Reactive metabolic byproducts contribute to antibiotic lethality under anaerobic conditions

Highlights
- Central carbon metabolite concentrations are altered by antibiotic treatment
- Reactive metabolic byproducts accumulate in antibiotic-treated cells
- DNA and membrane damage occur in antibiotic-treated cells
- Reactive metabolic byproduct scavengers protect against antibiotic lethality

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In brief
Wong et al. show that, under anaerobic conditions, reactive metabolic byproducts (RMBs) accumulate in *Escherichia coli* after treatment with bactericidal antibiotics and are associated with macromolecular damage. RMB scavengers protect against antibiotic lethality, suggesting that RMBs contribute to the cell death induced by bactericidal antibiotics under anaerobic conditions.
Reactive metabolic byproducts contribute to antibiotic lethality under anaerobic conditions

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SUMMARY

Understanding how bactericidal antibiotics kill bacteria remains an open question. Previous work has proposed that primary drug-target corruption leads to increased energetic demands, resulting in the generation of reactive metabolic byproducts (RMBs), particularly reactive oxygen species, that contribute to antibiotic-induced cell death. Studies have challenged this hypothesis by pointing to antibiotic lethality under anaerobic conditions. Here, we show that treatment of Escherichia coli with bactericidal antibiotics under anaerobic conditions leads to changes in the intracellular concentrations of central carbon metabolites, as well as the production of RMBs, particularly reactive electrophilic species (RES). We show that antibiotic treatment results in DNA double-strand breaks and membrane damage and demonstrate that antibiotic lethality under anaerobic conditions can be decreased by RMB scavengers, which reduce RES accumulation and mitigate associated macromolecular damage. This work indicates that RMBs, generated in response to antibiotic-induced energetic demands, contribute in part to antibiotic lethality under anaerobic conditions.

INTRODUCTION

Elucidating how bactericidal antibiotics kill bacteria remains a central problem in microbiology (Kohanski et al., 2010a). Although the primary binding targets of antibiotics have been characterized, how bacterial cell death occurs as a result of primary binding interactions remains unclear. We have previously hypothesized that the induction of stress response pathways to alleviate the deleterious consequences of the initial target corruption leads to increased energetic demands that heighten metabolic activity (Kohanski et al., 2007; Lobritz et al., 2015; Stokes et al., 2019). Increased metabolic flux results in the production of reactive metabolic byproducts (RMBs), including but not limited to reactive oxygen species (ROS), that contribute to antibiotic lethality by reacting with and damaging cellular components downstream of antibiotic binding to the primary target (Kohanski et al., 2007; Belenky et al., 2015; Foti et al., 2012). The feedback between damage to macromolecules, increased metabolic flux, and production of RMBs persists until bacterial cell death occurs (Stokes et al., 2019).

This hypothesis has been supported by multiple laboratories using independent lines of evidence (Gusarov et al., 2009; Wang and Zhao, 2009; Davies et al., 2009; Yeom et al., 2010; Shatalin et al., 2011; Nguyen et al., 2011; Luo and Helmann, 2012; Dwyer et al., 2012; Grant et al., 2012; Brynildsen et al., 2013; Moronez-Ramirez et al., 2013; Hong et al., 2019; Drlica and Zhao, 2021). It also has been challenged by other studies based on the argument that antibiotic killing of Escherichia coli (E. coli) in aerobic and anaerobic conditions can be similar (Keren et al., 2013; Liu and Imlay, 2013), and ROS are ostensibly not generated under anaerobic conditions. However, it is important to point out that the above mechanistic model does not assume or purport that ROS are the sole arbiters of antibiotic lethality. Moreover, the induction of stress response pathways as a result of corruption of the primary drug target and its associated cellular processes, as well as the resulting energetic demands that heighten metabolic activity, may not be exclusive to aerobic conditions.

Building on the foregoing points, we hypothesize that, under anaerobic conditions, bactericidal antibiotics corrupt target-specific cellular processes and result in increased energetic...
demands similar to those previously described under aerobic conditions (Kohanski et al., 2007). As a result of increased metabolic flux through anaerobic energy-producing processes (e.g., fermentation and anaerobic respiration with non-oxygen terminal electron acceptors, including nitrate, sulfate, and ferric iron) and additional processes such as free radical interactions, diverse RMBs are produced, and these RMBs react with, and disrupt, cellular components, leading to macromolecular damage that contributes in part to antibiotic lethality. This hypothesis implies that, although specific aspects—including which metabolic pathways experience altered flux and which RMBs are subsequently generated—may vary between anaerobic and aerobic conditions, RMBs contribute in part to antibiotic lethality under both conditions.

Here, we aim to test this hypothesis by determining whether RMBs may contribute to antibiotic lethality under anaerobic conditions. To this end, we used a combination of single-cell assays, bulk-culture measurements, and biochemical perturbations to examine how antibiotics affect the metabolic states of treated cells and how RMBs may contribute to antibiotic lethality under anaerobic conditions. As in previous work (Kohanski et al., 2007; Keren et al., 2013; Liu and Imlay, 2013), we focus here on three main classes of bactericidal antibiotics: aminoglycosides, which bind to the 30S ribosomal subunit and induce protein mislocalization; fluoroquinolones, which bind to DNA gyrase or topoisomerase to perturb DNA replication and transcription; and β-lactams, which bind penicillin-binding proteins to disrupt peptidoglycan cell wall biosynthesis. We show that treatment of E. coli with these three classes of antibiotics under anaerobic conditions leads to alterations in the cellular concentrations of central carbon metabolites. By employing single-cell microscopy and specific gas chromatography-mass spectrometry (GC-MS) experiments, we also find that RMBs, particularly reactive electrophilic species (RES), accumulate in antibiotic-treated E. coli under anaerobic conditions. At the single-cell level, we show that antibiotic treatment results in DNA double-strand breaks (DSBs) and membrane damage, consistent with the hypothesis that macromolecular damage can be induced in part by RES. Importantly, we demonstrate that antibiotic lethality is decreased by three different RMB scavengers, which reduce RES accumulation and mitigate the macromolecular damage induced by antibiotics under anaerobic conditions. This work indicates that under anaerobic conditions, antibiotic-induced changes in cellular metabolism result in the production of RMBs that react with cellular components, leading to macromolecular damage that contributes in part to antibiotic lethality.

RESULTS

Antibiotic lethality occurs under anaerobic conditions

To determine the extent to which antibiotic killing occurs under anaerobic conditions, we treated bulk cultures of log phase E. coli with the aminoglycoside kanamycin, the fluoroquinolone ciprofloxacin, and the β-lactam mecillinam at concentrations ranging from 0.1× to 50× the anaerobic minimum inhibitory concentrations (MICs; Table S1). Cells were grown, treated, plated, and incubated under anaerobic conditions in LB medium containing resorufin, an oxygen-sensitive dye that is colorless below a redox potential of ~110 mV and used here to validate that environmental oxygen is largely absent. The addition of resorufin is important, as an anaerobic environment by itself can be insufﬁcient to deprive culture media of oxygen. Under these conditions, time-kill assays and colony-forming unit (CFU) quantitation of antibiotic-treated cultures revealed that killing occurred for all three antibiotics, leading to between 0 and ~6 logs of decreased survival, depending on antibiotic class, antibiotic concentration, and treatment time (Figure 1A). We observed similar levels of decreased survival under aerobic conditions, with the exception of treatment with mecillinam, for which at most ~1 log of decreased survival occurred (Figure S1). These ﬁndings indicate the extents to which antibiotic killing occurs in LB medium under anaerobic and aerobic conditions, which are similar to the ﬁndings of previous studies (Keren et al., 2013; Liu and Imlay, 2013).

Antibiotic treatment results in altered central carbon metabolite concentrations under anaerobic conditions

As antibiotic lethality occurs under anaerobic conditions, we hypothesized that, under such conditions, bactericidal antibiotics corrupt target-specific cellular processes and result in increased energetic demands similar to those previously described under aerobic conditions (Kohanski et al., 2007). Under anaerobic conditions, E. coli can generate ATP through glycolysis, which converts glucose into pyruvate, resulting in the generation of ATP and NADH, and anaerobic respiration with non-oxygen terminal electron acceptors, which oxidizes NADH to replenish NAD⁺ (Figure 1B). In order to determine whether intracellular concentrations of central carbon metabolites are altered after antibiotic treatment, we measured glucose, pyruvate, and NADH/NAD⁺ concentrations in antibiotic-treated cells using enzymatic assays. First, we found that endogenous concentrations of glucose were increased in log phase, antibiotic-treated cell cultures relative to untreated controls (Figure 1C). Cells treated with kanamycin, ciprofloxacin, and mecillinam across a range of low (1× MIC) and high (5× to 50× MIC) concentrations exhibited average glucose concentrations that were as large as 4-fold that of untreated cells. Intriguingly, cells treated with mecillinam at a high (50× MIC; 10 μg/mL) concentration exhibited lower glucose concentrations, compared with treatment at a low mecillinam concentration; why this occurs is unclear, but it is possible that extensive primary target corruption at high mecillinam concentrations may dominate the cellular response to the antibiotic under these conditions.

Building on our measurements of glucose concentrations, we next measured pyruvate and NADH/NAD⁺ concentrations. As pyruvate is an end-product of glycolysis, pyruvate accumulation may indicate increased glycolytic flux (Zhu et al., 2008). Additionally, alterations to NADH/NAD⁺ concentration ratios may lead to redox imbalance and increased RMB production (Kohanski et al., 2007). We found increased concentrations of pyruvate similar to glucose (Figure 1D): average pyruvate levels in antibiotic-treated cells were between ~1.5- to ~4-fold that of untreated cells, and treatment with mecillinam at a high concentration resulted in lower pyruvate accumulation, compared with treatment at a low concentration (similar to what was found for glucose). Measuring NADH/NAD⁺ concentration ratios, we found that average NADH/NAD⁺ ratios in antibiotic-treated cells
were between ~0.5- to ~1.2-fold that of untreated cells (Figure 1E). Treatment with kanamycin at a high (5x MIC; 250 μg/mL) concentration and mecillinam at a low (1x MIC; 0.2 μg/mL) concentration increased average NADH/NAD⁺ concentrations relative to non-treatment conditions, whereas average NADH/NAD⁺ concentrations were decreased for all other treatment conditions. These measurements suggest that, under different treatment conditions, NADH might be differentially altered by processes such as anaerobic respiration and fermentation on the timescale of interest. In general, NADH/NAD⁺ concentrations changed to a lesser extent than those of glucose and pyruvate, which may reflect better NADH/NAD⁺ homeostasis (Figure 1B).

Taken together, these results indicate largely altered glucose, pyruvate, and/or NADH/NAD⁺ concentrations in antibiotic-treated E. coli relative to untreated controls. These findings suggest that antibiotic treatment may result in alterations to cellular metabolism, wherein increased ATP demand as a result of primary target corruption leads to increased central carbon metabolism through catabolic processes including glycolysis. Importantly, increased metabolism may result in the formation of RMBs, including RMBs that arise as a consequence of the oxidation of NADH to NAD⁺ and the reduction of terminal electron acceptors (Figure 1B).

