Identification of Gip as a novel phage-encoded gyrase inhibitor protein of Corynebacterium glutamicum

Larissa Kever | Max Hünnefeld | Jannis Brehm | Ralf Heermann | Julia Frunzke

1Institute of Bio- und Geosciences, IBG-1: Biotechnology, Forschungszentrum Jülich, Jülich, Germany
2Institut für Molekulare Physiologie, Biozentrum II, Mikrobiologie und Weinforschung, Johannes-Gutenberg-Universität Mainz, Mainz, Germany

Correspondence
Julia Frunzke, Institute of Bio- und Geosciences, IBG-1: Biotechnology, Forschungszentrum Jülich, 52425 Jülich, Germany.
Email: j.frunzke@fz-juelich.de

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Abstract

By targeting key regulatory hubs of their host, bacteriophages represent a powerful source for the identification of novel antimicrobial proteins. Here, a screening of small cytoplasmic proteins encoded by the CGP3 prophage of Corynebacterium glutamicum resulted in the identification of the gyrase-inhibiting protein Cg1978, termed Gip. Pull-down assays and surface plasmon resonance revealed a direct interaction of Gip with the gyrase subunit A (GyrA). The inhibitory activity of Gip was shown to be specific to the DNA gyrase of its bacterial host C. glutamicum. Overproduction of Gip in C. glutamicum resulted in a severe growth defect as well as an induction of the SOS response. Furthermore, reporter assays revealed an RecA-independent induction of the cryptic CGP3 prophage, most likely caused by topological alterations. Overexpression of gip was counteracted by an increased expression of gyrAB and a reduction of topA expression at the same time, reflecting the homeostatic control of DNA topology. We postulate that the prophage-encoded Gip protein plays a role in modulating gyrase activity to enable efficient phage DNA replication. A detailed elucidation of the mechanism of action will provide novel directions for the design of drugs targeting DNA gyrase.

KEYWORDS
bacteriophages, DNA gyrase, gyrase inhibitors, prophage induction, topoisomerase II inhibitors

1 | INTRODUCTION

Bacteriophages represent the “dark matter” of the biological world (Hatfull, 2015; Ofir & Sorek, 2018; Rohwer & Youle, 2011). With the recent massive expansion of the genomic sequence space, the number of functionally unknown open reading frames (ORFs) in phage genomes is continuously increasing (Yin & Fischer, 2008). By targeting diverse cellular processes and regulatory hubs in their host cell, bacteriophages represent a rich source for the identification of novel antibacterial proteins as well as for the establishment of highly efficient molecular tools (De Smet et al., 2017; Nobrega et al., 2018; Roach & Donovan, 2015; Schroven et al., 2021). Especially, small cytoplasmic phage proteins have been shown to influence and reprogram a variety of key cellular processes, including transcription, translation, cell division, and central metabolism (De Smet et al., 2017; Orr et al., 2020; Storz et al., 2014).

DNA gyrase represents a type IIA topoisomerase present in all bacteria and plays a crucial role in the homeostatic control of DNA topology. Because of its unique ability to introduce negative supercoils into covalently linked double-stranded DNA (dsDNA), the
activity of DNA gyrase is indispensable for bacterial growth and a key target of antibacterial agents. The heterotetrameric enzyme consists of two GyrA and two GyrB subunits (GyrA<sub>2</sub>GyrB<sub>2</sub>). While the GyrA subunit of the enzyme catalyzes the breakage and resealing of dsDNA, the GyrB subunit exhibits ATPase activity (Bush et al., 2015; McKie et al., 2021; Vanden Broeck et al., 2019). Currently, two major classes of small molecules drugs targeting the bacterial gyrase are known: the aminocoumarins and the quinolones (Collin et al., 2011). Besides a range of small molecules, also some proteinaceous, bacterial toxins were found to inhibit the activity of the gyrase, including Microcin B17 (Pierrett & Maxwell, 2003), a glycine-rich peptide found in *Escherichia coli* strains carrying the mcb operon as well as the CcdB toxin as part of the ccd toxin-antitoxin system encoded by the F-plasmid (Dao-Thi et al., 2005; Miki et al., 1992).

*Corynebacterium glutamicum*—a member of the phylum Actinobacteria—is an important industrial platform organism used for the industrial production of a wide range of value-added compounds, including amino acids, organic acids, and proteins (Wendisch et al., 2016). The genome of the model organism *C. glutamicum* ATCC 13032 contains four cryptic prophages (CGP1-4) (Frunzke et al., 2008; Ikeda & Nakagawa, 2003). The largest prophage CGP3 (~219 kb, containing also prophage CGP4) was shown to be inducible in an SOS-dependent manner as well as in an SOS-independent manner (Helfrich et al., 2015; Nanda et al., 2014; Pfeifer et al., 2016). Recently, the Lsr2-type protein CgpS was identified as a prophage-encoded nucleoid-associated protein involved in the silencing of phage gene expression maintaining the lysogenic state of the large CGP3 prophage (Pfeifer et al., 2016). Interference with CgpS binding was shown to result in prophage activation and consequently cell death.

In this study, a systematic screening of small cytoplasmic proteins encoded by the CGP3 prophage of *C. glutamicum* resulted in the identification of phage proteins causing severe growth defects and prophage induction. The small phagic protein Cg1978 was further shown to directly target the DNA gyrase enzyme by interacting with the GyrA subunit and inhibiting the supercoiling activity of the *C. glutamicum* DNA gyrase in vitro. Cg1978 was therefore termed Gip for gyrase inhibiting protein. A detailed elucidation of the mechanism of action may point to novel directions for the design of drugs targeting DNA gyrase.

