The AraC-Type Transcriptional Regulator GliR (PA3027) Activates Genes of Glycerolipid Metabolism in Pseudomonas aeruginosa

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Abstract: Pseudomonas aeruginosa encodes a large set of transcriptional regulators (TRs) that modulate and manage cellular metabolism to survive in variable environmental conditions including that of the human body. The AraC family regulators are an abundant group of TRs in bacteria, mostly acting as gene expression activators, controlling diverse cellular functions (e.g., carbon metabolism, stress response, and virulence). The PA3027 protein from P. aeruginosa has been classified in silico as a putative AraC-type TR. Transcriptional profiling of P. aeruginosa PAO1161 overexpressing PA3027 revealed a spectacular increase in the mRNA levels of PA3026-PA3024 (divergent to PA3027), PA3464, and PA3342 genes encoding proteins potentially involved in glycerolipid metabolism. Concomitantly, chromatin immunoprecipitation-sequencing (ChIP-seq) analysis revealed that at least 22 regions are bound by PA3027 in the PAO1161 genome. These encompass promoter regions of PA3026, PA3464, and PA3342, showing the major increase in expression in response to PA3027 excess. In vitro DNA binding assay confirmed interactions of PA3027 with these regions. Furthermore, promoter-reporter assays in a heterologous host showed the PA3027-dependent activation of the promoter of the PA3026-PA3024 operon. Two motifs representing the preferred binding sites for PA3027, one localized upstream and one overlapping with the −35 promoter sequence, were identified in PA3026p and our data indicate that both motifs are required for full activation of this promoter by PA3027. Overall, the presented data show that PA3027 acts as a transcriptional regulator in P. aeruginosa, activating genes likely engaged in glycerolipid metabolism. The GliR name, from a glycerolipid metabolism regulator, is proposed for PA3027 of P. aeruginosa.

Keywords: Pseudomonas aeruginosa; AraC family; transcriptional regulator; PA3027; regulon; glycerolipid metabolism

1. Introduction

The abundance of transcriptional regulators in bacteria allows for the optimization of gene expression and response to stimuli. Nineteen prokaryotic transcriptional regulator families have been evaluated thus far [1]. The most abundant groups in bacterial genomes include the LysR, TetR/AcrR, AraC/XylS, and Lrp families. The ample repertoire of transcriptional regulators (TRs) is a characteristic feature of free-living microorganisms and opportunistic pathogens, in contrast to intracellular organisms.

Pseudomonas aeruginosa is a ubiquitous bacterium, also known for being a facultative human pathogen, especially dangerous for immunocompromised patients, mainly due to its genetic plasticity and intrinsic antibiotic resistance mechanisms [2]. The P. aeruginosa genome encodes complex regulatory systems, which include more than 500 known and potential transcriptional regulators or two-component system proteins [3], altogether constituting almost 10% of all its genes. The regulatory network of this bacterium allows it to
modulate and manage cellular metabolism to survive in variable environmental conditions including the human body, causing infections of lungs, wounds, blood, and urinary tracts [4]. Recent studies performed by Huang and co-workers [5] showed how 20 key virulence-related transcriptional regulators from different TR families work, crosstalk, and affect transcription of target genes in *P. aeruginosa*. It highlighted how complicated the regulatory network is and how many factors influence the regulation of gene expression at any given moment.

The AraC family regulators are an abundant group of TRs in bacteria, mostly acting as gene expression activators [6]. The representatives of this family possess a highly conserved C-terminal domain containing two helix-turn-helix (HTH) motifs responsible for DNA-binding, and a variable N-terminal domain, called the ligand-binding domain (LBD) or response domain, involved in interactions with cognate ligands as well as participating in protein oligomerization.

The well characterized archetype of this family is the AraC activator of the *araBAD* operon, which is involved in the metabolism of L-arabinose in *Escherichia coli* [7]. Its action strongly depends on arabinose binding [8]. There are three AraC binding sites in the *araBAD* promoter: I1 and I2, located next to RNA polymerase binding sites, and the distant upstream site named O2. In the absence of arabinose, AraC binds to I1 and O2 and forms a dimer, which causes DNA looping. This conformation prevents RNA polymerase from binding to the *araBAD* promoter, so the AraC acts as a repressor. When arabinose is present, AraC binds the ligand and changes conformation. As a result, the DNA loop breaks and AraC monomers bind to I1 and I2 sites and help to recruit RNA polymerase to the promoter to initiate transcription [9].

The mode of action of AraC-type regulators was shown to be controlled by different mechanisms (e.g., the oligomeric state and/or conformation can be changed after ligand binding but also upon interaction with a partner protein) [6]. In the absence of ligands, they can act as repressors, while after ligand binding and conformation changes, they can act as the activators [6]. The ligands of AraC-type regulators are usually small organic compounds like sugars (e.g., arabinose for *E. coli* AraC [7], lactose for *Clostridium perfringens* BgaR) [10], amino acids, and their derivatives (e.g., sarcosine for SouR (PA4184) from *P. aeruginosa*) [11], or other small molecules (e.g., urea for UreR in *Providencia stuartii*) [12]. The AraC-type regulators control diverse cellular functions including carbon metabolism (e.g., GapR activates the transcription of the gap gene encoding glyceraldehyde-3-phosphate dehydrogenase in *S. aureofaciens*) [13]; type III secretion systems (e.g., VirF in *Shigella* spp., HilC/HilD in *Salmonella* spp., SPI-1, or ExsA in *P. aeruginosa*) [14]; quorum sensing and virulence (e.g., QsvR in *Vibrio parahaemolyticus*) [16].

*P. aeruginosa* PAO1 encodes 16 known and 41 potential TRs from the AraC/XylS family (based on NCBI and pseudomonas.com databases), which makes this group one of the largest among TR families in *P. aeruginosa*. Most of the AraC family TRs thus far characterized in this bacterium are described as involved in the regulation of metabolism: OruR (PA0831) is an ornithine degradation activator [17], ArgR (PA0893) controls arginine biosynthesis and aerobic catabolism [18], AntR (PA2511) is an activator of anthranilate degradation [19], MmsR (PA3571) is a positive regulator of amino acid biosynthesis [20], SouR (PA4184) is essential for growth on sarcosine [11], PchR (PA4227) regulates pyochelin biosynthesis [21], GbdR (PA5380) controls choline metabolism [22], CdhR (PA5389) regulates carnitine metabolism [23], and PruR (PA0780) is a proline utilization regulator important for virulence [24]. Some of the TRs from the AraC/XylS family play different roles in *P. aeruginosa* virulence, for example, ChpD (PA0416) was required for spreading and colonizing the liver [25]; ExsA (PA1713) played a role in colonization of the corneal epithelium [26]; VqsM (PA2227), a positive regulator of quorum sensing (QS), was also shown to be involved in modulation of antibiotic resistance and biofilm formation [27,28]; CdpR (PA2588), a positive regulator of pyocyanin and biofilm production [29]; SphR (PA5324), an activator of sphA gene important for survival in the murine lung as well as...
for resistance to the antimicrobial effect of the pulmonary surfactant sphingosine [30], or CmrA (PA2047) involved in activation of mexEF–oprN and increased resistance of *P. aeruginosa* to the pump substrates such as chloramphenicol, fluoroquinolones, or trimethoprim [31].

The PA3027 protein from *P. aeruginosa* has been classified in silico as a putative AraC/XylS-type transcriptional regulator. The PA3027 gene was previously identified as downregulated in *P. aeruginosa parA* and *parB* mutants, which showed disturbed chromosome segregation [32–34]. The genes with altered expression in *par* mutants represented different functional categories, however, a significant number of genes encoding transcriptional regulators, often of unknown functions, was noted [32,35]. This study aimed to decipher the function of one of them, the AraC-type transcriptional regulator PA3027.

2. Results

2.1. Overview of the PA3027 from *P. aeruginosa* PAO1161

The PA3027 gene in *P. aeruginosa*, transcribed divergently to the PA3026–PA3022 cluster, encodes a protein classified in silico as a putative AraC-type TR with two predicted domains: the N-terminal ligand-binding domain (LBD) and the C-terminal DNA binding domain with a HTH motif (Figure 1A). The region of PA3027 with the highest similarity to well-described members of the AraC family: AraC, Rob, and MarA from *E. coli*, was identified in the C-terminal part of the protein encompassing the putative HTH containing region (Figure 1A).

![Figure 1](image-url)

**Figure 1.** Properties of PA3027 protein from *P. aeruginosa*. (A) Genomic context of the PA3027 gene in the *P. aeruginosa* genome and domain structure of the PA3027 protein. The gene names from PAO1 and PAO1161 strains are presented. Alignment represents comparison of PA3027 HTH domain with corresponding regions of *E. coli* AraC (GenBank: CAA23508.1), Rob (GenBank: CAD6017604.1), and MarA (GenBank: AAK21293.1). Sequences were aligned using Clustal Omega.
Identical residues in all proteins were marked with yellow, in three sequences with blue and in two with grey. The secondary structure elements are marked with boxes based on MarA secondary structure [37]. (B) Structure of PA3027 monomer bound with DNA predicted using COACH and HDOCK [38–40]. LBD—ligand binding domain; HTH—helix-turn-helix. (C) Bacterial two-hybrid (BACTH) analysis of PA3027 self-interactions. E. coli BTH101 cya− was transformed with the pairs of vectors allowing expression of the indicated fusion proteins. Interactions between proteins were assayed by analysis of the β-galactosidase activity in cell extracts and analysis of colony color upon growth on McConkey medium with 1% maltose. Data indicate mean β-galactosidase activity from at least three replicates ±SD. (D) Size exclusion chromatography (SEC) with multi-angle static light scattering (MALS) analysis for His−PA3027. Left axis—UV absorption and light scattering (LS), right axis—molecular weight of protein (MW). (E) Oligomerization state of purified His−PA3027 assayed by crosslinking with increasing concentration of glutaraldehyde. Samples were used in Western blot analysis with anti-His antibodies. For (D,E), one red dot indicates a monomer and two dots indicate a dimer.

