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A transcriptomic Analysis of the Activity and Mechanism of Action of a Ruthenium(II)-Based Antimicrobial That Induces Minimal Evolution of Pathogen Resistance

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A transcriptomic Analysis of the Activity and Mechanism of Action of a Ruthenium(II)-Based Antimicrobial That Induces Minimal Evolution of Pathogen Resistance

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KEYWORDS antimicrobials, pathogens, resistance, ruthenium, transcriptomics.

ABSTRACT Increasing concern over rising levels of antibiotic resistance amongst pathogenic bacteria has prompted significant research into developing efficacious alternatives to antibiotic

1
2
3 treatment. Previously, we have reported on the therapeutic
4 activity of a dinuclear ruthenium (II) complex against pathogenic,
5 multidrug resistant bacterial pathogens. Herein, we report that
6 the solubility properties of this lead are comparable to those
7 exhibited by orally available therapeutics, that - in comparison
8 to clinically relevant antibiotics - it induces very slow evolution
9 of resistance in the uropathogenic, therapeutically resistant, *E.*
10 *coli* strain EC958 and this resistance was lost when exposure to
11 the compound was temporarily removed. With the aim of further
12 investigating the mechanism of action of this compound, the
13 regulation of nine target genes relating to the membrane, DNA
14 damage and other stress responses provoked by exposure to the
15 compound was also studied. This analysis confirmed that the
16 compound causes a significant transcriptional downregulation of
17 genes involved in membrane transport and the tricarboxylic acid
18 cycle. By contrast, expression of the chaperone protein-coding
19 gene, *spy*, was significantly increased suggesting a requirement
20 for repair of damaged proteins in the region of the outer membrane.
21 The complex was also found to display activity comparable to that
22 in *E. coli* in a range of other therapeutically relevant Gram-
23 negative pathogens.
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5 Over recent years, there has been a significant increase in
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7 bacterial infections that display multidrug resistance, leading to
8
9 a concomitant increase in mortality rates.¹⁻³ As last-line
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11 treatments such as carbapenems increasingly fail,³⁻⁶ antimicrobial
12
13 resistance is rapidly becoming a global threat to public health
14
15 and the economy.⁷⁻⁹ It has been estimated that by 2050 ten million
16
17 lives per year and a cumulative \$100 trillion of economic output
18
19 will be lost due to the rise of drug-resistant infections.¹⁰
20
21

22
23 *Escherichia coli* strains are a significant cause of infection
24
25 within clinical settings and are linked to high morbidity and
26
27 mortality globally causing a wide range of infections including
28
29 meningitis, pneumonia and bacteraemia.¹¹ *E. coli* is highly
30
31 prevalent in urinary tract infections and accounts for 80 % of all
32
33 community acquired urinary tract infections.^{12,13} High antimicrobial
34
35 resistance within uropathogenic *E. coli* (UPEC) strains is common;
36
37 a study of antibiotic resistance in cases of urinary tract
38
39 infections in Nigeria found that from a total of 137 *E. coli*
40
41 isolates 36 % were resistant to ten out of 11 urine line
42
43 antibiotics.¹⁴
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48 In this context, UPEC sequence type ST131 is an emerging pathogen
49
50 of particular concern. Apart from being commonly resistant to
51
52 fluoroquinolones, this strain produces the CTX-M extended spectrum
53
54 beta lactamase that confers resistance to oxyimino-cephalosporins
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3 and monobactams.¹⁵⁻¹⁸ Furthermore, CTX-M encoding genes are found
4 on plasmids which frequently carry additional resistance genes. As
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6 on plasmids which frequently carry additional resistance genes. As
7
8 a consequence, geographical variants commonly possess quinolone
9
10 modifying enzymes that provide fluroquinolone resistance, as well
11
12 as the enzymes carbapenemases and cephamycinases.¹⁹
13

14 As global concern increased, the USA surveillance programs SENTRY
15
16 and MYSTIC estimated through extrapolation that ST131 accounted
17
18 for approximately 17 % of all *E. coli* isolates, 44 % of all
19
20 antimicrobial resistant isolates and around 68 % of
21
22 fluoroquinolone resistant isolates.^{16,20} This is problematic as
23
24 within the US fluoroquinolones are prescribed as a first line
25
26 treatment against urinary tract infections. Given these facts, it
27
28 is unsurprising that - even when treated with standard antibiotic
29
30 regimes - urinary tract infections caused by ST131 can dangerously
31
32 progress into pyelonephritis and sepsis.^{21,22}
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37 With one third of women having a course of antibiotics to treat
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39 a urinary tract infection by the age of 26²³ multidrug resistant
40
41 uropathogenic bacteria are clearly a significant threat, with
42
43 ST131 being at the forefront of concern. It is therefore apparent
44
45 that urine line antibiotics are a pivotal tool in healthcare and,
46
47 as they are quickly becoming less effective, it is crucial that
48
49 novel treatment options are identified.²⁴
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52 Metal complexes are a class of compounds that demonstrated
53
54 significant early promise as therapeutic leads but are
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3 underdeveloped. As early as the 1950s, the Dwyer group reported
4
5 that Ru^{II} polypyridyl complexes had potential as antimicrobials.^{25,26}
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7 Their work demonstrated that increasing the lipophilicity of the
8
9 parent [Ru(phen)₃]²⁺ cation resulted in enhanced antimicrobial
10
11 action, leading to a derivative that displayed promising activity
12
13 against Gram-positive bacteria. However, due to the wide range of
14
15 effective conventional antibiotics clinically available at that
16
17 time, no further development of these distinctive leads occurred
18
19 for decades.
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23 In the last decade or so, due to the growing antibiotic resistance
24
25 crisis, the use of metals as antimicrobials has been revisited,²⁷⁻³²
26
27 with particular focus on the Ru^{II} systems,^{28,33} although most of
28
29 these newly reported systems still exhibit higher activities
30
31 against Gram-positive bacteria. As part of a program to develop
32
33 novel metal-complex-based imaging probes,³⁴⁻³⁶ therapeutics³⁷⁻⁴⁰ and
34
35 phototherapeutics,⁴¹⁻⁴⁵ we recently identified a series of dinuclear
36
37 ruthenium (II) complexes that exhibit higher activity against
38
39 Gram-negative species.⁴⁶ Subsequent detailed studies involving
40
41 several strains of *Staphylococcus aureus* indicated that the
42
43 lowered activity against Gram-positive bacteria is due to the
44
45 complexes binding to cell wall teichoic acids residues, leading to
46
47 reduced internalization.⁴⁷
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53 Although these reports provided preliminarily insights into the
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55 mechanism of action of these new potential therapeutics, in this
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3 study a deeper and more focused understanding of the antimicrobial
4 activity of the main lead complex, $\mathbf{1}^{4+}$ (Figure 1) is developed. The
5 complex was synthesized as a PF_6^- , for biological studies anion
6 metathesis was used to convert the complex into the Cl^- salt form.
7
8 Using the multidrug resistant uropathogenic strain of *E. coli* as
9 a model Gram-negative pathogen quantitative PCR (qPCR) was used to
10 monitor the regulation of key genes thought to be responsible for
11 sensing and reacting to the presence of $\mathbf{1}^{4+}$, providing further
12 insights into its mechanism of action. As we also find the complex
13 displays solubility that is comparable with established orally
14 available therapeutics, is active against a range of Gram-negative
15 pathogens, and resistance towards it emerges only very slowly,
16 these data further underlines its therapeutic potential.
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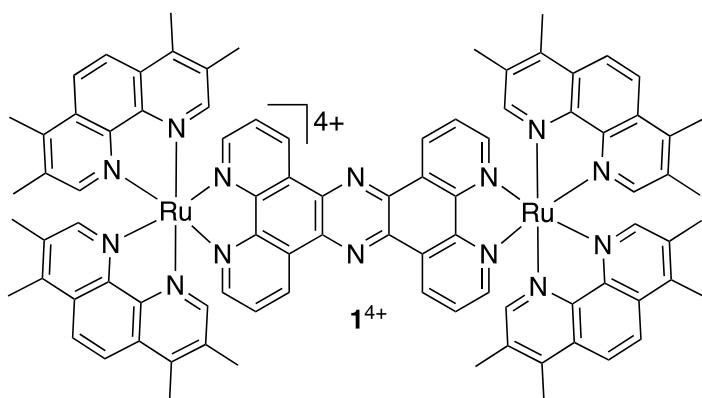
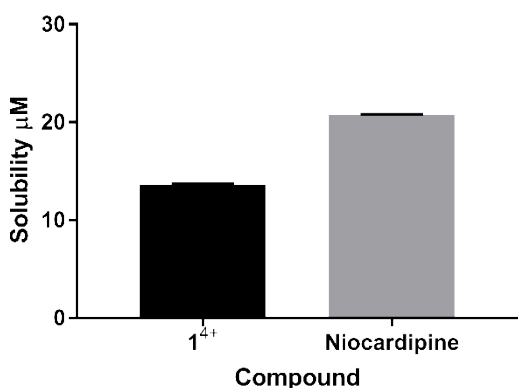


Figure 1. Structure of the ruthenium (II) complex relevant to this report.

