



**EMORY**  
LIBRARIES &  
INFORMATION  
TECHNOLOGY

**OpenEmory**

# Overproduction of the MtrCDE Efflux Pump in *Neisseria gonorrhoeae* Produces Unexpected Changes in Cellular Transcription Patterns

Elizabeth A. Ohneck, *Emory University*

Maira Goytia, *Emory University*

Corinne Loughlin, *Emory University*

Sandeep J. Joseph, *Emory University*

[Timothy Read](#), *Emory University*

Ann E. Jerse, *Uniformed Services University of the Health Sciences*

[William Shafer](#), *Emory University*

---

**Journal Title:** Antimicrobial Agents and Chemotherapy

**Volume:** Volume 59, Number 1

**Publisher:** American Society for Microbiology | 2015-01-01, Pages 724-726

**Type of Work:** Article | Final Publisher PDF

**Publisher DOI:** 10.1128/AAC.04148-14

**Permanent URL:** <https://pid.emory.edu/ark:/25593/pqsw3>

---

Final published version: <http://dx.doi.org/10.1128/AAC.04148-14>

## Copyright information:

© 2015, American Society for Microbiology

This is an Open Access work distributed under the terms of the Creative Commons Attribution-Noncommercial 3.0 Unported License

(<http://creativecommons.org/licenses/by-nc/3.0/>).



Accessed December 3, 2021 4:22 AM EST

# Overproduction of the MtrCDE Efflux Pump in *Neisseria gonorrhoeae* Produces Unexpected Changes in Cellular Transcription Patterns

Elizabeth A. Ohneck,<sup>a</sup> Maira Goytia,<sup>a</sup> Corinne E. Rouquette-Loughlin,<sup>a</sup> Sandeep J. Joseph,<sup>b</sup> Timothy D. Read,<sup>b</sup> Ann E. Jerse,<sup>c</sup> William M. Shafer<sup>a,d</sup>

Departments of Microbiology and Immunology<sup>a</sup> and Medicine,<sup>b</sup> Emory University School of Medicine, Atlanta, Georgia, USA; Department of Microbiology and Immunology, F. Edward Herbert School of Medicine, Uniformed Services University of the Health Sciences, Bethesda, Maryland, USA<sup>c</sup>; Laboratories of Bacterial Pathogenesis, VA Medical Center, Decatur, Georgia, USA<sup>d</sup>

**The global consequence of drug efflux gene overexpression in bacteria has not been specifically analyzed because strains showing high-level expression typically have mutations in genes encoding regulatory proteins that control other genes. Results from a transcriptional profiling study performed with a strain of *Neisseria gonorrhoeae* that is capable of high-level transcription of the *mtrCDE* efflux pump operon independently of control by cognate regulatory proteins revealed that its overexpression has ramifications for systems other than drug efflux.**

Bacteria use efflux pumps to resist the action of antibiotics during treatment of infection or to survive in the presence of antimicrobials in their environment (1). The level of expression of efflux pump-encoding genes is modulated by a complex regulatory system of repressors and activators, but derepression due to mutation in repressor-encoding genes or their promoters can significantly decrease bacterial susceptibility to antimicrobials (2). As an example, mutation in the *mtrR* repressor gene of *Neisseria gonorrhoeae* enhanced transcription of the *mtrCDE*-encoded efflux pump and resulted in both increased resistance of gonococci to structurally diverse antimicrobials (3–5) and greater *in vivo* fitness in an experimental mouse model of lower genital tract infection (6, 7). MtrR also has more global action, however, as it directly or indirectly impacts expression of >65 genes involved in diverse systems such as peptidoglycan synthesis, pilin secretion, the general stress response, and glutamine metabolism (8–10).

The question asked in this study was this: does overexpression of a drug efflux pump *per se* have a potential impact on bacteria? Heretofore, this question could not be addressed because bacteria that overexpress efflux pump genes typically contain mutations that impact cognate gene regulators (e.g., MtrR), which would make interpretation of results difficult. Taking advantage of a unique promoter that drives high-level expression of *mtrCDE* independently of MtrR control (11) (as well as activation by MtrA [12]), we performed a transcriptional profiling study. Here, we provide data that support the hypothesis that overexpression of a bacterial efflux system may have unexpected global ramifications with respect to microbial physiology.