The secondary structure prediction showed two potential HTH DNA binding motifs in the C-terminus of PA3027 (Figure 1A) [37]. The first HTH motif was predicted to be located at position 256–284, while the second one was predicted to be formed by residues 298 to 335. Similarly, PA3027 protein structure prediction using COACH and HDOCK [38–40] also suggested the existence of two possible HTH motifs involved in contact with DNA (Figure 1B). These data indicate the sequence similarity between PA3027 and other AraC-type TRs.

To analyze the oligomeric state of PA3027, various methods were applied. The bacterial two-hybrid analysis revealed that PA3027 can self-assemble in vivo, but only in the case when the C-terminal part of the protein is free in at least one of the tested fusions (variants T18–PA3027/T25–PA3027 and PA3027–T18/T25–PA3027) (Figure 1C). Concomitantly, size-exclusion chromatography combined with multi-angle light scattering (SEC-MALS) analysis with purified His−PA3027 demonstrated that this protein existed preferentially as a monomer, but it could also form dimers in solution under the tested conditions (Figure 1D). Similarly, glutaraldehyde crosslinking of purified His−PA3027 followed by Western blot analysis also showed the presence of protein dimers (Figure 1E). These data indicate that like other AraC-type regulators, PA3027 may exist as a monomer or dimer.

### 2.2. Effect of Increased PA3027 Level on Gene Expression

AraC-type TRs may act as gene expression activators but also as repressors. To identify genes regulated by PA3027 in P. aeruginosa, RNA-sequencing was performed under conditions of slight PA3027 overproduction, not affecting the growth of the cells (Appendix A Figure A1A). The rationale behind the analysis of cells with PA3027 excess rather than the ΔPA3027 mutant was based on (1) relatively low expression of PA3027 under standard growth conditions (LB or M9 medium, data not shown); (2) using protein excess could possibly mimic the induced, activated state of the protein; and (3) the effector for this regulator is unknown. Comparison of the transcriptomes of PAO1161 cells carrying pKKB1.11 (tacp−PA3027, hereafter called PA3027+) and PAO1161 cells with pAMB9.37 (tacp, empty vector control, hereafter called EV+), grown under selection in L broth with 0.05 mM IPTG, demonstrated 539 loci with altered expression (fold change (FC) ≤ −2 or ≥2, FDR adjusted p-value ≤ 0.01) (Figure 2A; Table S1). A total of 306 loci were downregulated, while 233 showed increased expression. The genes with altered mRNA levels were assigned to PseudoCAP functional categories [41] and grouped arbitrarily into six more general classes, as described previously [32,42]. The majority of identified genes belonged to classes II, IV, and V, and the highest enrichment was observed for the following categories: energy metabolism (18%, mostly downregulated genes), cell wall/LPS/capsule (18%, mostly upregulated genes), and transport of small molecules (16%) (Figure 2A). These data indicate that PA3027 excess in P. aeruginosa influenced the expression of genes from different functional categories.
Figure 2. Effect of increased PA3027 level on gene expression in *P. aeruginosa* PAO1161 cells. (A) Enrichment of PseudoCAP functional categories [41] for 539 genes (306 downregulated; 233 upregulated) showing changes in mRNA level in response to PA3027 abundance (fold change ≤ −2 or ≥ 2, FDR adjusted *p*-value ≤ 0.01). The numbers in brackets show the number of all genes in the PAO1 genome in the indicated PseudoCAP category. One gene could be classified into more than one category or class. Numbers in red or blue bars represent the number of up- or downregulated genes, respectively, in each category. The PseudoCAP categories were grouped into six more general classes. Genes annotated only in PAO1161 strain but not in PAO1 are described as non classified. (B) Volcano plot visualization of the results of differential expression analysis between transcriptomes of PA3027 overproducing cells and control cells. Each dot represents one gene and genes with the most significant changes are colored in red. For clarity genes with *p*-value < 0.1 are not shown. (C) Validation of RNA-seq results by RT-qPCR analysis. The same RNA used for RNA-seq analysis was used for cDNA synthesis and RT-qPCR analysis. Data represent mean fold change for three samples of PA3027 overproducing cells relative to the mean of the control samples ± SD.

The volcano plot visualization of the results of differential expression analysis highlighted genes with the most significant changes in mRNA level (Figure 2B). A spectacular increase in expression (>40) in response to PA3027 excess was observed for the PA3026–PA3023 gene cluster, transcribed divergently to the PA3027 (Figure 1A) as well as PA3464 gene (Figure 2B, Table 1). These genes encode proteins, with predicted functions in glycerolipid metabolism: an oxireductase acting on CH–OH group of donors (PA3026), a
glycerol-3-phosphate dehydrogenase (PA3025), a carbohydrate kinase (PA3024), a diacylglycerol/lipid kinase (PA3023), and a phospholipase C (PA3464). Concomitantly, a high decrease in expression in PA3027+ cells relative to EV+ cells was observed for the genes encoding cytochrome c (PA0523) as well as the regulatory protein NosR; both involved in the regulation of expression of the nitrous oxide reductase gene nosZ [43] (Table 1). Essentially, RT-qPCR (reverse transcription followed by quantitative PCR) analysis, using the material used for RNA-seq analysis validated the observed changes in mRNA level of chosen genes, confirming the influence of PA3027 on their expression (Figure 2C).

Table 1. Genes with altered expression in response to PA3027 excess (fold change < −10 or >10). PseudoCAP categories description as in Figure 1A.

| PA01 ID | PA01161 ID (D3C65) | Fold Change in RNA-seq | PseudoCAP Category | Gene Product |
|---------|--------------------|------------------------|---------------------|--------------|
| PA3024  | 10155              | 173.40                 | EM; PE              | carbohydrate kinase |
| PA3025  | 10150              | 163.31                 | CCC; EM             | glycerol-3-phosphate dehydrogenase/oxidase |
| PA3026  | 10145              | 135.03                 | HUU                 | FAD-binding oxidoreductase |
| PA3464  | 07800              | 129.57                 | HUU                 | phospholipase |
| PA3027  | 10140              | 107.00                 | TR                  | AraC family transcriptional regulator |
| PA3023  | 10160              | 47.32                  | HUU                 | lipid kinase YegS |
| PA2202  | 14525              | 15.56                  | MP; TSM             | amino acid ABC transporter permease |
| PA3342  | 08435              | 15.38                  | MP                  | DUF2804 domain-containing protein |
| PA2203  | 14520              | 14.14                  | MP; TSM             | amino acid ABC transporter permease |
| PA5024  | 27340              | 13.07                  | HUU                 | sulfite exporter TauE/SafE family protein |
| PA4191  | 04005              | 13.06                  | PE                  | isopenicillin N synthase family oxygenase |
| PA4193  | 03995              | 12.57                  | MP; TSM             | amino acid ABC transporter permease |
| PA4192  | 04000              | 10.44                  | TSM                 | amino acid ABC transporter ATP-binding protein |
| PA0282  | 01495              | 10.01                  | MP; TSM             | sulfate ABC transporter permease subunit CysT |
| PA3392  | 08180              | −11.34                 | EM                  | TAT-dependent nitrous-oxide reductase |
| PA0519  | 02740              | −11.50                 | EM                  | nitrite reductase |
| PA0524  | 02765              | −23.76                 | EM                  | nitric-oxide reductase large subunit |
| PA3391  | 08185              | −44.43                 | EM; MP              | regulatory protein NosR |
| PA0523  | 02760              | −62.05                 | EM                  | cytochrome c |

2.3. Identification of PA3027 Binding Sites in P. aeruginosa

To identify the PA3027 binding sites in the PAO1161 genome and hence the direct targets of the regulator, a chromatin immunoprecipitation-sequencing (ChIP-seq) analysis was performed using PAO1161 ΔPA3027 pKKB1.12 (tacp–flag–PA3027) strain expressing flag–PA3027 under the control of tacp (hereafter called F–3027+). The addition of the FLAG tag to PA3027 did not alter the ability of the protein to retard bacterial growth in medium with a high (0.5 mM) concentration of IPTG (Figure A1), suggesting that the fusion protein is functional. The PAO1161 ΔPA3027 (pABB28.1 tacp–flag) strain was used as an empty vector control (F–EV+). ChIP-seq was performed using cells grown under selection in LB broth with 0.05 mM IPTG to OD600 of 0.5 and anti-FLAG antibodies. Sets of ChIP-seq peaks called separately for each of the three F–3027+ ChIP replicates using the FDR adjusted p-value cut-off of 0.05 and fold enrichment cut-off 2 (Table S2A–C) were compared and peaks in genome regions also showing enrichment in the F–EV+ ChIP samples were discarded (Table S2D; Figure 3A). Overall, this analysis pointed out 22 high confidence PA3027 binding sites in the P. aeruginosa PAO1161 genome (Figure 3B). Two FLAG-PA3027 bound regions, 4/5 and 13/14, showed ChIP-seq peaks with two clearly separated summits, hence these were considered as separate binding sites in subsequent analyses.
Interestingly, these sites mapped to the promoter region as well as the terminator region of the flanked gene (Figure 3B). (Table 2).

