Escherichia coli enterobactin synthesis and uptake mutants are hypersensitive to an antimicrobial peptide that limits the availability of iron in addition to blocking Holliday junction resolution

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The peptide wrwycr inhibits Holliday junction resolution and is a potent antimicrobial. To study the physiological effects of wrwycr treatment on Escherichia coli cells, we partially screened the Keio collection of knockout mutants for those with increased sensitivity to wrwycr. Strains lacking part of the ferric-enterobactin (iron-bound siderophore) uptake and utilization system, parts of the enterobactin synthesis pathway, TolC (an outer-membrane channel protein) or Fur (an iron-responsive regulator) were hypersensitive to wrwycr. We provide evidence that the ΔtolC mutant was hypersensitive to wrwycr due to its reduced ability to efflux wrwycr from the cell rather than due to its export of newly synthesized enterobactin. Deleting ryhB, which encodes a small RNA involved in iron regulation, mostly relieved the wrwycr hypersensitivity of the fur and ferric-enterobactin uptake mutants, indicating that the altered regulation of a RyhB-controlled gene was at least partly responsible for the hypersensitivity of these strains. Chelatable iron in the cell, measured by electron paramagnetic resonance spectroscopy, increased dramatically following wrwycr treatment, as did expression of Fur-repressed genes and, to some extent, mutation frequency. These incongruous results suggest that while wrwycr treatment caused accumulation of chelatable iron in the cell, iron was not available to bind to Fur. This is corroborated by the observed induction of the suf system, which assembles iron–sulfur clusters in low-iron conditions. Disruption of iron metabolism by wrwycr, in addition to its effects on DNA repair, may make it a particularly effective antimicrobial in the context of the low-iron environment of a mammalian host.

INTRODUCTION

Holliday junctions are DNA structures with four double-stranded arms that serve as intermediates in recombination-dependent repair and rearrangement of DNA in both prokaryotes and eukaryotes. We have identified synthetic hexapeptides that bind Holliday junctions as disulfide-bridged dimers and inhibit Holliday junction resolution (Boldt et al., 2004; Kepple et al., 2005). One of these peptides, with the sequence wrwycr (we use the D-amino acid form to limit biological degradation), is a potent, broad-spectrum antimicrobial with the ability to inhibit, in vivo, mechanisms of DNA recombination and damage repair that proceed through a Holliday junction intermediate (Gunderson & Segall, 2006; Gunderson et al., 2009; L. Marcusson and other authors, unpublished data). Because mammalian host defences such as gastric acid and the oxidative burst in macrophages induce DNA damage and thus DNA repair in bacterial invaders, use of wrwycr in antimicrobial therapy may increase the efficacy of these natural defence systems. Recently, wrwycr was shown to inhibit the growth of Salmonella enterica serovar Typhimurium inside macrophages (Su et al., 2010) and significantly enhance acid-induced killing of Shiga toxin-producing strains of Escherichia coli associated with haemolytic uraemia syndrome (Lino et al., 2011).

While it is known that wrwycr targets Holliday junctions, it is not known how wrwycr enters cells, what other cellular targets it may have and what defence mechanisms cells employ against wrwycr. To address these questions, we screened a knockout library of E. coli for mutants that are resistant or hypersensitive to wrwycr. We expected mutations that conferred wrwycr resistance to inform us of cellular targets of wrwycr or mechanisms of wrwycr entry into the cell. We expected wrwycr-hypersensitive mutants to inform us of wrwycr efflux or other detoxification mechanisms as
An additional layer of iron regulation in E. coli

Iron is an essential element for almost all organisms, but can also be toxic. In what is known as Fenton chemistry, iron catalyses the decomposition of hydrogen peroxide (Fenton, 1894), resulting in hydroxyl radical and hydroxyl iron catalyses the decomposition of hydrogen peroxide can also be toxic. In what is known as Fenton chemistry, their iron-responsive regulatory system.

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Iron-responsive transcriptional regulator, plays a key role in the regulation of iron-responsive genes. Incongruously, they import from their environment. Fur, an iron-responsive regulatory system, is well as pathways that, when perturbed, increase targets for the peptide. From a set of wrwycr-hypersensitive mutants described here, we have discovered that wrwycr also affects the regulation of iron-responsive genes. Incogruously, wrwycr appears to simultaneously increase the measurable amount of desferrioxamine-chelatable iron and decrease the pool of bioavailable iron in the cell. This latter trait makes wrwycr especially detrimental to bacteria in iron-deficient environments or to bacteria that, through mutation, have lost components of their high-affinity iron-uptake system or their iron-responsive regulatory system.

An additional layer of iron regulation in E. coli is mediated through the small RNA RyhB, which, as a repressor, acts in concert with the RNA-binding protein Hfq to facilitate target transcript degradation by RNase E (Massé et al., 2003). While Fur regulates the amount of incoming iron, RyhB aids degradation of certain transcripts to direct available iron away from non-essential uses (such as iron storage or incorporation into non-essential proteins) in low-iron conditions, thus ensuring that the limited amount of iron is put to the most beneficial use (Massé et al., 2005). RyhB has also been shown to increase the levels of some transcripts (Massé et al., 2005), perhaps by preventing their degradation by RNases, and to activate translation of the shiA (shikimate transport) transcript by allowing access to the transcript’s ribosome-binding site (Prévost et al., 2007). In high-iron conditions, Fur represses transcription of ryhB (Massé & Gottesman, 2002; Vassinova & Kozyrev, 2000), leading to an increase in iron storage and a decrease in shikimate uptake.