Phenotypic changes and fluorescence of reactive metabolic byproduct-sensitive dyes occur under anaerobic conditions

As a result of increased metabolism in antibiotic-treated cells, we hypothesized that the production of RMBs is associated with antibiotic lethality. To address this, we characterized cellular phenotypes at the single-cell level using fluorescent dyes that...
are sensitive to RMBs and RMB-mediated cellular damage. We have recently shown that cytoplasmic condensation, a phenotype in which discrete portions of the cytoplasm become phase light when imaged under phase contrast, is associated with cell death in a fraction of cells treated with aminoglycosides or fluoroquinolones (Wong et al., 2021a). By contrast, β-lactams induce well-studied phenotypes of membrane bulging and lysis (Yao et al., 2012; Wong and Amir, 2019; Wong et al., 2021b). Under aerobic conditions, cytoplasmic condensation and/or cellular lysis are associated with membrane damage, cessation of growth at the single-cell level, and the accumulation of ROS, nitric oxide, and lipid peroxidation adducts, as measured by the fluorescent dyes carboxy-H2DCFDA, DAF-FM, and C11-BODIPY, respectively (Wong et al., 2021a). We have previously observed that kanamycin- or ciprofloxacin-treated cells irreversibly cease to elongate as soon as they exhibit cytoplasmic condensation (Wong et al., 2021a). Furthermore, the fraction of cells exhibiting cytoplasmic condensation in antibiotic-treated populations is reduced in the presence of glutathione, an antioxidant that also decreases antibiotic lethality (Wong et al., 2021a). Here, under anaerobic conditions, we found that treatment with kanamycin or ciprofloxacin induced cytoplasmic condensation after 4 h of treatment, a timescale corresponding to decreases of at least 4 logs in survival in bulk culture (Figures 1A, 2A, and 2B). Moreover, treatment with all three antibiotics (kanamycin, ciprofloxacin, and mecillinam) induced significant, 

Figure 2. Phenotypic change and fluorescence of reactive metabolic byproduct-sensitive dyes occur under anaerobic conditions.

(A) Cellular phenotypic changes induced by antibiotic treatment at bactericidal concentrations corresponding to 1 × to 50 × the anaerobic MICs (Table S1). Concentrations used were as follows: kanamycin, 50 μg/mL; ciprofloxacin, 1 μg/mL; mecillinam, 10 μg/mL. Images are shown in phase contrast, and colored arrows highlight specific phenotypes (bottom of figure). Cells were imaged 4 h after antibiotic treatment, and images are representative of three different fields of view from two biological replicates. Scale bars, 3 μm.

(B) Census of single-cell phenotypes in populations treated with antibiotics at the concentrations indicated in (A) after 4 h. Data are from four fields of view from four biological replicates with at least 150 cells per group. Colors refer to the legend at the bottom of the figure. Data are presented as mean values ± SEM, and error bars indicate one standard deviation.

(C) Summary of a panel of RMB-sensitive dyes and their primary reactive species targets.

(D–F) Fluorescence intensities of RMB-sensitive dyes in single cells after treatment at the concentrations indicated in (A), which exhibit different posttreatment phenotypes, as determined by epifluorescence microscopy after 4 h of treatment. Values indicate fluorescence intensities relative to background levels. Cells were cultured, treated, and imaged under anaerobic conditions. White points indicate mean values, and each distribution represents at least 10 cells. Two-sample Kolmogorov-Smirnov tests for differences from untreated, turgid cells: NS, not significant; *p < 10−2, **p < 10−4, and ***p < 10−10. Columns with no data, indicating that the specified phenotypes were not observed, are not shown.
phenotype-dependent increases in fluorescence of carboxy-
H$_2$DCFDA, DAF-FM, and C11-BODIPY in most cells, with typical
fluorescence increases being associated with condensation and/or lysis in kanamycin- and ciprofloxacin-treated cells and
with membrane bulging and/or lysis in mecillinam-treated cells (Figures 2C–2F). More limited fluorescence increases, especially
of C11-BODIPY, occurred as soon as 30 min after antibiotic
treatment and well before cells exhibited cytoplasmic condensa-
tion, membrane bulging, and lysis, suggesting that the genera-
tion of certain RMBs precedes cellular phenotypic changes (Figure S2). Importantly, these fluorescence increases did not
arise from oxygen contamination, as resorufin remained strictly colorless throughout our experiments. Taken together, these
single-cell assays suggest that RMBs accumulate in antibiotic-
treated cells under anaerobic conditions.

Antibiotics induce the production of reactive
electrophilic species under anaerobic conditions
In addition to arising from molecular oxygen, RMBs, including
ROS and non-ROS free radicals, may arise under anaerobic condi-
tions from endogenous sources, including NADPH oxidases
(Hajjar et al., 2017) and reactive nitrogen species (RNS) syn-
thases (Crane et al., 2010); from anaerobic respiration with termi-
nal electron acceptors, including nitrate, sulfate, and ferric iron;
and from glycation reactions involving amino acids (Yin et al.,
1995). Our observations of cytoplasmic condensation (indicative
of membrane damage) and lipid peroxidation (as assayed by
C11-BODIPY) suggest that RMBs might directly react with mem-
brane lipids, a process known to produce highly promiscuous and
deleterious RES (Yin et al., 2011). Therefore, we focused on
RES, which may be produced anaerobically and aerobically by processes such as glycolysis (Ferguson et al., 1999) and lipid
peroxidation in bacteria (Yin et al., 2011). RES are biomarkers of
RNS-induced, ROS-induced, and other free radical-induced damage, and RES also stimulate the production of other reactive byproducts, including advanced glycation end-products (AGE) and advanced lipoxidation end-products (ALE), through reac-
tions with nucleic acids, proteins, and lipids (STAR Methods).
Examples of RES include 4-hydroxynonenal (4-HNE), a byproduct of lipid peroxidation, and methylglyoxal (MGO), a byproduct of
glycolysis and lipid peroxidation. These aldehydes bind cellular
components, contribute to mutagenesis, and in the case of
MGO, accumulate in millimolar quantities within cells during unbalanced sugar metabolism (Ferguson et al., 1999). Under
anaerobic conditions, we have previously shown that bactericidal
antibiotics elevate central carbon metabolism (Yang et al.,
2019), and a recent study has suggested that glycolysis contrib-
utes to β-lactam killing of Gram-positive bacteria (Kawai et al.,
2019). Consistent with these and other works (Dwyer et al.,
2014; Wong et al., 2021a), we hypothesized that bactericidal an-
tibiotics kill bacteria under anaerobic conditions in part through the
generation of RES and the subsequent reactions of RES with cellular components.

To investigate this hypothesis in anaerobic bulk culture, we measured RES concentrations in log phase cells treated with antibi-
totics. Using GC-MS, we measured 4-HNE and MGO concen-
trations after derivatization at the time of harvest (Luo et al.,
1995) (Figure S3). Cells were harvested after treatment with antibiotics
across a range of concentrations, from 1 x to 50 x the anaerobic
MICs, at an antibiotic treatment endpoint of 2 h. We found that
4-HNE and MGO were present in lysates of cells treated with
kanamycin, ciprofloxacin, or mecillinam under anaerobic condi-
tions (Figures 3A and 3B). Furthermore, our measurements sug-
gested that increased antibiotic concentrations resulted in
increased average concentrations of 4-HNE and MGO for kana-
mycin and mecillinam. Ciprofloxacin treatment at higher concen-
trations (5 x MIC; 0.1 μg/mL) did not induce elevated levels of
either RES after 2 h relative to treatment at lower concentrations.
It is possible that extensive primary target damage and/or simul-
taneous inhibition of protein synthesis may affect the production of
RMBs, including RES, at high ciprofloxacin concentrations, as
previously suggested (Drlica and Zhao, 2021). Nevertheless, for
all other treatment groups, these measurements indicate that
RES are present and may accumulate in antibiotic-treated cells
under anaerobic conditions (Figures 3A and 3B). Additionally,
increased 4-HNE (but not MGO) concentrations were detectable
in cell lysates 30 min after antibiotic treatment (Figure S3), sug-
gesting that the generation of certain RES may occur rapidly af-
ter antibiotic treatment.

Next, to understand the physiological effects of RES on cells, we measured cellular survival after exogenous treatment with
millimolar concentrations of 4-HNE and MGO (Figure S4). We
found that addition of either 4-HNE or MGO led to between 0 to ~6 logs of decreased survival in log phase cells, depending
on concentration and treatment time, and lethality occurred un-
der both anaerobic and aerobic conditions (Figures S4 and S5).
Strikingly, consistent with our phenotypic observations for kana-
mycin and ciprofloxacin treatments (Figures 2A and 2B), single-
cell observations reveal that bactericidal concentrations of
4-HNE or MGO also induce cytoplasmic condensation and
cellular lysis (Figure S4). Together, these observations qualita-
tively indicate that, under anaerobic conditions, RES are dele-
tious to bacterial cells and could contribute in part to the cell
death phenotypes observed after antibiotic treatment.

Antibiotic-treated cells display macromolecular
damage consistent with damage induced by reactive
electrophilic species under anaerobic conditions
Previous studies have shown that, under aerobic (Belenky et al.,
2015; Foti et al., 2012; Dwyer et al., 2012, 2014; Hong et al.,
2019; Wong et al., 2021a) and anaerobic (Wong et al., 2021a;
Giroux et al., 2017) conditions, antibiotics induce macromolec-
ular damage distinct from the damage induced by their primary
drug-target binding activity. Here, we asked whether macromo-
lecular damage also occurs under anaerobic conditions, and if
so, whether it is consistent with our finding that antibiotic-treated
cells generate RES. We focused specifically on two types of
damage, DNA DSBs and membrane damage. We note that
kanamycin and mecillinam can damage cellular membranes
through non-specific ionic interactions (Martin and Beveridge,
1986) and primary target binding to penicillin-binding proteins
(Yao et al., 2012), respectively, whereas ciprofloxacin damages
DNA by binding to its primary targets of DNA gyrase and
Topoisomerase. Nevertheless, RES may further contribute to
DNA and membrane damage in antibiotic-treated cells. We
measured the frequencies of DSBs and membrane damage in
antibiotic-treated cells using a fluorescent protein-based probe (GamGFP; Shee et al., 2013) and SYTOX Blue, a membrane damage-sensitive dye, respectively.

Using a GamGFP strain of E. coli, we found that cells exhibited DSBs—as manifested by fluorescent GFP foci—after treatment with kanamycin, ciprofloxacin, or mecillinam under anaerobic conditions (Figures 3C and 3D). Although GFP fluorescence is typically quenched under anaerobic conditions, our observations of GamGFP foci suggest that under the anaerobic conditions used here, there remains enough oxygen for fluorophore maturation. This suggestion is supported by previous work that has shown GFP fluorophore maturation even under conditions of 0.1 PPM dissolved oxygen (Hansen et al., 2001), a level below the higher limit of atmospheric oxygen (5 PPM) used in our setup. Here, we found that the fraction of cells exhibiting at least one fluorescent GFP foci after 2 h of treatment depends on antibiotic class and concentration, and this fraction varies in a dose-dependent manner (Figure 3D). As expected from the primary binding target, higher (0.1 μg/mL) doses of ciprofloxacin induced the largest increase in the fraction of positive cells, and nearly all cells exhibited DSBs; additionally, increases of ~10% to 40% in the fraction of positive cells were found in cells treated with kanamycin and high concentrations of mecillinam (50x MIC; 10 μg/mL). Low doses of mecillinam, near the respective MIC, did not induce substantially higher levels of DNA damage, compared with non-treatment, on the timescale of our assay.