### Table 2. *P. aeruginosa* loci with PA3027 binding sites identified in the ChIP-seq analysis.

| Region ID | Binding Site Position | Gene in PAO1161 (D3C65_ | Gene in PAO1 | Fold Change (RNA-seq) | Fold Enrichment (ChIP-seq) | Position of Summit | PseudoCAP Category | Gene Product |
|-----------|------------------------|---------------------------|-------------|------------------------|---------------------------|-------------------|-------------------|--------------|
| 1 term    | 01245 PA0234           | −1.71                     | 5.73        | 264,723                | HUU nucleoside-binding protein |
| 2 term    | 04760 PA4045           | 1.54                      | 3.76        | 987,296                | HUU cobalamin-binding protein |
| 3 intra   | 07240 PA3572           | −2.18                     | 7.06        | 1,510,957              | HUU hypothetical protein |
| 4 prom    | 07800 PA3464           | 129.57                    | HUU         | phosholipase           |
| 5 term    | 07800 PA3464           | 129.57                    | 11.84       | 1,641,558              | HUU phosholipase |
| 6 prom    | 08155 PA3397           | 1.38                      | 3.72        | 1,712,366              | BCPGC; EM ferredoxin-NADP reductase |
| 7 intra   | 08185 PA3391           | −44.43                    | 4.30        | 1,720,198              | EM; MP regulatory protein |
| 8 prom    | 08435 PA3342           | 15.38                     | 5.68        | 1,762,086              | MP DUF2804 domain-containing protein |
| 9 intra   | 08440 PA3341           | −1.16                     | 2.42        | 1,763,471              | TR MarR family transcriptional regulator |
| 10 intra  | 08605 PA3390           | −3.36                     | 9.83        | 1,805,406              | HUU universal stress protein |
| 11 intra  | 09845 PA3083           | 1.51                      | 8.26        | 2,055,622              | TPTMD aminopeptidase |
| 12 intra  | 11,685 PA2742          | −1.20                     | 3.10        | 2,411,617              | TPTMD 50S ribosomal protein L33 |
| 13 prom   | 12,415 PA2601          | 1.04                      | 3.90        | 2,570,649              | TR LysR family transcriptional regulator |
| 14 term   | 12,415 PA2601          | 1.04                      | TR          | 3-mercaptopropionate dioxygenase |
| 15 intra  | 14,830 PA2147          | −1.73                     | 5.28        | 3,153,105              | AP catalase HP II |
| 16 intra  | 18,720 PA1414          | −2.19                     | 9.03        | 3,976,577              | HUU hypothetical protein |
| 17 prom   | 19,855 PA1196          | −4.91                     | 3.16        | 4,218,075              | TR sigma-54-dependent Fis family transcriptional regulator |
| 18 intra  | 21,295 PA0928          | −1.04                     | 4.07        | 4,499,850              | TCRS hybrid sensor histidine kinase/response regulator |
| 19 intra  | 25,030 PA4610          | −3.03                     | 2.56        | 5,276,490              | HUU copper transporter |
| 20 term   | 26,025 PA4772          | 1.83                      | 3.09        | 5,480,812              | EM FAD-binding oxidoreductase |
| 21 intra  | 28,310 PA5208          | −3.54                     | 3.28        | 5,982,827              | HUU TIGR00153 family protein |
| 22 intra  | 28,490 PA5243          | −1.56                     | 3.49        | 6,024,153              | BCPGC porphobilinogen synthase |
| 23 prom   | 28,760 PA5294          | −1.60                     | 3.02        | 6,078,786              | HUU multidrug resistance protein NorM |
| 24 prom   | 28,755 PA5293          | −1.17                     | 3.02        | 6,078,786              | TR probable transcriptional regulator |

PA3027 ChIP-seq peaks identified in promoters (prom), terminators (term), or gene body (intra). Underlined regions (4 and 14) are second peaks in regions encompassing PA3027 binding sites with two clearly separated signals. PseudoCAP category descriptions as in Figure 1A.
Figure 3. PA3027 binding sites in *P. aeruginosa* genome. (A) Venn diagram for ChIP-seq peaks obtained for samples of FLAG-PA3027 overproducing cells (F–PA3027+) and negative control (F–EV+). (B) ChIP-seq signal over regions encompassing PA3027 binding sites. The plots show coverage with reads for indicated positions in the PAO1 genome (kb), normalized per genome coverage (RPGC), and averaged for ChIP replicates. Genes are presented as grey arrows, only names of PAO1 orthologs are shown for clarity. (C, D) The consensus sequence logos of predicted PA3027 binding sites, obtained by MEME software [44,45] using 200 bp around 24 PA3027 peak summits (A) as well as the same 24 PA3027 peak summit regions with an extended to 500 bp region encompassing *PA3026* upstream sequences (B). The height of an individual letter represents the relative frequency of that nucleotide at that position. The consensus sequence (up line) and the most common nucleotide at each position (down line) are presented for each motif. The reverse complement presentation of sequence logos are shown below.

The identified PA3027 binding sites were compared with RNA-seq data to define genes possibly directly regulated by PA3027 binding in their vicinity (Table 2). Six PA3027 binding sites mapped to the promoter regions, five in potential terminator regions and 12 were identified in the gene bodies (Table 2). The PA3027 binding site with the highest fold enrichment encompasses the downstream region/terminator as well as the promoter region of the *PA3464* gene, which also showed major upregulation in response to PA3027.
excess (Figure 3B, region 4/5). PA3027 binding sites were also detected in regions preceding PA3026 and PA3342 genes, similarly, demonstrating increased expression in RNA-seq analysis (Figure 3B, region 24 and 8/9). Concomitantly, PA3027 binding regions were also identified within gene bodies, and some of these seemed to exert an effect on neighboring gene expression, as shown in the RNA-seq analysis for PA3391 (nosR), PA3572, PA3309, PA1414, PA1196, PA4610, PA5208, all exhibiting a decreased expression in response to PA3027 excess (Table 2; Figure 3B).

To identify recurrent DNA sequences in ChIP-seq peaks and define the PA3027 consensus-binding site, the MEME tool [45] was applied on 24 sequences corresponding to the PA3027 ChIP-seq peak summits ±100 bp sequences as well as the same 24 PA3027 peak summit regions with an extended to 500 bp region 24 encompassing PA3026 upstream sequences. Two sequences called here motif A (15 bp) and motif B (11 bp) were identified with proposed consensus sequences YYGGCGHTDTYSGMC and GGAYAWCGCCG, respectively (Figure 3C,D). Interestingly, in the reverse complement orientation, a part of motif B resembles a part of motif A. The localization of identified motifs within promoter regions of activated genes (PA3026, PA3364, and PA3342) showed their presence upstream, or even overlapping the predicted −35 promoter region, a position preferred for binding by transcriptional activators (e.g., AraC, MarA) [9,46] (Figure 4A–C). The localization of identified motifs in 24 regions detected as bound by PA3027 is presented in Appendix A Table A1. Additionally, in the promoter region of the PA3026 gene, next to motifs A and B, partial palindromes CCGGCCGTCGCTGGCGC and GGCCGGCGCCGCCGGCC as well as inverted repeat TCGGCCCTTTA-N29-TCCAGGCGCGA had been noticed (Figure 4A). Partially, they resemble the identified PA3027 binding motifs and potentially might be involved in DNA recognition and binding by PA3027.
Figure 4. PA3027 interaction with DNA assayed in vivo and in vitro. (A–C) Putative PA3027 binding motifs in PA3026 (A), PA3464 (B), and PA3342 (C) promoter fragments, used in β-galactosidase activity assays and electrophoretic mobility shift assay (EMSA) analysis. Blue—motif A, violet—motif B, green—pseudopalindrome, grey—"–10" and "–35" promoter regions. (D) The scheme of variants of the PA3026–PA3024 promoter used in the analysis and cloned to pCM132 upstream of a promoter-less lacZ reporter gene. (E) Influence of PA3027 on the activity of PA3026p, PA3464p, and PA3342p. β-galactosidase activity in extracts from E. coli DH5α Δlac cells bearing pCM132 derivatives containing the indicated promoters upstream of lacZ as well as pKKB1.11 (tap-PA3027), allowing IPTG inducible PA3027 production or empty vector pAMB9.37 (EV). Strains were cultured in medium with or without 0.1 mM IPTG. Data indicate mean ±SD from five cultures. * p-value < 0.05 in student’s two tailed t-test. (F) EMSA using His6–PA3027 and PCR amplified DNA of the indicated promoter regions. The 100 ng Cy5 tagged DNA was incubated with an increasing amount of His6–PA3027. Samples were separated using 10% acrylamide gel and Cy5 fluorescence was visualized. The 331 bp pCM132 fragment was used as a control to rule out non-specific DNA binding.

2.4. Regulatory Properties of PA3027

To evaluate the regulatory properties of PA3027, the PA3026 promoter region was selected to which PA3027 showed strong binding, and a spectacular increase in PA3026 expression was observed in response to PA3027 excess. Three variants of the PA3026 promoter region were cloned in the probing vector pCM132 [47], carrying a promoter-less lacZ (Figure 4D). All tested variants were able to act as promoters for the lacZ reporter gene, as manifested by higher β-galactosidase activity in cells carrying PA3026p-lacZ fusions in comparison to the promoter-less lacZ control (Figure 4E). No change of lacZ expression driven from PA3026 variants was observed in the absence or the presence of IPTG, when no PA3027 was delivered (EV), or in the cultures without PA3027 induction (–IPTG). Essentially, an addition of IPTG promoting PA3027 expression led to a major increase in the PA3026 promoter activity (Figure 4E). The lowest induction, approximately 2-fold, was observed for the shortest PA3026pA fragment. For fragment PA3026pB, the highest β-galactosidase activity and the highest induction (almost 12-fold) was observed, in comparison to promoter activity without PA3027 induction. A slightly lower, (10-fold) increase was observed for the PA3026pC-lacZ fusion. Importantly, the identified PA3027 binding motifs (Figure 3C,D), were present in all tested fragments of PA3026p, however, in different numbers. Only one PA3027 binding motif A, encompassing the ~35 region of PA3026p, was present in the shortest tested fragment A. The pseudopalindrome located 38 bp upstream of −35 box of PA3026p was additionally present in fragment B, and an additional distal putative PA3027 binding motif, resembling motif B, located 162 upstream to the −35 promoter region was present in PA3026pC (Figure 4A).

Essentially, all the DNA fragments used in the regulatory experiments were bound by His6–PA3027 in the electrophoretic mobility shift assay (EMSA) (Figure 4F), whereas no DNA shift of the control fragment (part of pCM132) was observed. The binding to the shortest version of the PA3026p promoter (fragment A) was somewhat weaker than for other tested PA3026 variants (Figure 4F). These data demonstrate the ability of PA3027 to activate PA3026p in a heterologous host and highlights the requirement of sequences upstream of the ~35 promoter region for full activation of this promoter by PA3027.