Many bacteria, both environmental and pathogenic species, have specialized systems for taking up iron from their environments, which can be very low in soluble iron (Neilands, 1981). Enterobactin is a siderophore synthesized non-ribosomally in a number of Gram-negative bacteria and in a few Gram-positive bacteria (reviewed by Raymond et al., 2003). It is exported to the extracellular milieu by EntS (Furrer et al., 2002) and TolC (Bleuel et al., 2005) and, once it has bound iron to become ferric enterobactin, is imported by the cell (Supplementary Fig. S1, available with the online version of this paper). FepA is an outer-membrane protein that transports extracellular ferric enterobactin into the periplasm; the TonB–ExbB–ExbD complex associated with the cytoplasmic membrane provides the energy for the activity of FepA (Guerinot, 1994). FepB binds to ferric enterobactin in the periplasm and directs it to the FepD–FepG cytoplasmic membrane transporter; FepC is an ATPase that assists FepD and FepG in transporting ferric enterobactin into the cytoplasm (Guerinot, 1994). Fes is the ferric enterobactin esterase that degrades ferric enterobactin into three 2,3-dihydroxybenzoyl-L-serine units to release iron, which is then reduced by an unknown mechanism (Brickman & McIntosh, 1992).

In our screen for E. coli mutants with altered sensitivity to wrwycr, we identified several physiologically related hypersensitive mutants: ΔfepB, ΔfepC, ΔfepD, ΔfepG, Δfes, ΔtolC and Δfur. In the course of the work presented here, we found that ΔentB, ΔentC and ΔmenF mutants were also wrwycr-hypersensitive. Fur represses expression of the fep genes in high-iron conditions, EntB and EntC are involved in synthesizing enterobactin (Supplementary Fig. S1) in addition to other compounds (Bleuel et al., 2005; Koronakis, 2003), and MenF synthesizes isochorismate for menaquinone production (Darwala et al., 1996). We present evidence that wrwycr limits iron availability (although it actually increases chelatable iron in the cell) or otherwise alters iron regulation in the cell, causing hyper-repression or hypo-activation of an unidentified gene by RyhB and thus hypersensitivity in the fur and enterobactin synthesis and uptake mutants.

METHODS

Strains, media and chemicals. Strains (Supplementary Table S1, available with the online version of this paper) were routinely cultured in LB [5 g NaCl l−1 (Davis et al., 1980)] at 37 °C with shaking (225 r.p.m.). The medium used for screening the Keio collection, no citrate E salts (NCE)/glucose/Casamino acids/vitamins (NGCV), was 1 × NCE minimal medium base (Davis et al., 1980) with 0.2% glucose, 0.1% autoclaved Casamino acids and 1 × vitamin mix 1 (500 × stock of vitamin mix 1 is an autoclaved mixture of 0.025% folic acid, 0.025% pantothenic acid, 0.025% nicotinamide, 0.025% pyridoxal-HCl, 0.025% thiamine-HCl, 0.0025% riboflavin and 0.05% biotin) and no added iron. The MOPS/glucose/Casamino acids/vitamins (MGCV) medium was based on a component of a ‘culture medium for enterobacteria’ (Neidhart et al., 1974) and consisted of 40 mM MOPS, 4 mM Tricine (the 10 × MOPS/Tricine mixture was adjusted to pH 7.4 with KOH), 9.5 mM NH4Cl, 0.276 mM K2SO4, 0.5 μM CaCl2, 0.525 mM MgCl2, 5 mM NaCl, 1.32 mM K2HPO4, 0.2% glucose, 1% Casamino acids and 1 × vitamin mix 2 (1000 × stock of vitamin mix 2 is a filter-sterilized mixture of 0.05% folic acid, 0.05% pantothenic acid, 0.05% nicotinamide, 0.05% nicotinic acid, 0.05% pyridoxine-HCl, 0.05% thiamine-HCl, 0.005% riboflavin and 0.1% biotin, with NaOH added to assist solubilization). FeCl3 was added to assist solubilization). FeCl3 was added to 2.4 or 6 mM (as indicated) for low-iron conditions and up to 150 μM (as indicated) for high-iron conditions. Mueller–Hinton broth powder was commercially prepared (Becton Dickinson). Chemicals and media components were purchased from Sigma-Aldrich or Fisher Scientific unless indicated otherwise. The peptide wrwycr consisted of d-amino acids.
acid residues with an amide group at the C-terminus, and was synthesized and purified to >95% purity by BioSynthesis or Sigma-Genosys. See Supplementary Material for full details on D-wrwycr.

Screening of Keio collection for hypersensitive mutants. To screen the Keio collection of 3985 mutants (Baba et al., 2006), a 96-place replicator was used to inoculate microtiter wells containing the tolerance-testing medium (100 μl) with overnight cultures. The tolerance-testing medium was NGCV with either DMSO (solvent-only control; final concentration 0.5%) or wrwycr (20 μM with 0.2% final DMSO concentration) added. The inoculated plates were sealed with paraffin film to reduce evaporation, incubated at 30 °C and the OD<sub>600</sub> was read at 24 and 72 h. To factor out the different wrwycr-free growth abilities of the various mutant strains, the optical density reading in the presence of wrwycr for each time point was divided by the optical density for the same strain in the presence of DMSO alone to get a ratio. Typically, mutants with a ratio of <0.3 were deemed to be candidate hypersensitive strains if the majority of strains on the plate had a higher ratio. For verified hypersensitive strains and additional Keio collection strains of interest, the mutated locus was transduced into <i>E.coli</i> MG1655 (Supplementary Table S1) or another desired genetic background using bacteriophage P1 (Miller, 1972). Where indicated, the kanamycin-resistance cassette (FRT-kan<sup>R</sup>-FRT) was removed via recombination between the flanking FRT sites using Flp recombinase supplied from plasmid pCP20 (Cherepanov & Wackernagel, 1995); the strains were subsequently cured of pCP20 by growth at 42 °C. The deletion/insertion site of all listed strains (Supplementary Table S1) was verified by PCR.

Other strain construction. The ΔryhB mutant strain was constructed by replacing the <i>ryhB</i> locus with a FRT-kan<sup>R</sup>-FRT cassette from plasmid pKD13 into strain EDT1160 and transducing the Δ<i>ryhB</i>:FRT-kan<sup>R</sup>-FRT locus into strain G652 to create strain EDT1571. The Δ<i>fur</i>Δ<i>ryhB</i> double mutants were constructed by transducing the Δ<i>ryhB</i>:FRT-kan<sup>R</sup>-FRT locus into the appropriate single mutants (Supplementary Table S1) using P1. To create strain EDT1689, the FRT-kan<sup>R</sup>-FRT cassette was removed from EDT1571; to create strain EDT1712, the Δ<i>fur</i>:FRT-kan<sup>R</sup>-FRT locus was transduced into strain EDT1689. Full details of these methods can be found in the Supplementary Material.