**Bacterial strains, culture conditions, and antimicrobial testing.** *N. gonorrhoeae* strains FA19 and FA19*mtr*<sub>120</sub> were the main strains used (Table 1). “*mtr*<sub>120</sub>” signifies a single nucleotide change (C to T) 120 nucleotides upstream of *mtrCDE* (7) that generates a new promoter for *mtrCDE* transcription (11) and increases gonococcal resistance to diverse antimicrobials (Table 1), including host defense antimicrobials (7, 13). Transformants of these strains bearing an insertional inactivated *ccpR* gene due to the presence of the nonpolar *aphA-3* cassette were constructed by previously described methods (12). Gonococci were routinely grown as non-piliated, opacity-negative variants on gonococcal medium base (GCB) agar or in broth, each with defined supplements I and II, as

previously described (4). The MIC of antimicrobials recognized by the MtrCDE efflux pump was determined by agar dilution (4). To test gonococcal susceptibility to peroxides, a microtiter plate assay was used. Briefly, this involved exposing 10<sup>5</sup> gonococci in GCB broth to various concentrations of hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>) or tert-butylhydroperoxide (tBuOOH) at 37°C for 45 min before spotting 5 μl onto GCB agar plates for assessment of viability; H<sub>2</sub>O<sub>2</sub> and tBuOOH are not substrates of the MtrCDE pump (unpublished observations).

**RNA-seq and qRT-PCR studies.** A transcriptional-profiling comparison study using transcriptome sequencing (RNA-seq) and three independent RNA samples prepared from broth-grown, late-logarithmic cultures of isogenic strains FA19 and FA19*mtr*<sub>120</sub> was performed using previously described methods (14). We defined differentially expressed genes as having (i) a fold change value of ≥2, (ii) a total read number larger than 5, and (iii) a Bonferroni-corrected *P* value of ≤0.05. For quantitative reverse transcriptase PCR (qRT-PCR) analysis of gene expression, RNA samples from RNA-seq experiments were used to synthesize cDNA using random hexamers. Transcripts of the genes of interest (*ccpR* and *rpS15*) were quantified using qRT-PCR and oligonucleotide primers (see Table S1 in the supplemental material) essentially as described previously (14); *rpS15* was used as a reference for normalization of the results.

**Global gene expression consequences of overexpression of *mtrCDE*.** As expected from previous studies (7, 11), RNA-seq

Received 24 August 2014 Returned for modification 13 October 2014

Accepted 29 October 2014

Accepted manuscript posted online 3 November 2014

Citation Ohneck EA, Goytia M, Rouquette-Loughlin CE, Joseph SJ, Read TD, Jerse AE, Shafer WM. 2015. Overproduction of the MtrCDE efflux pump in *Neisseria gonorrhoeae* produces unexpected changes in cellular transcription patterns. *Antimicrob Agents Chemother* 59:724–726. doi:10.1128/AAC.04148-14.

Address correspondence to William M. Shafer, wshafer@emory.edu.

Supplemental material for this article may be found at <http://dx.doi.org/10.1128/AAC.04148-14>.

Copyright © 2015, American Society for Microbiology. All Rights Reserved. doi:10.1128/AAC.04148-14

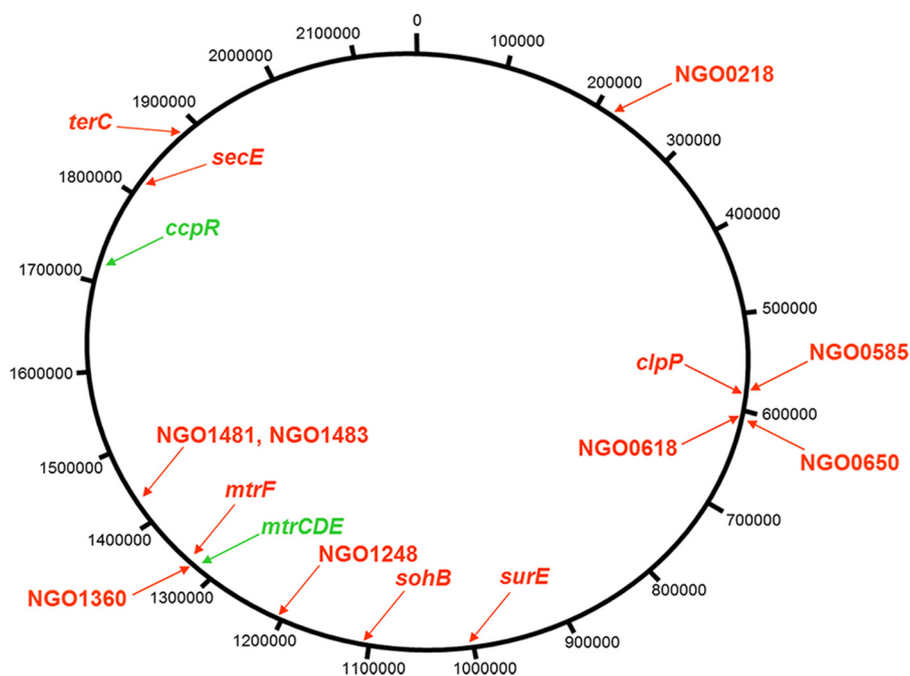
**TABLE 1** Transcriptional response of gonococci to overexpression of *mtrCDE* efflux pump operon