Results

Kinetic Turbidimetric Solubility

Over the last two decades, largely as a consequence of high through-put screening and demand for structurally complex drug leads, poor aqueous solubility has increasingly become a limiting development factor in drug development; compounds with poor solubility present high attrition risks and increased drug development costs.⁴⁸⁻⁵⁰ As complex **1⁴⁺** displays activity against Gram-negative pathogens, and it is known that solubility is a key criterion for oral availability of any drug lead,⁵¹ its solubility was assessed through kinetic turbidimetric stability assays (Figure 2).



Compound	LogS	Solubility/μM	Result
1 ⁴⁺	1.138	13.7	Pass
Nicardipine	1.318	20.8	Pass

Figure 2. Kinetic turbidimetric solubility. The kinetic solubility of **1⁴⁺** measured through turbidimetry and compared with a positive control nicardipine. Turbidimetry was measured at 7 controls (0.2 – 100 μM) in DMSO (1 %). Samples were incubated for 5 minutes at

1
2
3 25 °C. Absorbance determined at 620 nm. N = 4 ± SD. Any complex
4
5 with a solubility of <1 µM is considered insoluble and therefore
6
7 fails the solubility assay.
8
9

10 Nicardipine - a drug used to treat high blood pressure and angina,
11
12 that is frequently employed in drug metabolism and pharmacokinetic
13
14 studies - was used as a positive control and it was found that **1**⁴⁺
15
16 exhibits solubility in the range of this control - see SI for
17
18 details. Under the Biopharmaceutics Classification System provided
19
20 by the FDA, which is used to predict intestinal drug absorption,
21
22 [**1**]Cl₄ is seen to be freely soluble⁵². As aqueous solubility is a
23
24 major factor in the bioavailability of antimicrobial compounds;
25
26 the high solubility of **1**⁴⁺ indicates that it is a good orally
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28 delivered drug candidate.
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33 **Resistance of *Escherichia coli* EC958 to clinical antibiotics**

34 The uropathogenic *E. coli* strain EC958 used in this study is a
35
36 sequence type ST131 isolate. Due to the presence of the extended-
37
38 spectrum β-lactamase gene *bla*CTX-M-15 on its pEC958 virulence
39
40 plasmid it has been designated as a Priority 1: Critical Pathogen
41
42 by the World Health Organization that urgently requires new
43
44 treatments.¹⁸ To confirm the categorization and antibiotic
45
46 resistance profile of this EC958 clinical isolate, phenotypic
47
48 testing of its sensitivity to several β-lactam antibiotics was
49
50 carried out. These studies - which included a monobactam and
51
52 different generations of cephalosporins, as well as various other
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major groups of antibiotics - were accomplished through a standardized EUCAST disc diffusion assay (Table 1).⁵³

The tests revealed that the strain shows resistance to all tested cephalosporins and the monobactam; aztreonam, thus confirming its Priority 1 categorization; however, it is still sensitive to carbapenems, tigecycline, fosfomycin and nitrofurantoin.

Table 1. EUCAST disc diffusion antibiotic sensitivity testing against *E. coli* strain EC958

Antibiotic	Disk content (µg)	Mean diameter ± SEM	zone (mm)	Sensitivity*
Cephalosporins				
Cefotaxime	3	10 ± 0		Resistant
Ceftazidime	30	9.5 ± 0.47		Resistant
Cefuroxime	30	0 ± 0		Resistant
Tetracyclines				
Tigecycline	15	23 ± 0		Sensitive
Monobactams				
Aztreonam	30	19.5 ± 0.47		Resistant

Carbapenems			
Doripenem	10	29 ± 0	Sensitive [#]
Meropenem	10	31.5 ± 0.47	Sensitive
Ertapenem	10	26 ± 0.47	Sensitive
Imipenem	10	29.7 ± 0.47	Sensitive
Aminoglycosides			
Gentamicin	10	15 ± 0	Resistant
Fluoroquinolones			
Ciprofloxacin	5	0 ± 0	Resistant
Levofloxacin	5	0 ± 0	Resistant
Miscellaneous			
Rifampicin	2	0 ± 0	Resistant
Fosfomycin	50	28.5 ± 0.82	Sensitive
Nitrofurantoin	100	22 ± 0	Sensitive

*Sensitivity or resistance determined by EUCAST breakpoint figures, 2019⁵⁴.#Sensitivity or resistance determined by EUCAST breakpoint figures, 2018⁵⁵

Growth and viability of growing *E. coli* strain EC958 cultures is diminished upon exposure to **1**⁴⁺

Having established the multidrug resistance properties of the model EC958 strain, we investigated its sensitivity to complex, **1**⁴⁺. Previously, we determined that the minimal inhibitory concentration (MIC) of **1**⁴⁺ was 2.8 μM⁴⁶, however the MIC is measured using very low turbidity cultures recently diluted from stationary phase. As treatment of most infections occurs when the bacterial load is already high and actively growing, we investigated what effects exposure to the compound had on actively growing cultures. Growth assays in which cultures in the early-exponential growth