Growth assays. Because phosphate, a major component of NCE and thus the NGCV medium, co-precipitates with other iron cations, a MOPS-based medium (MGCV, above) was used for growth studies involving the addition of iron, except when noted. For wrwycr sensitivity and iron complementation assays, overnight cultures were diluted to 1% in MGCV, or NGCV for the Δ<i>fur</i> and Δ<i>fur</i> Δ<i>ryhB</i> mutants, containing the concentration of wrwycr indicated in figure legends or the same volume of 100% DMSO, and FeCl<sub>3</sub> (dissolved in 0.1 M HCl) added at the concentrations indicated in the figure legends or the same volume of 0.1 M HCl in a 96-well plate; the plates were sealed with paraffin film, grown at 37 °C with 1 min agitation every 10 min, and the OD<sub>600</sub> was periodically assayed in a VersaMax or SpectraMax plate reader (Molecular Devices). The data are presented with the <i>y</i>-axis on a log<sub>2</sub> scale, as appropriate for cell growth data.

Determining dimer : monomer ratio and cell-associated concentration of wrwycr. Reverse-phase (RP)-HPLC was used to visualize (by absorbance at 280 nm) and separate the monomer and dimer peaks of wrwycr in order to monitor the extent of dimerization visualized (by absorbance at 280 nm) and separate the monomer and dimer peaks of wrwycr in order to monitor the extent of dimerization (protocol adapted from an unpublished protocol by E. Ozyamak and I. Booth, University of Aberdeen, Scotland). Following centrifugation, 35 μl of the formic acid solution of each sample was mixed with 215 μl water and 100 μl of the mix was analysed by RP-HPLC as described above. The area under the monomer and dimer wrwycr peaks was used to determine the relative amount of wrwycr in each sample. Bacteria not treated with wrwycr were treated and analysed similarly. The RP-HPLC results were compared with those of formic acid only to identify a cell-dependent peak that eluted from the column at approximately 6–7 min, the area of which correlated with the number of untreated cells used in the extraction up to a maximum of ~109 cells.

The area under the cell-dependent peak or the OD<sub>600</sub> of the treated cultures was used to normalize the results for total wrwycr concentration to account for differences in the number of sampled cells. Data for each normalization method are shown separately. Full details are given in the Supplementary Material.

Quantitative PCR (qPCR) of Fur regulon genes. Overnight cultures (200 μl) from three independent colonies per strain grown in MHB were used to inoculate 20 ml MHB in 50 ml flasks. After 1 h growth, 10 μM wrwycr or 0.1% DMSO was added to the cultures. After 1.5 h further growth, RNA was extracted and used to synthesize cDNA, which was then analysed by qPCR. The qPCR conditions are given in the Supplementary Material and the primers used are listed in Supplementary Table S2.

4-Methylumbelliferyl-b-D-glucuronide (MUG) assays for β-galactosidase activity. Overnight cultures in MGCV supplemented with 6 μM FeCl<sub>3</sub> were subcultured 1:500 in fresh MGCV + 6 μM FeCl<sub>3</sub> and incubated at 37 °C until the cultures reached OD<sub>600</sub> ~0.3. Cells were treated in a 96-well microtitre plate by adding 100 μl of the cultures to 100 μl of the MUG (0.4 mg ml<sup>-1</sup>) buffer (200 mM, pH 10.3). The fluorescence intensity of the product, 4-methylumbelliferone, was measured at an excitation of 360 nm and an emission of 450 nm in a Molecular Devices SpectraMax GEMINI plate reader (Molecular Devices). The fluorescence was normalized by the OD<sub>600</sub> of the culture at the time the 10 μl aliquots were taken. See Supplementary Material for full details.

Catalase assays Catalase activity in extracts of various bacterial strains was determined using an assay described previously (Li & Schellhorn, 2007). Briefly, MG1655 and the Δ<i>fepB</i> strain were grown for 1.5 h, then DMSO or wrwycr (to 10 μM) was added to the culture and cells were sampled after a further 1.5 h or 3 h incubation. Cell pellets from overnight or treated cultures were resuspended in 50 mM potassium phosphate buffer, pH 7.0, lysed by sonication and the cell debris was removed by centrifugation. Total protein concentration in the cell extracts was determined by using a Bradford-type assay (Bio-Rad). Immediately before assay readings were taken, 250 μl 5 mM H<sub>2</sub>O<sub>2</sub> in 50 mM potassium phosphate buffer, pH 7.0, was added to samples and standards in a 96-well UV-transparent plate (Costar 3635, Corning). Catalase activity was assayed by monitoring the OD<sub>600</sub> increase in the presence of wrwycr for each time point was divided by the OD<sub>600</sub> of the culture at the time the 10 μl aliquots were taken. See Supplementary Material for full details.
absorbance at 240 nm, which decreases as H₂O₂ is degraded. The A₅₇₀ was recorded in a SpectraMax plate reader (Molecular Devices) for 5 min at ~8 s intervals, the slope of change in A₅₇₀ (calculated for each time point relative to t=0) was determined, and catalase activity (in units per mg total protein) was calculated from the standard curve and protein concentration determination. Full details are given in the Supplementary Material.

**Electron paramagnetic resonance (EPR) spectroscopy.** Intracellular chelatable iron concentrations were measured by EPR spectroscopy as described by Woodmansee & Imlay (2002). Full details are given in the Supplementary Material.