We next treated E. coli MG1655 with antibiotics in the presence of SYTOX Blue under anaerobic conditions (Figures 3E and 3F). SYTOX Blue is a nucleic acid stain that only penetrates cells with damaged membranes and is therefore a specific marker of membrane damage. Here, cells with compromised membranes are indicated by a largely uniform increase in fluorescence in the cellular cytoplasm. Similarly to GamGFP cells, the fractions of positive cells after 4 h of treatment were increased in a dose-dependent manner for all three antibiotics (kanamycin, ciprofloxacin, and mecillinam). Altogether, for treatment with all three antibiotics, the fractions of DSB- and membrane damage-positive cells were associated with relative levels of RES accumulation (Figures 3A and 3B), with the exception of high concentrations of ciprofloxacin, for which we measured lower levels of RES but for which more cells exhibited DNA and membrane damage. Although this may be expected for DNA damage due to ciprofloxacin’s primary binding targets, the observation for membrane damage suggests that ciprofloxacin-treated cells may be more susceptible to membrane damage in a manner that is not directly proportional to RES accumulation.

The foregoing results suggest that, under anaerobic conditions, antibiotics induce DNA damage different from the damage directly induced by primary target binding for treatment with kanamycin and mecillinam and membrane damage different from the damage directly induced by primary target binding for treatment with ciprofloxacin. It is important to note that our findings suggest that DNA damage is associated with—and may not necessarily cause—antibiotic lethality. Yet, DNA damage could induce the SOS response, which may result in membrane alterations through the activities of repair proteins such as RecA (Garvey et al., 1985). Additionally, these forms of damage may be also consistent with the antibiotic-induced accumulation of
Figure 4. Reactive metabolic byproduct scavengers protect against antibiotic lethality under anaerobic conditions, which is not explained by alterations to cellular growth rate, cellular metabolism, proton motive force, and intracellular antibiotic concentration

(A) RMB scavengers and their primary scavenging targets. Examples of reactive species targets are indicated in parentheses.

(B) Survival curves of *E. coli* MG1655 after kanamycin, ciprofloxacin, and mecillinam treatment at various concentrations, with and without exogenous supplementation of RMB scavengers (10 mM), as determined by plating and CFU counting. Cells were cultured, treated, and plated in LB under anaerobic conditions. Error bars indicate one standard deviation, and each point is representative of two biological replicates. Data are presented as mean values ± SEM; where SEM is small, error bars are present but are inside symbols. Where applicable, CFU/mL values less than 10^2 were truncated to a value of 10^2 to reflect the lower limit of quantification.

(C) OD_{600} measurements for anaerobic cell cultures supplemented with RMB scavengers (10 mM) or grown in dilute LB. Error bars indicate one standard deviation, and data are presented as mean values ± SEM. Where SEM is small, error bars are present but are inside symbols. Data from two biological replicates in each condition are shown.

(D) Ratios of ATP luminescence values to OD_{600} values, relative to cultures grown in LB without scavengers. Treated cells were grown, without antibiotics, anaerobically with RMB scavengers for 2 h, then harvested. Data from two biological replicates are shown (black points), and bars indicate average values.

(E) Single-cell measurements of DiBAC_4(3), ACMA, and DiSC_3(5) fluorescence, which respectively detect changes in the ΔΨ and ΔpH components of PMF and changes in membrane permeability. Data from cells treated with 1 mM carbonyl cyanide m-chlorophenylhydrazone (CCCP), a PMF uncoupler, are shown as a
RES and other RMBs. Indeed, 4-HNE and MGO participate in a variety of cellular reactions, including reactions with DNA to form adducts and generation of additional free radicals that may result in lipid peroxidation (STAR Methods). Treating antibiotic-free cells exogenously with RES, we found that both 4-HNE and MGO induced DSBs and membrane damage in a dose-dependent manner and resulted in a wide range of fractions of GamGFP- or SYTOX Blue-positive cells, from 20% to 90%, that are similar to the fractions of positive antibiotic-treated cells at high concentrations (Figure S4). These findings indicate that the generation of 4-HNE and MGO in antibiotic-treated cells may contribute in part to the DNA or membrane damage induced by antibiotic treatment. Importantly, we note that not all antibiotic-treated cells are GamGFP- or SYTOX Blue-positive, suggesting that additional forms of cellular damage may be needed to fully explain antibiotic lethality.

Chemical scavengers of reactive metabolic byproducts decrease antibiotic lethality under anaerobic conditions

As RMBs are produced in cells treated with antibiotics under anaerobic conditions, and RMBs, particularly RES, induce macromolecular damage consistent with that induced by antibiotics, we asked whether RMBs might contribute to antibiotic lethality. To address this question, we tested whether perturbations that rescue cells from RMBs also rescue cells from antibiotics under anaerobic conditions. We considered a panel of three chemical scavengers—glutathione, acetylcysteine, and pyridoxamine—which are known to detoxify various RMBs (Figure 4A; STAR Methods), and probed their effects on antibiotic killing under anaerobic conditions. Prior studies indicate that the non-enzymatic reactivity of glutathione or acetylcysteine with H₂O₂ is limited (Imlay, 2015; Winterbourn and Metdiewia, 1999; Zhitkovich, 2019), suggesting that their scavenging activities may predominantly target RES or RNS. Second-order rate constants have been measured to be ~0.1 to 1 M⁻¹ s⁻¹ for the non-enzymatic reaction of glutathione or acetylcysteine with H₂O₂ (Winterbourn and Metdiewia, 1999; Deponte, 2017), 1 to 10 M⁻¹ s⁻¹ for glutathione with a variety of electrophiles (Chen et al., 2008), and ~10⁸ M⁻¹ s⁻¹ for glutathione with nitric oxide (Deponte, 2017). Thus, application of these scavengers may support the general involvement of RMBs in antibiotic lethality, rather than the specific contribution of H₂O₂ and ROS alone, as previously suggested (Imlay, 2015).

We first focused on glutathione, which has been shown to increase the MICs of kanamycin and ciprofloxacin and to attenuate antibiotic lethality under aerobic conditions (Wong et al., 2021a; Lopatkin et al., 2019) (Figure S1). To test whether these observations hold under anaerobic conditions, we performed MIC and time-kill assays with exogenous supplementation of glutathione at a concentration of 10 mM, identical to that used in previous aerobic measurements (Wong et al., 2021a). We found that addition of glutathione resulted in a 2-fold increase in the anaerobic kanamycin MIC, an 8-fold increase in the anaerobic ciprofloxacin MIC, and no increase in the anaerobic mecillinam MIC (Table S1). Substantial (>2-fold) increases in MIC were specific to ciprofloxacin, as limited MIC changes were observed in cells treated with the bacteriostatic antibiotics rifampicin, chloramphenicol, and tetracycline (Table S1). Additionally, consistent with these MIC values, CFU quantitation revealed decreased antibiotic killing of cells in the presence of glutathione (Figures 4B and S6). After 4 h of treatment, anaerobic killing by kanamycin and ciprofloxacin was reduced by as much as 5 logs, and anaerobic killing by mecillinam was reduced by ~1 to 2 logs across a range of antibiotic concentrations (Figures 4B and S6). Aside from unsubstancial killing (≤1 log of decreased survival) observed during mecillinam treatment under aerobic conditions, the observed protection from antibiotic killing was largely similar to that under aerobic conditions (Figures 4B, S1, and S6). Moreover, while differences in MIC contribute to decreased lethality for kanamycin and ciprofloxacin, the observed increases in MICs were typically smaller than the shifts in concentration shown in the time-kill plots (Figure 4B; Table S1). This observation suggests that glutathione protects cells from kanamycin and ciprofloxacin killing only in part by increasing the MICs. In kanamycin- and ciprofloxacin-treated cells, glutathione protection was reflected at the single-cell level, where the frequency of cytoplasmic condensation was decreased (Figure S2). The frequency of membrane bulging in mecillinam-treated cells was not significantly changed by glutathione supplementation, suggesting that, in mecillinam-treated cells, glutathione protection occurs independently from membrane bulging. Moreover, we found that glutathione decreases antibiotic lethality irrespective of the availability of environmental oxygen and the known effects of glutathione on cellular efflux and detoxification processes (Figures S7–S10; Tables S2–S4; STAR Methods).

We next considered two other chemical scavengers of RMBs, acetylcysteine and pyridoxamine (Figure 4A). Acetylcysteine, a glutathione precursor, is believed to be a poor scavenger of ROS but a potent scavenger of RES, with which it forms less-reactive Michael adducts (Zhikoviev, 2019; Negre-Salvayre et al., 2008), and it can also serve as a precursor to H₂S (Shatalin et al., 2011; Mironov et al., 2017) whose anionic form (HS⁻) reacts with RES via direct sulfhydration (Nishida et al., 2012). Pyridoxamine is known to primarily scavenge RES and prevents the formation of other toxic species, including AGE and ALE, and may additionally scavenge ROS (Voziyan and Hudson, 2005; Amarnath et al., 2004). We found that acetylcysteine and positive control for DiBAC₄(3) and ACMA, and data from cells treated with 100 µg/ml valinomycin, an ionophore, are shown as a positive control for DiSC₃(5). Data are presented as mean values ± SEM, and error bars indicate 95% confidence intervals for the mean. Data representative of 20 cells in each group. Two-sample t tests for differences in mean values from LB only: ***p < 10⁻⁵, other bars not significant.
(F) Intracellular antibiotic concentration measurements. Cells were treated with kanamycin-Texas Red (50 µg/ml), ciprofloxacin (1 µg/ml), or mecillinam (10 µg/ml) for 1–4 h under anaerobic conditions, and intracellular antibiotic concentrations were assayed fluorometrically (kanamycin) or using LC-MS (ciprofloxacin and mecillinam). Single-cell counts for kanamycin-treated cells are indicated in parentheses, and 95% confidence intervals for the mean are shown. Two-sample t tests for differences in mean values of cells with no scavenger: **p < 10⁻³, ***p < 10⁻⁵, other bars not significant. For ciprofloxacin and mecillinam, data from two biological replicates in bulk culture are shown (black points), and bars indicate average values. Dashed lines indicate baseline values.
pyridoxamine, at concentrations (10 mM) comparable to glutathione supplementation, increased the anaerobic kanamycin and ciprofloxacin MICs similarly to glutathione, by 2- and 8-fold, respectively, and did not affect the mecillinam MIC (Table S1). Additionally, and consistent with these MIC values, we found that supplementation of acetylcycteine and pyridoxamine attenuated antibiotic killing both anaerobically and aerobically (Figures 4B, S1 and S6). Similarly to the observed protection against antibiotics for glutathione, we found that, at the single-cell level, supplementation of acetylcycteine and pyridoxamine reduced the frequency of cytoplasmic condensation in kanamycin- and ciprofloxacin-treated cells, but not the frequency of membrane bulging in mecillinam-treated cells (Figure S2).