## RESULTS

### 2.1 Systematic screening of small CGP3-encoded proteins influencing growth and prophage induction

Most of the proteins encoded by the cryptic CGP3 prophage are of unknown function. Phage proteins were shown to frequently target key regulatory hubs to shut down bacterial metabolism (Roach & Donovan, 2015). In this study, we screened the impact of overall 11 small (<75 amino acids), cytoplasmic phage-encoded proteins on cellular growth and prophage induction. For this purpose, plasmid-based overexpression (pAN6-GOI) of the selected genes of interest was performed in the prophage reporter strain *C. glutamicum* ATCC 13032::P<sub>lyS-eyfp</sub>. In previous studies, this chromosomal reporter (P<sub>lyS-eyfp</sub>) was successfully established to translate prophage activation into a fluorescent output (Helfrich et al., 2015). During cultivation, biomass was measured as a function of backscattered light intensity with an excitation wavelength of 620 nm. Following this approach, the overproduction of 9 out of 11 phage proteins (Cg1902, Cg1910, Cg1924, Cg1925, Cg1971, Cg2026, Cg2035, Cg2045, and Cg2046) displayed comparable phenotypes as the empty vector control regarding backscatter signal and fluorescence output measured via flow cytometry (Figure 1c and Figure S1a,b). By contrast, overproduction of Cg1914 and Cg1978 showed a significant effect on growth and prophage induction in the presence of 50 µM IPTG. Cg1914 overproduction resulted in a reduced growth rate (µ = 0.23 ± 0.01 hr<sup>−1</sup>) and a reduced final backscatter (Figure 1a, blue line) compared with the empty vector control pAN6 (µ = 0.38 ± 0.01 hr<sup>−1</sup>). In the case of Cg1978, overproduction led to an elongated lag-phase, but the final backscatter as well as the growth rate in the exponential phase (µ = 0.36 ± 0.01 hr<sup>−1</sup>) (Figure 1a, red line) were comparable with those of the empty vector control.

A comparable impact on cell growth due to cg1914 or cg1978 overexpression was also detected in the prophage-free strain MB001, indicating that the observed growth defect was independent of the presence and/or activity of the CGP3 island (Figure 1b).

For both target proteins, Cg1914 and Cg1978, overproduction resulted in an increased fluorescence signal after 24 hr of cultivation (cg1914: 7.4 ± 2.6% induced cells, cg1978: 3.5 ± 0.2% induced cells) indicating CGP3 prophage induction in the respective subpopulation (Figure 1c). As a positive control, we expressed an N-terminally truncated variant of the prophage silencer CgpS (CgpS-N), which was previously shown to trigger prophage induction (Pfeifer et al., 2016). Since overproduction of Cg1914 and Cg1978 showed a high impact on prophage induction, we tested the inducibility of the CGP3 prophage in mutants lacking the respective genes using a plasmid-based prophage reporter (P<sub>lyS-ys-venus</sub>). Remarkably, the corresponding strains *C. glutamicum* ATCC 13032 Δcg1914 and Δcg1978 featured no difference—neither in cell growth nor in prophage inducibility—upon treatment with the DNA-damaging antibiotic mitomycin C (Tomasz, 1995), which was used to trigger SOS-dependent prophage induction (Figure S2). These results indicated that both proteins are not essentially involved in SOS-dependent CGP3 induction. Prophage induction, therefore, appeared to be an indirect effect of Cg1914 or Cg1978 overproduction. Based on further results described in the following, we focused on the role and cellular target of the small phage protein Cg1978.

### 2.2 Overproduction of Cg1978 triggers the activation of SOS response and RecA-independent prophage induction

As the CGP3 prophage was already characterized to be inducible in an SOS-dependent manner as well as in an SOS-independent
manner (Helfrich et al., 2015; Nanda et al., 2014; Pfeifer et al., 2016), we determined the SOS dependency of Cg1978-mediated prophage induction. To this end, the fluorescent outputs of different reporter strains were measured via flow cytometry in a time-resolved manner during cg1978 overexpression. Besides the prophage reporter strain (C.g. ATCC 13032::P<sub>lys</sub>-eyfp), an SOS reporter strain (C.g. ATCC 13032::P<sub>recA</sub>-venus) as well as an SOS-deficient prophage reporter strain (C.g. ATCC 13032 Δ<sub>recA</sub>::P<sub>lys</sub>-eyfp) lacking the coprotease RecA—required for the induction of the host SOS response (Janion, 2008)—were used.

As described above, overexpression of cg1978 resulted in a similar growth phenotype of all reporter strains characterized by an elongated lag-phase (marked in gray) with subsequent wild type-like growth (Figure 2a). The impaired cell growth under cg1978 overexpression conditions was confirmed by time-lapse fluorescent microscopy of a C. glutamicum microcolony of the prophage reporter strain, which was grown in a microfluidic chamber. Increased levels of Cg1978 led to elongated cell morphology and a small fraction of cells featuring a strongly increased output of the prophage reporter (Figure 2c and Videos S1 and S2).

Measurement of the reporter output over time not only confirmed CGP3 induction but also revealed an induction of the cellular SOS response (Figure 2b). Interestingly, the wild type-like and the RecA-deficient prophage reporter strain showed nearly the same percentage of induced cells upon cg1978 overexpression throughout the entire measurement, reaching a peak value after 9.5 hr of cultivation (Δ<sub>recA</sub>::P<sub>lys</sub>-eyfp: 4.6 ± 0.7, P<sub>lys</sub>-eyfp: 4.8 ± 0.9). These results emphasize RecA-independent CGP3 induction as an indirect effect of cg1978 overexpression.

Remarkably, all reporter strains revealed an increasing fluorescence during the lag-phase, which decreased again upon transition into the exponential growth phase (Figure 2b), suggesting the growth of a subpopulation resistant to Cg1978 overproduction effects.