**Determination of mutation frequency using the medium method.** Sixteen overnight cultures, each inoculated from an independent colony, were subcultured 1:500 in NGCV media with HPLC for all samples. A two-tailed and dimer : monomer ratio of cell-associated wrwycr detected by RP-multiple comparison test was performed to compare the total amount wrwycr to achieve a final concentration of 8 or 16 µM FeCl₃ with wrwycr. We found that, in the MG1655 background, we reasoned that if wrwycr hypersensitivity in the Δfep and Δfes mutants was caused by hindrance of ferric enterobactin transport across the cytoplasmic membrane rather than hindrance of transport of another compound through the same system, the impairment of enterobactin synthesis would also render E. coli hypersensitive to wrwycr. We thus transduced the mutated loci of entB and entC, two genes whose products are required for enterobactin synthesis, from the corresponding Keio collection strains to the MG1655 background and measured the growth of the resulting strains in wrwycr. While both the enterobactin synthesis mutants tested were hypersensitive to wrwycr, the ΔentB strain was more hypersensitive than the ΔentC mutant strain (Fig. 1c). These results suggest that enterobactin synthesis is required for wild-type-level resistance to wrwycr but that there is an EntC-independent way of making or taking up the dihydroxybenzoate substrate for enterobactin synthesis under our test conditions. EntC is an isochorismate synthase and E. coli encodes another isochorismate synthase, MenF (Daruwala et al., 1996), which is needed to produce menaquinone for electron transport in anaerobic conditions. As our growth assays were done in sealed, predominantly static 96-well microtitre plates with glucose as the carbon source, it is likely that the dissolved oxygen content in the cultures was very low. Therefore, we tested whether MenF could substitute for EntC in

**RESULTS**

**Identification of wrwycr-hypersensitive E. coli mutants**

The Keio collection of E. coli deletion–insertion mutants was partially screened to find mutations that increased or decreased the tolerance of E. coli for the peptide wrwycr (Methods). Here we describe our investigation of a subset of strains that appeared hypersensitive in this screen, a group of mutants related to enterobactin synthesis and transport, namely ΔfepA, ΔfepC, ΔfepD, ΔfepG and Δfes. FepB (a periplasmic binding protein), FepC, FepD and FepG are involved in the uptake of the iron-bound siderophore molecule ferric enterobactin across the cytoplasmic membrane, and Fes breaks down imported ferric enterobactin in the cytoplasm to release iron (Guerinot, 1994). We verified the hypersensitivity of these mutants by transducing the mutations into E. coli MG1655 and testing the response of the MG1655-based strains to wrwycr compared with that of the MG1655 parent strain. We also measured the effect of two other ferric enterobactin uptake-related mutations transduced from the Keio collection, ΔfepA and ΔtonB, on sensitivity of MG1655 to wrwycr. We found that, in the MG1655 background, ΔfepB, ΔfepC, ΔfepD, ΔfepG, ΔtonB and Δfes showed no growth (OD₆₀₀ ratio <0.3) at 30 °C in 15 µM wrwycr in NGCV medium, which has no added iron other than that introduced by other media constituents or from the glassware, even after 72 h incubation. The ΔfepA mutant showed hypersensitivity only at the 24 h and not at the 48 and 72 h observation points. MG1655 is resistant (OD₆₀₀ ratio >0.8) to 15 µM wrwycr in the same conditions (for comparison, it is sensitive to 25–30 µM wrwycr). We removed the kanamycin resistance cassette from the MG1655-based strains and followed their growth when challenged with 22 µM wrwycr in MOPS-based MGCV medium (Methods) containing 2.4 µM FeCl₃; this amount of iron was chosen because the enterobactin uptake mutants were unable to grow in lower concentrations of FeCl₃ in MGCV in the absence of wrwycr. This medium contains a low level of phosphate, insufficient to precipitate the added iron, unlike the greater amount of phosphate present in NGCV. All the mutants grew essentially the same as the parent strain in MGCV/2.4 µM FeCl₃ without wrwycr but with the DMSO solvent (Fig. 1a). In contrast, the growth of the mutants, except ΔfepA and ΔtonB, in MGCV/2.4 µM FeCl₃ with wrwycr was delayed relative to M1655 (open symbols in Fig. 1b, c), verifying the original phenotype; the ΔfepA and ΔtonB mutants showed very little hypersensitivity to wrwycr in this assay (Fig. 1b). In all cases, the mutants recovered, often with a similar growth rate; we show below that this recovery is likely to be largely dependent on the TolC-dependent efflux, since the efflux-defective ΔtolC strain is hypersensitive to wrwycr and loses much of its ability to recover.

We reasoned that if wrwycr hypersensitivity in the Δfep and Δfes mutants was caused by hindrance of ferric enterobactin transport across the cytoplasmic membrane rather than hindrance of transport of another compound through the same system, the impairment of enterobactin synthesis would also render E. coli hypersensitive to wrwycr. We thus transduced the mutated loci of entB and entC, two genes whose products are required for enterobactin synthesis, from the corresponding Keio collection strains to the MG1655 background and measured the growth of the resulting strains in wrwycr. While both the enterobactin synthesis mutants tested were hypersensitive to wrwycr, the ΔentB strain was more hypersensitive than the ΔentC mutant strain (Fig. 1c). These results suggest that enterobactin synthesis is required for wild-type-level resistance to wrwycr but that there is an EntC-independent way of making or taking up the dihydroxybenzoate substrate for enterobactin synthesis under our test conditions. EntC is an isochorismate synthase and E. coli encodes another isochorismate synthase, MenF (Daruwala et al., 1996), which is needed to produce menaquinone for electron transport in anaerobic conditions. As our growth assays were done in sealed, predominantly static 96-well microtitre plates with glucose as the carbon source, it is likely that the dissolved oxygen content in the cultures was very low. Therefore, we tested whether MenF could substitute for EntC in
enterobactin synthesis in the low-oxygen environment of microtitre plates. A ΔmenF mutant had a hypersensitivity to wrwycr intermediate between that of the wild-type strain and the ΔentC mutant strain (Supplementary Fig. S2a), while a ΔentC ΔmenF double mutant was more sensitive to wrwycr than either of the single mutants (Supplementary Fig. S2a), suggesting that either MenF activity can compensate for the lack of EntC for enterobactin synthesis in our assay conditions or menaquinone synthesis itself is important for resistance to wrwycr in low-oxygen environments.