As differences in cellular growth, metabolism, and intracellular concentration of antibiotics may affect antibiotic lethality, we performed experiments measuring these variables. As detailed further in the STAR Methods, measurements of bacterial growth (Figure 4C) and metabolic (Figures 4D and S11) rates, in addition to antibiotic killing in dilute LB (Figures S12 and S13), indicate that scavenger protection does not arise from suppressive alterations to bacterial growth or metabolic state. Furthermore, dye-based measurements of the proton motive force (PMF; Figure 4E) and fluorescence-based and liquid chromatography-mass spectrometry (LC-MS)-based measurements of intracellular antibiotic concentrations (Figure 4F) indicate that scavenger protection is not explained by differences in PMF or intracellular antibiotic concentration.

Scavenger protection is associated with decreased accumulation of reactive electrophilic species and reduced macromolecular damage under anaerobic conditions

To test the hypothesis that RMB scavengers protect cells from antibiotic action in part by scavenging RES, we directly measured the effects of scavenger supplementation on RES levels using GC-MS. We treated cells with antibiotics at approximately the same multiples of their corresponding MIC values in the presence and absence of glutathione, acetylcycteine, or pyridoxamine (10 mM) to account for MIC changes induced by scavengers (Table S1). In cells treated with similar concentrations of antibiotics relative to multiples of the respective MICs, we found that 4-HNE and MGO concentrations after 2 h of treatment were largely decreased in the presence of scavengers (Figure 5A). The only exception was for kanamycin-treated cells in the presence of glutathione, which exhibited average MGO values larger than those of kanamycin-treated cells without scavenger. Notably, RES levels were decreased >2-fold under nearly all conditions with acetylcycteine and pyridoxamine, and similar decreases were observed for the application of scavengers to positive controls, in which cells were treated with exogenous MGO (Figure S14). Additionally, we found that these decreases in RES were often associated with decreases in DNA and/or membrane damage (Figure 5B) and reflected decreases in antibiotic lethality (Figure 5C). Indeed, scavengers also mitigated DNA and membrane damage induced by exogenous RES and rescued cells from RES lethality when RES were administered exogenously (Figures S14–S16). We note that these scavenger-mediated decreases in RES levels—along with cellular responses such as filamentation and the induction of the SOS response—may, in addition to decreasing antibiotic lethality (Figure 4C), also contribute to the observed increases in antibiotic MICs (Table S1). Together, these findings support the hypothesis that glutathione, acetylcycteine, and pyridoxamine decrease antibiotic lethality in part by reducing RES levels. As antibiotics are still lethal to scavenger-treated cells for which reduced RES levels were measured after 2 h of treatment, our findings further highlight the possibilities that substantial increases in RMB levels may occur post-plating (Hong et al., 2019) and that, in addition to RES, other RMBs or cellular pathways affected by primary target binding may also contribute to antibiotic lethality.

DISCUSSION

Understanding how bactericidal antibiotics kill bacteria remains an unresolved problem. Notwithstanding that the field has classified primary antibiotic binding targets, how such binding leads to bacterial cell death remains unclear. We have previously proposed that, downstream of primary target binding, the induction of stress response pathways in response to corrupted cellular processes leads to energetic demands that heighten metabolic activity (Kohanski et al., 2007; Lobritz et al., 2015; Stokes et al., 2019). Increased metabolism results in the accumulation of RMBs, which react with, and damage, cellular macromolecules, contributing to antibiotic lethality. This hypothesis was initially supported by our observations that treatment with bactericidal, but not bacteriostatic, antibiotics under aerobic conditions increased hydroxyl radical formation via the Fenton reaction; that the iron chelator 2,2’-dipyridyl and the hydroxyl radical scavenger thiourea reduce antibiotic lethality; and that genetic perturbations to metabolic pathways—for instance, deletion of the E. coli iscS gene, which decreases iron-sulfur cluster abundance—affect antibiotic lethality (Kohanski et al., 2007). Subsequent studies, from multiple laboratories using independent lines of evidence, have supported this hypothesis. Downstream of treatment with bactericidal antibiotics, these studies have measured increases in energetic demands, cellular respiration rate, and cellular metabolism (Lobritz et al., 2015; Dwyer et al., 2014; Mathieu et al., 2016; Yang et al., 2019); investigated associated processes such as the stringent response, which activates when protein synthesis is increased and amino acid pools are depleted, and in turn increases ROS production (Nguyen et al., 2011; Khakimova et al., 2013; Mathieu et al., 2016); studied redox-related alterations to bacterial cell physiology (Dwyer et al., 2007, 2012, 2014; Bilenky et al., 2015; Hong et al., 2019; Wang and Zhao, 2009; Wong et al., 2021a) and cellular mutagenesis (Kohanski et al., 2010b; Gutierrez et al., 2013; Pribis et al., 2019); and demonstrated that RMBs—including ROS—and related metabolism-associated changes impact antibiotic lethality (Brynioldsen et al., 2013; Hong et al., 2019; Gusarov et al., 2009; Dwyer et al., 2014; Wong et al., 2021a; Lopatkin et al., 2019; Goswami et al., 2016; Giroux et al., 2017). This common mechanism, as a working hypothesis for how bactericidal antibiotics in part kill bacteria...
has generated testable predictions and motivated potential strategies and targets for antimicrobial therapies.

In the present study, we have shown that bactericidal antibiotic treatment of E. coli under anaerobic conditions is associated with alterations in the concentrations of central carbon metabolites (Figure 1) and that RMB-sensitive dyes, including those sensitive to lipid peroxidation, RNS, and ROS, fluoresce in antibiotic-treated cells (Figure 2). Our observations of cytoplasmic condensation—indicative of membrane damage and cell death (Wong et al., 2021a)—and lipid peroxidation in antibiotic-treated cells suggest that RMBs might directly react with membrane lipids, producing RES. Using GC-MS, we measured endogenous concentrations of two RES, 4-HNE and MGO, in antibiotic-treated cells and found that they are present under anaerobic conditions (Figure 3). We also showed that antibiotic treatment under anaerobic conditions induces DNA and membrane damage, consistent with possible contributions of RES damage to these macromolecules (Figure 3). Further work should address whether these or other forms of damage are direct causes of antibiotic lethality. Importantly, three diverse RMB scavengers—glutathione, acetylcysteine, and pyridoxamine—reduce antibiotic lethality without decreasing cellular growth rate, cellular ATP levels, PMF, and intracellular antibiotic concentration (Figure 4).

Based on these results, we propose that RMBs, particularly RES, contribute to antibiotic-induced cell death under anaerobic conditions according to the following mechanism (Figure 6). Upon binding of the primary drug target, antibiotics corrupt target-specific cellular processes, leading to increases in energetic demands and metabolic fluxes. RMBs, including but not limited to RES, RNS, and ROS, are generated by increased metabolic flux through anaerobic energy-generating processes and additional processes such as free radical interactions under anaerobic conditions. These RMBs react with, and damage, cellular components including nucleic acids, proteins, and lipids, contributing in part to antibiotic lethality.

The hypothesis that RMBs contribute to antibiotic-induced bacterial cell death has been a model that has evolved and expanded, and it should continue to be tested by additional experiments and analyses. Considerable evidence, from both the present study and others, is consistent with a contribution of RMBs, including RES, RNS, and ROS, to antibiotic-induced cell death under anaerobic and aerobic conditions (Kohanski et al., 2007; Dwyer et al., 2007, 2012, 2014; Gusarov et al.,...
Bacterial cell death is a biologically complex process, and we may expect that additional RMBs and pathways contributing to antibiotic-induced cell death remain to be uncovered, understood, and exploited by our evolving understanding of how antibiotics work.

Limitations of the study

In this work, we have combined single-cell and bulk-culture approaches to study the metabolic and molecular pathways contributing to antibiotic-induced cell death in *E. coli*. Building on previous studies (Kohanski et al., 2007; Dwyer et al., 2014; Wong et al., 2021a), we have focused on *E. coli* as a model organism. Our findings suggest that RMBs contribute to antibiotic lethality under anaerobic conditions. This suggestion applies to different strains of *E. coli*, as shown here, and RMB scavengers protect against gentamicin lethality as well for a multidrug-resistant clinical *E. coli* isolate (Figure S17). However, the extent to which similar findings generalize to other bacteria and antibacterial treatments will require further study. Additionally, as detailed further in STAR Methods, our experimental setup limits the concentration of O₂ in our experiments to below ~30 μM, using a hydrogen in nitrogen gas mix. Although similar experimental setups have been used to study the effects of environmental oxygen on antibiotic lethality (Keren et al., 2013; Liu and Imlay, 2013; Dwyer et al., 2014; Wong et al., 2021a), we note that micromolar concentrations of O₂ have been shown to accommodate terminal oxidase function, and ROS formation, in plants (Blokhina et al., 2001). Furthermore, hydrogen gas may possess antioxidant properties, which could alter RMB generation and its downstream deleterious effects (Ohsawa et al., 2007). These limitations are consistent with our hypothesis that RMBs may arise from trace amounts of oxygen, in addition to endogenous sources including anaerobic respiration with alternate terminal electron acceptors and free radical interactions. We expect future work to further address these limitations, for instance, using anaerobic environments containing noble gases such as argon.

**STAR METHODS**

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QUANTIFICATION AND STATISTICAL ANALYSIS

SUPPLEMENTAL INFORMATION

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## STAR METHODS

### KEY RESOURCES TABLE

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RESOURCE AVAILABILITY

Lead contact
Further information and request for resources and reagents should be directed to and will be fulfilled by the lead contact, James J. Collins (jimjc@mit.edu).

Materials availability
The strains of E. coli used in this study are available with a materials transfer agreement (MTA).

Data and code availability
- All data reported in this paper will be shared by the lead contact upon request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL DETAILS

Bacterial strains
All experiments were performed with E. coli K-12 MG1655 unless otherwise noted, and all strains used in this work are summarized in Table S3. The ΔkefBΔkefC double knockout strain was constructed from the Keio collection (Baba et al., 2006) ΔkefB and ΔkefC strains (ΔkefB, JW3131-1; ΔkefC, JW0046-1) by lambda-red recombineering and kanamycin resistance was subsequently cured, both according to published methods (Datsenko and Wanner, 2000); a list of primers used in the construction and PCR verification is provided in Table S4. The ΔgstA single knockout strain is from the Keio collection (ΔgstA, JW1627-1) and was validated using PCR. For time-kill experiments involving the ΔkefBΔkefC and ΔgstA strains, the Keio collection parent strain, E. coli BW25113, was used as the control strain. SMR14334, a GamGFP strain, is a derivative of strain MG1655 and has been previously described (Shee et al., 2013). E. coli CDC 541, a multidrug-resistant strain, was obtained from the Centers for Disease Control and Prevention AR Isolate Bank (Atlanta, GA).

METHOD DETAILS

Antibiotics
Kanamycin sulfate (product 60615, Sigma-Aldrich, St. Louis, MO) and gentamicin sulfate (Sigma-Aldrich G1914) were dissolved in ultrapure Milli-Q water to make working stock solutions of up to 25 mg/mL. Ciprofloxacin (Sigma-Aldrich 17850) was dissolved in dilute acid (0.1 M HCl, Sigma-Aldrich H1758) to make working stock solutions of up to 10 mg/mL. Mecillinam (Sigma-Aldrich 33447) was dissolved in dimethyl sulfoxide (DMSO, Sigma-Aldrich D5879) to make working stock solutions of up to 10 mg/mL. Rifampicin (Sigma-Aldrich R3501), chloramphenicol (Sigma-Aldrich C0378), and tetracycline (Sigma-Aldrich 87128) were dissolved in
DMSO, ethanol, and dilute acid, respectively, to make working stock solutions of 3.2 mg/mL. All antibiotics were freshly prepared immediately before each experiment.