| Gene and category   | Common name | Fold change | Functional classification <sup>a</sup>                                   |
|---|-------------|-------------|--|
| Upregulated in the presence of the <i>mtr</i> <sub>120</sub> mutation   |             |             |  |
| NGO1363   | <i>mtrE</i> | 4.44        | Mtr efflux pump protein component: outer membrane channel protein (MtrE) |
| NGO1364   | <i>mtrD</i> | 5.63        | Mtr efflux pump protein component: RND family transporter (MtrD)         |
| NGO1365   | <i>mtrC</i> | 5.42        | Mtr efflux pump protein component: periplasmic fusion protein (MtrC)     |
| NGO1769   | <i>ccpR</i> | 2.52        | Cytochrome <i>c</i> peroxidase   |
| Downregulated in the presence of the <i>mtr</i> <sub>120</sub> mutation |             |             |  |
| NGO0218   | NGO0218     | -15.02      | Hypothetical   |
| NGO0585   | NGO0585     | -2.02       | Hypothetical integral membrane protein                                   |
| NGO0593   | <i>clpP</i> | -2.03       | ATP-dependent Clp protease subunit                                       |
| NGO0618   | NGO0618     | -2.14       | Hypothetical   |
| NGO0650   | NGO0650     | -2.05       | ATP-dependent RNA helicase   |
| NGO1058   | <i>surE</i> | -2.14       | Stationary-phase-survival protein  |
| NGO1246   | <i>sohB</i> | -2.72       | Periplasmic peptidase, S49 family  |
| NGO1248   | NGO1248     | -5.23       | Hypothetical   |
| NGO1360   | NGO1360     | -3.56       | FadR/GntR family transcriptional regulator                               |
| NGO1368   | <i>mtrF</i> | -2.67       | Mtr efflux pump accessory protein  |
| NGO1481   | NGO1481     | -4.59       | Putative SAM-dependent methyltransferase                                 |
| NGO1857   | <i>secE</i> | -4.83       | Protein translocase channel subunit                                      |
| NGO1917   | <i>terC</i> | -3.06       | Transmembrane transporter, tellurium resistance                          |

<sup>a</sup> RND, resistance-nodulation-division; SAM, S-adenosylmethionine.

analysis revealed that expression of *mtrC*, *mtrD*, and *mtrE* was highly upregulated in strain FA19*mtr*<sub>120</sub> compared to wild-type strain FA19, which, based on earlier reports (4, 5, 9), served as an internal control for the results of the transcriptome comparison (Table 1); this upregulation was confirmed by qRT-PCR (see Table S1 in the supplemental material). Importantly, analysis of the RNA-seq results revealed changes in the expression levels of 13

non-*mtr* genes which were widely distributed across the genome (Fig. 1) and represented a variety of functional classes (Table 1) based on annotation of the FA1090 genome ([www.genome.ou.edu](http://www.genome.ou.edu)). Only one of these non-*mtr* genes (*ccpR*) was overexpressed in strain FA19*mtr*<sub>120</sub>. Proteins encoded by the 13 underexpressed genes included four hypothetical proteins; a protease (encoded by *clpP*); a periplasmic peptidase (*sohB*); a stationary-



**FIG 1** Shown are chromosomal-map positions of genes differentially expressed in strain FA19*mtr*<sub>120</sub> compared to wild-type strain FA19 as identified by RNA-seq analysis (Table 1). The circular map is from the annotated FA1090 genome ([www.genome.ou.edu](http://www.genome.ou.edu)). Upregulated genes are shown in green, while downregulated genes are shown in red.

