\beta$ -D-thiogalactopyranoside [IPTG] [Cas: 367-93-1], L-arabinose [Cas: 5328-37-0], or salicylic acid [Cas: 69-72-7] [Merck KGaA]), respectively. Stock solutions were prepared in 100x concentration in water or 70% ethanol. The dynamic range is defined as the ratio of the fluorescence intensity of the highest signal and the signal obtained with the uninduced control.

## Plasmid isolation

Plasmids were isolated from bacterial cells by alkaline lysis using the 'innuPREP Plasmid Mini Kit' (Analytik Jena) according to the manufacturer's instructions. For elution, nuclease-free water warmed to 65°C was used. Isolated plasmid DNA was then stored at -20°C. DNA sequences were determined by Eurofins Genomics.

## Transformation of bacteria with plasmid DNA

Chemically competent *E. coli* cells prepared with the calcium-chloride/magnesium-chloride method were transformed using the heat-shock method (Hanahan, 1983) with slight modifications. 1–2  $\mu$ L of the plasmid DNA were added to 100  $\mu$ L chemically competent cells and then incubated on ice for 30 min. Subsequently, the heat shock was performed at 42°C for 90 s and 700  $\mu$ L of LB medium was added. Afterwards, the cells were incubated in a rotator wheel for 90 min ( $Km^R$ , or  $Gm^R$ ) at 37°C. Then, 100  $\mu$ L of the cells were plated on LB agar plates and incubated at 37°C overnight under respective selection pressure.

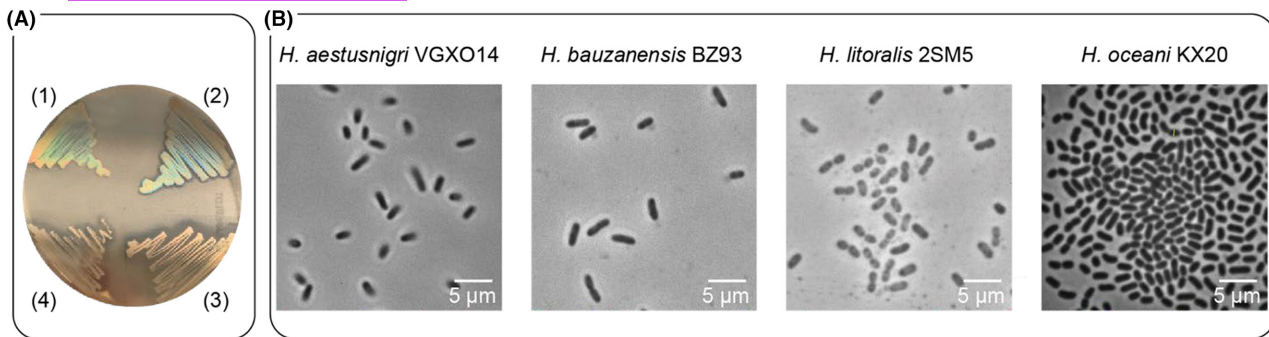
To transform *H. aestusnigri* VGXO14, *H. bauzanensis* BZ93, *H. litoralis* 2SM5, and *H. oceani* KX20 with plasmid DNA, a slightly modified version of the room temperature protocol for electroporation was used (Datsenko & Wanner, 2000; Tu et al., 2016). An overnight pre-culture was harvested via centrifugation (1 min, 21,000 g) and then washed thoroughly with 1 mL of sterile water. After another centrifugation step for 1 min, the cell pellet was resuspended in 80  $\mu$ L of sterile water, and 1  $\mu$ L (~100 ng) of the external DNA was added. Afterwards, the sample was transferred to an electroporation cuvette (electrode spacing 1 mm: Bio-Budget Technologies GmbH). The electroporation was performed by using the 'EC1' program (25  $\mu$ F, 200  $\Omega$ , 4.5–5 ms, 20 kV/cm) of MicroPulser (Bio-Budget Technologies GmbH). After that, 700  $\mu$ L of  $LB_{suc}$  medium was added to the cell suspension and then cultivated under agitation for 2 h at 30°C. Afterwards, 100  $\mu$ L of the cells were plated on LB-agar containing 3% (w/v) NaCl with glass beads and incubated at 30°C for 48 h under respective selection pressure.

## Tn7 transposition

For the Tn7 transposition, the miniTn7 tools pUC18R6KT-miniTn7T-Km, which was a gift from Herbert Schweizer (Addgene plasmid # 64969; <http://n2t.net/addgene:64969>, RRID: Addgene\_64969) (Choi et al., 2005) and the pBG-13 (Zobel et al., 2015) were used. Triparental conjugational gene transfer (Elhai & Wolk, 1988) was applied to transform the *Halopseudomonas* with these plasmids. To this end, *E. coli* S17-1  $\lambda$ pir was transformed with the plasmid of interest. A second transformation of *E. coli* S17-1  $\lambda$ pir with the helper plasmid pTNS2 was performed. Subsequently, 250  $\mu$ L of each overnight pre-culture of these *E. coli* strains, as well as 500  $\mu$ L of a *H. litoralis* 2SM5R pre-culture, were combined and mixed gently. After centrifugation (1 min, 21,000 g), the obtained cell pellet was resuspended in 100  $\mu$ L  $LB_{suc}$  and transferred onto a membrane filter, which was placed on LB-agar without any additives. After incubation for at least 5 h at 30°C in the dark, the cells were washed off the filter with 1 mL  $LB_{suc}$ . Finally, the cell suspension was centrifuged for 1 min at 21000 g and resuspended in 100  $\mu$ L of the supernatant. The suspension was plated on LB-agar (3% (w/v) NaCl, 25  $\mu$ g/mL Rif) using glass beads, incubated for two days at 30°C under the required selection pressure, and seeded on agar plates again. Further, the iridescent phenotype of *Halopseudomonas* colonies and their polyester hydrolyase activity (Figure 1B) can be utilised to confirm the isolation of transformed *Halopseudomonas* after the conjugation.

A PCR protocol was established to evaluate the successful genomic integration, including primers, each binding specifically at one of the two *glmS* genes (oligos





**FIGURE 1** Phenotypes of selected *Halopseudomonas* spp. (A) Cell lawns of *Halopseudomonas* strains after cultivation on LB-agar containing Impranil® DLN-SD. (1) *H. aestusnigri* VGXO14, (2) *H. bauzanensis* BZ93, (3) *H. litoralis* 2SM5, (4) *H. oceani* KX20. The agar plate was transilluminated with daylight while the picture was taken. (B) Phase contrast microscopy images, taken of motile bacteria from stationary phase cultures of *H. aestusnigri* VGXO14, *H. bauzanensis* BZ93, *H. litoralis* 2SM5, and *H. oceani* KX20 grown in LB<sub>suc</sub> medium.