2.3 | Cg1978 directly interacts with gyrase subunit A (GyrA)

To identify the direct cellular target of Cg1978, we performed an in vitro pull-down assay. For this purpose, the small protein Cg1978 containing a C-terminal Strep-tag was overproduced in E. coli BL21 (DE3) and purified via affinity purification. The purified target protein was incubated with C. glutamicum cell extract and this sample was again passed over a Strep-Tactin column to identify proteins copurifying with Cg1978. SDS-PAGE analysis of proteins coeluting with Cg1978 revealed an additional protein band at ~100 kDa (Figure 3a). Analysis of the copurified protein via MALDI-TOF as well as LC-MS/MS analysis of the whole elution fraction indicated the copurification of Cg1978 with C. glutamicum (C.g.) DNA gyrase subunit A (~95 kDa), a subunit of the heterotetrameric ATP-dependent DNA gyrase complex (A<sub>2</sub>B<sub>2</sub>). The DNA gyrase belongs...
FIGURE 2  Time-resolved measurement of reporter outputs upon Cg1978 overproduction in Corynebacterium glutamicum showed activation of SOS response and RecA-independent prophage induction. Cultivation of an SOS reporter strain ATCC 13032::P_recA-venus, a prophage reporter strain ATCC 13032::P_{lys}eyfp and an SOS-deficient prophage reporter strain ATCC 13032 ΔrecA::P_{lys}eyfp carrying the plasmids pAN6 or pAN6-cg1978 was performed in the BioLector® microcultivation system in CGXII-Kan25 medium with 2% (w/v) glucose and 50 µM IPTG. All data represent mean values with SDs from three independent biological replicates (n = 3). (a) Growth curves based on the backscatter measurements in the BioLector® microcultivation system. The elongated lag-phase of the Cg1978 overproducing strain is marked in gray. (b) Percentage of induced cells based on the flow cytometric measurements of eYFP or Venus fluorescence of the reporter strains. (c) Time-lapse fluorescence imaging of the C. glutamicum prophage reporter strain ATCC 13032::P_{lys}eyfp carrying the pAN6-cg1978 plasmid. Cells were cultivated in PDMS-based microfluidic chip devices (Grüninger et al., 2015) using CGXII-Kan25 medium with 2% (w/v) glucose. The medium was continuously supplied with a flow rate of 300 nl/min. Overexpression of cg1978 was induced by the addition of 50 µM IPTG. Fluorescent images represent cutouts from Videos S1 and S2.
to the subclass of type II topoisomerases and plays a key role in DNA metabolism as it is able to introduce negative supercoiling to double-stranded DNA in an ATP-dependent manner. Gyrase subunit B was also detected in the elution fraction, but in a significantly lower amount compared to GyrA rather suggesting unspecific copurification.

As a next step, surface plasmon resonance spectroscopy was used to examine the binding affinity of Cg1978 to GyrA. The sensorgram revealed a stable and specific 1:1 interaction between Cg1978 and GyrA with a high association rate ($k_a = 9.7 \times 10^4$ M$^{-1}$ s$^{-1}$) and a slow dissociation rate ($k_d = 5.2 \times 10^{-4}$ s$^{-1}$) resulting in an overall affinity ($K_D$) of 5.4 nM (Figure 3b). Purification of Cg1978-C-His and GyrA-N-Strep for SPR analysis are shown in Figure S3.

**2.4 | Cg1978 inhibits DNA supercoiling via interaction with the DNA gyrase in vitro**

Due to its essential role for cell survival, DNA gyrase represents an important drug target of antibiotics and protein-based inhibitors (Collin et al., 2011). Based on the observed growth defect upon Cg1978 overproduction and the interaction with GyrA, we
further assessed the effect of Cg1978 on gyrase activity by performing in vitro supercoiling inhibition assays with the purified enzyme.

For this purpose, Cg1978-C-His, GyrA-C-Strep, and GyrB-N-Strep from C. glutamicum were purified separately using affinity chromatography (Figure 4a). The formation of the heterotetrameric enzyme complex was obtained by incubating equimolar amounts of both gyrase subunits on ice for 30 min. In the first step, the activity of the purified C.g. DNA gyrase was measured by incubating 0.5 µg relaxed pBR322 plasmid DNA with different C.g. gyrase concentrations. The addition of increasing C.g. gyrase concentrations led to an increase in supercoiling of the plasmid DNA resulting in maximal supercoiling using 50 nM of C.g. gyrase. This concentration was defined as 1 U (corresponding to a specific activity of 1.88 × 10^3 U/mg) and was used for testing the potential inhibitory effect of Cg1978 (Figure 4b).

As shown in Figure 4c, incubation of increasing concentrations of Cg1978 (0.2–60 µM) with 1 U C.g. DNA gyrase and relaxed plasmid DNA resulted in a decreased supercoiling activity of the C.g. DNA gyrase. Complete inhibition of the introduction of supercoils by DNA gyrase was achieved by the addition of 10 µM Cg1978 leading to an accumulation of the relaxed/nicked conformation. Therefore, we named the gene product of cg1978 Gip (gyrase inhibiting protein).

As a negative control, Gip was incubated with relaxed plasmid DNA in absence of DNA gyrase to visualize any potential nuclease activity in the elution fraction of Gip. A slight band reflecting linear DNA could be detected when adding 40 µM Gip. However, incubating increasing concentrations of Gip with the C.g. DNA gyrase did not lead to an accumulation of linear DNA. As a positive control, the known fluoroquinolone-based gyrase-inhibitor ciprofloxacin (CFX) stabilizing the gyrase–DNA cleavage complex (Drlica & Malik, 2003) was used showing significant inhibition of the supercoiling activity of the C.g. gyrase at 50 µM (16.6 µg/ml). This was in line with already published data for the DNA gyrase of Mycobacterium smegmatis, which showed 50% inhibition of the supercoiling activity of 1 U gyrase by the addition of 10 µg/ml CFX (Manjunatha et al., 2002). As for Gip, inhibition of the DNA gyrase via CFX led to an accumulation of the relaxed/nicked plasmid conformation.