A ΔtolC mutant is hypersensitive to wrwycr

Another wrwycr-hypersensitive strain identified in the original screen was the ΔtolC mutant. This strain was significantly more sensitive to wrwycr than were the other strains reported here; it did not grow in 10 μM wrwycr even after 48 h (Fig. 1d), while the other mutants all grew at least somewhat in 22 μM wrwycr (Figs 1b and 2c). TolC not only is essential for the export of newly synthesized enterobactin across the outer membrane (Bleuel et al., 2005) but also assists in the efflux of many other compounds, including antibacterial drugs (reviewed by Koronakis et al., 2004). EntS is involved in transport of enterobactin across the cytoplasmic membrane (Furrer et al., 2002) and thus an entS mutant could be expected to have the same phenotype as a tolC mutant. However, the E. coli ΔentS strain was not hypersensitive to wrwycr (Supplementary Fig. S2b). This suggests that enterobactin export by EntS is not essential for the ability...
of *E. coli* to withstand wrwycr treatment, probably because *entS* mutants can export components of enterobactin that are themselves siderophores (Furrer et al., 2002), and that the *tolC* mutant must have an additional defect that made it hypersensitive to wrwycr. Below, we present evidence that the additional defect is a diminished capacity to export wrwycr.

**Fig. 2.** The *E. coli* ΔtolC mutant, but not the ΔfepB mutant, has an increased level of cell-associated wrwycr and a lower ratio of dimer to monomer wrwycr relative to the MG1655 parent strain, as measured in an RP-HPLC assay. (a) Overlay of typical RP-HPLC traces of formic acid extracts from *E. coli* MG1655 (G652) treated with wrwycr (solid line) or DMSO (short-dashed line) and of DMSO in formic acid (long-dashed line) to show identification of cell-dependent peaks and monomer and dimer wrwycr, as indicated. (b–d) Exponential-phase MG1655 (G652, ■), ΔtolC (EDT1311, ▲) or ΔfepB (EDT1316, ◆) cells were treated with 10 μM wrwycr for 1.5 h, after which cell-associated peptides were extracted using formic acid. The formic acid extracts were analysed by RP-HPLC and the detected monomer and dimer wrwycr peak areas were normalized to (b) the area of a cell-dependent peak, the area of which correlated with the number of DMSO-treated cells subjected to formic acid extraction (data not shown), or to (c) the OD₆₀₀ of the cultures at the time of formic acid extraction. (b, c) The normalized total peptide amounts expressed as monomer equivalents (b, c) and dimer to monomer ratios (d) are shown. For (b–d), each data point represents the result for an independent culture and the horizontal lines indicate the means of the samples. Data points are from (b) six, (c) three or (d) five separate experiments with three independent cultures each. A one-way analysis of variance with Tukey’s multiple comparison test determined that the difference in the means (horizontal bars) within each panel (b–d) is statistically significant (*P*<0.05) in bars labelled with a different lower-case letter, but not in those labelled with the same letter.

**Added iron reduces the antimicrobial effects of wrwycr by causing its precipitation**

The enterobactin synthesis and uptake mutants described above are deficient in the cell’s high-affinity iron- (as ferric enterobactin) uptake system. Our initial screening medium, NGCV, had no added iron and the phosphate buffer
in the medium may have precipitated any inherent iron. Indeed, the addition of iron to NGCV improves the growth of the wild-type strain in the absence of wrwycr (data not shown). We thus hypothesized that the hypersensitivity of the enterobactin synthesis, export and uptake mutants to wrwycr may be due to their limited ability to get enough iron through their other, lower-affinity iron-uptake systems (for a review see Guerinot, 1994). To address this hypothesis, we added exogenous iron to a low-phosphate, low-iron growth medium, MGCV, and found that the mutants were better able to withstand wrwycr treatment, shown by shorter lag times in the presence of wrwycr (Supplementary Fig. S3). However, we subsequently found that high concentrations of iron (150 μM and above) precipitate the peptide in the media (Supplementary Table S3), thereby lowering the effective concentration of wrwycr to which cells are exposed (Supplementary Fig. S4). Note that we did not observe any significant growth differences between cultures supplemented with FeSO₄ versus FeCl₃, either at low (6 or 12 μM) or at high (120 μM) final concentration (data not shown).

The ΔtolC mutant, but not the ΔfepB mutant, has an increased level of cell-associated wrwycr relative to the MG1655 parent

Because the entS mutant was not hypersensitive to wrwycr, we hypothesized that the ΔtolC mutant is wrwycr-hypersensitive not because of its involvement in enterobactin export but rather because of its inability to efflux wrwycr. To investigate if the tolC mutant is defective for wrwycr efflux, we measured the levels of cell-associated wrwycr in the ΔtolC mutant and in the parent (MG1655) strains using RP-HPLC (Fig. 2a). ‘Cell-associated wrwycr’ refers to all the wrwycr that could be extracted by formic acid from the cells, be it in the cytoplasm, periplasm, chromosome or the membranes. We found that the ΔtolC mutant does accumulate wrwycr to a higher level than the TolC⁺ MG1655 parent strain (Supplementary Fig. S2b, c), suggesting that the ΔtolC mutant is hypersensitive to wrwycr at least in part due to reduced efflux of wrwycr. In contrast, the ΔfepB mutant’s hypersensitivity is not due to increased cell-associated wrwycr. Interestingly, the ΔtolC mutant also showed an increased level of monomer wrwycr relative to dimerized wrwycr (Fig. 2d), while the ΔfepB mutant has about the same concentration and dimer-to-monomer ratio of cell-associated wrwycr as MG1655 (Fig. 2b–d).