Similar analysis with cells expressing PA3464p-lacZ or PA3342p-lacZ only showed a minor increase in expression from the tested promoters in the presence of PA3027 (Figure 4E), whereas DNA fragments with these sequences were clearly bound in the EMSA experiments. Thus, it is not excluded that additional factors or upstream/downstream sequences are required for the control of the expression of PA3464 and PA3342 by PA3027 in P. aeruginosa.

Overall presented data confirm PA3027 interactions with promoter regions of PA3026, PA3464, and PA3342 and highlight its role as a DNA-binding protein involved in transcription control.

2.5. Toward the Biological Function of PA3027–PA3026–PA3023 Gene Cluster

In silico analyses and database mining suggested that the products of the PA3026–PA3024 operon are potentially involved in the transformation of glycerol to alkylglycerone phosphate (Figure 5). PA3024 is probably involved in the conversion of glycerol to...
sn-glyceryl-3-phosphate. In *P. aeruginosa*, this reaction could also be catalyzed by GlpK, PA1487, and PA3579. The second protein, PA3025, is a homolog of GlpD, and possibly catalyzes the conversion of sn-glyceryl-3-phosphate to glycerone phosphate. The enzymes catalyzing the next reaction, converting glycerone phosphate to acylglycerone phosphate are unknown in *P. aeruginosa*, but the potential candidates could be genes encoding GNPAT–glyceronephosphate O-acyltransferases such as PA3673 (PlsB), PA0581, or PA4636. The last protein encoded in the PA3026–PA3024 operon could possibly facilitate conversion of acylglycerone phosphate to alkyglycerone phosphate. This analysis suggests that there might be a functional redundancy of enzymes encoded by PA3026–PA3024 and other proteins in *P. aeruginosa*.

Figure 5. Schematic representation of the part of central carbon metabolism in *P. aeruginosa*. The pathways were drawn based on the Pseudomonas database [48] and the literature [49–51]. Red and blue indicate increased or decreased gene expression in response to PA3027 overproduction, respectively.

In an attempt to assign a biological function to PA3027, the chromosomal mutants of PA01161 in PA3027 and PA3026–PA3024 genes (the whole operon deleted) were obtained using the allele exchange method. The ability to generate the ΔPA3027 and ΔPA3026–PA3024 strains indicates that the genes are not essential for the growth of *P. aeruginosa*. Lack of PA3027 or PA3026–PA3024 did not affect the growth of the cells on rich L broth medium, or minimal medium M9 supplemented with citrate and/or glycerol as the carbon source (Figure A2A–D), or in osmotic stress medium, a minimal medium with 0.5 M NaCl or 0.7 M sucrose (data not shown). Moreover, a BIOLOG microarray phenotype screening with plate no. 1, 2, 3, 4, 9, 10, 15, 16, and 18 did not show any effect of the gene deletion(s) on growth (data not shown) [52]. Additionally, no changes in bacterial motility or biofilm formation were observed between ΔPA3027 mutant and the WT strain (Figure A2E–G). These data indicate that under the conditions tested, the analyzed genes do not play a major role in cell fitness or that there is a backup pathway compensating the lack of these genes.

A search of genes encoding orthologs of proteins encoded by the PA3027–PA3023 gene cluster in 1748 representative and reference bacterial genomes included in the RefSeq database (release 91) using MultiGeneBlast revealed only four genomes encoding clustered orthologs of all analyzed proteins (Figure A3A; Table S3). Among these, two genomes possessed the gene encoding a PA3023 orthologue separated from the rest by an additional gene: a protein with a pirin domain for *Pseudomonas mendocina* and a potential
oxidoreductase for Pseudomonas resinovorans. Another eight strains demonstrated a similar arrangement of the PA3027 and PA3026–PA3024 orthologue encoding genes. The strains belong mostly to the genus Pseudomonas as well as other gamma-proteobacteria (e.g., Oblitimonas alkaliphila or Shewanella sedimentis) and one representative of the beta-proteobacteria (Rhodotherax ferrireducens). In R. ferrireducens, the PA3027 encoding orthologue is separated from the operon by genes encoding a glycerate 2-kinase, an unknown protein and a lipolytic enzyme. Comparison of the orthologs of the PA3027 transcriptional regulator from identified organisms showed the highest similarity among proteins encoded in Pseudomonas sp. genomes (Figure A3B). The most similar gene to PA3027 is the TR from P. citronellolis P3B5, which was isolated from ready-to-eat basil (Ocimum basilicum) phyllosphere material [53]. These data indicate that the presence of genes encoding PA3027–PA3024-like proteins is not unique to PAO1/PAO1161 or Pseudomonas and occurs in other bacteria, however, only in a very limited number of so far sequenced strains. Interestingly, among the identified strains outside the genus Pseudomonas, no homologs of glpD or glpK were identified, thus PA3025 and PA3024 may encode the only glycerol-3-phosphate dehydrogenase and carbohydrate kinase, respectively. These data suggest the involvement of PA3027 and PA3026–PA3024 proteins in glycerolipid metabolism, however, further studies are needed to assign a precise biological role to these proteins.

3. Discussion

The aim of this study was a multilayered analysis of PA3027 from P. aeruginosa, a previously uncharacterized AraC-type transcriptional regulator [54]. The sequence analysis showed that PA3027 possesses a tandem HTH motif in the C-terminal part of the protein (Figure 1A), indicating the existence of two surfaces involved in contact with DNA, which is typical for AraC-type regulators [6]. Additionally, SEC-MALS and glutaraldehyde crosslinking experiments showed that despite PA3027 being preferentially a monomer under the conditions tested, the protein may also form dimers (Figure 1D,E). Bacterial two-hybrid analysis revealed the lack of self-interactions for the PA3027-T18/PA3027-T25 pair, suggesting that the free C-terminus (containing the tandem helix-turn-helix) also contributes to PA3027 dimerization (oligomerization). This suggests the possibility of the head to head or tail to tail, but also head to tail PA3027 self-interaction, similar to the one previously observed, for example, for the AraC regulator ExsA from P. aeruginosa, bound to the promoters of regulated genes (exsC, exsD, exoT, pcrG) [26].

AraC-type regulators can work as monomers (e.g., Rns in E. coli) [55], however, some need to form dimers (e.g., ExsA in P. aeruginosa [56] or oligomers, like UreR in E. coli) [57]. The AraC from E. coli acts as a dimer, however, dimerization and the mode of DNA binding are largely affected by the ligand [58]. The state of oligomerization of PA3027 and the mode of action could also be regulated by the presence of the ligand, however, additional studies are required for its identification. Remarkably, PA3027 readily activated the PA3026p in heterologous host E. coli, which suggests that either a specific ligand is not required for activation or that the hypothetical ligand is present in E. coli cells, grown in L broth medium.

Comparison of transcriptomes of cells with a slight PA3027 overproduction and identification of PA3027 binding sites allowed defining the PA3027 regulon (Table 2). The ChIP-seq analysis pointed out 24 PA3027 binding sites, but summits of only six of these mapped to regions directly preceding genes. Similarly, among the 28 binding sites of P. aeruginosa CdpR, only three were located in intergenic regions including the divergently transcribed promoter of P. aeruginosa quinolone signal (PQS) [29]. Remarkably, some of the PA3027 ChIP-seq peaks had complex shapes, and sometimes clear sub-summits, suggesting PA3027 binding to multiple motifs in the analyzed regions. The most significant observation of this work was the direct binding of PA3027 to the promoters of PA3026–PA3023, PA3464, and PA3342 genes, also showing the spectacular increase of mRNA level in response to PA3027 excess, which indicates the role of PA3027 in the activation of these
genes via direct interactions with their promoter regions. Our data show that this activation can be readily observed for PA3026p.

Our analysis identified two putative PA3027 binding motifs, motif A (15 bp, consensus YYGGCGHTDTYSGMC) and motif B (11 bp, consensus GGAYAWCGC) (Figure 3C,D). Similar to other AraC-type regulators (e.g., AraC from *E. coli* or CuxR from *Sinorhizobium meliloti*), PA3027 binds to more than one motif [59,60]. The identified PA3027 binding sites showed some similarity to the motifs recognized and bound by other AraC representatives (e.g., VqsM (GGATSNNTYGCCA) or CdpR (RGWNNWNCGGCCA) from *P. aeruginosa*) [5]. The localization of PA3027 binding motifs within promoter regions of activated genes, upstream or overlapping the –35 promoter region position (Figure 4A–C), was similar to those observed previously for binding sites of transcriptional activators (e.g., AraC, MarA) [9,37]. A careful analysis of PA3026 also indicated the presence of partial palindromes CCGCGTCGCGTCGCG and GGCCGGCGCGCGCGGCGG, as well as inverted repeat TCGGCTTGA-N29-TCCAGGCCGA (Figure 4A), not directly resembling the PA3027 binding consensus for motif A or B, however, we might not rule out the involvement of these sequences in PA3027 action. The identified motifs are characterized by repeated stretches of GC pairs. The regulatory experiments and EMSA assays using truncated regions of the PA3026 promoter suggested that to act efficiently in transcription activation, PA3027 needs multiple sites, one in the vicinity of the -35 promoter sequence and at least one more located upstream, at a distance. The shortest tested fragment of PA3026, with one predicted PA3027 binding motif, exhibited the lowest increase in expression in response to PA3027, however, in EMSA analysis, this fragment was still bound by PA3027 (Figure 4F). Other tested variants of the PA3026 with more than one putative PA3027 binding motif were bound by PA3027 in EMSA and activated in response to PA3027 in the regulatory assays (Figure 4E). The presence of distal sites provides the possibility of modulating the action of the regulator in the presence or absence of a ligand, similar to the AraC regulator [9]; however, further studies are required to analyze the role of specific motifs in PA3026 activation by PA3027.