12 and 13; [Table S1](#)). The other primer binds within the integration cassette (oligo 9 or 10; [Table S1](#)). Thus, only a genomic integration led to a PCR product. To illustrate the localisation of *glmS1* (locus tag: [BLU11\\_RS05420](#)) and *glmS2* (locus tag: [BLU11\\_RS09655](#)) in relation to the origin of replication *oriC* of the chromosomes, Tubic Ori-Finder 2022 (Dong et al., 2022) was used to identify *oriC* of the chromosomes (GenBank: [LT629748.1](#) and [CP000010.1](#)). The region 3,507,165-3,507,739 nt of LT629748.1, and the regions 3,006,500-3,006,897 nt as well as 3,055,348-3,055,787 nt of CP000010.1 were identified as the possible origin of replication (the sequences can be found in the [Supplementary Material S1](#)).

## Phase-contrast microscopy

Approximately 10  $\mu$ L of the cells obtained from stationary phase cultures cultivated in LB<sub>suc</sub> medium in a BioLector® were placed on a microscope slide and covered by a coverslip. Images were taken on a motorised inverted microscope (Nikon Eclipse Ti, Nikon Europe B.V.) equipped with a high-resolution camera from Zyla (Andor Neo SCC-01462, Andor, Belfast) and a CFI Plan Achromat  $\lambda$ 100x Oil Ph3 DM oil-Immersion objective (NA 1.45; W.D.0.13; Nikon). Images were captured using the Nikon NIS-Elements AR software package. The analysis was performed using Fiji, ImageJ2 software (Schindelin et al., 2012).

## RESULTS AND DISCUSSION

We selected four strains from the genus *Halopseudomonas* as model organisms to establish standard cultivation conditions and probe their genetic accessibility using a set of replicative and integrative vectors. The strains were selected based on expected eurythermia and euryhalinity, presumably resulting from their natural habitats.

*H. aestusnigri* VGXO14 was isolated from a crude oil-contaminated sand sample of the intertidal zone (Sánchez et al., 2014). It harbours a remarkable number of lipolytic enzymes, some with astonishing substrate promiscuity and resistance to organic solvents (Bollinger, Molitor, Thies, et al., 2020; Coscolín et al., 2019). The soil bacterium *H. bauzanensis* BZ93 was isolated from an industrial site (Zhang et al., 2011). Both strains were thus expected to exhibit tolerance to various chemicals and may contribute to the biodegradation of pollutants as indicated by the degradation of poly(ethylene terephthalate) for enzymes from *H. aestusnigri* VGXO14 and *H. bauzanensis* BZ93 at ambient temperature (Avilan et al., 2023; Bollinger, Thies, Knieps-Grünhagen, et al., 2020). Additionally, we chose two marine water isolates, *H. litoralis* 2SM5 and *H. oceani* KX20. *H. litoralis* was also isolated from the tidal zone and is thus expected to cope with constantly changing environmental conditions. It can tolerate NaCl concentrations up to 15%, representing the highest reported osmotolerance among *Halopseudomonas* spp. (Bollinger, Thies, Katzke, & Jaeger, 2020; Pascual et al., 2012). In contrast, the deep-sea organism *H. oceani* was isolated from a habitat with high pressure and low temperature, reflected by the ability to grow at 4°C but tolerating up to 42°C (Wang & Sun, 2016).

## Growth of *Halopseudomonas* spp. under typical laboratory conditions

As a first step, we aimed to establish reliable cultivation conditions for the selected *Halopseudomonas* species and monitor growth. As previously reported, these strains form small beige, non-pigmented colonies on LB-agar plates (Pascual et al., 2012; Sánchez et al., 2014; Wang & Sun, 2016; Zhang et al., 2011). On LB-Impranil agar, the strains are known to form halos, indicating polyester-hydrolysing activity (Molitor et al., 2020). When cultivated on LB-Impranil agar, *H. litoralis* showed the largest halo, followed

by *H. oceanii*, whereas the smallest halo was observed around *H. aestusnigri* colonies (Figure 1A). Furthermore, bacteria grown in the form of dense lawns showed an iridescent, rainbow-coloured diffuse glow upon transillumination (Figure 1A). This observation indicated a coordinated movement and aggregation of cells in specific patterns within the lawn or the production of exopolysaccharides leading to structural colours (Kientz et al., 2012; Mizuno et al., 2022; Petruzzi et al., 2017), a phenomenon that, to our knowledge, has not been reported for any *Halopseudomonas* species before.

In order to access *Halopseudomonas* for biotechnological applications, suitable cultivation conditions need to be established. We first cultivated the strains in different cultivation vessels filled with LB medium to identify a suitable bioreactor system. The type and filling of cultivation vessels are decisive for oxygen transfer (by surface-to-volume ratio and shaking geometry and frequency) as well as shear forces (e.g. in the case of baffles). Growth in a 100 mL Erlenmeyer flask (10% filling volume) was compared with growth in two types of microbioreactors available for the BioLector<sup>®</sup> system (Beckman Coulter GmbH), i.e. a Round Well Plate<sup>®</sup> (RWP) (1 mL filling volume) and a FlowerPlate<sup>®</sup> (FP) (1 mL filling volume; oxygen transfer rate: 45 mmol/L/h) (Figure S1A). Here, the cultivation in FPs led to strong aggregate formation within all strains' cultures, hampering the growth monitoring using light scattering (see jagged curve in Figure S1B) or optical density. Aggregation of cells indicates stress that may here be caused by shear forces (Fakhrudin & Quilty, 2007; Trunk et al., 2018; Tsagkari et al., 2022). Comparable growth with less aggregate formation was observed when flasks and RWP were used as cultivation vessels. Next, we examined the cell morphology by phase contrast microscopy, revealing that the stationary phase cells of *Halopseudomonas* spp. maintained their rod-shaped morphology after 24 h of cultivation under these conditions (Figure 1B). The sampled cells appeared fully motile, but *H. litoralis* 2SM5 and *H. oceanii* KX20 still tended to aggregate (Figure S3). The number of colony-forming units (CFU) per mL determined for stationary phase cells was in the same order of magnitude after growing in all three cultivation vessels, as exemplarily assessed for *H. bauzanensis* (Figure S1C). Thus, we decided to use RWPs for all further cultivation experiments allowing to monitor growth of bacteria at higher throughput.