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3 phase were injected with a range of concentrations of 1^{4+} below and
4
5 above the MIC were performed. As Figure 3 illustrates, very little
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7 effect on growth is observed within the first 30 minutes of
8
9 exposure to the compound. Subsequently, growth in the presence of
10
11 1^{4+} becomes inhibited for at least 2.5 h and the extent of growth
12
13 inhibition correlates with increasing concentration of the
14
15 compound. After this period, there is some recovery at
16
17 concentrations up to 1 μM , but above this concentration little
18
19 recovery in growth is seen. To understand whether 1^{4+} was affecting
20
21 viability or causing bacteriostasis at these concentrations,
22
23 viability assays were performed (Figure 3B). From 2 h post-
24
25 injection, a significant difference in viability is observed
26
27 between untreated cultures and those exposed to 0.5 - 5 μM of 1^{4+} .
28
29 In agreement with this data, fluorescence microscopy shows
30
31 accumulation of the Ru^{II} complex between 20 and 120 minutes and a
32
33 significant increase in fluorescence at the 60- to 120-minute
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35 timepoints, indicating further accumulation of the compound
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37 (Figure 3C). Increase in fluorescence intensity was measured using
38
39 the integrated density measurement function on ImageJ. At 60
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41 minutes the average corrected total cell fluorescence (CTCF) for
42
43 the cells was 9308.9, at 120 minutes the average CTCF for the cells
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45 was 29956.9. This is in agreement with previous findings indicating
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47 that ruthenium accumulates within the cell during this time.⁴⁶
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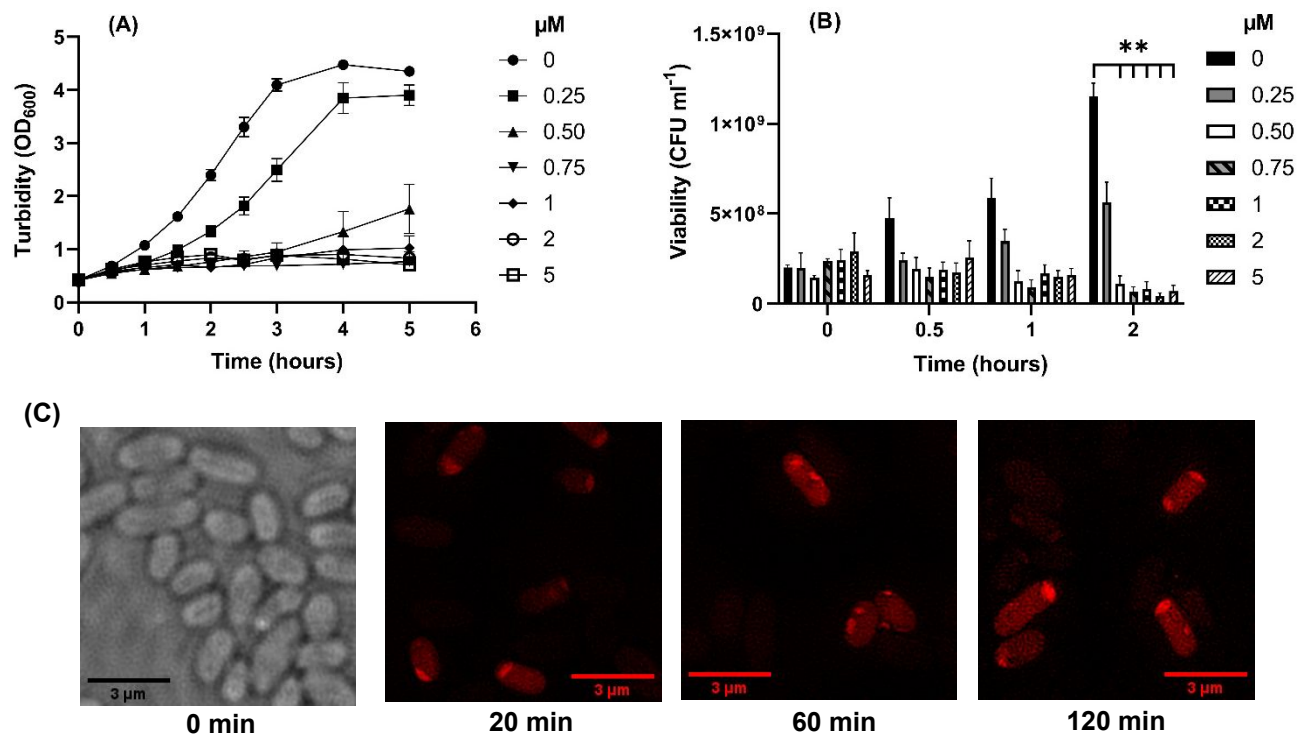
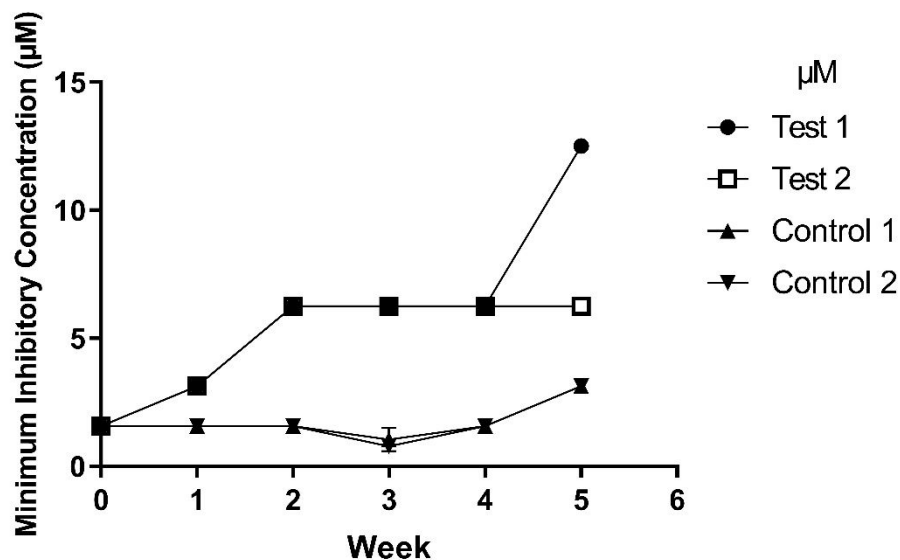


Figure 3. Effect of 1^{4+} concentration on the growth and viability of *E. coli* strain EC958 in glucose defined minimal medium (A) Cultures were grown to early-exponential phase at 37 °C and 150 rpm shaking in 250 ml flasks with defined minimal medium containing glucose as the sole carbon source. Upon reaching an OD₆₀₀ of ~0.4 differing concentrations of 1^{4+} (0 - 5 μM) were added to the cultures and growth was monitored at regular intervals. N ≥ 3 ± SEM. (B) Samples of the cultures described in (A) were removed and viable counts performed to determine the bactericidal effect of 1^{4+} at differing concentrations between 0 - 5 μM. N ≥ 3 ± SEM. Statistical significance (**) was determined with two-way ANOVA and Tukey's multiple comparisons (p < 0.05). (C) Localization of 1^{4+} in *E. coli* EC958 cells was visualized through structured

illumination microscopy at 0-, 20-, 60- and 120-min. Cells were imaged using the emission of 1^{4+} on excitation at 450 nm using A568 filter. After treatment with $0.8 \mu\text{M}$ 1^{4+} cells were washed with nitric acid before fixing with paraformaldehyde (4 %). Images were taken using a 1516 oil and SlowFadeTM Gold Antifade Mountant.

Evolution of resistance of *E. coli* strain EC958 to 1^{4+}

The potential of 1^{4+} as a putative antimicrobial lead would be enhanced if therapeutic resistance does not develop rapidly, nor to a high degree. To investigate this, we serially passaged EC958 cultures containing concentrations of 1^{4+} at half the minimal inhibitory concentration for five weeks. The WT MIC in minimal media ($1.5 \mu\text{M}$) was used for initiation of the experiment, new MIC assays were undertaken each week and changes to 1^{4+} concentrations were made if an MIC increase was observed to maintain the 0.5 x MIC concentration in the growing cultures (Figure 4).



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3 **Figure 4.** Evolution of 1^{4+} resistance of *E. coli* strain EC958 in
4 glucose defined minimal media. *E. coli* cultures were grown in
5 liquid GDMM containing 0.5 x MIC of 1^{4+} , cultures were passaged
6 every 24 h and weekly MIC's were determined. Experiments were
7 performed as biological duplicates and results displayed are
8 weekly mean MIC results ($N \geq 3 \pm SD$ for each biological repeat).
9 Control samples (upward and downward facing triangles) were
10 treated the same as test samples (\bullet , \square), except no compound was
11 added. Two-Way ANOVA with Tukey's multiple comparison test showed
12 significant differences between the original WT cultures and both
13 test cultures by week five and between the test and control
14 cultures in week five ($p < 0.0001$).

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31 Both independent cultures that were passaged in the presence of
32 1^{4+} showed a small increase in resistance over the first two weeks
33 of exposure with the MIC increasing from 1.5 μM to 6.1 μM . A
34 further increase in resistance was observed in one of the test
35 cultures after five weeks of exposure (MIC: 12.5 μM) whereas
36 resistance in the other test culture remained constant between
37 weeks two and week five. Significant differences were found between
38 both test cultures after five weeks exposure to 1^{4+} compared with
39 the initial MIC of the WT strain ($p < 0.001$). We observed a four-
40 to eight-fold increase in MIC over the course of five weeks
41 constant exposure to 1^{4+} . Additionally, tests were performed to
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3 determine whether resistance would evolve to 1^{4+} due to serial
4 passage in the absence of the ruthenium complex. Figure 4 shows a
5 minor increase in resistance to the compound in week five where
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10 1^{4+} was omitted, however this is a small increase, suggesting
11 exposure to the ruthenium complex is required to cause significant
12 resistance to 1^{4+} .
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17 Samples from all cultures were streaked weekly to check for
18 morphological changes or contaminants. Cultures exposed to the Ru^{II}
19 complex consistently showed different colony size and morphology
20 when subsequently streaked onto rich media in the absence of 1^{4+}
21 (Figure S-1). This effect was transient, as re-streaking of the
22 different sized colonies subsequently produced normal growth (data
23 not shown).
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33 All cultures in the evolution experiment depicted in Figure 4
34 were cryopreserved weekly. When cultures that had demonstrated a
35 four- to eight-fold increase in MIC were revived from these frozen
36 stocks and 1^{4+} susceptibility was retested, the MIC was reduced to
37 WT levels. This indicates that the strains may not have become
38 resistant to 1^{4+} via mutational change, instead it suggests that
39 altered expression of resistance genes such as those coding for
40 efflux pumps and modulated membrane permeability may be the cause
41 of the small increase in resistance after prolonged exposure to
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53 1^{4+} . Upon re-streaking from resistant cryopreserved stocks, these
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3 genes appear to revert to their pre-exposure expression levels,
4 resulting in the reduction in MIC to the original value.
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7 8 **Transcriptomic analysis reveals that membrane repair plays a significant role** 9 **in the response to 1⁴⁺ exposure**

10 To further study the effects of 1⁴⁺ on *E. coli* strain EC958 its
11 mechanism of action was probed through quantitative PCR. From the
12 study on bacterial growth after exposure to a range of
13 concentrations of 1⁴⁺ illustrated in Figure 3, we determined that
14 a final Ru^{II} complex concentration of 1.5 μ M would allow gene
15 expression to be accurately assessed. Significant cell death
16 occurs within 60-minutes of treatment when a lethal dose of the
17 compound is administered. We therefore monitored changes in gene
18 expression at timepoints ranging from 20 to 120 minutes to
19 determine whether early changes in the response to the compound
20 are altered after continued exposure.
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35 Cultures in glucose defined minimal media were grown in
36 triplicate to early-exponential phase, after which samples of the
37 culture were removed to act as the pre-exposure control.
38 Subsequently, at each timepoint, post-exposure samples were
39 removed from the culture, RNA was extracted, and the expression of
40 selected genes was compared to a reference.
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49 Prompted by our initial experiments indicating a dual mechanism
50 of action of 1⁴⁺,⁴⁶ the nine target genes selected for study have
51 functional roles in membrane permeability/stability and DNA repair
52 and a qPCR analysis was performed at all timepoints to assess
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expression of these genes after exposure to 1^{4+} . The reference gene *hcaT* was selected for this study as it is a well-defined, constitutively expressed, gene under many conditions and showed no significant change in expression in whole-transcriptome analyses of *E. coli* exposed to other ruthenium-based compounds.⁵⁶ In these experiments, it was found that three of the nine genes tested showed significant changes in expression upon exposure to 1^{4+} (Figure 5).