A spectacular increase of expression under PA3027 overproduction conditions was also observed for the PA3464 gene encoding PlcA, a phospholipase C acting on phosphatidylcholine (PC), phosphatidylethanolamine (PS), and phosphatidylserine (PE) [61]. Divergently transcribed PA3465 gene encoding conserved hypothetical protein classified as the membrane MSF (major facilitator superfamily domain) transporter was downregulated. Interestingly, according to ChIP-seq data, there were two PA3027 binding regions upstream and downstream of the PA3464 gene (Figure A2). In PA3464p, motif A is located 23 bp upstream to the -35 sequence of PA3464p, while motif B overlaps it. The PA3027 binding sites in the promoter and terminator regions were also observed for the PA3342 gene encoding a protein with an uncharacterized DUF2804 domain (Figure 3B). Only motif A located 133 bp to the start codon was identified in PA3342p. The cloned fragments preceding PA3464 or PA3342 were however not sufficient to promote PA3027 dependent gene activation in *E. coli*. Potentially like Rns, CfaR, VirF, AggR, and CsvR [62,63], PA3027 needs to have a second binding motif downstream of the promoter binding site to activate expression. In this case, the second motif would be in the gene terminator. These analyses point out that other genetic/proteic components are needed for the PA3027 dependent activation of these genes.

The PA3026–PA3024 operon potentially encodes proteins involved in glycerolipid metabolism. PA3024 encodes a putative FGGY carbohydrate kinase carrying out ATP-dependent phosphorylation of glycerol (Figure 5). Three other enzymes with this activity are encoded in the PAO1 genome: GlpK, PA1487, and PA3579, but only one, GlpK, has been characterized so far as a part of the GlpR regulon together with GlpDFK [64]. The GlpD homolog, PA3025, potentially catalyzes the next step in the pathway, oxidizing glycerol-3-phosphate to dihydroxyacetone phosphate (DHAP). *E. coli* encodes two glycerol 3-phosphate dehydrogenases, the *glpD* and *glpABC* genes. GlpD is required for aerobic growth with glycerol or glycerol 3-phosphate, while GlpABC is required for anaerobic growth with glycerol (or glycerol 3-phosphate) and fumarate [65,66]. PA3026 encodes a
potential oxidoreductase acting on the CH–OH group of donors. The next gene, PA3023, transcribed in the same orientation as the PA3024, encodes a probable diacylglycerol kinase, similar to the lipid kinase YegS from Salmonella typhimurium [67].

The ability to generate the ∆PA3027 and ∆PA3026–PA3024 strains indicates that the genes are not crucial for the growth of P. aeruginosa. The ability of the strains to grow in minimal media with different carbon sources (e.g., citrate, glycerol) was tested as several bacterial AraC transcription factors are known to control genes that are responsible for the degradation of complex carbon sources [68]. We cannot rule out that these genes play a role in the metabolism of glycerol and the fact that the ∆PA3026–PA3024 operon mutant could grow on glycerol might be explained by the functional redundancy of the glpD homologs in P. aeruginosa [69]. Another gene that is potentially regulated directly by PA3027 is PA3464, which encodes a phospholipase C, cleaving phospholipids just before the phosphate group [70]. PA3464 is active on PC, PS, and PE [61], however, further studies are needed to elucidate the role of this enzyme in P. aeruginosa biology.

Engagement of the products of the regulated genes in glycerol metabolism could be a first step in the production of glycerolipids [66], biosurfactants [71], biofilm formation [72], or as an energy source [73]. The link between PA3027 target genes (PA3026–PA3024, PA3464) is phospholipid metabolism (Figure 5). Phospholipids are membrane components that could contain sn-glycerol-3-phosphate esterified with fatty acids [74]. The PE, phosphatidylglycerol, and cardiolipin are the major phospholipids in bacteria. All membrane phospholipids include phosphatidic acid (PA), which is derived from glycerol-3-P via action of glycerol-3-P acyltransferases [75]. The enzymes that catalyze the turnover of phospholipids include both phospholipases and lipid phosphatases. It is important to elucidate the complex regulatory mechanisms that control the connected and coordinated pathways involved in the synthesis of glycerophospholipids as important components of each cell. The work presented here is the first characterization of the PA3027 transcriptional regulator involved in the regulation of glycerolipid metabolism in P. aeruginosa, thus we propose the name GliR (glycerolipid metabolism regulator) for PA3027.

4. Materials and Methods

4.1. Growth Conditions Bacterial Strains and Plasmids

Bacterial strains, plasmids, and primers used in this study are listed in Table A2 and A3. E. coli and P. aeruginosa strains were grown at 37 °C in Luria-Bertani (LB) broth or on LB plates containing 1.5% (w/v) agar [76] as well as in M9 minimal medium [77] supplemented with 0.25% citrate ions or 1% glycerol as a carbon source and leucine (10 mM) for PAO1161 leu− strain. For the selection of plasmids, LB medium was supplemented with appropriate antibiotics: kanamycin (50 μg mL−1 for E. coli, 500 μg mL−1 in solid media, and 250 μg mL−1 in liquid media for P. aeruginosa); benzylpenicillin sodium salt (300 μg mL−1 in solid media and 150 μg mL−1 for E. coli); carbenicillin (300 μg mL−1 for P. aeruginosa); rifampicin (300 μg mL−1 for P. aeruginosa); and chloramphenicol (10 μg mL−1 for E. coli, 150 μg mL−1 for P. aeruginosa).

Bacterial cells were routinely grown in flasks closed with a cotton plug, with shaking 200 rpm at 37 °C. Growth kinetics was monitored by measurements of optical density at 600 nm (OD600) in 96-well plates at 37 °C using a Varioskan Lux Multimode Microplate Reader and SkanIt RE 5.0 software (Thermo Fisher Scientific, Waltham, MA, USA). Motility assays were performed as described previously [33,78]. Plates were standardized by using the same volume of medium. Biofilm analysis was performed according to the previously described method [77] using cultures in LB or minimal medium with citrate grown at 37 °C for 24 h or 48 h.

Competent cells of E. coli were prepared by the CaCl2 method [77] and P. aeruginosa according to the method using MgCl2 [79]. E. coli strain DH5α was used for plasmid manipulations and S17-1 was used to mate pAKE600 [80] derivatives into P. aeruginosa. Standard DNA manipulations were performed as described [77].
4.2. Vectors and Strains Construction

The ΔPA3027 or ΔPA3026–PA3024 deletion mutants were constructed with the use of pAKE600 suicide vector derivatives [80]: pKKB1.61 and pKKB1.62 (Table A2). Upstream and downstream DNA fragments of mutated regions were amplified using primer pairs #1/#2 and #3/#4 for PA3027 or #5/#6 and #7/#8 for PA3026–PA3024 (Table A3). The PCR fragments were digested with BamHI, HindIII, and HindIII, EcoRI, respectively, and ligated with EcoRI, BamHI digested pAKE600. The allele exchange procedure was performed as described previously [33]. The E. coli S17 strain carrying suicide plasmid was conjugated with the recipient strain P. aeruginosa PAO1161 (RifR). Putative cointegrants were selected on LB agar with rifampicin and carbenicillin. Colonies were used to inoculate LB with 10% sucrose and checked for the Cbs phenotype to select clones without the vector. Allele exchange was screened by PCR (Table A3).

The PA3027 gene was cloned in pET28a to obtain the His6–tagged version of the protein at the N-terminus. The gene was amplified using PCR with primers #9/#10 and PAO1161 genomic DNA as a template. The purified fragment was then digested with EcoRI, SacI and ligated with EcoRI, SacI digested pET28a to obtain pKKB1.21.

To obtain an expression vector able to propagate in P. aeruginosa (pKKB1.11 and pKKB1.12), the PA3027 gene was excised from pKKB1.21 using EcoRI, SalI and inserted into pAMB9.37 or pABB28.1, a derivative of pBBR1MCS-1 [81,82], containing tacp and lacIQ or lacIQ, tacp, and flag, respectively.

To test the promoter activity of selected DNA fragments, the RK2 derivative pCM132 plasmid with a promoter-less lacZ reporter gene was used [47]. PCR products corresponding to PA3026pA (120 bp), amplified using primers #14/#13), PA3026pB (176 bp amplified using primers #12/#13), PA3026pC (330 bp, amplified using primers #15/#13), PA3027pX (167 bp, amplified using primers #18/#19), were digested with EcoRI, BamHI, and ligated with EcoRI, BamHI digested pCM132 to obtain pKKB1.305, pKKB1.303, pKKB1.304, and pKKB1.309, respectively (Table A2). pKKB1.307 (PA3464p-lacZ) was constructed by amplification of the 228 bp fragment using primers #16/#17, followed by digestion with BglII and ligation with BglII digested pCM132. The scheme of cloned and analyzed variants of the PA3026 promoter region is shown in Figure 4.

For bacterial two-hybrid analysis, DNA fragment encoding PA3027 was cloned into derivatives of pKT25 (pLKB2), pKNT25, pUT18, and pUT18C (pLKB4) [83]. To obtain pKKB1.51 and pKKB1.81 (pKNT25, pUT18 derivatives, respectively with PA3027 lacking STOP codon), the PCR fragments amplified using #9/#11 primers were digested with EcoRI, SacI and ligated with vectors digested with the same enzymes (Table A2). To construct pKKB1.52, PA3027 was excised from pKKB1.11 using EcoRI/SalI and ligated with pLKB2 digested with EcoRI, Ecl136II. The PA3027 gene was transferred from pKKB1.21 to pLKB4 as an EcoRI, SacI fragment to obtain pKKB1.82 (Table A2).