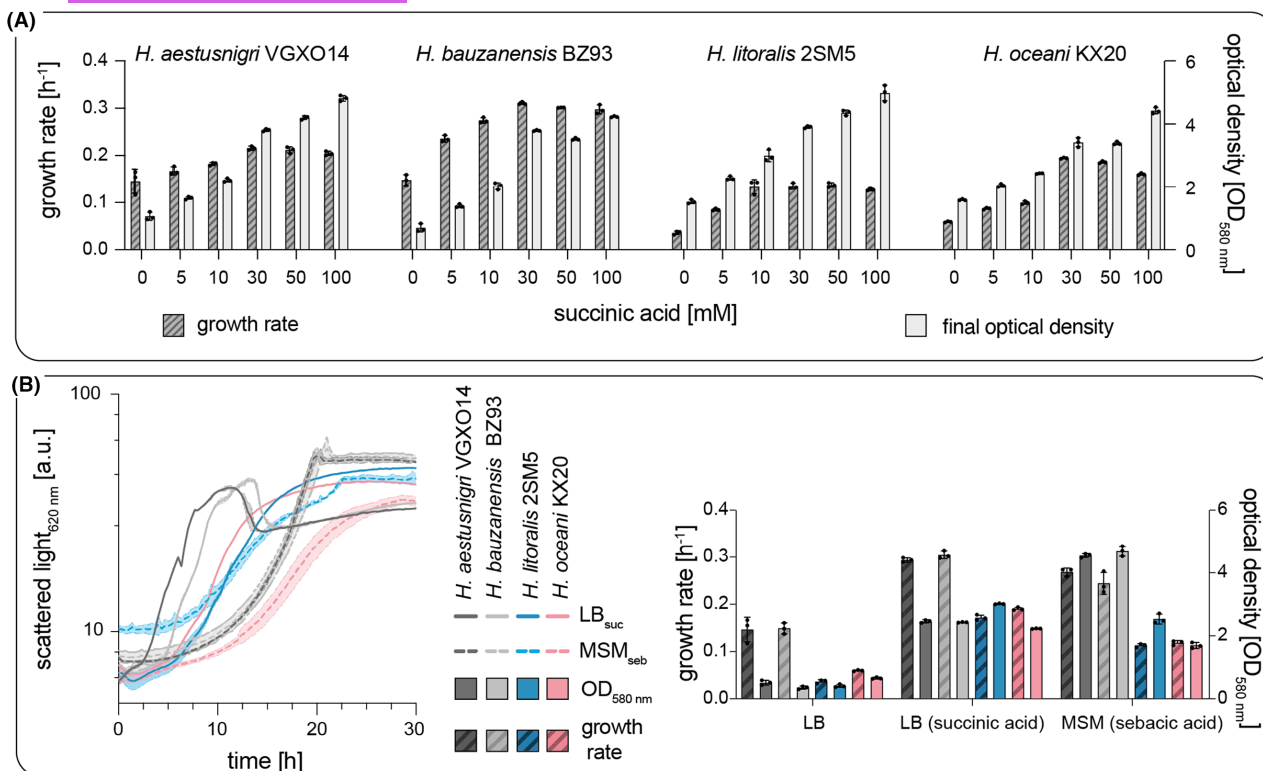
We observed multiphasic growth curves upon cultivation in LB medium indicating that different carbon sources are used up sequentially (Figure S1D, Figure S2A–D). After 24 h of growth in flasks and RWP, the cells reached a final cell density  $OD_{580\text{ nm}}$  of approximately 1, which is low when compared with cultures of *Pseudomonas* sp. or *E. coli* sp. grown in the same medium, indicating that the nutrients provided

by this medium were not depleted. Hence, we tried to identify additives and set up a suitable mineral salt medium (MSM). Previous studies reported that these strains hardly metabolised carbohydrates, but instead used dicarboxylic acids (Figure S4) (Pascual et al., 2012; Sánchez et al., 2014; Wang & Sun, 2016; Zhang et al., 2011). We therefore supplied the complex media with succinic acid ( $C_4$ ) and assessed their use as sole carbon source in MSM (Figure 2).

Supplementation of LB medium with succinic acid at the beginning of the logarithmic growth phase (4.5 h) improved the growth of all *Halopseudomonas* strains during batch cultivations (Figure 2A). Here, highest biomass densities (determined as  $OD_{580\text{ nm}}$ ) were observed with concentrations of 50 mM and 100 mM succinic acid. Maximal growth rates were measured upon the addition of 30 mM succinic acid. A shift from pH 7 to pH 10, presumably evoked by the metabolism of the added acid, was ruled out as main reason for the observed growth characteristics, as an elevated pH per se did not improve the growth of *Halopseudomonas* (Figure S5).

Succinic acid is an intermediate of the citrate cycle and is known as a suitable carbon source for many bacteria including *Pseudomonas* species (Collier et al., 1996; Dhamale et al., 2022; Mendonca et al., 2020). According to previous reports, *Halopseudomonas* are also able to grow on less common dicarboxylic acids (de Witt et al., 2023; Pascual et al., 2012; Sánchez et al., 2014; Wang & Sun, 2016; Zhang et al., 2011). Hence, we assessed the growth of *H. aestusnigri* VGXO14, *H. bauzanensis* BZ93, *H. litoralis* 2SM5, and *H. oceanii* KX20 upon supplementation of both LB and MSM medium with short-chain ( $C_2$  and  $C_3$ ) and long chain ( $C_5$ – $C_{10}$ ) dicarboxylic acids in C-equimolar concentrations (Table 1; Figure S6). In general, the addition of dicarboxylic acids with shorter chain length, such as oxalic acid ( $C_2$ -dicarboxylic acid), malonic acid ( $C_3$ -dicarboxylic acid) as well as glutaric acid ( $C_5$ -dicarboxylic acid), and pimelic acid ( $C_7$ -dicarboxylic acid) led to minor growth improvements compared to LB medium or inhibited growth.  $C_6$ - and  $C_8$ -dicarboxylic acids supplemented as additional (LB) and as sole carbon source (MSM) evoked a prolonged lag phase, but the cultures reached comparable final biomasses to those supplemented with succinic acid ( $C_4$ -dicarboxylic acid). Notably, supplementation of MSM with long-chain acids azelaic acid ( $C_9$ -dicarboxylic acid) and sebacic acid ( $C_{10}$ -dicarboxylic acid) resulted in comparable or even better growth performance of all species, indicated by a shorter lag phase, higher growth rate, and/or higher optical densities. (Figure S6).