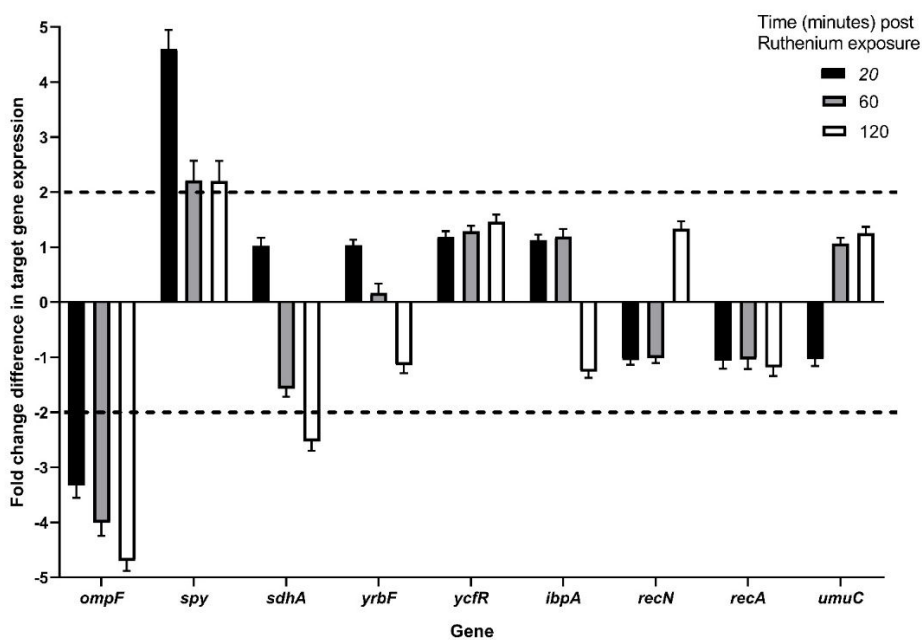


Figure 5. Relative *E. coli* gene expression levels of nine target genes after exposure to 1^{4+} show effects on membrane permeability and protein repair. The qPCR expression profiles of nine genes after exposure to $1.5 \mu\text{M}$ 1^{4+} are shown over a time course of 20- (black bars), 60- (grey bars) and 120-minutes (white bars). $N \geq 3 \pm \text{SEM}$. Expression levels were normalised against reference gene

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3 *hcaT*. Dashed horizontal lines represent significant expression
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5 changes (≥ 2 -fold change).
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9 A steady decrease in expression of the *ompF* gene was observed
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11 across the time-course. This gene encodes a non-specific porin
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13 found within *E. coli*⁵⁷ and its primary function is facilitate
14
15 passive diffusion of small hydrophilic molecules, including
16
17 tetracycline and fluoroquinolones, across the cell membrane.^{57,58}
18
19 Therefore, downregulation of this gene upon exposure to **1**⁴⁺
20
21 suggests an attempt by the cell to prevent uptake of the compound
22
23 by reducing membrane permeability. However, exposure of a wild
24
25 type *E. coli* strain and several porin knock-out mutants, including
26
27 *ompF* and *ompF-ompC* deletions⁵⁹ to **1**⁴⁺ showed no significant change
28
29 in the minimal inhibitory or bactericidal concentrations (Figure
30
31 S-2).
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36 Raised expression of the chaperone gene *spy* was observed over the
37
38 time-course, with initial rapid growth in expression slowly
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40 plateauing at a constant, increased level. These observations
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42 suggest that the Spy protein is rapidly required after exposure to
43
44 **1**⁴⁺. In Gram-negative bacteria, changes in external environment can
45
46 potentially affect the periplasmic space, resulting in unfavorable
47
48 conditions that cause proteins to aggregate and/or unfold and in
49
50 these circumstances chaperones like Spy are vital in maintaining
51
52 protein folding homeostasis.^{60,61}
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3 A significant decrease in expression of the *sdhA* gene was observed
4 after 120-minutes exposure to **1⁴⁺**. The succinate dehydrogenase
5 flavoprotein subunit, encoded by the *sdhA* gene, is a key protein
6 involved in the tricarboxylic acid cycle; generation of precursor
7 metabolites and aerobic respiration. Genes coding for key
8 components of aerobic respiration have previously shown to be
9 highly downregulated in *E. coli* when treated with another
10 ruthenium-containing compound.⁶²

11
12 To determine the wider effects of exposure to **1⁴⁺**, three other
13 genes were tested for changes in expression after exposure. These
14 were; the *yrbF* gene that encodes for a component of an ATP-binding
15 cassette transporter system that maintains lipid asymmetry in the
16 outer membrane which can be disrupted by chemicals or assembly
17 defects⁶³, *ibpA* that encodes for a small heat shock protein that
18 protects various proteins from thermal and oxidative stress⁶⁴, and
19 *ycfR* that encodes for a protein considered as both a biofilm
20 regulator and multi-stress response protein whose expression
21 increases as a result of multiple environmental changes⁶⁵. However,
22 the expression levels of none of these genes altered significantly
23 (by two-fold or greater) upon exposure to **1⁴⁺**.

24 **1⁴⁺ shows antimicrobial activity of against different strains of pathogenic** 25 **bacteria**

26 Given that antimicrobial resistance is a serious problem across
27 many species of pathogenic bacteria, and it is not always possible
28 to know the identity of the infecting pathogen prior to initiating

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3 treatment, it is important to understand the spectrum of activity
4
5 of **1⁴⁺**. As our previous work has indicated that this complex appears
6
7 to be more active in Gram-negative bacteria,⁴⁷ we explored its
8
9 activity against a wider spectrum of multidrug and pandrug
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11 resistant *E. coli* strains as well as other Gram-negative pathogenic
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13 species- Table 2. The *E. coli* strains include a KPC-producing
14
15 (*Klebsiella pneumonia carbapenase*) *E. coli*², an avian pathogenic
16
17 *E. coli*, the *E. coli* EC958 strain (for comparison) and an
18
19 antimicrobial testing control strain NCTC 12923. The other Gram-
20
21 negative pathogens included in this panel were clinical isolates
22
23 of *Pseudomonas aeruginosa*, *Salmonella kedougou*, *Shigella flexneri*,
24
25 *Enterobacter hormaechei*, *Citrobacter koseri* and *Acinetobacter*
26
27 *baumannii*. Interestingly, the potent activity of **1⁴⁺** against the
28
29 control strain is retained in all the tested drug resistant
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31 strains, including carbapenem resistant pathogens, indicating its
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33 broad spectrum of activity.
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42 **Table 2.** Minimum inhibitory concentration for **1⁴⁺** demonstrate
43
44 significant antimicrobial activity against a variety of bacterial
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46 pathogens
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Bacterial Species Strain	Type /	MIC (μ M)
<i>E. coli</i> EC958	Multidrug resistant, clinical isolate	1.7

<i>E. coli</i> NCTC12923	Antimicrobial strain	control	2.4
<i>E. coli</i> EC_160_KPC2	Carbapenem resistant		2.4
Avian pathogenic <i>E. coli</i>	Multidrug resistant		1.6
<i>P. aeruginosa</i>	Clinical isolate		1.9
<i>K. pneumoniae</i>	Clinical isolate		1.6
<i>S. kedougou</i>	Clinical isolate		1.4
<i>S. flexneri</i>	Clinical isolate		4.2
<i>E. hormaechei</i>	Clinical isolate		1.7
<i>C. koseri</i>	Clinical isolate		1.1
<i>A. baumannii</i>	Clinical isolate		1.6