To further investigate the activity profile of Gip, we determined its effect on the DNA gyrase of Mycobacterium tuberculosis and E. coli. Gyrase subunits A of the actinobacterial species C. glutamicum and M. tuberculosis (Mtb) share a sequence identity of ~71%, while GyrA of C. glutamicum and E. coli (E.c.) only show a ~45% sequence identity (Figures S4 and S5). As described previously for the DNA gyrase of C. glutamicum, 1 U of the E.c. and M.tb. gyrases were used to examine supercoiling inhibition via Gip. The supercoiling assay showed no significant change in the supercoiling activity of the respective gyrases when adding increasing concentrations of Gip. In case of the M.tb. gyrase, a slight shift from supercoiled plasmid DNA to linear plasmid DNA could be detected by addition of 40 µM Gip, which was even less pronounced for the E.c. gyrase. The corresponding assays are shown in Figure S6.

2.5 | Compensatory expression of gyrAB and topA as a response to gyrase inhibition via Gip

Supercoiling inhibition assays showed that Cg1978 (Gip) is a gyrase-inhibiting protein. As the activity of DNA gyrase is indispensable for bacterial growth, we investigated the impact of gip overexpression on the transcriptome of C. glutamicum. Since the gip overexpression strain revealed a wild type-like growth rate after an elongated lag-phase, we were especially interested to understand how the bacterial host counteracts gyrase inhibition. For this purpose, comparative transcriptome analysis of the C. glutamicum ATCC 13032 strain containing the empty vector control and the strain containing the overexpression plasmid pAN6-gip was performed using DNA microarrays. The shown transcriptomic changes are based on mRNA levels of cells harvested at an OD_600 of 6 in the mid-exponential growth phase.

The gip overexpressing strain showed a partial upregulation of CGP3 genes due to overexpression of gip (Table S4), confirming the prophage induction also revealed by the above-described reporter assays. Apart from the CGP3 region, overexpression of gip led to upregulation of 352 genes and downregulation of 333 genes reflecting the high impact of gyrase inhibition on overall cell metabolism.

Interestingly, both gyrase subunits gyrA and gyrB were markedly upregulated showing a more than 4-fold increase in expression levels (Table 1). In contrast, topA coding for topoisomerase I, which catalyzes the opposing reaction of DNA gyrase by removing negative supercoils, showed a reduced expression level. Moreover, the expression of further genes involved in DNA metabolism was influenced by gip overexpression including for example the reduced expression of genes coding for helicases (exemplarily cg0838, cg0842, cg0843, cg0845, cg0889, and cg1498). Additionally, 10 targets of the SOS key player LexA, for example, recN (DNA repair) and ftsK (cell division and chromosome segregation), showed an increased mRNA ratio, which was in line with the high fluorescent outputs of the SOS reporter strain upon Gip overproduction (Table 1, see Table S4 for the complete list of genes with altered expression levels).

3 | DISCUSSION

In this study, a screening of small cytoplasmic proteins encoded by the CGP3 prophage of C. glutamicum resulted in the identification of the novel gyrase inhibitor protein Gip (Cg1978, 6.8 kDa). Overproduction of Gip resulted in significant growth defects and prophage induction in a subpopulation. Further characterization of this small phagic protein confirmed a specific, stable, and high-affinity interaction with the GyrA subunit and inhibition of the supercoiling activity of the DNA gyrase in vitro.

DNA gyrase possesses the unique ability to catalyze the ATP-dependent negative supercoiling of double-stranded DNA by cleaving and rejoining it (Bush et al., 2015). Supercoiling inhibition assays showed that Gip-mediated gyrase inhibition resulted in the accumulation of the nicked/relaxed plasmid conformation, while no more
superciled plasmid DNA was detectable (Figure 4). Different molecular mechanisms of gyrase inhibition have been described so far. Fluoroquinolones and the well-characterized proteinaceous bacterial toxins Microcin B17 (MccB17, 3.1 kDa) and CcdB (11.7 kDa) poison the DNA gyrase by stabilizing the gyrase-DNA cleavage complex leading to double-strand DNA breaks (Bernard et al., 1993; Drlica & Malik, 2003; Pierrat & Maxwell, 2003). In contrast, aminocoumarins (e.g. novobiocin) inhibit ATP hydrolysis as the binding site overlaps with the ATP-binding pocket of the GyrB subunit (Maxwell & Lawson, 2003). Further proteins targeting DNA gyrase are pentapeptide repeat proteins (PRPs) like Qnr proteins or MfpA, whose inhibitory interaction is proposed to be based on DNA mimicry (Shah & Heddle, 2014). However, no conserved domains of Gip with other known proteinaceous gyrase inhibitors could be identified using the

Note: A genome-wide comparison of mRNA levels of the Corynebacterium glutamicum ATCC 13032 strain overexpressing gip and the wild-type strain carrying the empty vector control was performed. The shown mRNA ratios indicate mean values from three independent biological replicates (n = 3). The strains were cultivated in CGXII-Kan25 minimal medium with 2% (w/v) glucose and 50 µM IPTG and mRNA was prepared from cells harvested at an OD 600 of 6. The mRNA ratios were calculated by dividing the mRNA levels of the gip overexpressing strain by the mRNA levels of the strain carrying the empty vector control. The table includes selected genes from a larger set which showed a changed mRNA level in all experiments (mRNA ratio >2.0: upregulation [red] or <0.5: downregulation [green], p-value <0.05).
Investigations regarding the activity profile of Gip suggested that the inhibitory activity seems to be highly specific for the DNA gyrase of its bacterial host C. glutamicum, as the DNA gyrase of M. tuberculosis and E. coli were not significantly affected by Gip. Similar observations were made for the proteinaceous bacterial toxins Microcin B17 (MccB17, 3.1 kDa) and CcdB (11.7 kDa), which target the DNA gyrase of their host E. coli in vitro, while no inhibition of the DNA gyrase of M. smegmatis could be detected (Chatterji et al., 2001). Even though DNA gyrase is a conserved protein among bacteria, Gram-positive and Gram-negative bacteria show substantial differences in the amino acid sequence of GyrAB (Madhusudan & Nagaraja, 1996; Manjunatha et al., 2000). Accordingly, the absence of specific target residues potentially explains the different levels of susceptibility of the DNA gyrase to proteinaceous toxins (Chatterji et al., 2001).