Comparing the results among strains, we observed that cultures of *H. aestusnigri* VGXO14 and *H. bauzanensis* BZ93 grown in MSM achieved higher final biomass than complex media cultures, whereas *H. litoralis* 2SM5 and *H. oceanii* KX20 reached higher



**FIGURE 2** Growth of *H. aestusnigri* VGXO14, *H. bauzanensis* BZ93, *H. litoralis* 2SM5, and *H. oceani* KX20 in complex and mineral salt media compositions. (A) Growth in LB medium supplemented with increasing concentrations of succinic acid. The growth rate (dark grey, left y-axis) and the optical density at 580 nm after 24 h (light grey, right y-axis) are shown. (B) Comparison of the growth in LB medium with 45 mM succinic acid (continuous line) and MSM with 18 mM sebacic acid as sole carbon source (dotted line). Growth curves determined by light scattering are shown on the left, and growth rates (striated bars, left y-axis) and the optical densities after 24 h of cultivation (filled bars, right y-axis) are shown on the right. The shown data represent the mean of biological triplicates. The calculated standard deviations are either indicated by shadows or error bars.

final biomasses when using the complex medium (Figure 2B). *H. litoralis* and *H. oceani* metabolised a larger spectrum of dicarboxylic acids than *H. formosensis* FZJ (de Witt et al., 2023), whereas growth of *H. aestusnigri* and *H. bauzanensis* was even inhibited by  $\text{C}_6$  dicarboxylic acids (Table 1).

In summary, succinic acid and sebacic acid were the most beneficial supplements to LB medium, resulting in short lag phases, high growth rates, and biomass for all tested species. Interestingly, sebacic acid appeared superior to succinate when present as sole carbon source. The maximal growth rates achieved for the four species with the optimised media (MSM with sebacic acid) were  $0.267 \pm 0.08 \text{ h}^{-1}$ ,  $0.243 \pm 0.023 \text{ h}^{-1}$ ,  $0.112 \pm 0.004 \text{ h}^{-1}$ , and  $0.118 \pm 0.005 \text{ h}^{-1}$  for *H. aestusnigri*, *H. bauzanensis*, *H. litoralis*, and *H. oceani*, respectively. Reported growth rates of the Proteobacteria *Cupravidus necator* and *Pseudomonas nitroreducens* on sebacic acid were within a similar range (Janota-Bassalik & Bohdanowicz-Strucinska, 1974; Lang et al., 2007; Strittmatter et al., 2022). From Sullivan et al. (2022), approx. growth rates can be deduced for *Acinetobacter baylyi* ADP-1 of  $0.27 \text{ h}^{-1}$  growing on sebacic acid and of  $0.43 \text{ h}^{-1}$  on adipic acid.

In relation to well-established model organisms growing under the respective standard conditions, the growth rates obtained here with  $>1$  division/hour were 2–6 times lower but within the range of those reported for batch cultivations of *P. putida* ( $0.57 \text{ h}^{-1}$ ), *B. subtilis* ( $1.5 \text{ h}^{-1}$ ), *E. coli* ( $1.1 \text{ h}^{-1}$ ), or *C. glutamicum* ( $0.75 \text{ h}^{-1}$ ) growing on glucose (Blank et al., 2008; Wittgens et al., 2011). The growth rate of engineered *P. putida* for growth on adipic acid was determined with  $0.35 \pm 0.01 \text{ h}^{-1}$  (Ackermann et al., 2021).

Metabolism of long-chain dicarboxylic acid is not widespread among bacteria. It requires a set of specialised enzymes catalysing the respective analogue of fatty acid, which is best described for *A. baylyi* ADP1 (Ackermann et al., 2021; Parke et al., 2001). BLASTp homology search uncovered homologues of all the mentioned proteins in *H. litoralis* and *H. bauzanensis* (Table S2) and homologues of most of them in *H. aestusnigri* and *H. oceani*. The lack of some of the expected proteins may be attributable to the fact that only draft genomes are available for those strains that do not represent closed chromosomes but sets of scaffolds, indicating that parts of the genomes are lacking.



**TABLE 1** Overview of the metabolism of various dicarboxylic acids as an additional or sole carbon source by *H. aestusnigri* VGXO14, *H. bauzanensis* BZ93, *H. litoralis* 2SM5, and *H. oceani* KX20.

Dicarboxylic acids	LB medium				MSM medium			
	<i>H. aestusnigri</i>	<i>H. bauzanensis</i>	<i>H. litoralis</i>	<i>H. oceani</i>	<i>H. aestusnigri</i>	<i>H. bauzanensis</i>	<i>H. litoralis</i>	<i>H. oceani</i>
C <sub>2</sub>	-	-	-	-	-	-	-	-
C <sub>3</sub>	-	++	++	+	-	-	-	-
C <sub>4</sub>	+++	+++	+++	++	++	++	+++	+++
C <sub>5</sub>	+	-	+	+	-	-	-	-
C <sub>6</sub>	+	-	++	+++	+	+	+++	+++
C <sub>7</sub>	+	-	+	+	+	-	-	-
C <sub>8</sub>	+++	+++	+++	+++	++	+	+	++
C <sub>9</sub>	+++	+++	+++	+++	+++	+++	+++	++
C <sub>10</sub>	+++	+++	+++	+++	+++	+++	+++	++

The C<sub>2</sub>-C<sub>10</sub> dicarboxylic acids were tested in concentrations C-equimolar to 45mM succinic acid and were used as additional carbon sources in LB medium and as sole carbon sources in MSM medium. The indicators correlate to relative biomass values after 48h. The values determined for LB medium with additives were related to the biomass achieved for a specific strain with LB medium without additives (100%): - growth inhibiting; + a range up to 150%; ++ a range up to 180%; +++ a range greater than 180%. The highest biomass achieved for a specific strain in MSM with a respective carbon source was set to 100%. - growth inhibiting; + a range up to 50%; ++ a range up to 80%; +++ a range up to 100%. The respective growth curves are shown in [Figure S4](#).

Unlike to *A. baylyi*, the identified *dca*-homologues are distributed among several loci in the genomes of the selected Halopseudomonads, which may indicate functions of these enzymes in other metabolic pathways like terpene/branched amino acid degradation.

Organisms naturally evolved to utilise C<sub>6</sub>-C<sub>10</sub> dicarboxylic acids are of interest in the context of plastic degradation and upcycling of synthetic polymers, as those acids are building blocks for polyesters (like polyethylene adipate terephthalate (PBAT), polyester polyurethane foams and coatings, e.g. Impranil® DLN-SD, and Nylon 6,6,) as well as part of their hydrolysates and different plasticisers (Ackermann et al., 2021; de Witt et al., 2023; Howard et al., 2012; Sullivan et al., 2022). As it is known that *Halopseudomonas* spp. secrete enzymes that can depolymerise polyesters (Avilan et al., 2023; Bollinger, Thies, Knieps-Grünhagen, et al., 2020; de Witt et al., 2023; Haernvall et al., 2017; Molitor et al., 2020), it may be speculated that Halopseudomonads play a role in the breakdown of such polymers present in different habitats. On the other hand, these bacteria might represent interesting targets for establishing biotechnological recycling strategies, which can serve as alternatives to engineered or laboratory-evolved strains (Ackermann et al., 2021; Sullivan et al., 2022).