Transcription of Fur-repressed genes is upregulated in wild-type E. coli in response to wrwycr treatment

The hypersensitivity of the Fep system mutants to wrwycr suggested the possibility that wrwycr restricts the availability of iron to the bacteria. To determine the availability of intracellular iron during wrwycr treatment using a biological assay, we monitored the expression level of two genes, entB and fhuF, that are repressed in high-iron conditions by Fur. Compared to the expression level in MG1655 treated with DMSO, the expression level of entB and fhuF in MG1655 treated with 10 μM wrwycr for 1.5 h was increased by 111- and 161-fold, respectively, on average, as measured by qPCR (Fig. 3). This increased level of expression was nearly as high as that seen in a Δfur mutant not treated with wrwycr, in which fhuF and entB expression should be completely derepressed (Fig. 3). This suggests that wrwycr treatment
reduces the amount of bioavailable iron in \textit{E. coli} MG1655 and that the Fur regulon responds accordingly.

We anticipated that \textit{entB} and \textit{fhuF} transcript levels in the \Delta fur strain would appear insensitive to \textit{wrwycr} treatment compared to DMSO treatment. Instead, we did see changes (albeit relatively small, just over threefold), but \textit{entB} and \textit{fhuF} transcript levels responded in opposite directions. For example, while the level of \textit{fhuF} transcripts in \textit{wrwycr}-treated \Delta fur cultures was about half of that in DMSO-treated \Delta fur cultures at 1.5 h, the level of \textit{entB} transcripts was about threefold higher for the same comparison. The \Delta fur strain is itself hypersensitive to \textit{wrwycr} (see below); however, we do not know why \textit{wrwycr}-treated \Delta fur cultures had different levels of \textit{entB} and \textit{fhuF} transcripts from DMSO-treated \Delta fur cultures, or the reason for the differential response of \textit{entB} vs \textit{fhuF} transcripts in the \Delta fur background (see Discussion).

\textbf{Growth in NGCV medium and \textit{wrwycr} treatment decrease catalase activity in cells}

Low bioavailable iron levels lead to a decrease in the cellular concentration of the enzymically active form of non-essential iron-containing enzymes, such as the catalase hydroperoxidase I (KatG or haemoprotein b-590) (Hubbard \textit{et al.}, 1986); thus, catalase activity may indicate iron availability in the cell. Using a \textit{H}_2\textit{O}_2 depletion assay, we quantified the amount of total catalase activity in cell-free extracts of the enterobactin uptake mutant \Delta \textit{fepB} and in the MG1655 parent strain after overnight growth in the screening medium NGCV. Consistent with reduced levels of available iron in the conditions of our screen, we found that extracts of \Delta \textit{fepB} grown in NGCV overnight without \textit{wrwycr} treatment had 29\% of the catalase activity of the parent strain MG1655 in the same conditions (Supplementary Fig. S5a). Treatment of MG1655 in MGCV medium with 10 \mu M \textit{wrwycr} for 1.5 h resulted in only 15\% of the catalase activity of the DMSO-treated samples (Supplementary Fig. S5b).

\textbf{\textit{wrwycr} treatment increases the chelatable iron in the cell}

The pattern of expression of Fur-regulated genes, observed by qPCR, and the decrease in catalase activity suggested that bioavailable iron levels in the cell are reduced by \textit{wrwycr} treatment. We used EPR spectroscopy to measure the amount of desferrioxamine-chelatable iron in \textit{wrwycr}-treated cells. Surprisingly, EPR spectroscopy revealed that levels of chelatable iron in the cell actually increased by six- or sevenfold after 50 min of treatment with 8 or 16 \mu M \textit{wrwycr}, respectively, relative to cells treated with the same volume of DMSO (Table 1). Combined with our other findings, this suggests that while \textit{wrwycr} treatment increases chelatable iron in the cell, this iron is not available to Fur or KatG and that additional iron must be imported to rescue cells from the effects of \textit{wrwycr}.

A possible source of the excess chelatable iron in \textit{wrwycr}-treated cells may be \textit{wrwycr}-related damage to proteins that contain iron–sulfur clusters. Kohanski \textit{et al.} (2007) have suggested that many antibiotics may damage such proteins, causing them to release iron into the cell. We explored this possibility by testing the LacZ levels expressed from the \textit{suf} operon promoter, which encodes proteins involved in iron–sulfur cluster assembly under iron-limiting or oxidative stress conditions (Outten \textit{et al.}, 2004). Indeed we observed dose-dependent induction of \textit{β}-galactosidase activity, which decreased over time at the lower doses of \textit{wrwycr} tested (Fig. 4a). The \textit{suf} promoter is activated independently by apo-IscR and OxyR (Yeo \textit{et al.}, 2006). We found that deletion of \textit{iscR} prevented the rise in \textit{wrwycr}-dependent LacZ activity, whereas deletion of \textit{oxyR} did not affect induction (Fig. 4b). This suggests that iron–sulfur clusters are being damaged and are unavailable to bind to apo-IscR following \textit{wrwycr} treatment.

Free intracellular iron in conjunction with oxygen has been linked to increased DNA damage, leading to higher mutation frequencies (Imlay \textit{et al.}, 1988). To determine if the additional chelatable iron in \textit{wrwycr}-treated cells leads to an increased DNA mutation frequency, we treated cultures with either 8 or 16 \mu M \textit{wrwycr} for 3 or 24 h and measured the mutation frequency compared with that in untreated cultures. To minimize the effect of exogenous iron, cultures were grown in NGCV without any added iron. After 3 h, there was no difference in mutation frequency in treated versus untreated cultures. After 24 h of treatment, the mutation frequency was less than 1.6 \times 10^{-10} cells in untreated cultures, but increased 38-fold to 6.2 \times 10^{-9} cells in cultures treated with 8 \mu M \textit{wrwycr}, and increased almost ninefold to 1.4 \times 10^{-9} cells in cultures treated with 16 \mu M \textit{wrwycr}. The reduced mutation frequency observed after 16 \mu M \textit{wrwycr} treatment compared with 8 \mu M \textit{wrwycr} treatment may be due to lesser fitness of the mutants and/or a reduced replication rate at the higher peptide concentration.