Reporter outputs of the RecA-dependent SOS reporter strain and transcriptional analysis of Gip overproducing cells revealed an induction of the SOS response (Figure 2b, Table 1). These findings are in agreement with already published data describing activation of the SOS response as one of the pleiotropic effects of gyrase inhibition (Jeong et al., 2006). Stabilization of the gyrase-cleaved DNA complex results in arrested replication forks and widespread DNA damage by stimulating the formation of DNA breaks triggering the SOS response (DeMarini & Lawrence, 1992; Dwyer et al., 2007).

Gip overproduction was further shown to activate the induction of the CGP3 prophage. However, the fact that the observed growth defect of Gip overproducing cells is independent of the presence of the CGP3 prophage (Figure 1b) and that deletion of gip did not result in altered inducibility of CGP3 (Figure S2) emphasize prophage activation as an indirect effect of Gip overproduction. Recent studies already confirmed that CGP3 is inducible in an SOS-dependent manner as well as in an SOS-independent manner (Helfrich et al., 2015; Nanda et al., 2014; Pfeifer et al., 2016). As the wild type and the RecA-deficient prophage reporter strain revealed nearly identical fluorescent outputs, we propose that prophage induction occurred mainly in a RecA-independent manner. Here, influencing the introduction of supercoils due to gyrase inhibition might be a possible reason for CGP3 induction. The lysogenic state of CGP3 is maintained by the Lsr2-type silencer protein CgpS, which was shown to target more than 35 AT-rich regions within the CGP3 element (Pfeifer et al., 2016). The formation of this dense nucleoprotein complex was shown to be crucial for efficient CgpS-mediated silencing (Wiechert et al., 2020). Especially in the case of proteins-targeting AT-rich DNA sequences, the topologic state of DNA can affect protein–DNA interactions (Dorman & Dorman, 2016). Apart from that, different studies already demonstrated an influence of DNA supercoiling on the \( \lambda \) repressor and the lysogenic-lytic decision of phage \( \lambda \) (Ding et al., 2014; Norregaard et al., 2013, 2014).

In general, it is conceivable that the CGP3 prophage could have an advantage from encoding a gyrase inhibitor as it might allow a more efficient phage DNA replication by modulating host gyrase activity. Similar assumptions were recently made for the topo I inhibitor protein gp55.2 encoded by the T4 phage of E. coli. It was hypothesized that modulating DNA relaxation activity of topo I is required for an optimal phage yield during infection (Mattenberger et al., 2015). Another example of a phage-encoded protein altering DNA topology is represented by the gyrase-inhibiting peptide Igy encoded by phage LUZ24 infecting Pseudomonas aeruginosa (De Smet et al., 2021). Interaction of Igy with GyrB, possibly by functioning as a DNA mimicry protein, inhibits the DNA gyrase and LUZ24 infection seems to be independent of a functioning host DNA gyrase.

Global topological alterations caused by Gip overproduction were also reflected by the transcriptome analysis revealing a marked impact on global gene expression patterns (Table 1). As DNA gyrase is indispensable for replication and transcription by changing the topological state of DNA, its inhibition was previously described to globally affect the gene expression profile (Guha et al., 2018; Jeong et al., 2006). Particularly noteworthy in this context are the significantly increased mRNA levels of gyrA and gyrB as well as the down-regulation of topA. The DNA topology modulatory proteins, gyrase and topoisomerase I (topo I), catalyze opposing reactions of DNA supercoiling and relaxation (McKie et al., 2021). Previous studies already revealed that expression of the gyrAB and topA is controlled in a supercoiling-sensitive manner: While increasing DNA relaxation stimulates gyrAB expression (Menzel & Gellert, 1983), it represses expression of topoA allowing homeostatic maintenance of DNA topology (Ahmed et al., 2016; Tse-Dinh, 1985). As gyrase inhibition blocks the introduction of negative supercoils, increased expression levels of gyrAB and a decreased expression level of topA upon Gip overproduction are most probably used to counteract gyrase inhibition. The adaptation at the level of gene expression could then explain the resumed growth of the gip expressing strain—reaching almost wild type-like growth rates after a pronounced lag phase (Figure 1).

In summary, we identified Gip as a novel gyrase inhibitor protein encoded by the CGP3 prophage of C. glutamicum. Gip was shown to specifically inhibit the gyrase of its bacterial host C. glutamicum, but further studies are required to decipher its impact on the phage life cycle.

## 4 | EXPERIMENTAL PROCEDURES

### 4.1 | Bacterial strains and growth conditions

All bacterial strains and plasmids used in this study are listed in Tables S1 and S2, respectively. Corynebacterium glutamicum ATCC 13032 (NCBI reference: NC_003450.3) was used as a wild-type strain (Ikeda & Nakagawa, 2003). For growth studies and fluorescence measurements as well as for transcriptome analysis, C. glutamicum cells were precultivated in BHI (brain heart infusion, Difco BHI, BD, Heidelberg, Germany) at 30°C for 8 hr. The preculture was used to inoculate an overnight culture in CGXII minimal medium with 2% (w/v) glucose (Keilhauer et al., 1993), which was cultivated under...
the same conditions. The next day, the overnight culture was used to inoculate the main culture in CGXII minimal medium with 2% (w/v) glucose to an OD$_{600}$ of 1. All media contained kanamycin in a concentration of 25 µg/mL. Gene expression was induced using 50 µM IPTG (Isopropyl β-D-1-thiogalactopyranoside). For standard cloning applications, E. coli DH5α was cultivated in Lysogeny Broth (Difco LB, BD, Heidelberg, Germany) medium containing 50 µg/mL kanamycin (LB Kan$_{50}$) at 37°C and 170 rpm. For protein overproduction and following purification, the E. coli BL21 (DE3) strain was used. Precultivation was performed in LB Kan$_{50}$ medium, which was incubated overnight at 37°C and 120 rpm. The main culture was inoculated in LB Kan$_{50}$ medium to an OD$_{600}$ of 0.1 using the pre-culture. At an OD$_{600}$ of 0.6 gene expression was induced using 100 µM IPTG. Cells were harvested after additional 24 hr incubation at 16°C.