## Unlocking the genome of Halopseudomonads for engineering purposes

Genetic accessibility requires, among other traits, reliable selection of transformed clones, e.g. after conjugational transfer experiments. Halopseudomonads cannot grow on cetrinide agar as the common *Pseudomonas* selection medium (Goto & Enomoto, 1970) nor share any antibiotic resistance. In order to enable counterselection, we isolated spontaneously occurring rifampicin (Rif) resistant clones after spreading several colonies of each species on LB-agar containing 25 µg/mL rifampicin (Rif). The resulting strains were termed *H. aestusnigri* VGXO14R, *H. bauzanensis* BZ93R, *H. litoralis* 2SM5R, and *H. oceani* KX20R, respectively. To counteract spontaneously occurring Rif-resistant cells of the *E. coli*-donor strain, we exploited the pronounced osmotolerance of *Halopseudomonas* spp. by using LB agar plates containing Rif and 3% (w/v) NaCl, which impaired the donor's growth ([Figure S7](#)). Remarkably, the Rif-resistant strains formed no aggregates in liquid culture as they were observed for the respective wild-type strains.

A recent study reported the successful transformation of *H. aestusnigri* and *H. oceani* with a pBBR-derived plasmid by electroporation (Chan et al., 2023). Based on these results, we established an electroporation method using a pJT'Tmcs plasmid harbouring Gm<sup>R</sup>

the broad-host range pRO1600 *oriV* but no *oriT*. The washing protocol and the electroporation procedure did not impact the viability of cells obtained from R-strains (data not shown). For the four selected strains, transformation efficiencies were determined as the number of Gm/Rif-resistant CFU per mass of DNA: *H. aestusnigri* VGXO14R –  $3 \times 10^7$  CFU/ $\mu$ g; *H. bauzanensis* BZ93R –  $3 \times 10^4$  CFU/ $\mu$ g; *H. litoralis* 2SM5R –  $7 \times 10^7$  CFU/ $\mu$ g; and *H. oceani* KX20R –  $1 \times 10^8$  CFU/ $\mu$ g. Table S3 shows that all strains allowed for the replication of a set of broad-host-range plasmids established for *Pseudomonas* spp. (Martin-Pascual et al., 2021), harbouring *ori's* pRO1600, pBBR1, and RSF1010 as well as the respective Rep proteins, except for R6K-vectors. Interestingly, these strains also appeared to maintain the typical *E. coli* origins of replication ColE1 / pMB1 / pBR322 that are not applicable in *Pseudomonas* spp. This observation was confirmed by testing the antibiotic resistance of clones after transformation and the isolation of the plasmids from the *Halopseudomonas* strains (Figure S8A,B). Nevertheless, curing from such plasmids could be obtained by cultivating the strains in liquid cultures at 37°C for 24 h (Figure S8A).

Next, we evaluated in *H. aestusnigri* VGXO14R, *H. bauzanensis* BZ93R, *H. litoralis* 2SM5R, and *H. oceani* KX20R for the use of constitutive promoters, namely  $P_{em7}$  and  $P_{tac}$ , for the expression of genes encoding the fluorescent proteins GFP and mCherry (Figure S9). Expression of reporter genes from promoter  $P_{tac}$  in *H. aestusnigri* and *H. bauzanensis* carrying a plasmid with a RO1600 origin resulted in similar fluorescence yields, *H. litoralis* and *H. oceani* harbouring the respective plasmid showed only low fluorescence intensities. The highest fluorescence signal was observed with *H. litoralis* carrying  $pYT-P_{em7}$ -eYFP.

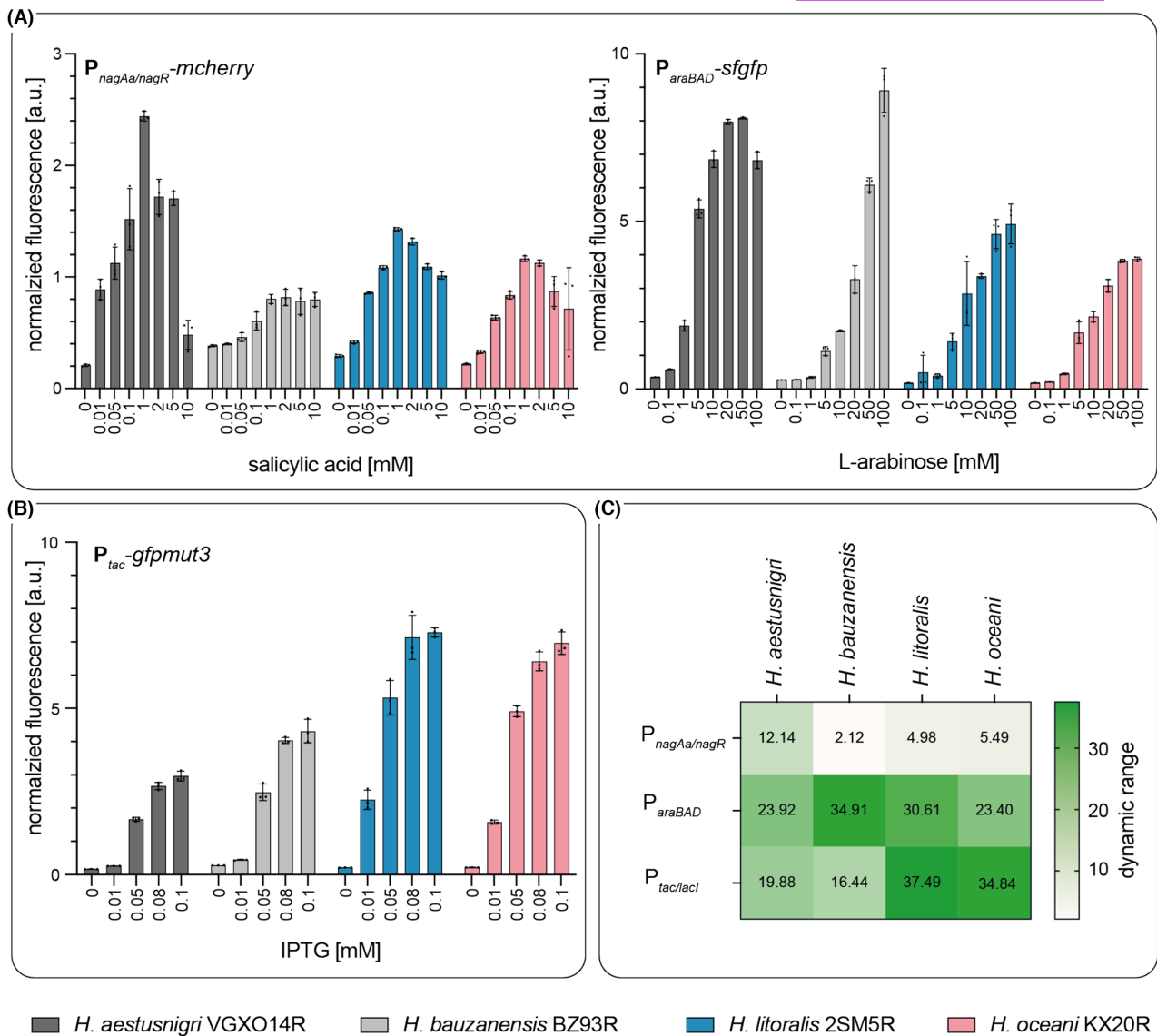
We then tested inducible promoter systems that either lead to an activation (Figure 3A) or de-repression (Figure 3B) of the gene expression. For all R-strains evaluated, a gradual induction profile was obtained. Nevertheless, we observed differences in the normalised fluorescence intensities between the different species. On the one hand, *H. aestusnigri* VGXO14R showed the highest fluorescence intensity compared to the other R-strains when gene expression was induced with L-arabinose or salicylic acid. L-arabinose at a concentration of 20 mM was sufficient to activate expression and reach maximum fluorescence intensity. For *H. litoralis* 2SM5R and *H. oceani* KX20R in contrast, 50 mM L-arabinose was required for maximum intensity, and at least 100 mM for *H. bauzanensis* BZ93R. Notably, as all strains cannot grow with arabinose as sole carbon source, they probably lack an efficient import mechanism. The highest dynamic range (Figure 3C) of the  $P_{araBAD}$  system was achieved in *H. bauzanensis* BZ93R and *H. litoralis* 2SM5R, matching the range previously determined for *P. putida* (Chan et al., 2023). Interestingly, the dynamic ranges achieved here with