\textbf{\textit{ryhB} overexpression is a cause of the \textit{wrwycr} hypersensitivity of a \Delta fur mutant and the enterobactin uptake mutants}

Because of the role Fur plays in regulation of the enterobactin uptake, we tested the sensitivity of the \Delta fur

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Chelatable intracellular iron (\mu M)*</th>
<th>Fold difference with respect to DMSO-treated cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>DMSO</td>
<td>34.2 ± 5.5</td>
<td>1</td>
</tr>
<tr>
<td>8 \mu M wrwycr</td>
<td>201.7 ± 17.2</td>
<td>5.9</td>
</tr>
<tr>
<td>16 \mu M wrwycr</td>
<td>238.2 ± 31</td>
<td>7</td>
</tr>
</tbody>
</table>

*Mean ± SEM.
mutant to wrwycr and found that it was hypersensitive to 20 μM wrwycr (Fig. 5). When Fur is not bound to iron, as is typical in low-iron media conditions, it does not repress ryhB transcription. In turn, the additional RyhB regulatory RNA molecules produced may decrease the expression of a gene that is important for wrwycr tolerance and/or increase expression of a wrwycr target during growth in the low-iron NGCV and MGCV media. In fact, the hypersensitivity of the Δfur mutant was alleviated by the deletion of ΔryhB (Fig. 5).

To further test the hypothesis that a RyhB-regulated gene is important for wrwycr resistance in the enterobactin-uptake mutants. However, deletion of ryhB in the wild-type strain does not significantly increase its resistance to wrwycr, thus the altered expression of ryhB due to its reduced repression by Fur in the iron-limited conditions following wrwycr treatment is not the sole cause of the sensitivity of the wild-type strain to wrwycr.

**DISCUSSION**

In this work, we showed that *E. coli* mutants defective in enterobactin biosynthesis or ferric enterobactin uptake
Fig. 6. Deletion of ryhB reduces the hypersensitivity to wrwycr of the enterobactin uptake mutants. Open symbols indicate RyhB⁺ strains and closed symbols indicate RyhB⁻ strains. The mean ± SEM OD₆₀₀ of three independent cultures of (a) E. coli MG1655 (G652, □), ΔryhB (EDT1571, ■), ΔfepC (EDT1686, △), ΔfepC ΔryhB (EDT1700, ▲), ΔfepG (EDT1687, ○) and ΔfepG ΔryhB (EDT1701, ●) or (b) E. coli MG1655 (G652, □), ΔryhB (EDT1571, ■), ΔfepB (EDT1343, ◊), ΔfepB ΔryhB (EDT1683, ◆), ΔfepD (EDT1344, ○), ΔfepD ΔryhB (EDT1684, ●), Δfes (EDT1345, △) and Δfes ΔryhB (EDT1685, ▲) measured during growth in MGCV with 20 μM wrwycr at 37 °C. Growth measurements shown in (a) and (b) were recorded simultaneously but are presented in two panels for clarity. All strains grew equally well in DMSO (data not shown).

across the cytoplasmic membrane were hypersensitive to the synthetic antimicrobial peptide wrwycr. We infer from these results that the ability to take up ferric enterobactin or another compound that uses this same transport system is important for E. coli to withstand wrwycr treatment. Mutants lacking FepA or TonB, which are responsible for ferric enterobactin transport across the outer membrane, were not significantly hypersensitive to wrwycr. Our recent analysis of the E. coli proteome found that FepA protein levels decreased upon wrwycr treatment (J. E. Roston and other authors, unpublished data). Since FepA levels in the wild-type strain are reduced by wrwycr treatment, it is not surprising that the absence of FepA in the ΔfepA mutant did not significantly increase sensitivity of E. coli to wrwycr. Ferric enterobactin may cross the outer membrane more easily in wrwycr-treated cells because the peptide increases the permeability of cells, as shown by greater fluorescence of lipophilic probes such as NPN (Jonnalagadda, 2009).

In support of ferric enterobactin limitation causing wrwycr hypersensitivity, we found that mutants lacking either EntB or EntC were also hypersensitive to wrwycr, although less so than some of the uptake mutants (perhaps the reason they were not identified in the original screen). The ΔentC strain was less hypersensitive to wrwycr than was the ΔentB strain, but a ΔmenF ΔentC strain was more sensitive than the ΔentC MenF⁺ strain. This suggests that either the second isochorismate synthase in E. coli, MenF (Daruwala et al., 1996), can make isochorismate for the synthesis of enterobactin in the absence of EntC and/or menaquinone synthesis by MenF is itself important for resistance to wrwycr. Indeed, a ΔmenF EntC⁺ strain was hypersensitive to wrwycr. The literature suggests that MenF cannot compensate for the lack of EntC for enterobactin synthesis unless the corresponding genes are overexpressed in trans, as most of the isochorismate synthesized by MenF and EntC is thought to be channelled to the next enzyme in their respective pathways, MenD or EntB (Buss et al., 2001). Sufficient isochorismate may escape the menaquinone pathway to make dihydroxybenzoate (and therefore enterobactin) in the absence of EntC in the semi-anaerobic conditions of the microtitre plate assays.

E. coli entS and tolC mutants are known to excrete little if any enterobactin but they do excrete monomer, dimer and trimer 2,3-dihydrobenzoylserine molecules (Bleuel et al., 2005; Furrer et al., 2002). These Fes-mediated enterobactin breakdown products are siderophores in their own right (in their iron-bound state they are taken back into the cell by the Fiu and Cir receptors) and thus entS and tolC mutants can achieve comparable growth to that of enterobactin-excreting strains (Hantke, 1990). As such, it is not surprising that the ΔentS mutant tested here was not hypersensitive to wrwycr. To explain the hypersensitivity of the ΔtolC mutant to wrwycr, we propose that TolC is responsible for efflux of wrwycr. Supporting this hypothesis, the ΔtolC mutant had higher levels of cell-associated wrwycr as well as a reduced ratio of dimer to monomer wrwycr compared to MG1655 and the ΔfepB mutant. In the ΔtolC mutant, wrwycr may enter and remain in the cytoplasm long enough for a greater fraction to become reduced to the monomer form. Exactly how wrwycr enters bacteria to achieve the observed steady-state level of cell-associated wrwycr is unknown.