**Bacterial culture and growth**

Cells were grown in liquid LB medium (product 244620, Becton Dickinson, Franklin Lakes, NJ). LB media containing 1.5% agar (Becton Dickinson 244520) was used to grow individual colonies. Cells were grown anaerobically or aerobically from single colonies at 37°C in 14 mL Falcon tubes with shaking at 300 rpm.

In anaerobic experiments, as a redox indicator, resazurin (Sigma-Aldrich R7017) was added to LB to a final concentration of 1 mg/L before autoclaving; the heat-activated form of resazurin, resorufin, is colorless below a redox potential of ~110 mV (Reddy, 2007). We note that resorufin remained strictly colorless in all anaerobic time-kill experiments, with the exception of those with exogenous H2O2. Addition of H2O2 resulted in rapid and persistent color changes of resorufin to pink, indicating increases in redox potential.

**Determination of MICs**

We determined MICs anaerobically for all antibiotics and RES considered in this work against *E. coli* MG1655 by diluting 1:10,000 from an anaerobically-grown overnight culture into 96-well plates (product 9018, Corning Inc., Corning, NY) capped with plate lids, with no shaking and two-fold dilutions of antibiotic or RES across wells. The MIC was determined as the minimum concentration of antibiotic or RES at which no visible growth (optical density at 600 nm, OD600 < 0.1) occurred overnight while incubated anaerobically at 37°C. A summary of the MIC values determined in this way is provided in Tables S1 and S2. All OD600 measurements in this work were performed using between 100 and 200 µL of culture volume in 96-well plates with a SpectraMax M3 plate reader (Molecular Devices, San Jose, CA).

**Anaerobic chamber and experiments**

Experiments were performed in an anaerobic chamber (Type B Vinyl, Coy Labs, Grass Lake, MI) equipped with twin palladium catalysts and a Coy Oxygen/Hydrogen Analyzer (Coy Labs) and maintained at 37°C. A 5% hydrogen in nitrogen gas mix (product NI HY5300, AirGas, Radnor, PA) was used to maintain the steady-state anaerobic environment at less than 5 PPM oxygen. Additionally, a BD BBL GasPak anaerobic indicator (Becton Dickinson), and growth media containing 1 mg/L resorufin as an anaerobic indicator, were used to validate anaerobic conditions. The minimum sensitivity of the oxygen indicators used is ~1 PPM; hence the concentration of O2 in the experiments was below ~30 µM. While previous studies have assumed that similarly anaerobic conditions were stringent enough to prevent ROS formation (Keren et al., 2013; Liu and Imlay, 2013), we note here that, inconsistent with this assumption, micromolar concentrations of O2 have been shown to accommodate terminal oxidase function, and ROS formation, in plants (Blokchina et al., 2001). These results are consistent with our hypothesis that RMBs can arise from trace amounts of molecular oxygen, in addition to various endogenous sources including anaerobic respiration with alternate terminal electron acceptors and free radical reactions.

For all anaerobic experiments, starting cultures were taken from cultures grown overnight inside the anaerobic chamber. Cells were grown with shaking at 300 rpm in 14 mL Falcon tubes and treated with antibiotics or RES as described above. For time-kill assays, cells were serially diluted, plated, and grown overnight inside the anaerobic chamber. For microscopy experiments, cells were plated and imaged with a Zeiss Axioscope A1 microscope inside the anaerobic chamber, as detailed further below. Plating and microscopy were performed strictly inside the anaerobic chamber.

To allow for deoxygenation, all materials used in our experiments were brought into the anaerobic chamber at least 24 h before the start of the experiment, with the exception of antibiotics or RES, fluorescent dyes and reagents, RMB scavengers, the BacTiter-Glo solution for ATP measurements, and cell lysis solution containing B-PER for preparation of cell lysates. To ensure freshness of these reagents, these reagents were prepared immediately before each experiment, brought into the anaerobic chamber, and equilibrated in open-cap tubes for 1 to 2 h before usage. After addition of these reagents to growing cultures, with the exception of ROS-treated cultures, we observed that resorufin in the growth media remained strictly colorless, indicating that these reagents did not introduce significant sources of environmental oxygen to our experiments.

**Time-kill assays and CFU measurements**

For all time-kill assays in bulk culture, cells were diluted 1:100 from an overnight culture (grown either anaerobically or aerobically, corresponding to the oxygen conditions of the time-kill experiment) into 14 mL Falcon tubes containing 2 mL of growth media. Following previous work (Keren et al., 2013) in which cells were grown for a fixed incubation time under both anaerobic and aerobic conditions, here cells were grown for ~1.5 h to early exponential phase, OD600 = 0.02 (anaerobic) and 0.1 (aerobic), in the conditions described above; we note here that the starting densities between this work and previous work from our lab (Dwyer et al., 2014) differ, and this difference may contribute to differences in survival relative to previous work from our lab (Dwyer et al., 2014). Antibiotics were added to the final concentrations indicated, and cultures were re-incubated with shaking at 300 rpm at 37°C. At the indicated times, cells were aliquoted and serially diluted in LB, and between 5 to 100 µL of cell culture was plated or spotted on LB agar.

LB-agar petri dishes were incubated at 37°C at least overnight (16-24 h) under the same oxygen conditions as the time-kill experiment. CFUs were determined by manual counting, and all measurements are based on counts containing at least five colonies; we note here that a typical lower limit of quantification in our time-kill assays is between 1 to 10^2 CFU/mL, and, where applicable, we have
truncated CFU/mL values according to the lower limit of quantification. Survival was determined by dividing all CFU/mL measurements to that immediately before antibiotic treatment at time 0 h.

**Metabolic measurements**

We treated log-phase bulk cultures with kanamycin, ciprofloxacin, and mecillinam at concentrations ranging from 1 × to 50 × the anaerobic MICs under anaerobic conditions and harvested cells after a treatment time of 20 min, similar to previous work examining changes in metabolic flux in antibiotic-treated cells under aerobic conditions (Kohanski et al., 2007; Belenky et al., 2015). For all metabolic measurements in bulk culture, cells were diluted 1:100 from an overnight culture grown anaerobically into 50 mL Falcon tubes with working volumes of 50 mL. Cells were grown anaerobically for ~1.5 h at 37°C without shaking to early exponential phase, OD600 = 0.02. Antibiotics were added to the final concentrations indicated, and cultures were incubated without shaking at 37°C. After 20 min of treatment, cells were centrifuged at centrifuged at 3720 × g for 5 min, and washed with 500 μL PBS. Cells were centrifuged again at 1500 × g for 5 min, the supernatant was discarded, and 1 mL B-PER II (product 78260, Thermo Fisher Scientific, Waltham, MA) containing 100 μg/mL lysozyme (Sigma-Aldrich L6876) and 2.5 U/mL Dnase I (Thermo Fisher 90083) was added to each cell pellet for harvesting. Cells were incubated with vortexing for 10 min. Cells were then centrifuged at 1500 × g for 5 min, and 800 μL of supernatant was aliquoted and removed from the anaerobic chamber for further analysis. Protein concentrations in the sample supernatants were determined using the Coomassie Plus protein assay reagent, as described below. Metabolite concentrations in the sample supernatants were determined using enzymatic assays. For these enzymatic assays, the supernatant was deproteinized by aliquoting 500 μL of supernatant into a 10 kDa molecular weight cut-off (MWCO) spin filter (Thermo Fisher 88513), centrifuging at 12,000 × g for 30 min, and collecting the flow-through for analysis. The flow-through from B-PER II with lysozyme and DNAase (no cell lysis) was also collected for background determination during analysis.

**Glucose concentration measurements**

Glucose concentrations were measured using an enzymatic assay kit (Sigma-Aldrich MAK263). Standards of 0, 0.004, 0.008, 0.012, 0.016, and 0.02 nmole/μL glucose were generated, and 50 μL of standard or sample was added to each well of a 96-well flat-bottom plate. 50 μL of the reaction mix comprising glucose assay buffer, glucose probe, and glucose enzyme mix were then added to each well following the manufacturer’s instructions. The plate was incubated for 30 min at 37°C, protected from light. The fluorescence intensity at Ex/Em = 535/587 nm was measured using a SpectraMax M3 plate reader. Glucose concentration values were inferred by linearly interpolating fluorescence intensity values with respect to the standard curve.

**Pyruvate concentration measurements**

Pyruvate concentrations were measured using an enzymatic assay kit (Sigma-Aldrich MAK332). Standards of 0, 5, 10, 15, 20, 30, 40, and 50 μM pyruvate were generated, and 10 μL of standard or sample was added to each well of a 96-well flat-bottom plate. 90 μL of the reaction mix comprising enzyme mix and dye reagent were then added to each well following the manufacturer’s instructions. The plate was incubated for 30 min at room temperature. The fluorescence intensity at Ex/Em = 530/585 nm was measured using a SpectraMax M3 plate reader. Pyruvate concentration values were inferred by linearly interpolating fluorescence intensity values with respect to the standard curve.

**NADH and NAD+ concentration measurements**

NADH and total NADH and NAD+ concentrations were measured using an enzymatic assay kit based on NAD cycling (product MET-5014, Cell Biolabs, San Diego, CA). In this assay, the concentrations of NADH, NAD+, or total NADH and NAD+ can be determined using an enzymatic cycling reaction in which NAD+ is reduced to NADH, then NADH reacts with a probe that produces a colored product. NADH was specifically extracted, and NAD+ was destroyed, by addition of NaOH. Briefly, 27.5 μL of each sample was added to a microcentrifuge tube, 5.5 μL of 0.1 N NaOH was added, and the tube was mixed. Tubes were incubated at 80°C for 1 h, protected from light. Tubes were centrifuged briefly to pool all sample solution, and 22 μL of the provided assay buffer was added to shift the pH back to neutral. The incubation with base was skipped for measurements of total NADH and NAD+. pH in all samples was confirmed to be ~7.0 using pH test indicator strips (Sigma-Aldrich P4786). Standards of 0, 0.004, 0.008, 0.015, 0.031, 0.063, 0.125, 0.25, 0.5, and 1 μM NAD+ were generated, and 50 μL of standard or sample was added to each well of a 96-well flat-bottom plate. 50 μL of the NAD cycling reagent comprising NAD cycling substrate, NAD cycling enzyme, colorimetric probe, and assay buffer was added to each well following the manufacturer’s instructions. The plate was incubated for 3 h at room temperature, protected from light. The absorbance at 450 nm was measured using a SpectraMax M3 plate reader. NADH and total NADH and NAD+ concentration values were inferred by linearly interpolating absorbance with respect to the standard curve, and NAD+ values were determined by subtracting NADH values from total NADH and NAD+ values.