### 4.2 Recombinant DNA work and cloning techniques

All plasmids and oligonucleotides used in this study are listed in Tables S2 and S3, respectively. Standard cloning techniques like PCR and restriction digestion were performed according to standard protocols (Sambrook & Russell, 2001). In all cases, Gibson assembly was used for plasmid construction (Gibson, 2011). DNA regions of interest were amplified via PCR using the chromosomal DNA of C. glutamicum ATCC 13032 as a template. The plasmid backbone was cut using the indicated restriction enzymes. Sequencing and synthesis of oligonucleotides were performed by Eurofins Genomics (Ebersberg, Germany). Genomic deletions were constructed using the pK19mobsacB plasmid and the two-step homologous recombination method (Niebisch & Bott, 2001). The 500 bp up- and downstream regions of the respective gene were amplified using the indicated oligonucleotides listed in Table S3. Both PCR products and the digested pK19mobsacB plasmid (with HindIII, EcoRI) were assembled via Gibson assembly (Gibson, 2011). The correct deletion was verified by sequencing of the colony PCR product with the indicated oligonucleotides (Table S3).

### 4.3 Microtiter cultivation and reporter assays

For growth experiments and fluorescence assays, the BioLector® microcultivation system of m2p-labs (Aachen, Germany) was used (Kensy et al., 2009). The main cultivation was executed in FlowerPlates (MTP-48-B, m2p-labs) at 30°C and 1,200 rpm with a starting OD$_{600}$ of 1 using 750 µl of CGXII minimal media with 2% (w/v) glucose containing 50 µM IPTG and 25 µg/mL kanamycin. During cultivation, biomass was measured as a function of backscattered light intensity with an excitation wavelength of 620 nm (filter module: λ$_{EM}$/λ$_{EM}$: 620 nm/620 nm, gain: 15). Data for biomass measurements were baseline-corrected by subtracting the $t_0$ value from all data points. The measurements of backscatter were taken at 15 min intervals.

### 4.4 Protein purification via affinity tags

For heterologous protein overproduction, E. coli BL21 (DE3) cells containing the pET-cg1978-C-strep plasmid, the pET-gyrA-C-strep plasmid, the pET-gyrA-N-strep plasmid, the pET-gyrB-N-strep plasmid, or the pET-cg1978-C-his plasmid were cultivated as described in "Bacterial strains and growth conditions."

Cell harvesting and disruption were performed as described by Pfeifer et al. (2016). In case of Cg1978-C-Strep, buffer A (100 mM Tris–HCl, pH 8.0) with cComplete™ Protease inhibitor (Roche, Basel, Switzerland) was used for cell disruption and buffer B (100 mM Tris–HCl, 250 mM NaCl, pH 8.0) for purification. Purification of Strep-tagged Cg1978 was conducted by applying the supernatant to an equilibrated 2 ml Strep-Tactin®-Sepharose® column (IBA, Göttingen, Germany). After washing with 20 ml buffer B, the protein was eluted with 6 ml buffer B containing 15 mM d-desthiobiotin (Sigma–Aldrich, St. Louis, MO, USA). Purification of GyrA-C-Strep, GyrA-N-Strep, and GyrB-N-Strep was conducted in the same way using an adjusted buffer B$_{gyr}$, for cell disruption and purification (buffer B$_{gyr}$, 20 mM Tris–HCl, 500 mM NaCl, 10% (w/v) glycerol, 5 mM EDTA, 1 mM DTT, pH 7.9).

For purification of Cg1978-C-His, the cell pellet was resuspended in 50 ml TNI20 buffer (20 mM Tris–HCl, 300 mM NaCl, 20 mM imidazole, and pH 8.0) with cComplete™ Protease inhibitor (Roche, Basel, Switzerland), and cells were disrupted as described above. Purification of His-tagged Cg1978 was performed by applying the supernatant to an equilibrated 2 ml Ni-NTA Agarose column (Invitrogen, California, USA). After washing with 30 ml TNI20 buffer, the protein was eluted with increasing imidazole concentrations using TNI buffer (20 mM Tris–HCl, 300 mM NaCl, pH 8.0) containing 50 mM, 100 mM, 200 mM, or 400 mM imidazole.

After purification, the elution fractions with the highest protein concentration were pooled and analyzed with SDS-PAGE (Laemmli, 1970) using a 4%–20% Mini-PROTEAN® gradient gel (Bio-Rad, Munich, Germany).

### 4.5 In vitro pull-down assay and MALDI-TOF analysis

Protein purification of Cg1978-C-Strep was conducted as described above. The elution fractions showing the highest protein concentration in a Bradford assay (Bradford, 1976) were pooled and purified with size-exclusion chromatography using PD10 desalting columns (GE Healthcare, Freiburg, Germany) and buffer B (100 mM Tris–HCl, 250 mM NaCl, pH 8.0) according to manufacturer’s manual to remove excess desthiobiotin.

For the detection of possible interaction partners of the target protein on a protein-protein level, C. glutamicum ATCC 13032 wild-type cells were cultivated in a BHI medium. At an OD$_{600}$ of 5 to 6, the cells were harvested at 11,325 g and 4°C for 15 min and cell pellet of 100 ml cell culture was resuspended in 25 ml buffer A (100 mM Tris–HCl, pH 8.0) with cComplete™ Protease inhibitor (Roche, Basel, Switzerland). Cell disruption was performed using the French Press cell with a pressure of 172 mPA for five passages followed by a centrifugation step at 5,000 g for 50 min.
For copurification of possible protein interaction partners, the purified target protein Cg1978 was incubated with the C. glutamicum crude extract at RT for 1 hr. After loading the mixture to the StrepTactin column, the purification was performed as described above. The elution fractions with the highest protein concentration were precipitated by the addition of 100% (w/v) trichloroacetic acid (TCA) in a volume ratio of four units of protein to one unit TCA (Sivaraman et al., 1997). After incubation at 4°C overnight, the precipitation approach was centrifuged for 5 min at 14,000 g. The supernatant was discarded and the pellet was washed with 200 µl cold acetone twice. Afterward, the pellet was dried for 10 min at 95°C and resuspended in 30 µl 1.5 x SDS loading buffer for gel electrophoresis or in 30 µl trypsin reaction buffer provided by the Trypsin Singles, Proteomics Grade kit (Sigma–Aldrich) for LC-MS/MS sample preparation. Analysis of elution fractions via SDS-PAGE (Laemmli, 1970) was performed using a 4%–20% Mini-PROTEAN® gradient gel (Bio-Rad, Munich, Germany). After staining the gel with Comassie dye-based RAPIDstain solution (G-Biosciences, St. Louis, MO, USA) MALDI-TOF-MS measurements were performed with an Ultraflex III TOF/TOF mass spectrometer (Bruker Daltonics, Bremen, Germany) for identification of the co-purified proteins (Bussmann et al., 2010). Elution fractions were further analyzed via LC-MS/MS.