*H. aestusnigri* VGXO14R and *H. oceani* KX20R were higher than previously reported for the respective wild-type strains (Chan et al., 2023). In *E. coli*, an arabinose concentration of 13.3 mM was necessary to fully induce gene expression, whereas 133 mM (2% w/v) was required in *P. putida* (Cook et al., 2018; Guzman et al., 1995). Using the  $P_{nagAa/nagR}$ -system, a maximum of the normalised fluorescence intensity was reached after adding 1 mM salicylic acid. Notably, the growth of *H. aestusnigri* was impaired at this concentration, whereas *H. bauzanensis*, *H. litoralis*, and *H. oceani* appeared tolerant to up to at least 10 mM (Figure S10). *P. putida* KT2440, for comparison, tolerates up to 5 mM salicylic acid; the maximum protein production was achieved when supplementing with 2 mM salicylic acid (Weihmann et al., 2023). However, 10  $\mu$ M salicylic acid was sufficient to induce expression of the reporter gene *mcherry* in *H. aestusnigri* VGXO14R, whereas five times the amount was necessary for *H. litoralis* 2SM5R and *H. oceani* KX20R, and ten times the amount for *H. bauzanensis* BZ93R. These observations may indicate the presence of more efficient mechanisms to keep the inducer out of the cell, matching the observed higher tolerance of those strains. However, fluorescence was also detected when no inducer molecule was added, indicating a leakiness of the promoter system. On the other hand, *H. aestusnigri* VGXO14R showed the weakest fluorescence intensity when the gene expression was derepressed by the addition of IPTG. Here, *H. litoralis* 2SM5R and *H. oceani* KX20R showed the highest sensitivity towards IPTG, since the de-repression of *gfpmut3* expression occurred upon induction with 10  $\mu$ M IPTG, whereas 50  $\mu$ M IPTG were needed in *H. aestusnigri* VGXO14R, and *H. bauzanensis* BZ93R. An inducer concentration of 80–100  $\mu$ M IPTG needed for Halopseudomonads represents one-tenth of concentrations commonly used in Pseudomonads or *B. subtilis* and is only slightly higher than used for *E. coli* Tuner (DE3) (Binder et al., 2014, 2016; Hogenkamp et al., 2021; Li et al., 2019). Species-specific differences may be caused by variances in inducer import, inducer tolerance, but also plasmid copy numbers, as reported for, among others, *H. aestusnigri* and *H. oceani* (Chan et al., 2023).

## Chromosomal integration of reporter genes by Tn7 transposition

Expression of a chromosomally integrated gene of interest is often favourable as compared to a plasmid-based gene since it enables the cultivation without the addition of antibiotics and factors such as plasmid loss and plasmid copy number do not play a role (Jahn et al., 2016). Chromosomal integration can occur either at randomised positions, e.g. via Tn5 transposition, or site-specific, via homologous recombination or Tn7





**FIGURE 3** Evaluation of heterologous gene expression in *Halopseudomonas* strains using different promoters. Induction profiles of different promoters requiring activation ( $P_{araBAD}$ ,  $P_{nagAa/nagR}$ ) (A) or de-repression ( $P_{tac}$ ) (B). Expression of reporter genes was induced by L-arabinose, salicylic acid, or IPTG at the beginning of the logarithmic growth phase after 4.5 h of cultivation. Cells treated with 70% EtOH served as negative controls. The data represent the means of biological triplicates, and error bars indicate the calculated standard deviations. (C) The dynamic range is shown as a heatmap, increasing from white to dark green.

transposition into the *attTn7*-site. The *attTn7*-site is located in a neutral region around 27 bp downstream of the gene *glmS* encoding a transaminase involved in cell wall biosynthesis (Mitra et al., 2010; Peters & Craig, 2001). Interestingly, the here investigated *Halopseudomonas* carry two copies of this essential gene instead of one. Therefore, their genomes apparently provide two sites for Tn7 integration: *H. aestusnigri* VGXO14 (WP\_088273794.1, WP\_088276615.1); *H. bauzanensis* BZ93 (WP\_074778335.1, WP\_036992462.1); *H. litoralis* 2SM5 (WP\_090273138.1, WP\_090272410.1); and *H. oceani* KX20 (WP\_104738233.1, WP\_104738216.1). Even though the two copies of these genes in each strain are not identical in sequence, the TnsD recognition sites (gene sequences that translate to

PRNLAKSVTVE [Mitra et al., 2010]) were found to be highly conserved. *H. litoralis* 2SM5 is the only strain with a closed genome sequence available that allows for precise localisation of these sites; we, therefore, chose to investigate this strain for the accessibility of both potential sites for Tn7 transposition. To this end, we constructed two mini-Tn7 vectors differing in selection markers and fluorescence reporter genes, namely pUC18-R6KT-miniTn7T-Km- $P_{tac/lacI}$ -*mcherry*, and pBG-13 (Zobel et al., 2015) harbouring *sfgfp* under the control of the constitutive promoter  $P_{em7}$ .