**Description of antibiotic-induced phenotypes**

We have previously studied the cellular phenotypes induced by bactericidal antibiotics at the single-cell level (Wong and Amir, 2019; Wong et al., 2021a). In brief, salient phenotypes induced by aminoglycosides and fluoroquinolones include cytoplasmic condensation and lysis (Wong et al., 2021a), while salient phenotypes induced by β-lactams include membrane bulging and lysis (Yao et al., 2012; Wong and Amir, 2019; Wong et al., 2021b). Aminoglycoside- and fluoroquinolone-treated cells, treated with antibiotics for a...
timescale of hours, experience cytoplasmic condensation wherein discrete portions of the cellular cytoplasm become phase-light and the cell shrinks. This shrinkage occurs over a timescale of minutes, irreversibly halts cellular elongation, and has been evidenced to arise from membrane damage (Wong et al., 2021a). Furthermore, we have previously shown that increases in the fluorescence intensities of carboxy-H$_2$DCFDA, DAF-FM, and C11-BODIPY occur coincident with cytoplasmic condensation and/or lysis in amino-glycoside- and fluoroquinolone-treated cells (Wong et al., 2021a). These cells remain condensed over a timescale of hours, until sudden lysis occurs (Wong et al., 2021a).

β-lactam-treated cells, treated with antibiotics for a timescale of tens of minutes, experience membrane bulging, wherein micron-sized, phase-dark membrane extrusions appear over the course of seconds. Cells remain bulged over a timescale of seconds to minutes, until sudden lysis occurs (Yao et al., 2012; Wong and Amir, 2019; Wong et al., 2021b). We have previously shown that membrane bulging can be explained by the formation of cell wall defects and the resulting elastic response of the cellular envelope (Wong and Amir, 2019; Wong et al., 2021b).

**Microscopy**

Microscopy experiments were performed with cells sandwiched between cover glasses and glass slides unless otherwise stated. Cells were concentrated by centrifugation at 2350 $\times$ g for 5 min and resuspended in a smaller volume of supernatant. We plated 1 to 2 µL of the resuspended bacterial culture on 3” $\times$ 1” $\times$ 1” microscope slides (product 125444, Fisher Scientific, Hampton, NH) containing LB-agarose pads for cell immobilization, and sealed the slides using 18 mm square cover glasses (product 48366, VWR, Radnor, PA). Cells were imaged immediately afterward. We used a Zeiss Axioscope A1 upright microscope equipped with a Zeiss AxioCam 503 camera and a Zeiss 100x NA 1.3 Plan-neofluar objective (Zeiss, Jena, Germany) located inside the anaerobic chamber. Images were recorded using Zen Lite Blue (Zeiss), and processed and analyzed using ImageJ (NIH, Bethesda, MD). When possible, epifluorescence exposure times were limited to a maximum of 300 ms to avoid photobleaching. All microscopy experiments were replicated at least twice, and we verified that the absolute values of all fluorescence intensities were comparable across experiments performed using this microscopy setup.

**RMB detection with fluorescent dyes**

All dyes below were added directly to cell cultures and incubated for at least 30 min before subsequent analyses. For general detection of oxidative stress and ROS (H$_2$O$_2$, ROO·, and NOO·), we used the cell-permeant dye carboxy-H$_2$DCFDA (Invitrogen C400), which was dissolved in DMSO and added to incubating liquid cultures to a final concentration of 10 µM. As a positive control, cells were treated with 10 mM H$_2$O$_2$ (Sigma-Aldrich H1009) for the same durations as with antibiotics. For detection of RNS, in particular nitric oxide (NO), we used DAF-FM diacetate (Invitrogen D23844), dissolved in DMSO and added to incubating liquid cultures to a final concentration of 10 µM. As a positive control, we treated cells with 1 mM diethylamine NONOate (Sigma-Aldrich D184), which was dissolved in ethanol to prepare stock solutions, for the same durations as with antibiotics. C11-BODIPY 581/591 (Invitrogen D3861), a fluorescent dye-based lipid peroxidation sensor whose fluorescence emission peak shifts from red to green upon lipid peroxidation, was dissolved in DMSO and added to incubating liquid cultures to a final concentration of 10 µM. As a positive control, cells were treated with 10 mM H$_2$O$_2$ for the same durations as with antibiotics. For all samples, cellular phenotypes were manually determined with respect to corresponding phase contrast images. We note here that lysed cells may exhibit fluorescence, as indicated in Figure 2 of the main text: in such cells, despite large decreases in cytoplasmic phase contrast intensity accompanying cellular lysis, cytoplasmic material remained after lysis, and the remaining material retained some fluorescence.

**Exogenous RES**

4-HNE (product 32100, Cayman Chemical, Ann Arbor, MI) was supplied as a solution in ethanol, and 4-HNE controls were performed by adding ethanol to cells. MGO (Sigma-Aldrich M0252) was supplied as an aqueous solution. All of these reagents were freshly prepared or diluted before each experiment.

**DNA double-strand break detection with GamGFP**

DNA double-strand breaks (DSBs) were detected using an engineered fluorescent protein-based probe (Shee et al., 2013). This probe uses a fusion of GFP to the Gam protein from phage Mu that binds to DSBs when expressed in *E. coli*. We performed this experiment under anaerobic conditions as follows. An overnight culture of *E. coli* SMR14334, grown anaerobically, was diluted 1:100 in LB media containing 100 ng/mL doxycycline for GamGFP induction. At OD$_{600}$ = 0.02, where applicable, cells were treated with the indicated compounds at the indicated concentrations. Cultures were incubated for the indicated times, and cells were analyzed using fluorescence microscopy as detailed above.

**Membrane damage detection with SYTOX Blue**

Membrane damage was detected using a membrane damage-sensitive dye, SYTOX Blue Nucleic Acid Stain (Invitrogen S11348), following previous work (Wong et al., 2021a). We performed this experiment under anaerobic conditions as follows. An overnight culture of *E. coli* MG1655, grown anaerobically, was diluted 1:100 in LB media. At OD$_{600}$ = 0.02, cells were treated with SYTOX Blue at a final concentration of 10 µM and, where applicable, any additional indicated compounds at the indicated concentrations. Cultures were incubated for the indicated times, and cells were analyzed using fluorescence microscopy as detailed above.
Image analysis

Cells were counted manually to determine phenotypes, such as condensation and lysis. For RMB dye experiments, fluorescence was quantitatively analyzed with ImageJ (National Institutes of Health, Bethesda, MD). Closed cell contours were delineated based on phase contrast images and ImageJ was used to measure the mean fluorescence pixel intensity within a given contour. This was done both manually and using the MicrobeJ plugin (Ducret et al., 2016) for reproducible automation. The average background fluorescence intensity was subtracted from all measured values. Fractional fluorescence enhancement was calculated as the ratio of fluorescence intensities (1) inside a cell contour and (2) averaged over background regions with no cells. We were not blinded to allocation in the image analysis, and all cells for which we could reliably determine phenotypes or measure fluorescence intensities were used. For GamGFP and SYTOX Blue experiments, closed cell contours were delineated based on phase contrast images, and the respective quantification of foci-containing and fluorescently stained cells were performed manually and semi-automatically using ImageJ. We were not blinded to allocation in the image analysis, and cells for which we could reliably determine the presence or absence of foci or measure fluorescence intensities were used.

Description of RMB scavengers used

Previous work has shown that thiourea, a general scavenger of RMBs, quenches RMB-sensitive fluorescent dye oxidation and protects against antibiotic killing both anaerobically (Keren et al., 2013; Liu and Imlay, 2013) and aerobically (Kohanski et al., 2007), consistent with our finding of RMB accumulation under anaerobic conditions and the suggestion that thiourea may scavenge RMBs under different oxygen conditions. Contrasting arguments have posited that thiourea acts independently of scavenging RMBs and rescues cells by reducing growth or metabolic rate (Liu and Imlay, 2013). Here, we demonstrate that protection against antibiotic lethality is general across different RMB scavengers and show that these scavengers typically reduce RES accumulation.

Glutathione is a major antioxidant that has previously been shown to attenuate antibiotic killing under aerobic conditions (Dwyer et al., 2014; Wong et al., 2021a; Lopatkin et al., 2019). Glutathione may protect against antibiotic lethality by direct scavenging of ROS and RNS, as well as by enzymatic reactions catalyzed by glutathione peroxidase (Lo et al., 1994). In addition to ROS and RNS detoxification, glutathione is also a potent detoxifier of RES through direct scavenging and the activity of glutathione-dependent enzymes, including glutathione S-transferases (Allocati et al., 2009). Spontaneous RES reactions with MGO can occur with the sulphydryl group of the cysteinyl residue of glutathione, generating hemithiolacetal (HTA; Ferguson et al., 1999). Moreover, glutathione can react spontaneously with 4-HNE through Michael addition (Negre-Salvayre et al., 2008; Aquilano et al., 2014). In addition to its detoxification functions, glutathione and its adducts regulate efflux, predominantly of K+, from the cell through the Kef system (Ferguson et al., 1999, 2000).

Previous studies have shown that glutathione acts to: (1) detoxify RMBs by direct scavenging; and (2) regulate the KefB/KefC potassium efflux channels, which activate when RMBs accumulate within cells (Ferguson et al., 2000), as well as glutathione S-transferases (GSTs), which catalyze the conjugation of glutathione to electrophiles and other xenobiotics (Allocati et al., 2009). As mentioned in the main text, we asked whether glutathione protection occurred mainly through quenching of RMBs or the effects of glutathione on KefB/KefC-mediated potassium efflux—resulting in cytoplasmic acidification, which protects cellular DNA from RES attack—and GST-based detoxification. To address this question, we first constructed a ΔkefBΔkefC strain and found that these two genetic deletions suffice to confer sensitivity to killing by exogenous MGO, consistent with a prior phenotypic study (Ferguson et al., 2000) of this operon under aerobic conditions (Figure S7; Tables S2–S4). We then tested for glutathione protection. Under both anaerobic and aerobic conditions, we found that ΔkefBΔkefC cells displayed glutathione protection against antibiotic treatment at comparable levels to wild-type (Figures S8 and S9), indicating that glutathione does not solely act on KefB/KefC-mediated potassium efflux to protect cells from antibiotic killing. We also found that the ΔkefBΔkefC strain was not hypersensitive to antibiotic killing, suggesting that KefB and KefC do not, by themselves, alter antibiotic lethality. Similar results were recapitulated in a ΔgstA strain, which lacks a gene encoding a major GST (Figure S10). Together, these findings suggest that glutathione decreases antibiotic lethality irrespective of the availability of environmental oxygen, the known effects of glutathione on KefB/KefC-mediated potassium efflux, and GstA-mediated detoxification.

Acetylcysteine is a glutathione and H2S precursor that, by itself, is often used as an antioxidant. However, it has been noted that the reaction rates of acetylcysteine for major classes of ROS are slow and almost undetectable for superoxide radical (Zhitkovich et al., 2019), and the antioxidant properties of acetylcysteine could arise from its role as a glutathione precursor. The disulfide-reducing activity of acetylcysteine is higher than that of glutathione due to its stronger nucleophilicity, suggesting that acetylcysteine primarily scavenges RES (Zhitkovich et al., 2019). Consistent with this inference, acetylcysteine has been shown to exhibit protective properties against 4-HNE and MGO, as previously reported in eukaryotes (Negre-Salvayre et al., 2008) and demonstrated for E. coli in this study.