4.3. Protein Overproduction and Purification

Overproduction of His6-PA3027 was carried out in the E. coli BL21 strain carrying pKKB1.21. Overnight culture was diluted 1:50 in 1000 mL autoinduction LB (Formedium, Norfolk, UK) supplemented with 1% glycerol, 0.5% NaCl, and kanamycin and grown for 48 h at 18 °C. Cells were harvested by centrifugation and sonicated in Buffer S (20 mM MES pH 6.5, 200 mM (NH4)2SO4, 200 mM NaCl) with 1 mM phenylmethysulfonyl fluoride (PMSF) and 1 mg/mL lysozyme. The His-tagged PA3027 was purified on Ni-agarose columns (Ni-TED 1000 Protino, Marchel&Nagel) using Buffer S with 250 mM imidazole for elution. The purification procedure was monitored by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) using a Pharmacia PHAST gel system. Elution fractions were dialyzed using Buffer S with 1% β-mercaptoethanol and stored at −80 °C.

4.4. SEC-MALS Analysis
Size exclusion chromatography coupled to multi-angle light scattering (SEC-MALS) analysis was performed using a high-performance liquid chromatography (HPLC) instrument (1260 Infinity LC, Agilent Technologies Inc., Santa Clara, CA, USA) equipped with a UV detector, a MALS detector (DAWN HELEOS II, Wyatt Technology Santa Barbara, CA, USA), and a differential refractometer (Optilab T-rEX, Wyatt Technology, Santa Barbara, CA, USA). A total of 100 μL of 1 mg/mL samples, obtained as described in the above section, were loaded onto a Superdex 200 Increase 10/300 column (GE Healthcare, Milwaukee, WI, USA) equilibrated with Buffer S. Absorption at UV wavelengths of 280, 254, and 215 nm were monitored during SEC. Samples were run at room temperature at a flow rate of 0.5 mL/min. The results were analyzed using ASTRA v. 6 software (Wyatt Technology, Santa Barbara, CA, USA) in accordance with the manufacturer’s instructions.

4.5. Glutaraldehyde Crosslinking

The oligomerization state of purified His6–PA3027 was assayed by crosslinking as described previously [84].

4.6. RNA Isolation, RNA-Seq, and RT-qPCR Analysis

Strains were obtained by transformation of PAO1161 cells with pKKB1.11 (tacp–PA3027) or pAMB9.37 (tacp) plasmids (Table A2). Transformants were selected on L-agar plates supplemented with 150 μg mL⁻¹ chloramphenicol and were verified by isolation of plasmid DNA and its digestion. After overnight growth, each culture of P. aeruginosa PAO1161 carrying pKKB1.11 or pAMB9.37 vector was diluted 1:100 into fresh L-broth supplemented with 75 μg mL⁻¹ chloramphenicol and 0.05 mM IPTG as inducer. Cells were collected from 2 mL of cultures in the logarithmic phase of growth (optical density at 600 nm of 0.4–0.6) were mixed with 4 mL of RNAprotect Bacteria Reagent (Qiagen, Hilden, Germany). RNA was isolated using the Qiagen RNeasy Mini Kit, according to the manufacturer’s instructions. RNA was treated with DNA-free DNA Removal Kit (Invitrogen, Thermo Fisher Scientific, Waltham, MA, USA) and a lack of DNA contamination was confirmed by PCR. RNA concentration was determined using μDrop plate of Varioskan Lux Multimode Microplate Reader and quality was checked using Bioanalyzer.

Library preparation and sequencing were performed at the Laboratory of DNA Sequencing and Oligonucleotide Synthesis, Institute of Biochemistry and Biophysics, PAS (Warsaw, Poland). Ribosomal RNA was depleted using the RiboZero Bacteria Kit (Illumina, San Diego, CA, USA). Obtained mRNA was used for cDNA library construction using the KAPA Stranded RNASeq Kit. The library was further quality checked on 1% agarose gel and concentration was measured using the qPCR KAPA Library Quantification Kit (Roche Holding AG, Basel, Switzerland). Libraries were sequenced using standard Illumina protocols. Reads were quality-checked and filtered using FASTP version 0.20.0 [85]. Reads were mapped to the P. aeruginosa PAO1161 genome (CP032126.1) using Bowtie2 version 2.3.4.3 [86] with default settings. The number of reads mapping to individual genes was counted using FeatureCounts v 2.0.1 (part of the Subread) with the -s2 option [87]. Differential expression analysis was conducted using edgeR ver 3.28.0 [88]. Raw data are available in the NCBI’s Gene Expression Omnibus (GEO) database under accession number GSE163555.

For selected genes, the results were confirmed by RT-qPCR using RNA isolated from the same cultures. Reverse transcription on 4 μg RNA was performed using a TranScriba Kit (A&A Biotechnology, Gdansk, Poland). qPCR was performed on a LightCycler 480 II System (Roche Molecular Diagnostics, Mannheim, Germany) using 5x HOT FIREPol EvaGreen qPCR Mix Plus (Solis Biodyne, Tartu, Estonia). Each 18 μL reaction contained 3.6 μL 5x reaction mix, 1 μL of five times diluted cDNA and 1.5 μL of mixed 5 μM primers (Table A3). The relative expression was determined by comparison of crossing points (Cp) between the target and the reference gene (nadB). Three technical repetitions were used for each primer pair. The ratio was calculated using the Pfaffl formula [89].
4.7. Chromatin Immunoprecipitation with Sequencing

ChIP-seq analysis was performed using the PAO1161 ΔPA3027 strain with pKKB1.12 (strain overproducing FLAG–PA3027) and PAO1161 ΔPA3027 with pABB28.1 (lacIQ–tacp–flag). Cells were grown at 37 °C in LB with 50 μg mL⁻¹ chloramphenicol and 0.05 mM IPTG. Cells were collected at OD₆₀₀ of 0.5. ChIP protocol was based on a modified protocol using Dynabeads Protein A [90]. Lysate after sonication was thawed on ice and 150 μL of each strain variant was incubated with 20 μL of magnetic beads coupled with protein A (Dynabeads Protein A, 10001D, Invitrogen, Thermo Fisher Scientific, Waltham, MA, USA), separated from the original suspension using a magnetic separation stand. A pre-clearing step was performed for 1 h at 4 °C with rotation. A total of 50 μL of magnetic beads, separated from the suspension as above, was then mixed with 6 μL of anti-FLAG mouse polyclonal antibodies (DYKDDDDK Tag polyclonal antibodies 1 mg/mL, PA1-985B, Invitrogen, Thermofisher Scientific, Waltham, MA, USA) diluted in 200 μL of PBS with 0.05% Tween-20. Mixtures of magnetic beads and antibodies were incubated for 10 min at 4 °C with gentle rotation. Beads with bound antibodies were then separated from the supernatant, washed once with 200 μL of the PBS with 0.05% Tween-20 solution, and stored on ice. Pre-cleared lysate was separated from the beads used for pre-clearing and added to the beads coated with antibodies. A mixture containing lysate and magnetic beads with antibodies was incubated at 4 °C for 20 min with mixing on a rotator. Beads were then collected and washed as described earlier [35]. Elution was performed twice for 15 min in 50 μL at 65 °C in a thermostock (Thermomixer compact, Eppendorf, Hamburg, Germany) with 1400 rpm (7× g) mixing. Elutions from six parallel reactions were pooled. The obtained eluates were incubated with 8 μL of RNase A (100 mg/mL, 19101, Qiagen, Hilden, Germany) for 30 min at 65 °C. Then, 40 μL of Proteinase K (20 mg/mL, 19133, Qiagen, Hilden, Germany) was added and the samples were incubated for 1 h at 50 °C, followed by overnight incubation at 65 °C. Next, 40 μL of Proteinase K was added and the samples were again incubated for 1 h at 50 °C. Subsequently 24 μL of 3 M sodium acetate (pH = 5) was added and the volume was adjusted to 700 μL using water. DNA purification was performed using a Qiaquick Qiagen PCR purification Kit, according to the manufacturer’s instructions. The DNA was stored at −20 °C. Purified DNA from ChIP performed with the empty vector strain was included as a background control.

Sequencing of ChIP samples was performed in the Laboratory of DNA Sequencing and Oligonucleotide Synthesis of Institute of Biochemistry and Biophysics Polish Academy of Sciences in Warsaw, Poland. The NGS library was constructed using a QiaSeq Ultralow Input Library Kit (Qiagen, Hilden, Germany). Samples were quality checked on 1% agarose gel and concentration was measured using a qPCR KAPA Library Quantification Kit (Roche Holding AG, Basel, Switzerland). Libraries were sequenced using standard Illumina protocols.

Reads were quality-checked and filtered using FASTP version 0.20.0 [85]. Reads were mapped to the *P. aeruginosa* PAO11161 genome (CP032126.1) using Bowtie2 version 2.3.4.3 [86] using default settings. Obtained *.sam* files were sorted (samtools sort -n), run through samtools fixmate with the -m option, again sorted (samtools sort), and duplicates were marked with samtools markdup. Samtools ver. 1.9 was used [91]. The files were indexed and used to generate coverage *.bigwig* files, normalized to 1× sequencing depth (RPNC), without binning and smoothing using the bamCoverage tool version 3.3.0 included in deepTools [92].

ChIP-seq peaks were called separately for each ChIP sample using MACS2 version 2.1.2 [93] with default options for paired-end BAM files, and 0.05 as the false discovery rate (FDR). Subsequently, peak matching was performed using the findOverlapsOfPeaks function from ChIPpeakAnno [94]. Peaks overlapping and regions enriched in the background control were excluded. Visualization of the coverage data was performed using Integrated Genomics Viewer version 2.4.17 [95]. Peaks were annotated using a custom R script. Raw data are available in the NCBI’s Gene Expression Omnibus (GEO) database under accession number GSE163554.
4.8. Regulatory Experiments

The β-galactosidase activity was tested in extracts from exponentially growing *E. coli* DH5αΔlac cells bearing pCM132 derivatives (pKKB1.305 PA3026pA, pKKB1.303 PA3026pB, pKKB1.304 PA3026pC, pKKB1.309 PA3342p, pKKB1.307 PA3464p), and pKKB1.11 (*tacp–PA3027*) or pAMB9.37. To assay β-galactosidase activity, bacteria were grown at 37 °C in L-broth containing kanamycin and chloramphenicol with or without 0.1 mM IPTG. The β-galactosidase activity was calculated using the J. Keith Joung modified Miller Units equation [96].