The Tn7 transposition was performed via triparental conjugational gene transfer with the donor strains *E. coli* S17-1/pUC18R6KT-miniTn7T-Km- $P_{tac/lacI}$ -*mcherry* and *E. coli* S17-1/pTNS2 and the acceptor strain *H.*

*litoralis* 2SM5R. The success of Tn7 integration was determined using an analytic PCR with two site-specific forward primers that bind within the sequence of either *glmS1* or *glmS2*, and a reverse primer that binds within the integration cassette. The obtained PCR products (Figure 4A) were confirmed by sequencing analysis. Of 12 clones analysed after transposition with miniTn7T-Km-P<sub>tac/lacI</sub>-*mcherry*, five had the transposon integrated downstream of *glmS1*, and likewise five carried the transposon downstream of *glmS2*; for two, the PCR results were inconclusive. Hence, both copies of *glmS* were indeed addressable for Tn7 transposition. Only a limited number of bacterial genomes are reported to contain more than one copy of *glmS* accompanied by two *attTn7*-sites adjacent to these genes; among them are different *Burkholderia* spp.,  $\beta$ -proteobacteria with usually two chromosomes (Choi et al., 2006). *B. mallei*, which harbours two *glmS* copies on chromosome 1, was studied in more detail. Here, Tn7 transposition was found to occur at both sites; however, in contrast to our observations with *H. litoralis*, one *attTn7*-site was clearly preferred, with 92% of the clones having the transposon integrated downstream of *glmS1* only (Choi et al., 2006). In *B. mallei*, *glmS*-1 is located closer to the origin of replication. In *H. litoralis*, in contrast, both *glmS* genes are located at similar distances from *oriC* (Figure 4A). This might point to an explanation for a presumable more randomised mode of integration in the latter.

As the localisation of an integron in the chromosome often influences the expression strength (Chaves et al., 2020; Englaender et al., 2017; Sauer et al., 2016), we compared the output of the LacI/P<sub>tac</sub> expression systems of transposons integrated at each of the *attTn7*-sites of *H. litoralis* based on the mCherry fluorescence 20 h after induction of gene expression. Notably, no significant differences in fluorescence intensities were detected between both integration loci (Figure 4C). This observation matches previous reports regarding the correlation of protein production with the location of the genomic integration locus with regard to *oriC* (Chaves et al., 2020), independent of the strand polarity (Sauer et al., 2016).

Afterwards, we evaluated the simultaneous occupation of both *attTn7*-sites by transforming a strain carrying Tn7- LacI/P<sub>tac</sub>-*mcherry* in *attTn7.1* with a second Tn7 transposon harbouring a *sfgfp* gene behind a constitutively active promoter on plasmid pBG-13. Another reverse primer was designed to bind within the *sfgfp* sequence to verify the second integration. Thus, the presence of a PCR product indicated that the transposition occurred into the respective site (Figure 4B); sequencing of the PCR products confirmed subsequent transpositions into both sites. During cultivation and expression, it was observed that the fluorescence reporter genes of both sites were expressed, as confirmed by recording the specific emission spectra of

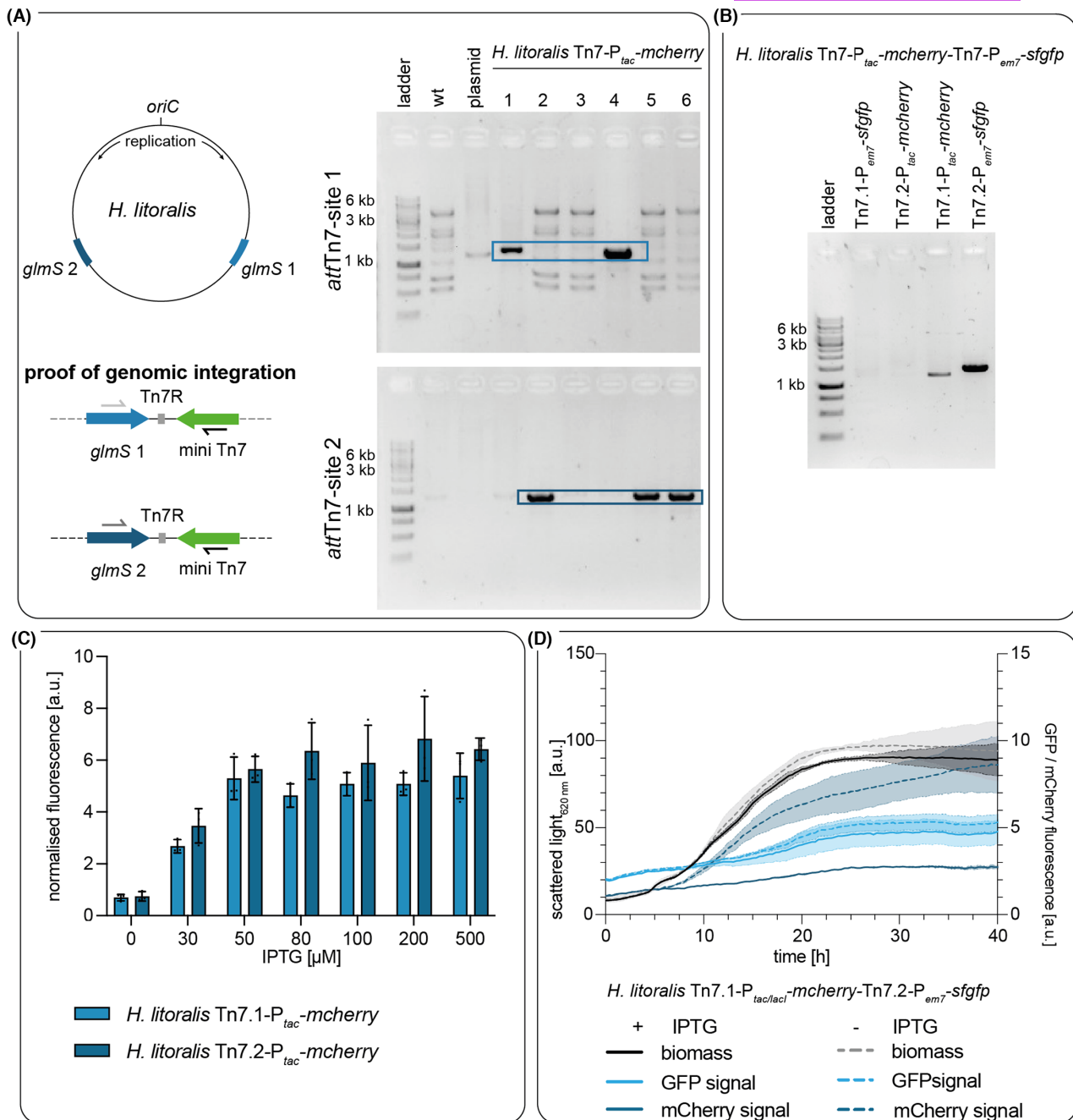
mCherry and sfGFP (Figure S11A). In the majority of the cultures, the mCherry signal appeared immediately after the addition of IPTG, whereas the sfGFP signal unexpectedly did not correlate with the biomass signal despite the presence of a constitutive promoter (Figure 4D). The growth of the cultures appeared not to be affected by the expression of both recombinant genes (Figure S11C).