Pyridoxamine, a form of vitamin B6, is known to prevent the formation of AGE such as CML and CEL. It inhibits lysine modifications by 4-HNE, and prevents the formation of ALE by scavenging 1,4-dicarbonyls (Amaranth et al., 2004). Pyridoxamine may also inhibit ROS production (Voziyan and Hudson, 2005) and can react directly with MGO to form a dimer, inhibiting AGE formation by MGO-driven modifications (Nagaraj et al., 2002).

RMB scavengers

L-glutathione reduced (Sigma-Aldrich G4251), n-acetyl-l-cysteine (Sigma-Aldrich A9165), and pyridoxamine dihydrochloride (Sigma-Aldrich P9380) were dissolved in ultrapure Milli-Q water. All scavengers were freshly prepared before each experiment.
Rationale for scavenger control experiments

As described in the main text, differences in cellular growth, metabolism, and intracellular concentration of antibiotics may affect antibiotic lethality. We performed additional experiments, as described below, to determine whether these factors could contribute to scavenger-mediated protection. Supplementation of each scavenger did not lead to significant differences in growth rate, as measured by optical density (Figure 4C). Previous studies from our laboratory and others (Lopatkin et al., 2019; Mathieu et al., 2016) have suggested that, under aerobic conditions, antibiotic lethality depends on the cellular metabolic state associated with ambient growth conditions, and that antibiotic lethality correlates with increased ATP levels (Lopatkin et al., 2019). As the BacTiter Glo assay has been used to measure ATP levels of bulk cultures (Lopatkin et al., 2019), we utilized this assay to determine ATP levels in scavenger-treated bulk cultures under anaerobic conditions. If decreased antibiotic lethality in the presence of scavengers (10 mM) arose due to decreased metabolism, then the measured ATP levels should be decreased in the presence of RMB scavengers. We found, on the contrary, that measured ATP levels were increased in the presence of RMB scavengers (Figures 4D and S11). These findings are inconsistent with the hypothesis that scavenger protection from antibiotic lethality arises from suppressive alterations to bacterial metabolic state. Importantly, time-kill experiments with cells cultured and treated in diluted LB (1:1000 in PBS) confirm that decreasing metabolism alone is insufficient to explain the protection observed: we find that there is essentially no growth, and between 0 to 2 logs of killing, occurring in antibiotic-treated cells in dilute LB across all antibiotic concentrations tested (Figures S12 and S13). In contrast, growth persists at lower antibiotic concentrations relative to the MICs in the presence of glutathione, acetylcysteine, or pyridoxamine in nutrient-replete media (Figure 4B). Together, these findings indicate that suppressive alterations to bacterial metabolic state do not explain the observed scavenger protection from antibiotic lethality.

A prior study (Ezraty et al., 2013) has linked iron chelator-mediated protection against aminoglycosides, ostensibly evidence for ROS-dependent aminoglycoside lethality (Kohanski et al., 2007), to changes in iron-sulfur clusters and electron transport chains resulting in altered proton motive force (PMF) and decreased aminoglycoside uptake. In E. coli, the PMF is generated by two components, the membrane potential, ΔΨ, and the pH gradient, ΔpH, across the membrane. As described below, we assayed for changes in ΔΨ and ΔpH using two fluorescent dyes, DiBAC4(3) and ACMA, as well as for changes in membrane permeability using DiSC3(5), in the presence and absence of RMB scavengers. DiBAC4(3) can enter depolarized cells, where it binds to intracellular proteins or the cell membrane and enhances exhibited green fluorescence. ACMA is a DNA intercalator that selectively binds to poly (d(A-T)) and membranes in the energized state, and becomes quenched if a pH gradient forms. DiSC3(5) accumulates on hyperpolarized membranes and can be translocated into the lipid bilayer. We found that the fluorescence intensities of DiBAC4(3), ACMA-, and DiSC3(5)-labeled cells, measured at the single-cell level, were not significantly different in the presence of RMB scavengers (Figure 4E). In contrast, as controls for these dyes, we used CCCP, a PMF decoupling protonophore, and valinomycin, an ionophore antibiotic. Consistent with the suppression of PMF in positive controls, we found that CCCP induced fluorescence of DiBAC4(3) and quenching of ACMA (Figure 4E). Consistent with changes in membrane permeability induced by valinomycin, we found that DiSC3(5)-labeled cells fluoresced in the presence of valinomycin (Figure 4E). These findings suggest that the observed scavenger protection from antibiotic lethality does not arise from suppressive alterations to PMF.

Lastly, we performed intracellular antibiotic concentration measurements using fluorimetry for kanamycin and liquid chromatography-mass spectrometry (LC-MS) for ciprofloxacin and mecillinam. We treated cells with antibiotics at a constant concentration in the presence of scavenger (10 mM), and found approximately equal antibiotic concentrations in cells with and without scavenger, as indicated by single-cell measurements for cells treated with a fluorescent derivative of kanamycin and LC-MS measurements for bulk-culture ciprofloxacin- and mecillinam-treated cell lysates (Figure 4F). These results indicate that intracellular antibiotic concentrations are not decreased in any of the scavenger-antibiotic pairs. Altogether, these findings suggest that the observed scavenger-mediated increases in MICs and protection from antibiotic killing are not explained by differences in cellular growth rate, cellular metabolic state, PMF, or intracellular antibiotic concentration.

ATP abundance assay

Intracellular ATP was quantified using the BacTiter-Glo Microbial Cell Viability Assay (product G8230, Promega, Madison, WI) according to the manufacturer’s instructions. Luminescence and optical density (OD600) of samples were measured with a SpectraMax M3 plate reader. For anaerobic cultures, cells were treated with the BacTiter-Glo assay solution inside the anaerobic chamber, incubated for at least 5 min for lysis, then removed from the anaerobic chamber for luminescence measurements performed immediately after sampling, using a SpectraMax M3 microplate reader located outside the anaerobic chamber. As an additional positive control used for ATP/OD600 determination, cells cultured in Neidhardt EZ Rich Defined Medium (Teknova Inc., Hollister, CA) were noted to exhibit substantially increased ATP/OD600 levels (ATP/OD600 ~ 7) relative to cells cultured in LB only (ATP/OD600 = 1), and the former culture condition was included and assayed in parallel in all experiments.

Changes in membrane permeability and PMF

DiBAC4(3) (Invitrogen B438), a fluorescent reporter of membrane potential (ΔΨ), was dissolved in DMSO and added to incubating liquid cultures to a final concentration of 10 μg/mL to stain cells with and without depolarized membranes. ACMA (9-amino-6-chloro-2-methoxyacridine; Invitrogen A1324), a fluorescent reporter which binds to membranes in the energized state and becomes quenched if a pH gradient forms, was dissolved in DMSO and added to incubating liquid cultures to a final concentration of 10 μg/mL to assay changes in ΔpH. DiSC3(5) (Invitrogen D306), a fluorescent reporter which accumulates on hyperpolarized membranes and is
translocated into the lipid bilayer, was dissolved in DMSO and added to incubating liquid cultures to a final concentration of 1 μM. As a positive control for DiBAC₄(3) and ACMA, CCCP (carbonyl cyanide 3-chlorophenylhydrazone; Sigma-Aldrich C2759), a protonophore which uncouples PMF, was dissolved in DMSO and added to incubating liquid cultures at least 5 min before imaging, to a final concentration of 1 mM. As a positive control for DiSC₃(S), valinomycin (Sigma-Aldrich V0827), an ionophore antibiotic, was dissolved in DMSO and added to incubating liquid cultures at least 5 min before imaging to a final concentration of 100 μg/mL.

**Intracellular kanamycin concentration measurements**

Intracellular kanamycin concentration measurements were performed using kanamycin-Texas Red analogously to previous work (Meylan et al., 2017; Sandoval et al., 1998). One milligram of Texas red sulfonyl chloride (Thermo Fisher T1905) was resuspended in 50 μL of anhydrous N,N-dimethylformamide (Sigma-Aldrich 227056) on ice. The solution was added to 2.3 mL of 100 mM K₂CO₃ (Sigma-Aldrich P5833) at pH 8.5, with or without 10 mg/mL kanamycin, on ice. To quantify kanamycin-Texas Red concentrations, 2 mL of cells was grown to log-phase anaerobically from a 1:100 dilution of an overnight culture to OD₆₀₀ = 0.02, as described above. Concentrated kanamycin-Texas Red was then added to achieve a working concentration of 50 μg/mL, scavenger (10 mM final concentration) was added where applicable, and samples were incubated anaerobically for 1 h. Samples were then analyzed using fluorescence microscopy, as detailed above.

**Preparation of samples for LC-MS**

For each sample used to measure ciprofloxacin or mecillinam concentration, 100 mL of cells was grown to log-phase anaerobically from a 1:100 dilution of an overnight culture to OD₆₀₀ = 0.02 without shaking and in 250-mL flasks. Cells were then treated with mecillinam or ciprofloxacin at a final concentration of 10 μg/mL or ciprofloxacin at a final concentration of 1 μg/mL, and scavenger (10 mM final concentration) was added where applicable. After 4 h, 50 mL of cell culture was aliquoted, centrifuged at 3720 × g for 10 min, and washed with 500 μL PBS. Cells were centrifuged again at 1500 × g for 10 min, the supernatant was discarded, and 500 μL acetonitrile (Sigma-Aldrich 271004) was added to the cell pellet. Mecillinam or ciprofloxacin concentrations in these samples were determined using LC-MS, as detailed below. The remaining 50 mL of cell culture was used for protein concentration determination, as detailed below.

**Description of RES measured**

RES, including MGO and 4-HNE, are known to participate in a variety of cellular reactions. MGO, a reactive dicarbonyl α-oxoaldehyde, is known to accumulate in millimolar concentrations within *E. coli* cells during unbalanced glycolysis, as previously studied in an *E. coli* K-12 strain with impaired glycerol kinase activity (Freedberg et al., 1971). In addition to lipid peroxidation and glycolysis, MGO can arise from the degradation of glucose, DNA, and L-threonine, and enzymatically from MGO synthase with the glycolytic intermediate dihydroxyacetone phosphate (DHAP) as an intermediate. MGO forms Schiff bases with amino acids, particularly nucleophilic residues including lysine, cysteine, and arginine (Lo et al., 1994). After Amadori rearrangement, AGE, including carboxymethyl- and carboxyethyl-lysines (CML and CEL), are generated. Furthermore, like 4-HNE (below), MGO forms adducts with DNA that lead to DNA damage and degradation (Negre-Salvayre et al., 2008; Lee and Park, 2017). MGO can also generate free radicals by glycating amino acids in a process which occurs independent of oxygen availability (Yim et al., 1995). Additionally, methylglyoxal has previously been suggested to increase persister frequency after ampicillin treatment as a consequence of loss-of-function mutations of *glpD* and *tktA*, two genes associated with cellular metabolism (Girgis et al., 2012). Our findings support the notion that methylglyoxal accumulates in antibiotic-treated cells and further suggest that methylglyoxal may be deleterious under the conditions considered here.