Discussion

Examination of the antibiotic susceptibility profile of this uropathogenic *E. coli* strain demonstrates its significant multidrug resistance. Current treatment recommendations for uncomplicated urinary tract infections where antibiotic therapy is indicated include the administration of nitrofurantoin, trimethoprim / sulfamethoxazole, fluoroquinolones, fosfomycin or oral β -lactam agents. Clinicians are guided in their choice of treatment by local susceptibility patterns of *E. coli* and other uropathogens as strain specific antimicrobial susceptibility profiles are usually not determined ⁶⁶. The EC958 strain tested here showed resistance to several β -lactam antibiotics and both fluoroquinolones tested, however both nitrofurantoin and fosfomycin were shown to be effective (Table 2). Much of the

1
2
3 antibiotic resistance profile demonstrated herein is expected when
4
5 interrogating the whole genome sequence of this organism ¹⁸.
6
7 Totsika *et al*, report the presence of the pEC958 plasmid in this
8
9 strain that contains multiple antibiotic resistance genes. The
10
11 presence of *tetA* and *tetR* found on the pEC958 plasmid provide
12
13 resistance to early members of the tetracyclines, however do not
14
15 confer resistance to the third-generation tigecycline tested here.
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19 ⁶⁷ ⁶⁸ Resistance to the fluoroquinolones and aminoglycosides can be
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21 explained by the presence of *aac(6')-Ib-cr*, encoding an
22
23 aminoglycoside acetyltransferase capable of causing resistance to
24
25 aminoglycosides and fluoroquinolones via modification of the drug.
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28 ⁶⁹ ⁷⁰ ⁷¹ However, interestingly no obvious genomic explanation for
29
30 rifampicin resistance could be found. Resistance to rifampicin
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32 commonly occurs via mutation of the *rpoB* gene and resistance is
33
34 easily evolved. ⁷² Therefore our strain of EC958 may be a variant
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36 of the one described previously^{18,73} that has acquired this
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38 mutation. With no apparent carbapenem resistance genes in the
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40 genome sequence and EC958 displaying sensitivity to all four types
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42 tested this drug class would provide a good treatment option in
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44 the case of disease progression from EC958 causing a urinary tract
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46 infection to pyelonephritis and bloodstream infection.
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51 Given the extensive multidrug resistance of this, and many other
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53 bacterial pathogens, new antimicrobial leads that do not readily
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55 succumb to evolving resistance are urgently needed. The rate at
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3 which resistance to an antimicrobial arises is dependent on its
4 mechanism of action and whether single or multiple changes are
5 required for significant resistance to arise. We compared the
6 resistance changes of EC958 when exposed to 1^{4+} with literature
7 reports of resistance evolution in *E. coli* exposed to commonly
8 used antibiotics. One study showed that continuous exposure to
9 levofloxacin caused *E. coli* to increase resistance to this
10 antibiotic by 16-fold within the first 24 h of exposure and 64-
11 fold after 14 days. The mechanism of resistance was identified as
12 mutation of targets of the fluoroquinolone and changes in membrane
13 permeability.⁷⁴ Resistance to fluoroquinolones such as levofloxacin
14 only requires a single point mutation in DNA gyrase to emerge, so
15 the rapid increase observed in this study is not surprising. A
16 separate study exposed *E. coli* to three antibiotics: trimethoprim,
17 chloramphenicol and doxycycline, in different evolution
18 experiments. Trimethoprim, an antibiotic commonly used to treat
19 urinary tract infections, showed a consistent increase in
20 resistance of around 1,680-fold after 20 days of exposure.
21 Increases of 870- and 10-fold were observed against
22 chloramphenicol and doxycycline respectively⁷⁵. One further study
23 found that clinical resistance could be evolved in previously
24 susceptible uropathogenic *E. coli* strains to ciprofloxacin,
25 amoxicillin and aminoglycosides by as little as one, two and three
26 to five day(s) of passage respectively⁷⁶. Thus, the evolution of
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3 resistance to $\mathbf{1}^{4+}$ is slow in comparison to many clinically available
4 antibiotics. The slow and low level resistance gains of EC958
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6 against $\mathbf{1}^{4+}$ may be due to its multi-mechanism mode of action which
7
8 targets both the cell membrane and intracellular targets^{46,47}.
9
10 Therefore, it seems likely that resistant isolates would take
11
12 significantly longer to arise, further adding promise to $\mathbf{1}^{4+}$ as a
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14 therapeutic tool.
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19 In addition to gaining further insight into the potential for
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21 resistance to the Ru^{II} complex, it is important to fully understand
22
23 how $\mathbf{1}^{4+}$ causes bacterial growth inhibition and cell death. Figure
24
25 2 demonstrates that very little impact on growth is observed within
26
27 the first 30 minutes post-injection of the compound. This is in
28
29 agreement with previous stimulated emission depletion imaging that
30
31 identified a 20-minute period when compound accumulation at cell
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33 membranes occurs before localisation at cell poles⁴⁶. Alexa Fluor
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35 NHS-ester-405 counterstaining experiments also demonstrated that
36
37 membrane damage is not present within at least the first five
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39 minutes of exposure to $\mathbf{1}^{4+}$ with subsequent bacterial inner and
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41 outer membrane damage identified 60 minutes post-exposure. $\mathbf{1}^{4+}$.
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46 To gain insight into how EC958 senses and responds to the presence
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48 of $\mathbf{1}^{4+}$ at a transcriptomic level, we selected nine genes associated
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50 with the proposed targets of the complex; the bacterial membranes
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52 and DNA, to better understand the bacterial response to membrane
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54 damage and to determine whether DNA is a secondary target.
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3 We observed a steady decrease in expression of *ompF* upon exposure
4 to **1**⁴⁺, OmpF is a non-specific porin found within *E. coli*⁵⁷. This
5 porin allows passive diffusion of small hydrophilic charged
6 molecules and antibiotics including tetracyclines and
7 fluoroquinolones across the cell membrane ⁵⁷. As downregulation of
8 *ompF* causes resistance to multiple antibiotics, this change in
9 gene expression may be a cause of the transient resistance gained
10 during prolonged exposure to **1**⁴⁺ (Figure 2)⁷⁷. However, with upper
11 limits for molecular weight; the cut-off for porins of *E. coli* is
12 around 600 Da.⁷⁸ As the molecular weight of **1**⁴⁺ is considerably
13 larger than this, OmpF mediated transport is unlikely to be a
14 mechanism of passage into the bacterial cell⁴⁶. Aside from the
15 generalized porins, there are solute-specific facilitated
16 diffusion channels through the outer membrane into the periplasm
17 that allow solutes to bypass the porin-specific size and charge
18 requirements, **1**⁴⁺ may utilize one of these routes to gain entry to
19 the cell⁷⁸.

20
21 An increase in expression of the *spy* gene was observed upon
22 exposure to **1**⁴⁺. Spheroplast protein Y (Spy) is a non-ATP dependent
23 periplasmic chaperone, vital in maintaining the homeostasis of
24 protein folding under cellular stress ^{60,61}. In Gram-negative
25 bacteria, such as EC958, changes in external environment impact
26 the periplasmic space, resulting in unfavorable conditions for
27 proteins leading to aggregation and unfolding. In these conditions

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3 *spy* expression is upregulated to help prevent unfolding and
4 aggregation and to re-fold substrates without ATP. Stresses that
5 have been found to cause induction in the *spy* gene include ethanol,
6 indole, tannins and metals such as zinc, copper and other
7 ruthenium-based complexes with differing mechanistic
8 actions^{62,79,80}. In this case, the upregulation of *spy* upon treatment
9 with **1**⁴⁺ is likely due to the compound damaging the bacterial
10 membrane and altering the conditions of the periplasmic space.
11 Therefore, this increase in expression may be an attempt by the
12 cell to regain the balance of protein aggregation and unfolding in
13 this region. The expression of *spy* is controlled by the two-
14 component systems CpxAR and BaeSR; both responsible for regulation
15 of the envelope stress response. Accumulation of **1**⁴⁺ at the poles
16 of the cell is demonstrated herein (Figure 3C) and supports
17 previous findings⁴⁶. This transcriptomic data provides further
18 evidence that the membrane is a major site of activity for this
19 compound, which appears to selectively target and damage bacterial
20 membranes but not to damage mammalian cells nor demonstrate
21 significant toxicity in animal models at similar concentrations⁴⁶.
22 This is an important finding in the selectivity and therapeutic
23 potential of **1**⁴⁺.