4.9. Electrophoretic Mobility Shift Assay (EMSA)

The electrophoretic mobility shift assay (EMSA) was performed using purified His6-PA3027 and the amplified, purified PA3026 promoter variants (Figure 4). To prepare DNA fragments, PCR was performed using pCM132 derivatives as a template and appropriate pair of primers as follows: pKKB1.305 for the PA3026pA fragment (#14/#20), pKKB1.303 for the PA3026pB fragment (#12/#20), pKKB1.304 for the PA3026pC fragment (#15/#20), pKKB1.309 for the PA3342p fragment (#18/#20), pKKB1.307 for the PA3464p fragment (#16/#20), and pCM132 for the control fragment (#21/#20). The reverse primer (#20), which binds to the plasmid sequence was coupled with Cy5 dye, enabling visualization using a FluorChemQ MultimageII Chemilumager. The images were captured using AlphaView software (Alpha Innotech, version 3.5.0). The binding reaction was performed in the presence of unspecific DNA (600 ng, salmon sperm DNA) and the complexes were analyzed on a 10% acrylamide gel.

4.10. Bacterial Two-Hybrid Analysis (BATCH)

Appropriate pairs of plasmids were co-transformed into the *E. coli* BTH101 cya strain [83]. The presence of interactions between the tested proteins was assayed by the analysis of the appearance of red colonies of transformants on the MacConkey medium and β-galactosidase activity in the extracts of transformants [96].

4.11. Bioinformatic Analysis

A comparison of the C-terminal domain of the chosen AraC-type regulators was performed using Clustal W. Structure of the PA3027 monomer bound with DNA was predicted using COACH and HDOCK [38-40]. Clustered orthologs of PA3024-PA3027 were identified in 1748 reference genomes from the Refseq database (Release 91) using MultiGeneBlast [97]. The DNA binding motifs (Table A1) were identified using MEME-ChIP version 5.3.0 [44,45] using sequences corresponding to 200 bp around 24 PA3027 peak summits (Supplementary Text S1).

5. Conclusions

The AraC-type regulators are involved in the control of various cellular functions, helping to adjust metabolism to efficiently use available resources, maintain homeostasis, and propagate. Here, we performed a thorough analysis of a representative of the AraC family PA3027 transcriptional regulator from *P. aeruginosa*. The regulon of PA3027 was identified, highlighting the role of PA3027, named GliR in the regulation of genes involved in glycerolipid metabolism. The significance of our research is in identifying the pathways regulated by GliR, which in turn allows for a better understanding of the complicated regulatory network of the human pathogen *P. aeruginosa*.

Supplementary Materials: The following are available online at www.mdpi.com/1422-0067/22/10/5066/s1, Text S1: Sequences of PA3027 ChIP-seq peak summits used for motif searching (peaks marked by numbers and gene ID identified in the vicinity, see Table 2). Table S1: Results of RNA-seq analysis. List of genes with altered expression identified by comparison of transcriptomes of *P. aeruginosa* PAO1161 cells overproducing PA3027 with transcriptomes of cells carrying the empty vector (fold change (FC) ≤ −2 or ≥ 2, FDR adjusted p-value ≤ 0.05). Genes annotated only in
PAO1161 strain but not in PAO1 are described as “not annotated” (NA). Table S2: Results of ChIP-seq analysis. PA3027-FLAG ChIP-seq peaks identified in each repeat—set A, B, C or in the negative control—D (Table S2A–D). Table S3: Distribution and evolutionary conservation of PA3027-PA3023 cluster in bacteria identified using MultiGeneBlast [97].

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Appendix A
**Figure A1.** Impact of PA3027 excess on the growth of *P. aeruginosa* PAO1161. PAO1161 or ΔPA3027 mutant strains carrying empty vector pAMB9.37 tacp or pKKB1.11-tacp-PA3027 (A,B) or empty vector tacp-flag or pKKB1.12 tacp-flag-PA3027 (C,D) were grown in L-broth under selection with the indicated concentration of inducer IPTG (0.05–0.5 mM). The red line indicates the growth in the presence of 0.05 mM IPTG, conditions selected for RNA-seq analysis. Data represent mean OD$_{600}$ from three independent replicates. Standard deviations are not shown for clarity.

**Figure 2.** Phenotypic characterization of *P. aeruginosa* PAO1161 cells lacking PA3027 or PA3026–PA3024 operon. (A–D) Growth of PAO1161 leu+ and ΔPA3027 mutant on (A) L broth, (B) minimal medium with citrate, (C) minimal medium with glycerol, or (D) minimal medium with citrate and glycerol. Data represent OD$_{600}$ mean from three independent replicates. (E) Growth of PAO1161 and ΔPA3027 mutant on swimming and swarming medium. (F,G) Biofilm formation of PAO1161 leu and ΔPA3027 mutant on (F) L broth or (G) minimal medium with citrate. Data represent mean from five independent replicates.
Figure 3. The occurrence of PA3023–PA3027 gene cluster in bacteria. (A) Clustered genes encoding orthologs of PA3023–PA3027 identified in 1748 representative and reference bacterial genomes included in the RefSeq database (release 91) [98] using MultigeneBlast [97]. (B) Phylogenetic tree of PA3027 and its identified orthologues, constructed using CoBaltDB [99].
Table A1. PA3027 DNA binding motifs in ChIP-seq peaks. The DNA binding motifs were identified using MEME-ChIP version 5.3.0 [44,45] and DNA corresponding to 200 bp around 24 summits of ChIP-seq peaks (Supplementary Text S1). Motif A was identified with the settings, allowing one occurrence per sequence, whereas motif B was searched using a setting allowing zero or one occurrence per sequence.

| Peak Number | Strand | Start | \( p \)-Value | Site |
|-------------|--------|-------|---------------|------|
| **Motif A** |        |       |               |      |
| 20          | +      | 71    | \( 9.60 \times 10^{-8} \) | CTGGCGCTATTCGCC |
| 2           | +      | 72    | \( 5.89 \times 10^{-7} \) | GTGCGGTATTGCC |
| 1           | +      | 10    | \( 2.01 \times 10^{-6} \) | CCGCGTATCGCC |
| 9           | +      | 47    | \( 5.80 \times 10^{-6} \) | TGCGGATTTGCC |
| 7           | +      | 37    | \( 5.80 \times 10^{-6} \) | CTGGCGAGTTGCC |
| 5           | +      | 2     | \( 7.42 \times 10^{-6} \) | CCGCGTATCGCC |
| 15          | +      | 34    | \( 1.32 \times 10^{-5} \) | ATCGCGGATCGCC |
| 11          | +      | 40    | \( 1.32 \times 10^{-5} \) | CCGCGAAATCGCC |
| 12          | -      | 28    | \( 1.48 \times 10^{-5} \) | TGCGGATTTGCC |
| 21          | +      | 60    | \( 2.78 \times 10^{-5} \) | TGCGGACTTTGCC |
| 16          | +      | 22    | \( 2.78 \times 10^{-5} \) | CCGCGTATGCGCT |
| 4           | +      | 36    | \( 4.53 \times 10^{-5} \) | ATCGCGATATCGCC |
| 17          | -      | 23    | \( 7.14 \times 10^{-5} \) | GCGCGATTTGCC |
| 10          | +      | 4     | \( 7.14 \times 10^{-5} \) | TGCGAATTTGCC |
| 23          | -      | 62    | \( 7.79 \times 10^{-5} \) | TGCGGATTTGCC |
| 18          | +      | 14    | \( 7.79 \times 10^{-5} \) | TGCGGCGTTGCC |
| 24          | -      | 17    | \( 7.79 \times 10^{-5} \) | TGCGGATTTGCC |
| 8           | +      | 39    | \( 1.62 \times 10^{-4} \) | TCGCGGATTTGCC |
| 13          | -      | 44    | \( 2.03 \times 10^{-4} \) | ATGGGTATTTGCC |
| 19          | +      | 54    | \( 2.18 \times 10^{-4} \) | TGCGGCGTTGCC |
| 3           | -      | 83    | \( 2.35 \times 10^{-4} \) | TGCGGCGTTGCC |
| 22          | -      | 16    | \( 3.31 \times 10^{-4} \) | TGCGGCGTTGCC |
| 6           | +      | 9     | \( 4.03 \times 10^{-4} \) | ATCGGATTTCC |
| 14          | -      | 70    | \( 4.29 \times 10^{-4} \) | TGCGGCGTTGCC |
| **Motif B** |        |       |               |      |
| 2           | +      | 61    | \( 7.20 \times 10^{-7} \) | GATAACGGCC |
| 1           | -      | 11    | \( 7.20 \times 10^{-7} \) | GATAACGGCC |
| 21          | -      | 40    | \( 1.90 \times 10^{-6} \) | GACATCGCC |
| 20          | +      | 60    | \( 3.50 \times 10^{-6} \) | GACATCGCC |
| 5           | -      | 3     | \( 4.83 \times 10^{-6} \) | GATAACGGCC |
| 24          | -      | 43    | \( 6.79 \times 10^{-6} \) | GACATCGCC |
| 16          | -      | 33    | \( 1.22 \times 10^{-5} \) | GAAACCGAC |
| 18          | +      | 49    | \( 1.48 \times 10^{-5} \) | GATAACGGCC |
| 17          | +      | 77    | \( 1.84 \times 10^{-5} \) | GAAACCGAC |
| 13          | +      | 83    | \( 2.31 \times 10^{-5} \) | GACATCGCC |
| 7           | +      | 24    | \( 3.02 \times 10^{-5} \) | GACATCGCC |
| 19          | +      | 19    | \( 3.88 \times 10^{-5} \) | GACATCGCC |
| 22          | +      | 51    | \( 7.02 \times 10^{-5} \) | GATAACGGCC |
| 4           | +      | 69    | \( 8.23 \times 10^{-5} \) | GACATCGCC |
| 3           | +      | 60    | \( 9.79 \times 10^{-5} \) | GAAAGTGAGC |
**Table A2.** Bacterial strains and plasmids used and constructed in this study.