TABLE 2 Sensitivity of isogenic gonococci to antimicrobials

| Strain  | Level of susceptibility to antimicrobials <sup>a</sup> |      |         |     |                               |        |
|---|--|------|---------|-----|-------------------------------|--------|
|   | Em   | CV   | TX-100  | PxB | H <sub>2</sub> O <sub>2</sub> | tBuOOH |
| FA19  | 0.25   | 0.31 | 125     | 100 | 0.002                         | 0.004  |
| FA19 <i>ccpR::kan</i>                           | 0.25   | 0.31 | 125     | 100 | 0.002                         | 0.004  |
| FA19 <i>mtr</i> <sub>120</sub>                  | 2  | 1.25 | >16,000 | 400 | 0.004                         | 0.016  |
| FA19 <i>mtr</i> <sub>120</sub> <i>ccpR::kan</i> | 2  | 1.25 | >16,000 | 400 | 0.002                         | 0.008  |

<sup>a</sup> Em, erythromycin; CV, crystal violet; TX-100, Triton X-100; PxB, polymyxin B; H<sub>2</sub>O<sub>2</sub>, hydrogen peroxide; tBuOOH, tert-butylhydroperoxide. For efflux pump substrates Erm, CV, TX-100, and PxB, the numbers refer to MICs (μg/ml), while the numbers for H<sub>2</sub>O<sub>2</sub> and tBuOOH are minimal bactericidal concentrations (MBCs) in percent (vol/vol). All determinations were performed in triplicate.

phase-associated survival protein (*surE*); a transcriptional regulator (NGO1360) involved in modulating expression of glutamate metabolism genes in meningococci (15) and located just downstream of the upregulated *mtrCDE* operon; an ATP-RNA helicase (NGO650); a component of the protein translocating system (*secE*); a tellurium resistance-associated protein (*terC*); and a putative methyltransferase (NGO1481).

We selected *ccpR*, which encodes cytochrome C peroxidase (16), as a model gene to further test if overexpression of *mtrCDE* could have distal effects on gonococcal gene expression; *ccpR* was also chosen as it may contribute to gonococcal resistance to oxidative stresses (16, 17). By qRT-PCR analysis, we confirmed enhanced expression of *ccpR* in strain FA19*mtr*<sub>120</sub> compared to wild-type strain FA19 (see Table S1 in the supplemental material). We next examined our test strains with or without a *ccpR::kan* mutation for differences in susceptibility to MtrCDE-substrate antimicrobials and two peroxides (H<sub>2</sub>O<sub>2</sub> and tBuOOH), since the peroxidase activity of CcpR would likely influence levels of gonococcal susceptibility to peroxides but not MtrR substrates. In three independent experiments, strain FA19*mtr*<sub>120</sub> showed an increase in resistance to efflux pump substrates (Table 2) and resistance to H<sub>2</sub>O<sub>2</sub> and tBuOOH at levels 2-fold- and 4-fold greater, respectively, than those seen with strain FA19; peroxide resistance levels were independent of the presence of a functional MtrCDE efflux pump (data not presented). While the susceptibility of strain FA19*mtr*<sub>120</sub> *ccpR::kan* to pump substrates was unchanged from that seen with FA19*mtr*<sub>120</sub>, strain FA19*mtr*<sub>120</sub> *ccpR::kan* was reproducibly 2-fold more sensitive to tBuOOH, which was not unexpected given the multiple ways gonococci resist peroxides (16, 17).

We propose that overexpression of a drug efflux pump has unexpected secondary effects on bacterial gene expression and associated metabolic processes. The mechanism(s) by which these expression changes occur is unclear but should be considered in drug efflux studies.

**Nucleotide sequence accession numbers.** The complete data set can be accessed through GEO accession number GSE47048 and SRA accession number SRA079863.

## ACKNOWLEDGMENTS

We thank Jacqueline Balthazar for excellent technical assistance and Yarahmah Zalucki for critical reading of early versions of the paper.

This work was supported by NIH grants AI021150-29 (W.M.S.), AI42053 (A.E.J.), AI096788 (T.D.R., D. Dean, and R. Selden), and AI031496-22 (to P. F. Sparling, University of North Carolina) and a VA

Merit Review Grant to W.M.S. W.M.S. is the recipient of a Senior Research Career Scientist Award from the VA Medical Research Service.