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After two hours of exposure to **1**⁴⁺ a significant decrease in the
expression of *sdhA* was observed. The succinate dehydrogenase
flavoprotein subunit A (SdhA) is part of the succinate

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3 dehydrogenase enzyme complex, bound to the inner surface of the
4 cytoplasmic membrane its primary functions are to catalyze the
5 oxidation of succinate to fumarate in the citric acid cycle and to
6 participate in the aerobic electron transport pathway to generate
7 energy for the cell by oxidative phosphorylation ⁸¹. At two hours
8 **1**⁴⁺ has penetrated the membranes of *E. coli* and entered the cell
9 where it could potentially cause significant intracellular damage,
10 therefore it is likely that treated bacteria downregulate various
11 metabolic pathways in order to conserve energy and prevent the
12 production of further potentially harmful species, causing the
13 reduction in *sdhA* expression.
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28 We previously showed that **1**⁴⁺ binds intracellular targets at 60-
29 minutes^{46,47}. To decipher whether this binding could cause damage
30 to bacterial DNA we monitored the regulation of three genes that
31 would be upregulated as a response to DNA damage and the SOS
32 response: *recA*, *recN* and *umuC*. No significant change was observed
33 in any of these genes suggesting that the bacteria did not produce
34 an SOS response. Therefore, at the low compound concentrations
35 used, DNA damage is unlikely to be a target for **1**⁴⁺ in *E. coli*. The
36 gene expression levels of three further proteins were examined
37 upon exposure to **1**⁴⁺. The protein expressed by *ycfR* is a biofilm
38 regulator. It is a multi-stress response protein, expression of
39 which increases as a result of environmental changes.⁶⁵ The gene
40 *yrbF* encodes for a component of an ATP-binding cassette transporter
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3 system, which is one mechanism that the cell may use to maintain
4 lipid asymmetry in the outer membrane when chemically disrupted⁶³.
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6 Finally, the heat shock protein gene, *IbpA* protects various
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8 substrates and proteins when the cell undergoes thermal and
9
10 oxidative stress⁶⁴. Interestingly, no significant change in gene
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12 expression was observed for these genes after exposure to **1**⁴⁺
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14 indicting that exposure to the complex results in a relatively
15
16 specific disruption to cellular function.
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21 This gene expression data provides further evidence to support
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23 the hypothesis that the bacterial cell membrane is a significant
24
25 target of **1**⁴⁺ with disruption to the stability of periplasmic
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27 proteins being a major stressor for the cell. The pathogen screen
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29 for sensitivity to this ruthenium complex showed significant
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31 activity against several Gram-negative species, each of these
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33 species also contain homologues of the *spy* and *sdhA* genes, so it
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35 is likely that these pathogens will elicit similar responses upon
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37 exposure (Table 2). However, the outer membrane proteins expressed
38
39 by the Gram-negative pathogens differ significantly between the
40
41 species tested, with *P. aeruginosa* containing a porin of unusually
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43 low permeability (OprF) and *A. baumannii* containing an *E. coli*
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45 OmpA homologue (OmpA_{AB}). These porins are thought to contribute to
46
47 the intrinsic antibiotic resistance of these species, but as shown
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49 in Table 2 they do not appear to contribute to resistance against
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51 **1**⁴⁺. Recently, Smitten et al examined the differing activity of **1**⁴⁺
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3 against Gram positive bacterial pathogens⁴⁷ demonstrating that the
4 level and type of teichoic acids present within the Gram positive
5 cell wall has a significant impact on the antimicrobial activity
6 of the compound. Testing antimicrobial activity of **1**⁴⁺ against
7 several Gram negative bacterial species has confirmed the broad-
8 spectrum activity of this compound against Gram negative
9 pathogens, including clinical isolates and multi-drug resistant
10 strains (Table 2) further demonstrating the potential of this
11 compound for future use as a therapeutic in the fight against
12 antimicrobial resistant infections.
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27 Conclusions

28 As with previous studies with **1**⁴⁺ and the EC958 pathogenic strain
29 of *E. coli*, it was found the compound is highly potent against
30 several pathogenic *E. coli* strains and other Gram-negative
31 pathogens. Although a small rise in resistance was observed over
32 five weeks of exposure, the slow rate and relatively low level of
33 evolution in comparison to that of clinically available organic
34 antibiotics makes **1**⁴⁺ a strong potential candidate for
35 antimicrobial therapeutics. In addition, the good kinetic
36 solubility indicates the compound will have good bioavailability
37 and could be used as an oral antimicrobial. Transcriptomic analysis
38 suggests that **1**⁴⁺ does cause damage both to the inner and outer
39 membrane, resulting in unfavorable, stressful conditions in the
40 periplasmic space. Together this work adds to the growing volume
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3 of research supporting the hypothesis that this class of Ru^{II}
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5 complexes can be developed into effective antimicrobial drugs.
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8 Future research will further consider the potential emergence
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10 of resistance to this lead with a particular focus on mechanisms
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12 of cellular entry and efflux and any proteins capable of binding
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14 to the complex making it unavailable to bind other more vulnerable
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16 targets within the cell. Such studies will facilitate the
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18 circumvention of these mechanism if they prove to be potential
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20 routes of resistance.
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24 **Materials and Methods**

25 **Bacterial strains and culture conditions**

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27 Strains used in this study are listed in Table 1, the primary
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29 strain used was a CTX-M-15 type extended spectrum β -lactamase
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31 (EBSL)-producing clinical isolate *E. coli* EC958 (ST131)⁷³. Bacteria
32
33 were routinely grown aerobically at 37 °C in either Mueller-Hinton
34
35 broth (Sigma Aldrich, UK) or defined minimal medium with glucose
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37 as the sole carbon source (GDMM) as described previously⁴⁶. Prior
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39 to experiments, bacterial starter cultures were prepared by
40
41 inoculating the appropriate liquid medium with colonies from a
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43 fresh agar plate and incubation at 37 °C with shaking at 180 rpm
44
45 for approximately 18 h. Where necessary cultures were
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47 cryopreserved in a sterile suspension of 25 % glycerol : 75 %
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49 Mueller-Hinton broth (v/v) at -80 °C.
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Preparation and storage of **1**⁴⁺

The **1**⁴⁺ compound was synthesised as previously described⁴⁶. Stocks solutions of **1**⁴⁺ were made to a concentration of 5 mg ml⁻¹ in sterile deionised water and were stored at room temperature protected from light.

Disc Diffusion Assay

Disk diffusion assays were performed in accordance with European Committee on Antimicrobial Susceptibility Testing (EUCAST) guidelines⁵³. After incubation the diameter of the zones of inhibition were measured and compared to EUCAST breakpoints.^{54,55}

Determination of minimal inhibitory concentrations

MICs of **1**⁴⁺ were determined via the standard broth-dilution method in 96-well microtiter plates in glucose defined minimal media. The MIC was evaluated using 2-fold decreasing concentrations of the compound between 50 to 0.09 µM against starter cultures diluted to an OD₆₀₀ of 0.05 - 0.1 (equivalent to a 0.5 McFarland standard). Plates were incubated at 37 °C for 16 - 20 h after which the presence or absence of growth in each well was observed to determine the minimal inhibitory concentration of **1**⁴⁺ for each of the strains tested.

Bacterial growth and viability measurements

Cultures were incubated in 250 ml conical flasks from 2.5 % starter cultures of *E. coli* EC958 in 30 ml GDMM media at 37 °C with shaking at 150 rpm. Once cultures reached an early exponential growth phase (OD₆₀₀ ~0.4) **1**⁴⁺ was added at the appropriate final

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3 concentrations and growth monitored at regular intervals for the
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5 subsequent five hours. Viability was measured for samples
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7 harvested at the relevant time points after serial dilution in
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9 phosphate-buffered saline by plating 10 μ l aliquots on Mueller-
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11 Hinton agar and incubation overnight at 37 °C.
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14 15 **Structured illumination microscopy**

16 Samples were prepared and analysed as previously described ⁴⁶.
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18 Briefly, *E. coli* cultures were grown in GDMM to an OD₆₀₀ of 0.3-
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20 0.4 and **1⁴⁺** was added to a final concentration of 0.8 μ M. Samples
21
22 were incubated for 0-, 20-, 60- and 120-min before washing with
23
24 nitric acid and fixation with paraformaldehyde (4 %). Localization
25
26 of **1⁴⁺** in *E. coli* EC958 cells was visualized through structured
27
28 illumination microscopy using a 1516 oil and SlowFade™ Gold
29
30 Antifade Mountant. Cell fluorescence was measured using ImageJ by
31
32 selecting a cell and measuring area, integrated density and mean
33
34 grey value. Background fluorescence was determined and corrected
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36 for in each image. Four cells were measured per image. The
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38 corrected total cell fluorescence was calculated by CTCF =
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40 Integrated Density - (Area of selected cell X mean fluorescence of
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42 background readings).
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48 **Evolution of resistance assays**