4.6 | LC-MS/MS sample preparation and measurement

LC-MS/MS was performed after TCA precipitation using the Trypsin Singles, Proteomics Grade kit (Sigma–Aldrich, St. Louis, MO, USA) according to the manufacturer’s instruction. The prepared tryptic peptide samples were separated chromatographically on a nanoLC Eksigent expert™ 425 LC system in microLC modus (Sciex) coupled with a 25 Micron ESI Electrode to a TripleTof™ 6600 mass spectrometer (Sciex). As a trap, a YMC-Triart C18 column with the dimension 5 x 0.5 mm ID, 3 µm, 12 nm (YMC) was used, combined with a YMC-Triart C18 column with 150 x 0.3 mm ID, 12 nm, 5-3 µm (YMC) as an analytical column. The column oven was set to 40°C.

For trapping, 2% acetonitrile in dd.H2O with 0.5% formic acid served as a loading solvent, whereas 0.1% formic acid was used as mobile phase A and acetonitrile with 0.1% formic acid (both LC-MS grade, ROTISOLV™, ≥99.9%, Carl Roth) as mobile phase B. First, 10 µl of each sample containing up to 8 µg of digested protein was loaded from the cooled autosampler onto the trap column using 100% loading solvent for 10 min at 10 µl/min for desalting and enrichment.

For the following separation of the peptides on the analytical column, a linear gradient with increasing concentrations of mobile phase B was used starting with 97% A and 3% B and a flow rate of 5 µl/min as an initial condition. During Information-Dependent Acquisition (IDA) and SWATH measurements, the following source and gas settings were applied: 5,500 V spray voltage, 35 psi curtain gas, 12 psi ion source gas 1, 20 psi ion source gas 2, and 150°C interface heater. Each sample was injected three times.

For shotgun measurements, the mass spectrometer was operated with a “top 50” method: Initially, a 250-ms survey scan (TOF-MS mass range m/z 400–1,500, high-resolution mode) was collected from which the top 50 precursor ions were automatically selected for fragmentation, whereby each MS/MS 97 Appendix event (mass range m/z 170–1,500, in high-sensitivity mode) consisted of a 40 ms fragment ion scan. For parent ion selection, the precursor ion intensity served as the main selection criterion. Ions with an intensity exceeding 100 counts/s and with a charge state of 2+ to 5+ were preferentially selected. Selected precursors were added to a dynamic exclusion list for 22 s and subsequently isolated using a quadrupole resolution of 0.7 amu and fragmented in the collision cell with a rolling collision energy (CE) of 10 eV. If <50 precursor ions fulfilling the selection criteria were detected per survey scan, the detected precursors were subjected to extended MS/MS accumulation time to maintain a constant total cycle time of 2.3 s.

For data analysis, the IDA data were processed with ProteinPilot™ (V5.01, Sciex, USA) using the ParagonTM Algorithm for peptide identification and the ProGroup™ Algorithm for protein identification.

4.7 | DNA microarrays

For a comparative transcriptome analysis of C. glutamicum ATCC 13032 carrying the empty pAN6 vector and cells carrying the pAN6-cg1978 vector, cultivation was performed as described in “Bacterial strains and growth conditions” using CGXII-Kan25 minimal media with 2% (w/v) glucose and 50 µM IPTG. For both strains, cells were harvested at an OD$_{600}$ of 6 in a reaction tube filled with ice (50 ml) for 5 min at 5,000 g and 4°C. RNA purification was carried out using the “RNeasy Mini”-Kit (QIAGEN, Hilden, Germany) according to the manufacturer’s manual. The preparation of labeled cDNA and DNA microarray analysis was performed as described previously (Donovan et al., 2015). The data processing was executed with in-house software according to (Polen & Wendisch, 2004). Genes with an mRNA ratio (sample/neg. control) of >2.0 (p-value <0.05) were classified as upregulated, whereas genes with an mRNA ratio of <0.5 (p-value <0.05) were classified as down-regulated. Array data were deposited in the GEO database (ncbi.nlm.nih.gov/geo) with accession number GSE151224.

4.8 | Flow cytometry

Analysis of fluorescent reporter outputs at the single-cell level was performed using the BD Accuri™ C6 flow cytometer (BD biosciences, Heidelberg, Germany). The chromophore of the yellow fluorescent protein eYFP or Venus was excited with a blue laser with an excitation wavelength of 488 nm. The fluorescence emission of eYFP and Venus was measured using a 530/30 nm standard filter. Particle size was detected using the forward light scatter (FSC). The flow cytometer was started up by flushing with filtered, dd.H2O for 10 min. For preparing flow cytometry samples, cell cultures were mixed with 1 ml flow cytometric fluid (BD™ 342003
4.9 | Cultivation and perfusion in microfluidic device

Single-cell analysis of cg1978 overexpressing cells was performed using an in-house developed microfluidic platform (Grüngerberger et al., 2013, 2015; Helfrich et al., 2015). Cultivation and time-lapse imaging were performed in CGXII minimal medium with 2% (w/v) glucose and 25 µg/ml kanamycin as described by (Pfeifer et al., 2016). Overexpression of cg1978 in the prophage reporter strain ATCC 13032::P_{lys}−eyfp carrying the pAN6-cg1978 vector was induced by adding 50 µM IPTG to the medium. An uninduced control served as a reference.