## REFERENCES

- Nikaido H. 1996. Multidrug efflux pumps of Gram-negative bacteria. *J Bacteriol* 178:5853–5859.
- Zalucki YM, Mercante AD, Cloward JM, Ohneck EA, Kandler JL, Goytia M, Johnson PJT, Shafer WM. 2013. Function and regulation of *Neisseria gonorrhoeae* efflux pumps, p 207–221. In Yu EW (ed), *Microbial efflux pumps: current research*. Horizon Press, Inc., Pittsburgh, PA.
- Pan W, Spratt BG. 1994. Regulation of the permeability of the gonococcal cell envelope by the *mtr* system. *Mol Microbiol* 11:769–775. <http://dx.doi.org/10.1111/j.1365-2958.1994.tb00354.x>.
- Hagman KE, Pan W, Spratt BG, Balthazar JT, Judd RC, Shafer WM. 1995. Resistance of *Neisseria gonorrhoeae* to antimicrobial hydrophobic agents is modulated by the *mtrRCDE* efflux system. *Microbiology* 141: 611–622. <http://dx.doi.org/10.1099/13500872-141-3-611>.
- Hagman KE, Shafer WM. 1995. Transcriptional control of the *mtr* efflux system of *Neisseria gonorrhoeae*. *J Bacteriol* 177:4162–4165.
- Warner DM, Folster JP, Shafer WM, Jerse AE. 2007. Regulation of the MtrC-MtrD-MtrE efflux-pump system modulates the *in vivo* fitness of *Neisseria gonorrhoeae*. *J Infect Dis* 196:1804–1812. <http://dx.doi.org/10.1086/522964>.
- Warner DM, Shafer WM, Jerse AE. 2008. Clinically relevant mutations that cause derepression of the *Neisseria gonorrhoeae* MtrC-MtrD-MtrE efflux pump system confer different levels of antimicrobial resistance and *in vivo* fitness. *Mol Microbiol* 70:462–478. <http://dx.doi.org/10.1111/j.1365-2958.2008.06424.x>.
- Folster JP, Dhulipali V, Nicholas RA, Shafer WM. 2007. Differential regulation of *ponA* and *pilMNOPQ* expression by the *Neisseria gonorrhoeae* MtrR transcriptional regulatory protein. *J Bacteriol* 189:4569–4577. <http://dx.doi.org/10.1128/JB.00286-07>.
- Folster JP, Johnson PJT, Jackson L, Dhulipali V, Dyer DW, Shafer WM. 2009. MtrR modulates *ropH* expression and levels of antimicrobial resistance in *Neisseria gonorrhoeae*. *J Bacteriol* 191:287–297. <http://dx.doi.org/10.1128/JB.01165-08>.
- Johnson PJT, Stringer VA, Shafer WM. 2011. Off-target gene regulation mediated by transcriptional repressors of antimicrobial efflux pump genes in *Neisseria gonorrhoeae*. *Antimicrob Agents Chemother* 55:2559–2565. <http://dx.doi.org/10.1128/AAC.00010-11>.
- Ohneck EA, Zalucki YM, Johnson PJT, Dhulipala V, Golparian D, Unemo M, Jerse AE, Shafer WM. 2011. A novel mechanism of high-level, broad-spectrum antibiotic resistance caused by a single base pair change in *Neisseria gonorrhoeae*. *mBio* 2:e00187–11. <http://dx.doi.org/10.1128/mBio.00187-11>.
- Rouquette C, Harmon JB, Shafer WM. 1999. Induction of the *mtrCDE*-encoded efflux pump system of *Neisseria gonorrhoeae* requires MtrA, an AraC-like protein. *Mol Microbiol* 33:651–658. <http://dx.doi.org/10.1046/j.1365-2958.1999.01517.x>.
- Shafer WM, Qu X-D, Waring AJ, Lehrer RI. 1998. Modulation of *Neisseria gonorrhoeae* susceptibility to vertebrate antibacterial peptides due to a member of the resistance/modulation/division efflux pump family. *Proc Natl Acad Sci U S A* 95:1829–1833. <http://dx.doi.org/10.1073/pnas.95.4.1829>.
- Vélez Acevedo RN, Ronpirin C, Kandler JL, Shafer WM, Cornelissen CN. 2014. Identification of regulatory elements that control expression of the *tbpBA* operon in *Neisseria gonorrhoeae*. *J Bacteriol* 196:2762–2774. <http://dx.doi.org/10.1128/JB.01693-14>.
- Pagliarulo C, Salvatore P, De Vitis LR, Colicchio R, Monaco C, Tredici M, Tala A, Bardaro M, Lavitola A, Bruni CB, Alifano P. 2004. Regulation and differential expression of *gdhA* encoding NADP-specific glutamate dehydrogenase in *Neisseria meningitidis* clinical isolates. *Mol Microbiol* 51:1757–1772. <http://dx.doi.org/10.1111/j.1365-2958.2003.03947.x>.
- Turner S, Reid E, Smith H, Cole J. 2003. A novel cytochrome c peroxidase from *Neisseria gonorrhoeae*: a lipoprotein from a Gram-negative bacterium. *Biochem J* 373:865–873. <http://dx.doi.org/10.1042/BJ20030088>.
- Falsetta ML, Steichen CT, McEwan AG, Cho C, Ketterer M, Shao J, Hunt J, Jennings MP, Apicella MA. 2011. The composition and metabolic phenotype of *Neisseria gonorrhoeae* biofilms. *Front Microbiol* 2:75. <http://dx.doi.org/10.3389/fmicb.2011.00075>.