49 After determination of the MIC for **1⁴⁺**, a 1 % inoculum of starter
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51 culture incubated in GDMM was added to 10 ml of GDMM. Compound was
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53 then added where appropriate to the culture at 0.5 times the MIC
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55 allowing growth of the strain in sub-MIC exposure. Control cultures
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3 were set up in the same manner in the absence of **1⁴⁺**. Cultures were
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5 incubated at 37 °C and 180 rpm for 24 h, after which a 1 % inoculum
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7 from this culture was used to inoculate a fresh tube of GDMM + /
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9 - **1⁴⁺**. This process was repeated every 24 h of the lifetime of the
10
11 experiment. Each week cultures were cryopreserved and re-tested
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13 for their MIC with the amount of compound added to subsequent
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15 cultures adjusted as required to maintain a 0.5 times MIC.
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19 Quantitative PCR analysis

20 Cells were grown as described above in GDMM. At the designated
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22 times pre- and post- injection of a final concentration of 1.5 µM
23
24 **1⁴⁺**, 5 ml samples of culture were removed into 10 ml volumes of RNA
25
26 Protect (Qiagen, UK) in triplicate for RNA stabilisation. RNA was
27
28 extracted using the RNeasy Mini Kit from Qiagen according to
29
30 manufacturer's instructions and qPCR performed as previously
31
32 described⁸² using a QuantStudio™ 3 Real Time PCR System (Applied
33
34 Biosystems). Primers used for PCR can be found in supplementary
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36 information.
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41 Kinetic Turbidimetric Solubility assays

42 Aqueous solubility was measured using a high throughput
43
44 turbidimetric assay. A 10 mM stock of each compound (nicardipine
45
46 hydrochloride and **1⁴⁺**) were made up in DMSO. This stock was diluted
47
48 in phosphate buffered saline (PBS pH 7.4) to give concentrations
49
50 (µM): 0.4, 2, 4, 20, 40, 100 and 200. The final DMSO concentration
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52 = 1 %. Each concentration was plated out in triplicate. The
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3 solutions were incubated for 2 h at 37 °C then the absorbance read
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5 on a plated reader at 620 nm.
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8 Statistical analyses

9 GraphPad Prism v7.05 software was used for statistical analysis
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11 tests include one-way ANOVA, two-way ANOVA With Turkey's multiple
12
13 comparison tests and Welch's T-test.
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16 Author contributions

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22 ‡These authors contributed equally to this manuscript
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25 Supporting Information

26 Supporting methods: chemistry methods for [Ru(3,4,7,8-tetramethyl-
27 1,10-phenanthroline)₂Cl₂]²⁺, [Ru(TMP)₂]₂(tpphz)]⁴⁺, Anion
28 metathesis.
29

30 Supporting data: S Table 1 - Primer sequences designed for
31 transcriptomic analysis, S Table 2 - Kinetic turbidimetric
32 solubility plate readings, Figure S-1 - Variation in colony size
33 of *Escherichia coli* after exposure to 1⁴⁺. Figure S-2 - MIC and MBC
34 assays of *Escherichia coli* porin mutants, compare to the wild type.
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45
46 that have greatly improved the quality of this report.
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52 strains for testing.
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3 ABREVIATIONS USED
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5	UPEC	uropathogenic <i>Escherichia coli</i>
6		
7	PCR	polymerase chain reaction
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9	qPCR	quantitative polymerase chain reaction
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11	DMSO	dimethyl sulfoxide
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13	FDA	Food and Drug Administration
14		
15	EUCAST	European committee on antimicrobial susceptibility tests
16		
17	MIC	minimal inhibitory concentration
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19	SEM	scanning electron microscopy
20		
21	ANOVA	analysis of variance
22		
23	WT	wild type
24		
25	GDMM	glucose defined minimal medium
26		
27	ATP	adenosine triphosphate
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29	KPC	<i>Klebsiella pneumoniae</i> carbapenemase
30		
31	NCTC	National Collection of Type Cultures
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33	PBS	phosphate buffered saline
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35	TMP	3,4,7,8-tetramethyl-1,10-phenanthroline
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37	TPPHZ	tetrapyridophenazine
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39	DMF	dimethylformamide
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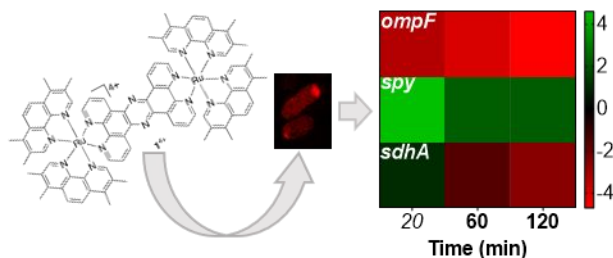
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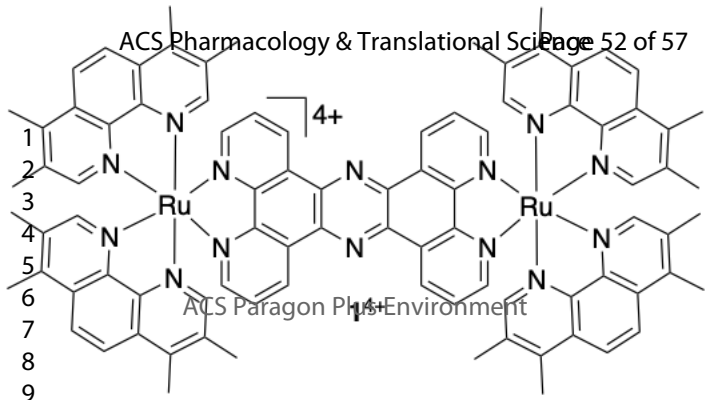
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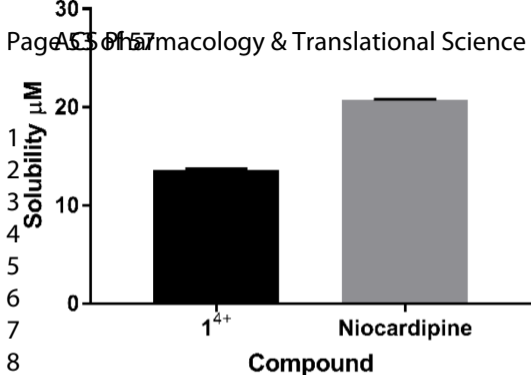
Title: A transcriptomic Analysis of the Activity and Mechanism of Action of a Ruthenium(II)-Based Antimicrobial That Induces Minimal Evolution of Pathogen Resistance