4.10 | Supercoiling inhibition assay

For the supercoiling inhibition assay, Cg1978 as well as both gyrase subunits (GyrA and GyrB) were purified by the means of a C-terminal His-Tag for Cg1978, a C-terminal Strep-Tag for GyrA, and an N-terminal Strep-Tag for GyrB as described above. Using PD10 desalting columns (GE Healthcare, Freiburg, Germany), the buffer of Cg1978 was exchanged to PBS (137 mM NaCl, 2.7 mM KCl, 8 mM Na₂HPO₄, 1.5 mM KH₂PO₄, pH 7.4). In case of GyrA and GyrB, the buffer was exchanged to 20 mM Tris-HCl, pH 7.9, 50% (w/v) glycerol, 0.5 M KCl, and 1 mM DTT. Formation of the heterotetramic gyrase complex was obtained by incubating equimolar amounts of GyrA and GyrB for 30 min on ice. The activity of the purified C. glutamicum (C.g.) gyrase as well as its inhibition by Cg1978 were determined using the M. tuberculosis Gyrase (HIS) Supercoiling Assay Kit (Inspiralis, Norwich, UK) according to the manufacturer’s manual. According to the assay conditions, the C.g. gyrase concentration of 50 nM resulting in full supercoiling of 0.5 µg relaxed plasmid DNA after 30 min of incubation at 37°C was determined as 1 U. Supercoiling inhibition of the C.g. gyrase was assayed by using 1 U of the C.g. gyrase with increasing concentrations of Cg1978-C-His (0.02–60.0 µM). Additionally, different concentrations of the known inhibitor ciprofloxacin (10 and 50 µM) were used as a positive control.

Moreover, the inhibitory effect of Cg1978 on the M.tb and E.c. gyrase was investigated according to the M. tuberculosis and E. coli Gyrase (HIS) Supercoiling Assay Kits (Inspiralis, Norwich, UK) using 1 U of the respective gyrase and the same Cg1978 concentrations as for the C.g. gyrase. All reactions were stopped by adding 30 µl STEB buffer (40% (w/v) sucrose, 100 mM Tris-HCl pH 8.0, 10 mM EDTA, 0.5 mg/ml Bromophenol Blue), and 30 µl chloroform/isooamy alcohol (v:v, 24:1).

4.11 | Surface Plasmon Resonance Spectroscopy (SPR)

For SPR analysis, Cg1978-C-His and GyrA-N-Strep were purified as described above. After purification, the buffer of both proteins was exchanged to PBS (137 mM NaCl, 2.7 mM KCl, 8 mM Na₂HPO₄, 1.5 mM KH₂PO₄, pH 7.4) using PD10 Desalting columns (GE Healthcare, Freiburg, Germany). The binding of His-tagged Cg1978-C-His to GyrA-N-Strep was analyzed by SPR analysis in a Biacore 3000 device (GE Healthcare, Freiburg, Germany) using a Sensor Chip CM5 (GE Healthcare, Freiburg, Germany). As the first step, an anti-histidine antibody (GE Healthcare, Freiburg, Germany) was immobilized to the chip matrix using amino coupling chemistry. All experiments were carried out in HBS-EP buffer (0.01 M HEPES, pH 7.4, 0.15 M NaCl, 3 mM EDTA, 0.005% w/v Surfactant P20) at 25°C. Following the standard coupling protocol for antibody immobilization, the mixture of 0.05 M N-Hydroxysuccinimide (NHS) and 0.2 M 1-Ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride (EDC) was injected for a total contact time of 420 s to activate the matrix. Then, the anti-histidine antibody (50 µg/ml) diluted in immobilization buffer (10 mM sodium acetate, pH 4.5) was injected for 420 s. To deactivate the unbound parts of the chip matrix, 1 M ethanolamine hydrochloride-NaOH (pH 8.5) was injected for 420 s. The flow rate was set to 10 µl/min during this immobilization procedure. Approximately 8,000–10,000 response units (RU) of the anti-histidine antibody were immobilized per flow cell. For the binding analysis, 180–250 RU of Cg1978-C-His was captured via injection of 40 µl (10 nM) at a flow rate of 5 µl/min followed by 10 min of HBS-EP buffer to remove unbound protein from the chip. The binding analysis between Cg1978-C-His and GyrA-N-Strep was then performed by injecting 90 µl of GyrA-N-Strep (10–250 nM) followed by a dissociation time of 300 s at a flow rate of 30 µl/min. After each cycle, the surface was regenerated by injection of regeneration buffer (10 mM Glycine-HCl, pH 1.5) for 30 s, at a flow rate of 30 µl/min. After the equilibration with three start up cycles without the analyte, this was repeated for various concentrations of GyrA-N-Strep (10–250 nM). Sensorgrams were recorded using Biacore 3000 Control Software 4.1.2 and analyzed with BIAevaluation software 4.1.1 (GE Healthcare, Freiburg, Germany). The surface of flow cell 1 immobilized with the anti-histidine antibody was used to obtain blank sensorgrams for the subtraction of the bulk refractive index background. The referenced sensorgrams were normalized to a baseline of 0. Peaks in the sensorgrams at the beginning and the end of the injection are due to the run-time difference between the flow cells for the chip.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.
AUTHOR CONTRIBUTIONS

LK, MH, and JF conceived the study; LK and JB performed the experiments; LK, MH, JB, RH, and JF analyzed the data; LK and JF wrote the manuscript. All authors reviewed and edited the manuscript.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available in the supplementary material of this article. Microarray data were deposited in the GEO database (ncbi.nlm.nih.gov/geo) with accession number GSE151224.

ORCID

Julia Frunzke https://orcid.org/0000-0001-6209-7950

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