Why is the ability to harvest ferric enterobactin protective against wrwycr in a low-iron medium? We hypothesized that wrwycr treatment reduced bioavailable iron in the cell, necessitating increased iron uptake by the ferric enterobactin
transporter. Supporting this hypothesis, the transcript level of the Fur:Fe\(^{2+}\)-repressed genes entB and fluF increased and catalase activity decreased after 1.5 h wrwycr treatment. These results are consistent with wrwycr treatment reducing the amount of bioavailable iron. In contrast, direct measurement of chelatable iron in the cell indicated that wrwycr treatment actually increased chelatable iron, perhaps by damaging iron–sulfur cluster proteins, as has been proposed for other antibiotics (e.g. Gu & Imlay, 2011; Kohanski et al., 2007). The induction of suf operon promoter activity by wrwycr treatment agrees with this idea, and implicates IscR (and not OxyR) as being necessary for this induction. A possible explanation for the apparent conflict between wrwycr's induction of genes that promote iron uptake and the cells' perceived shortage of iron is that the chelatable iron is not in fact available to bind to Fur or KatG and to rescue wrwycr-treated cells.

The increased iron uptake in the Δfur mutant should logically have provided protection against wrwycr due to derepression of genes encoding iron uptake systems, and thus the Δfur mutant should not have been hypersensitive to wrwycr, in conflict with our findings. However, perhaps the iron concentration of the test medium was not sufficient to counteract wrwycr challenge. Moreover, RyhB levels would have increased in the Δfur mutant, decreasing expression of RyhB-repressed genes and increasing expression of RyhB-activated genes. Deletion of ryhB mostly, but not completely, relieved the wrwycr hypersensitivity of the tested fep and fes mutants, indicating either that a RyhB-repressed function is critical for withstanding wrwycr treatment or that a RyhB-activated function somehow promotes the inhibitory effects of wrwycr. Deletion of the shiA gene, known to be activated by RyhB, does not render either wild-type or ΔfepB cells more sensitive than the ShiA\(^+\) isogenic strains (data not shown). Future studies will be aimed at identifying the RyhB-targeted transcript(s) that affects wrwycr sensitivity. As the altered regulation by RyhB does not fully explain the hypersensitivity of the enterobactin uptake mutants, wrwycr-dependent iron limitation probably plays a role in addition to the altered iron regulation.

A surprising result is the discordant transcription response to wrwycr of entB and fluF in the Δfur strain. One possibility for the observed differential response may be the additional positive regulation of entB by CRP-CAMP (Zhang et al., 2005). It is intriguing that in Salmonella, CRP-activated genes are induced about twofold in a Δfur strain by the presence of the iron chelating agent dipyridyl, similar to our observed induction of entB transcription in the presence of wrwycr (Campoy et al., 2002). The same study showed that the concentration of cAMP was higher in a Δfur mutant compared with the wild-type in iron-limiting conditions in the presence of dipyridyl. We thus note the parallel between the effects of dipyridyl and wrwycr on entB expression (and perhaps, by inference, on cAMP levels) in the Δfur mutant, although in the study by Campoy et al., the medium used was LB rather than the MHB media we used in our own qPCR experiments.

The observed Fur-regulon derepression in response to wrwycr indicates lower levels of bioavailable iron levels, but how does wrwycr treatment elicit this outcome? The most straightforward explanation would be that wrwycr chelates iron, albeit weakly. However, we have been unable to find any evidence for such an activity by using either a chrome azurol S colorimetric assay (Schwyn & Neilands, 1987) or by monitoring the spectrophotometric or fluorometric profile of wrwycr in the presence of iron. We may not have found the appropriate conditions for testing binding of wrwycr to iron. Alternatively, instead of binding iron directly, wrwycr may indirectly sequester iron by binding a siderophore and causing its aggregation, for example, but we have no clues as to the identity of such a molecule. Our results could also formally be explained by wrwycr binding to Fur and KatG directly, competing for iron binding intracellularly (see below). We will test these hypotheses in the future.

It is unclear what the relationship is between the different activities of peptide wrwycr in interfering with DNA repair and accumulating intracellular DNA damage and the iron-restricting activities described here. At least two DNA glycosylases involved in base excision repair, endonuclease III (Nth) and MutY protein, and the DinG helicase, induced by DNA damage, contain [4Fe–4S]\(^{2+}\)-type iron–sulfur centres (Boal et al., 2005; Cunningham et al., 1989; Lukianova & David, 2005; Ren et al., 2009). Impairment of the activity of MutY and Nth by wrwycr due to damage to their iron–sulfur centres may be one explanation for wrwycr-dependent increase in the mutation frequency in iron-limited conditions (NGCV media). Another explanation may be the presence of greater concentrations of intracellular iron after wrwycr treatment. Our best current hypothesis is that the effects of wrwycr on DNA repair (Gunderson et al., 2009) and on iron availability are independent, albeit potentially synergistic.

In summary, the antimicrobial properties of wrwycr are potentiated in low-iron conditions. Enterobactin has been thought not to be an important virulence factor because mammalian neutrophils produce siderocalin (lipocalin), which binds enterobactin in serum (Goetz et al., 2002). However, mammalian antimicrobial strategies include the export of iron from macrophages when they are infected with intracellular pathogens such as Salmonella, Mycobacteria and others (Forbes & Gros, 2001; Nairz et al., 2007). Moreover, several of the effects of wrwycr on iron metabolism described herein, including the accumulation of chelatable but likely unavailable iron in cells, are independent of enterobactin, and thus wrwycr almost certainly puts a strain on all high-affinity iron uptake systems. This could make it or similar molecules particularly effective against pathogens in the low-iron conditions of their hosts.

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Antimicrobial peptide that limits iron bioavailability


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