Authors: Adam M Varney, Kirsty L Smitten, Jim A Thomas and Samantha McLean

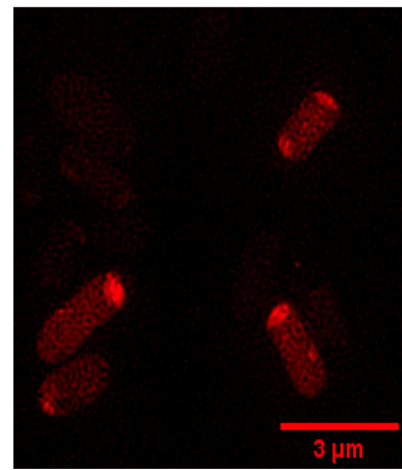
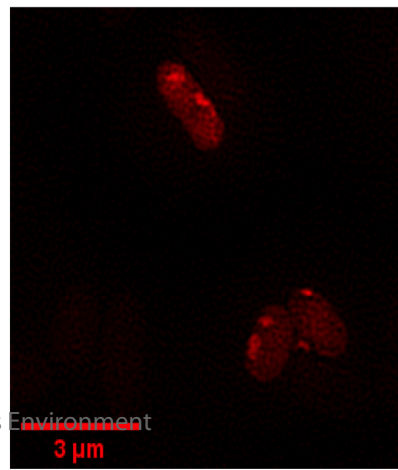
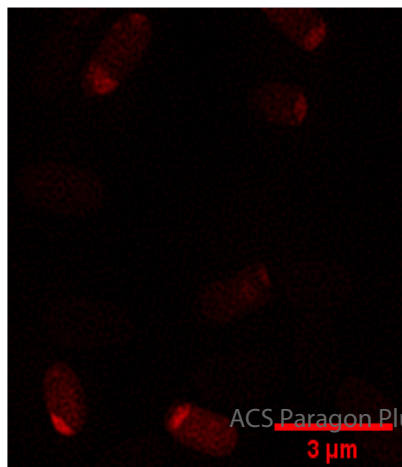
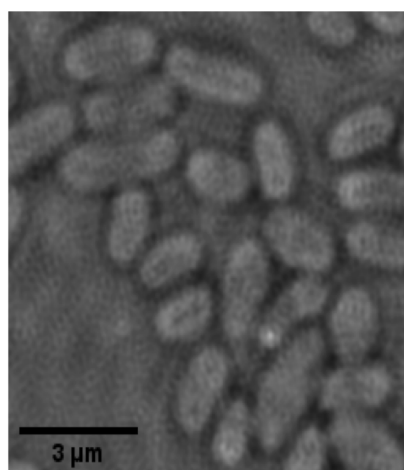
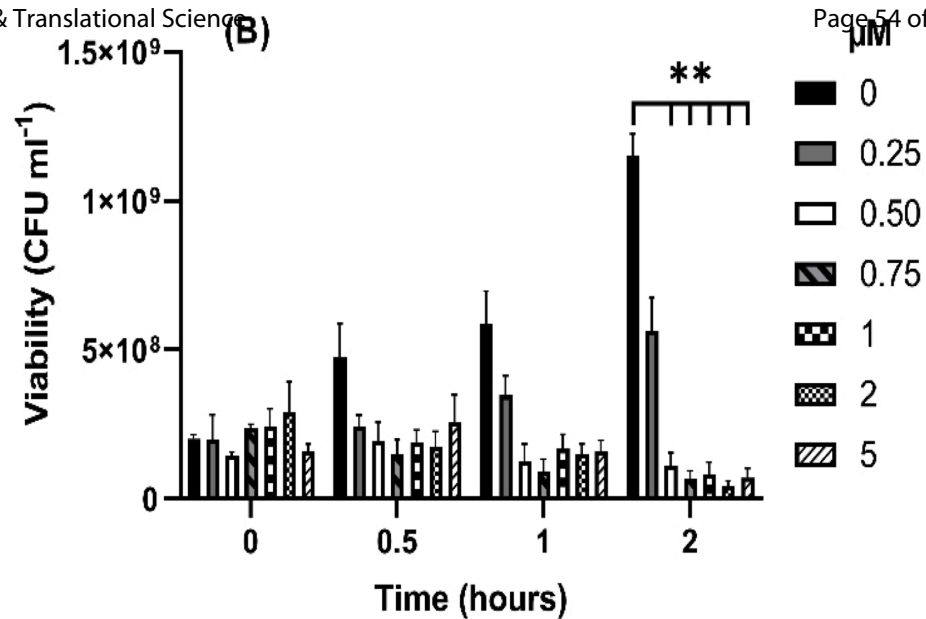
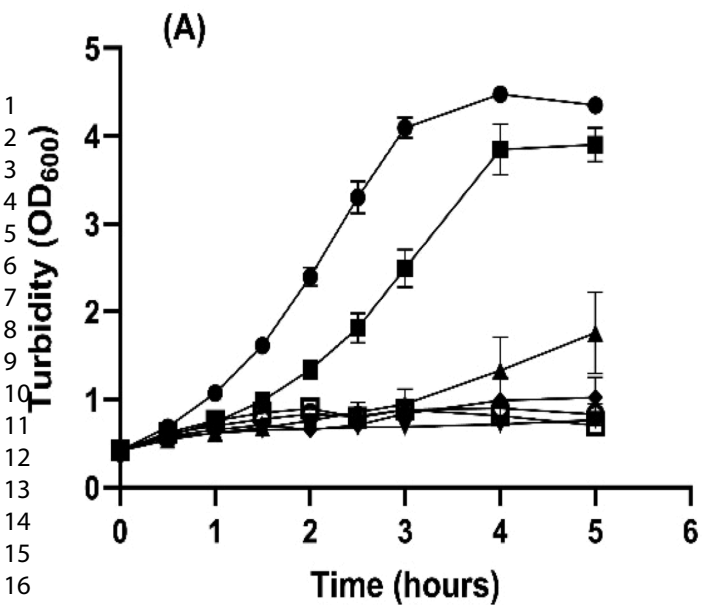
Brief synopsis: Addition of the lead compound to *Escherichia coli* EC958 causes changes in gene expression within the cell. Expression of membrane repair protein, *spy* is increased whilst expression of outer membrane protein, *ompF* and succinate dehydrogenase, *sdhA* is decreased.

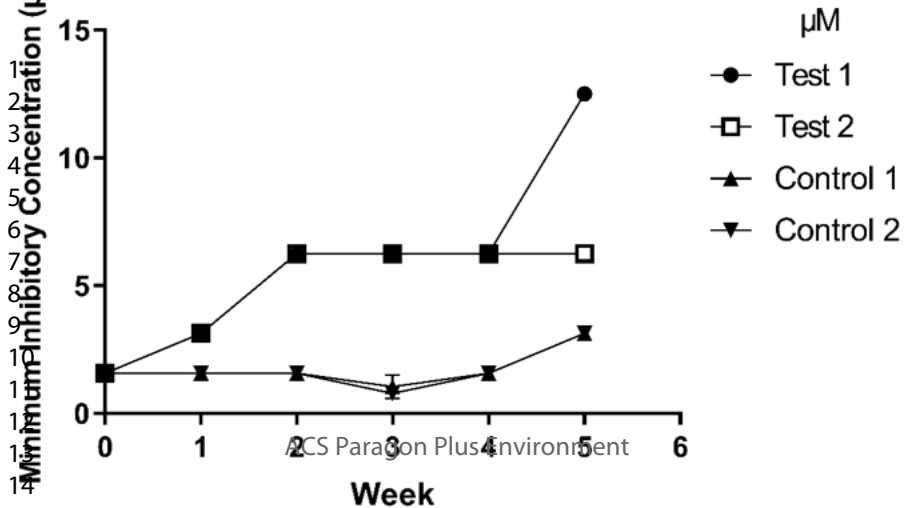


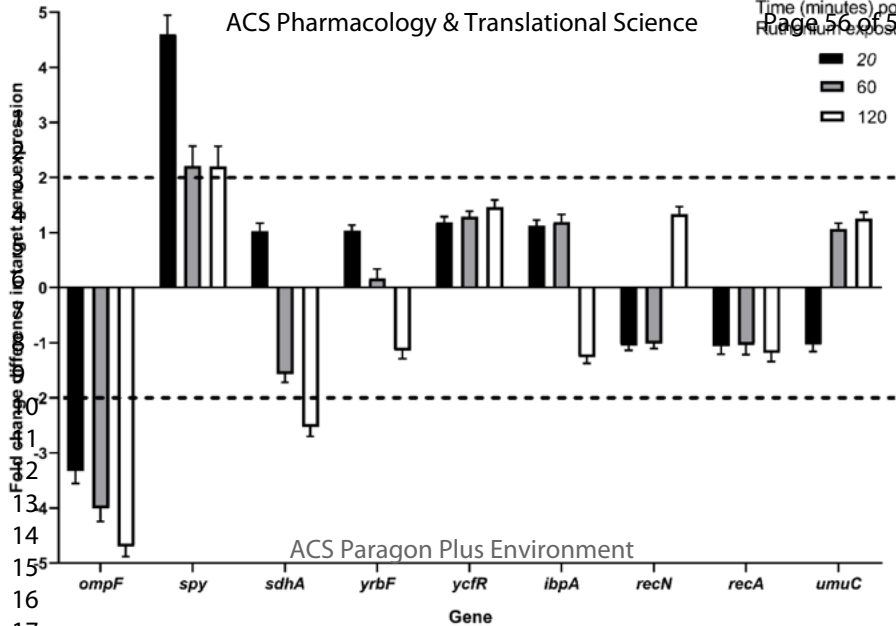




Compound	LogS	Solubility/µM	Result
1 ⁴⁺	1.138	13.7	Pass
Nicardipine	1.318	20.8	Pass

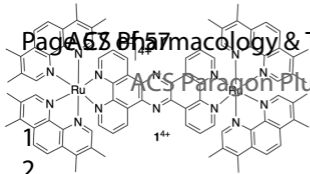






ACS Paragon Plus Environment

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Bacterial Species	Active?
<i>E. coli</i>	✓
<i>P. aeruginosa</i>	✓
<i>K. pneumoniae</i>	✓
<i>S. kedougou</i>	✓
<i>S. flexneri</i>	✓
<i>E. hormaechei</i>	✓
<i>C. koseri</i>	✓
<i>A. baumannii</i>	✓