4-HNE is a reactive α,β-unsaturated hydroxyalkenal that often arises from the oxidation of fatty acids. Unlike MGO, 4-HNE accumulation in *E. coli* has largely not been studied in previous work. While 4-HNE is commonly generated in eukaryotes by the lipid peroxidation of lipids containing polyunsaturated omega-6 acyl groups, 4-HNE can also arise from lipid peroxidation of lipids commonly found in bacteria including *E. coli*, such as cardiolipin (Dalleau et al., 2013; Liu et al., 2011; Dowhan et al., 1997). 4-HNE reacts with histidine, cysteine, or lysine residues of proteins to form stable Michael adducts, which eventually lead to the formation of protein aggregates that accumulate in cells (Negre-Salvayre et al., 2008). In eukaryotes, accumulation of 4-HNE-protein adducts has been evidenced to affect tyrosine kinase receptors, cell cycle progression, signaling, proteasome activity, oxidative stress responses, and induction of caspases resulting in apoptosis (Negre-Salvayre et al., 2008). Similar to MGO, 4-HNE can also form adducts with DNA, and these adducts have been evidenced to be potent mutagens (Csala et al., 2015).

We note here that additional RES, such as malondialdehyde (MDA), have previously been evidenced to accumulate in antibiotic-treated cells under aerobic conditions. MDA is a major dicarboxyl aldehyde byproduct of lipid peroxidation in fatty acids, particularly polyunsaturated fatty acids. While *E. coli* is not evidenced to synthesize polyunsaturated fatty acids under typical growth conditions, a non-enzymatic lipid peroxidation reaction can occur with monounsaturated fatty acids without the characteristic lipid peroxidation propagation step (Hong et al., 2012). This is supported by the detection of TBARS (thiobarbituric acid reactive substances) in *E. coli* exposed to oxidizing agents (Hong et al., 2012). Furthermore, as previously detailed (Belenky et al., 2015), MDA may arise in bacteria from deoxyribonuclease by an alternative pathway initiated by a Fenton oxidant abstracting a hydrogen atom at the 4'-position to form a carbon-centered radical that adds molecular oxygen at diffusion-controlled rates. Previous aerobic TBARS measurements for copper alloy-treated *E. coli* (Hong et al., 2012) and ELISA-based measurements of MDA adducts in antibiotic-treated *E. coli* suggest that, like 4-HNE and MGO, MDA can also accumulate in bacterial cells (Belenky et al., 2015; Hong et al., 2019).
Preparation of samples for GC-MS

For each sample used to measure RES concentration or protein concentration, at least 50 mL of log-phase bulk cultures of E. coli MG1655 grown in LB were treated with kanamycin, ciprofloxacin, and mecillinam at the indicated concentrations for the indicated times in 250-mL flasks. Cells were then aliquoted into 50 mL Falcon tubes. Tubes were centrifuged at 3720 × g for 10 min. Next, we discarded the supernatant in all samples. Each cell pellet was washed with 500 µL PBS, resuspended, and centrifuged at 1500 × g for 10 min. Afterward, the supernatant was discarded from each sample.

For samples used in RES concentration determination and protein concentration determination, 1 mL B-PER II containing 100 µg/mL lysozyme, 5 U/mL DNase I, and 25 mM O-(2,3,4,5,6-Pentafluorobenzyl)hydroxylamine (PFBHA; Sigma-Aldrich 76735) as a carbonyl derivatizing agent was then added to cell pellets for harvesting, and, as an internal standard, deuterated benzaldehyde (benzaldehyde-d₈, product D-0005, CDN Isotopes, Pointe-Claire, Quebec, Canada) was added to the lysis buffer to a final concentration of 100 µM immediately before dispensing of the lysis buffer to cell pellets. The same lysis buffer without PFBHA and deuterated benzaldehyde was used for protein concentration determination of samples from cell cultures that were processed using LC-MS to measure ciprofloxacin and mecillinam concentrations.

After addition of lysis buffer, all samples were vortexed and incubated at 37°C for 30 minutes. All samples were then centrifuged at 1500 × g for 10 min, and the supernatant was aliquoted from each sample for further analysis of RES or protein concentrations. For RES measurements, the supernatants were removed from the anaerobic chamber at this point, flash-frozen on dry ice, and stored at -80°C until processing at the Harvard Center for Mass Spectrometry.

Protein concentration assay

We used the Coomassie Plus protein assay reagent (Thermo Fisher 23236) following the manufacturer’s instructions. This measurement was performed under aerobic conditions using the supernatants prepared under anaerobic conditions. Briefly, for each sample, 10 µL of sample was pipetted directly into 300 µL of Coomassie Plus reagent in a 96-well plate. The plate was mixed and incubated at room temperature for 10 minutes, and the absorbance at 595 nm was read with a SpectraMax M3 spectrophotometer. Standard curves were generated from control samples with bovine serum albumin (BSA) concentrations of 2000, 1500, 1000, 750, 500, 250, 125, 25, and 0 µg/mL, where BSA was diluted in the same lysis buffer used to prepare samples. Protein concentration values were inferred by linearly interpolating absorbance values with respect to the standard curves.

GC-MS

GC-MS experiments were performed at the Harvard Center for Mass Spectrometry. Cell lysates were analyzed for target derivatives of PFBHA with 4-HNE and MGO. All samples were run on a Thermo GC-QE mass spectrometer using an Agilent DB-5MS (30 m, 0.25 mm diameter, 0.25 µm film) column with the following orbitrap parameters: polarity, pos; resolution, 60000; agc target, 1e6; scan range, 66.7 to 1000 m/z. Samples were injected in splitless mode. The GC oven was maintained at 50°C for 4 min, before increasing to 300°C at 15°C/min and finally maintained at 300°C for 24 min. Cell lysate samples were prepared as follows. Between 750 to 800 µL of each sample was transferred to a new tube, then 500 µL of methanol was added to each sample. Samples were transferred to 8 mL glass vials and incubated at room temperature for 30 min. 1 mL of hexane was added, and samples were vortexed for 1 min. Six drops of sulfuric acid were added, and then samples were again vortexed for 1 min. Samples were centrifuged at 2500 × g, the supernatant was transferred to glass autosampler vials while avoiding the aqueous phase, and the samples were dried under flow of N₂. Finally, the samples were resuspended in 50 µL hexane, and 45 µL of the suspension was transferred to microinserts (the remaining 5 µL being used to test pH). Standards were prepared as a 100 µM solution of 4-HNE and MGO in lysis buffer containing internal standard, and six 1/5 dilution series of each standard were subsequently prepared. 1 mL of each standard was then transferred to 8 mL vials, 500 µL of methanol was added to each standard, and standards were then treated like samples.

Benzaldehyde-d₈ was used as an internal standard, yielding a fragment at m/z = 181.0069. The following accurate m/z of fragment ions were used for quantification: 250.0287 for 4-HNE, 265.0396 for MGO, and 181.0069 for the internal standard. The quantification was based on the ratio with the internal standard area. Samples with low internal standard signals were not used. The limits of detection for 4-HNE and MGO in this protocol were 0.64 pmol and 16 pmol, respectively. Sample chromatograms for standards and chemical structures of target derivatives are shown in Figure S3. Most GC-MS measurements were performed on two independent occasions, and typical values are reported as fold changes to facilitate comparison.

Intracellular ciprofloxacin concentration LC-MS

In previous work (Gutierrez et al., 2017; Asuquo and Piddock, 1993), intracellular ciprofloxacin concentrations were assayed fluorometrically with Ex/Em in the range of 275/410 nm. However, the addition of exogenous scavengers may interfere with this fluorometric reading: pyridoxamine, for instance, has Ex/Em in the range of 335/400 nm (Sikorska et al., 2004). We therefore quantified intracellular ciprofloxacin using LC-MS experiments, which were performed at the Harvard Center for Mass Spectrometry. Deuterated ciprofloxacin, ciprofloxacin-d₈ (product 25466, Cayman Chemical Company, Ann Arbor, MI), was used as an internal standard. All solvents used are LC-MS grade. 100 µL of ciprofloxacin-d₈ in acetonitrile was added to thawed samples. Samples were vortexed for 1 min, until the cell pellets were fully resuspended: an ultrasound bath was used, if needed, to loosen the pellets. Samples were then incubated in an ultrasound bath for 30 min, centrifuged at maximum speed for 10 min, transferred to new microcentrifuge tubes, dried under N₂ flow, and resuspended in 100 µL of acetonitrile (50% in water).
A standard curve was prepared using eight 1/5 dilution series of a 100 μM solution of ciprofloxacin in acetonitrile containing internal standard. Standards were prepared similarly to the samples. The lower limit of quantification was found to be less than 1.28 nM.

All samples were run on a Thermo QE+. The column used was Phenomenex Kinetex C18, 1.7 μm, 100 Å, 150 × 2.1 mm. The source used was HESI+. MS parameters were as follows: full MS 200-600 m/z, resolution 70000, 100 ms max IT, 3e6 AGC. The mobile phases were A: water, 0.1% formic acid, 5 mM ammonium formate and B: acetonitrile. The following gradient was used: 3 min at 10% B, then to 100% B in 0 min, followed by 7 min at 100% B. The column was then equilibrated at 10% B for 3 min. The flow rate was 0.25 mL/min. Each sample was injected twice continuously, and the second injections were used for quantification.

**Intracellular mecillinam concentration LC-MS**

LC-MS experiments were performed at the Harvard Center for Mass Spectrometry. Deuterated ampicillin, ampicillin-d₅ (Cayman Chemical 25356), was used as an internal standard. All solvents used are LC-MS grade. 300 μL of acetonitrile and internal standard (ampicillin-d₅ at 0.33 μM) was added to frozen samples. 200 μL of H₂O was added, and samples were left to thaw. Samples were vortexed for 30 s, and then incubated in an ultrasound bath for 30 min. Samples were then centrifuged at max speed for 20 min, transferred to new tubes, dried under N₂ flow, and resuspended in 100 μL of acetonitrile (50% in water).

A standard curve was prepared using nine 1/5 dilution series of a 1 mM solution of mecillinam in LB. 50 μL of each standard was prepared similarly to the samples. The lower limit of quantification was found to be 2.5 nM.

All samples were run on an Agilent 6460 Triple Quad Mass Spectrometer coupled to a 1290 LC. The column used was Phenomenex Kinetex C18, 1.7 μm, 100 Å, 150 × 2.1 mm. The source used was ESI turbojet, with gas temperature 330°C, gas flow 8L/min, nebulizer 40 psi, sheath gas 375°C at 9L/min, capillary at +2800 V and nozzle at +300 V, and delta EMV of +100. The mobile phases were A: water, 0.1% formic acid and B: acetonitrile, 0.1% formic acid. The following gradient was used: 5 min at 0% B, then to 100% B in 10 min, followed by 5 min at 100% B. The column was then equilibrated at 0% B for 5 min. The flow rate was 0.2 mL/min. Each sample was injected twice continuously with the exception of two samples, and resulting measurements of mecillinam concentration were averaged. The two samples that were not injected twice continuously were injected only once each: these were both samples corresponding to treatment with mecillinam and pyridoxamine, and the mecillinam concentrations in these samples were found to be within 10% of variation.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Statistical parameters were reported either in individual figures or corresponding figure legends. Two-sample Kolmogorov-Smirnov tests or two-sample t-tests for differences in mean value, as indicated in each figure where applicable, were performed at the standard 5% significance level. We were not blinded to allocation in the statistical testing. All statistical analyses were performed using MATLAB.