| Strains                  | Genotype/Plasmid Details                                                                                       | Source     |
|--------------------------|---------------------------------------------------------------------------------------------------------------|------------|
| *E. coli DH5α*           | F<sup>−</sup> (φ80d lacZΔM15) recA1 endA1 gyrA96 thi-1 hsdR17 (r~m~c~) supE44 relA1 deoR Δ(lacZYA–argF)U196 | [100]      |
| *E. coli BL21*           | F~ompT~ hsdS (r~m~c~) gal dcm (λ DE3)                                                                            | [100]      |
| *E. coli S17-1*          | pro hsdR hsdM recA Tp<sup>+</sup> Sm<sup>+</sup>ΩRP4-Tc::Mu Km::Tn7                                              | [100]      |
| *E. coli BTH101*         | F~cya~ araD139 galE15 galK16 rpsL (Str<sup>+</sup>) hsdR2 mcrA1 mcrB1                                         | [83]       |
| *E. coli DH5α Δlac*      | F<sup>−</sup> (φ80d lacZΔM15) recA1 endA1 gyrA96 thi-1 hsdR17 (r~m~c~) supE44 relA1 deoR Δ(lacZYA–argF)U196 | lab collection |
| *P. aeruginosa PAO1161*  | leu r<sup>−</sup> m<sup>+</sup> Rif<sup>R</sup>                                                            | [101]      |
| *P. aeruginosa PAO1161 ΔPA3027* | deletion of 1015 bp fragment encompassing *PA3027* gene                                              | this work  |
| *P. aeruginosa PAO1161 ΔPA3026–PA3024* | deletion of 4711 bp fragment encompassing *PA3026-PA3024* operon                                  | this work  |
| **Plasmids**             |                                                                                                             |            |
| pAKE600                  | Ap<sup>R</sup>; ori<sub>MB1</sub>, ori<sub>T</sub>, sacB                                                  | [80]       |
| pKKB1.61                 | Ap<sup>R</sup>; pAKE600 derivative with 431 bp fragment encompassing up- and down- sequence of *PA3027*, cloned using BamHI-HindIII-EcoRI and 18/2# and 3#/4# primers | this work  |
| pKKB1.62                 | Ap<sup>R</sup>; pAKE600 derivative with 472bp fragment encompassing up- and down- sequence of *PA3026–PA3024*, cloned using BamHI, HindIII/HindIII, EcoRI EcoRI and 4#/5# and 6#/7# primers | this work  |
| pET28a(+)                | Km<sup>R</sup>; ori<sub>bamy221</sub>, ori<sub>vi</sub>; expression vector                               | Novagen    |
| pKKB1.21                 | pET28a(+) derivative with *his<sup>E</sup>-PA3027*, PA3027 cloned 1078 bp as EcoRI, SacI                  | this work  |
| pBRR1-MCS1               | Cm<sup>R</sup>; IncA/C broad-host-range cloning vector, lacZA–MCS, mob, T7p, T3p                           | [81]       |
| pAMB9.37                 | Cm<sup>R</sup>, *ori<sub>C</sub>N<sub>ACS</sub>*, pBRR1-MCS-1 derivative with laci<sup>E</sup>–lacz<sup>E</sup>, expression vector | [82]       |
| pABB28.1                 | Cm<sup>R</sup>, *ori<sub>C</sub>N<sub>ACS</sub>*, pBRR1-MCS-1 derivative with laci<sup>E</sup>–lacz<sup>E</sup>–flag expression vector | this work  |
| pKKB1.11                 | pAMB9.37 (laci<sup>E</sup>–lacz<sup>E</sup>) derivative with 1097 bp fragment, encoding PA3027 gene, cloned using EcoRI, Sall | this work  |
| pKKB1.12                 | pABB28.1 (laci<sup>E</sup>–lacz<sup>E</sup>–flag) derivative with 1097 bp flag–PA3027 fragment, cloned using EcoRI, Sall | this work  |
| pLKB2                    | Km<sup>R</sup>, *ori<sub>flu</sub>*, pKT25 modified with lacz<sup>E</sup>–cyaT25<sup>E</sup>–MCS          | [83,102]   |
| pKNT25                   | Km<sup>R</sup>; *ori<sub>flu</sub>*, lacz<sup>E</sup>–MCS–cyaT25<sup>E</sup>          | [83]       |
| pUT18                    | Ap<sup>R</sup>; *ori<sub>C</sub>, lacz<sup>E</sup>–MCS–cyaT18<sup>E</sup> | [83]       |
| pLKB4                    | Ap<sup>R</sup>, *ori<sub>C</sub>, pUT18C modified with lacz<sup>E</sup>–cyaT18<sup>E</sup>–MCS | [83,102]   |
| pKKB1.51                 | pKNT25 derivative with PA3027–cyaT25<sup>E</sup>; 1042 bp PA3027 fragment cloned using EcoRI, SacI enzymes | this work  |
| pKKB1.52                 | pLKB2 derivative with cyaT25–PA3027; 1092 bp PA3027 fragment cloned using EcoRI, Ecl136I/Smal enzymes | this work  |
| pKKB1.81                 | pUT18 derivative with PA3027–cyaT18<sup>E</sup>; 1042 bp PA3027 fragment cloned using EcoRI, SacI enzymes | this work  |
| pKKB1.82                 | pLKB4 derivative with cyaT18–PA3027; 1092 bp PA3027 fragment cloned using EcoRI, SacI enzymes | this work  |
| pCM132                   | Km<sup>R</sup>; *ori<sub>V</sub>–*ori<sub>T</sub>–*ori<sub>T</sub>; promoter-less lacZ reporter gene | [47]       |
| pKKB1.305                | pCM132 derivative with PA3026pa, 120 bp fragment amplified with primers 14#/13# cloned using EcoRI, BamHI/BglIII | this work  |
Table A3. List of primers used in this study.

| Nr  | Starter            | Used for:                  | Sequence                                                                 |
|-----|--------------------|-----------------------------|--------------------------------------------------------------------------|
| #1  | 3027mLF            | PA3027 gene deletion        | gcggatcCAATTCGACCACGGTGCTTTC                                              |
| #2  | 3027mLR            | gcaagcttGGTCTGCATGGTCGTTTC  |
| #3  | 3027mPF            | gcaagctttagTAATGAGAACCGGCACATCCG                                      |
| #4  | 3027mPR            | gcgaattcCGGTGCTCTATCCGAACCAGAGTTCC                                     |
| #5  | 3026-4mLF          | gcaagcttcCATGGCCGAGGTATTTCCAG                                       |
| #6  | 3026-4mLR          | gcaagtctAGACGACCATGAAACCTTCC                                          |
| #7  | 3026-4mPF          | gcgaattcTCTGGCTCTCCAGGTTAC                                              |
| #8  | 3026-4mPR          | gcgaatATGACACCTTGGCTCCAC                                               |
| #9  | 3027eF             | PA3027 expression          | gcgaattcATGCAGACCCTTGGCTCCAGG                                            |
| #10 | 3027eR             | gcgaGCTCTGGCGCCGACACTTCAG                                              |
| #11 | 3027eR2            | gcaagctttgtaAATGAGAACCGGCACATCCG                                      |
| #12 | 3027pdF            | cagaacctgtgatgAGGCTGGTGTCAATCTTCAG                                    |
| #13 | p3026R2            | cggCTGGCAGGACAGGAGTGGCC                                               |
| #14 | p3026F2            | gcaagcttcTCTGGCTCTGGCCAGG                                              |
| #15 | p3026F3            | gcaagcttcTCTGGCTCTGGCCAGG                                              |
| #16 | p3464R             | PA3464 promoter            | gcagatctCGGCTTGGTGCTCTGGTC                                               |
| #17 | p3464F             | gcagatctCAGATGGGCTGCTTGGATC                                            |
| #18 | p3342F             | PA3342 promoter            | cagaaccagctgatgAGGCTGGTGTCAATCTTCAG                                      |
| #19 | p3342R             | gcaagcttcTCTGGCTCTGGCCAGG                                              |
| #20 | CM132RCy5          | EMSA analysis              | Cy5—CTTCCACAGTATTCCACCC                                                  |
| #21 | CM132pF            | GCGACAGGCTTCTCCTGAGTAG                                                  |
| #22 | 3027qF             | CTGGATCAGGGCGATCAGGAG                                                 |
| #23 | 3027qR             | CCGGACAGGCGAAGAGGAGTGC                                                |
| #24 | 3026qF             | AATCAGCTATCTCCCGGCACTCC                                               |
| #25 | 3026qR             | ATGATCCACAGGGCTTCCACCC                                                |
| #26 | 3464qF             | CAACCTGTTTCGAGACCAACC                                                |
| #27 | 3464qR             | TACAGAGACGAACTGCGCTT                                                  |
| #28 | 3023qF             | CTACCTGTGTTGACGGGTTGAG                                                |
| #29 | 3023qR             | CCTTCCACACTGAGAATCCG                                                  |
| #30 | 3342qF             | GAGAACCGTTATGCGTCAACCC                                                |
| #31 | 3342qR             | TACAGAGACGAACTGCGCTT                                                  |
| #32 | 2203qF             | TCTTCTGGTACTTCCGGGTT                                                  |
| #33 | 2203qR             | CATATCCAGGCGAGGAGGC                                                   |
| #34 | 2204qF             | CCCCTGCAAAACTTCTCCT                                                   |
| #35 | 2204qR             | GATGAGCCGCTTGGCAATGTG                                                 |
| #36 | 1911qF             | AGATCTCTTCTGAGAAGCGCC                                                 |
| #37 | 1911qR             | CATCTCTGGCGGTACATTTG                                                  |
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