However, in four out of 45 cultures, the sfGFP signal correlated with the biomass signal (Figure S11B). Notably, the mCherry signal increased in these cultures only in the stationary growth phase despite the time point of IPTG supplementation (Figure S11B), which was not observed when only one locus harboured the transposon. Thus, interference between both loci acting either on the expression level or strain stability may be assumed, leading to repression or loss of the target genes. Studies regarding the simultaneous occupation of both Tn7-sites in *B. mallei* or exploitation of three artificial *attTn7*-sites in *Salmonella enterica* reported, in contrast to some extent a correlation of the expression signal with the integron copy number (Bruckbauer et al., 2015; Roos et al., 2015). At present, the underlying mechanism remains elusive; it may involve an unknown regulatory mechanism.

In summary, we have demonstrated that both *attTn7*-sites in the genome of *H. litoralis* are suitable for integrating expression modules. They may be exploited to carry homologous target or reporter genes, for complementation of mutants, or to establish novel biocatalytic functions or reporter genes (Choi et al., 2006; Norris et al., 2010).

## CONCLUSION

*Halopseudomonas* constitutes an only recently established phylum of Proteobacteria. These bacteria, discovered in various challenging habitats over the last two decades, possess only limited metabolic flexibility and apparently produce a variety of polyester hydrolases. Here, we have studied four selected species of *Halopseudomonas* and could demonstrate the metabolism of several dicarboxylic acids with sebacic acid and succinic acid as preferred carbon sources. We further isolated rifampicin-resistant strains designated *H. aestusnigri* VGXO14R, *H. bauzanensis* BZ93R, *H. litoralis* 2SM5R, and *H. oceani* KX20R and provide protocols for electroporation and conjugational gene transfer. Moreover, a genetic toolbox was successfully established, including plasmids with different origins of replication and constitutive and inducible promoter systems. The species-specific differences in tolerance and induction profiles observed for different *Halopseudomonas* species suggest a chassis-a-la-carte model (Czajka et al., 2017; Fatma et al., 2020; Riley & Guss, 2021), allowing to select the species



**FIGURE 4** Genomic integration of a reporter gene construct into two *attTn7*-sites of *H. litoralis* 2SM5R. (A) Identification of integration sites. The localisation of genes *glmS1* (light blue) and *glmS2* (dark blue) within the genome of *H. litoralis* 2SM5 is shown (illustration not to scale). Transposition events were verified by PCR using the primer pairs shown by convergent arrows with a *glmS1* or *glmS2* specific forward primer in combination with a primer binding in the transposon either within the antibiotic resistance gene (pUC18R6KT-miniTn7T-Km- $P_{tac}$ -*mcherry*) or within gene *sfgfp* (pBG-13). The PCR results after transposition into the different *attTn7*-sites are shown on the right-hand side for six strains with one transposon. Crude cell extract of *H. litoralis* wild type was applied as control. Blue boxes indicate PCR products of the expected size (1300 bp). (B) One of the identified strains was subsequently transformed with a second and different Tn7 transposon. The colony PCR was performed using a primer combination indicating the integration of  $P_{em7}$ -*sfgfp* either into the first (Lane 1) or the second (Lane 4) integration site and integration of  $P_{tac/lacI}$ -*mcherry* (lane 2 (*attTn7*-site 1); lane 3 (*attTn7*-site 2)) (C) Comparison of the gene expression between two strains with miniTn7T-Km- $P_{tac/lacI}$ -*mcherry* in either the first (light blue) or second (dark blue) *attTn7*-site. The gene expression was induced by adding IPTG after 4.5 h of cultivation at the beginning of the logarithmic growth phase. (D) Simultaneous expression of target genes integrated in different *attTn7* sites. The biomass signal (black) of *H. litoralis* Tn7.1- $P_{em7}$ -*sfgfp*-Tn7.2- $P_{tac}$ -*mcherry* as well as the sfGFP fluorescence (light blue, Tn7.1-site) and the mCherry fluorescence (dark blue, Tn7.2-site) were measured with a microbioreactor system (BioLector 1). The cultures were induced with 50  $\mu$ M IPTG (dashed line) after 4.5 h of cultivation or treated with 70% EtOH as negative controls (continuous line). The data represent the mean of biological triplicates, and error bars or shadows indicate the calculated standard deviations.



which best fits the desired properties rather than aiming for a one-fits-all *Halopseudomonas* model (Czajka et al., 2017; Fatma et al., 2020; Riley & Guss, 2021). We further demonstrated for *H. litoralis* the presence of two *attTn7*-sites suitable for stable chromosomal integration of expression modules. Hence, the results reported here will contribute to further exploring the physiology and biotechnological applications of *Halopseudomonas* species.

## AUTHOR CONTRIBUTIONS

**Luzie Kruse:** Conceptualization (supporting); formal analysis (lead); investigation (lead); methodology (lead); validation (lead); visualization (lead); writing – original draft (lead). **Anita Loeschcke:** Conceptualization (supporting); funding acquisition (equal); project administration (supporting); writing – review and editing (supporting). **Jan de Witt:** Investigation (supporting); methodology (supporting). **Nick Wierckx:** Conceptualization (supporting); funding acquisition (supporting); project administration (supporting); writing – review and editing (supporting). **Karl-Erich Jaeger:** Conceptualization (supporting); funding acquisition (supporting); project administration (lead); writing – review and editing (supporting). **Stephan Thies:** Conceptualization (lead); funding acquisition (equal); project administration (supporting); writing – review and editing (lead).

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## CONFLICT OF INTEREST STATEMENT

The authors declare that there are no competing interests associated with the manuscript.

## DATA AVAILABILITY STATEMENT

All data are available upon request.


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## SUPPORTING INFORMATION

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