Antibiotic Resistance Is Prevalent in an Isolated Cave Microbiome

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Abstract

Antibiotic resistance is a global challenge that impacts all pharmaceutically used antibiotics. The origin of the genes associated with this resistance is of significant importance to our understanding of the evolution and dissemination of antibiotic resistance in pathogens. A growing body of evidence implicates environmental organisms as reservoirs of these resistance genes; however, the role of anthropogenic use of antibiotics in the emergence of these genes is controversial. We report a screen of a sample of the culturable microbiome of Lechuguilla Cave, New Mexico, in a region of the cave that has been isolated for over 4 million years. We report that, like surface microbes, these bacteria were highly resistant to antibiotics; some strains were resistant to 14 different commercially available antibiotics. Resistance was detected to a wide range of structurally different antibiotics including daptomycin, an antibiotic of last resort in the treatment of drug resistant Gram-positive pathogens. Enzyme-mediated mechanisms of resistance were also discovered for natural and semi-synthetic macrolide antibiotics via glycosylation and through a kinase-mediated phosphorylation mechanism. Sequencing of the genome of one of the resistant bacteria identified a macrolide kinase encoding gene and characterization of its product revealed it to be related to a known family of kinases circulating in modern drug resistant pathogens. The implications of this study are significant to our understanding of the prevalence of resistance, even in microbiomes isolated from human use of antibiotics. This supports a growing understanding that antibiotic resistance is natural, ancient, and hard wired in the microbial pan genome.

Introduction

The question of whether the extensive presence of resistance elements in microbes is primarily the result of human activity is controversial. Investigation of antibiotic resistance in bacteria from the Galapagos, a remote environment with limited human occupation and presumably low anthropogenic antibiotic exposure, revealed that acquired antibiotic resistance genes were absent in bacteria isolated from terrestrial animals [1]. Similarly, plasmids from bacterial collections that predate the antibiotic era are largely devoid of resistance elements [2,3]. This suggests that the presence of antibiotics is an important selective force in evolution and spread of antibiotic resistance genes and can contribute significantly in altering the natural microbiota. In a survey of soil samples from the Netherlands spanning the pre-and post-antibiotic eras (1940–2008), an increase in the relative abundance of antibiotic resistance genes for major antibiotic families (ß-lactams, tetracyclines and macrolides) was observed in contemporary soil samples in comparison to pre-antibiotic era samples [4]. This body of evidence is consistent with the hypothesis that widespread resistance is a modern phenomenon linked to human use of antibiotics.

On the other hand, antibiotics and antibiotic biosynthetic pathways are believed to have evolved over millions of years suggesting that antibiotic resistance is an equally ancient phenomenon [5,6,7]. Indeed, we have recently shown that antibiotic resistance elements were abundant and diverse in ancient DNA dating from the Pleistocene (30,000 years ago) [8]. The concept of the antibiotic resistome predicts that resistance is the result of dynamic and competitive microbial interactions that pre-date human use of antibiotics [9,10]. Consistent with this notion is our survey of contemporary soil actinomycetes that reported widespread multidrug resistance even in the absence of obvious human sources of antibiotics [11]. There have also been reports of antibiotic resistance in microorganisms isolated from extreme natural habitats including the deep terrestrial subsurface [12] and the deep ocean [13]; environments presumably largely absent of human influence. These studies support a hypothesis that resistance is an ancient and genetically rich natural phenomenon, deeply embedded in the microbial pan genome.

One of the challenges in measuring contemporary environmental resistance is rigorously ensuring the absence of anthropogenic sources of antibiotics as a selective pressure for the acquisition of ‘modern’ resistance genes. A survey of a diverse
environmental bacterial population that has never been exposed to modern antibiotics and resistance genes would unequivocally establish the depth of the contemporary environmental resistome. Lechuguilla Cave offers a unique environment for such a study.

Lechuguilla Cave, located within Carlsbad Caverns National Park (USA), was formed in the Capitan Formation of the Delaware Basin by hypogenic (ascending water) sulfuric acid speleogenesis in the last 7 million years [14,15,16], resulting in the formation of a very large (>200 km) and deep (>500 m), maze-like cave system [17]. While the cave is still connected to the artesian aquifer that formed it (Figure 1), the majority of the cave became isolated as the aquifer dropped [18]. Due to its isolated hypogene nature, extensive cave development occurred primarily at −300 to −400 m, while the impermeable Yates Formation that overlies the cave limits vadose water from entering the system [14,17,19]. The deep recesses of Lechuguilla Cave, isolated from surface input for the past 4–7 million years, therefore provide a unique environment to study the presence and prevalence of antibiotic resistance elements.

We report here a sampling of the culturable Lechuguilla Cave resistome—the antibiotic resistance genes found in a culture collection of isolates from this cave environment. In addition to finding a wide array of resistance in this culture collection, we also describe a novel macrolide kinase variant and an inducible daptomycin hydrolase, demonstrating not only that resistance is prevalent in the absence of anthropogenic antibiotics, but hitherto undiscovered mechanisms of resistance that have potentially important clinical implications are prevalent in the environment.

![Figure 1. Plan and profile maps of Lechuguilla Cave, Carlsbad Caverns National Park, New Mexico.](image)

The sites where microbial strains were collected (LCECE, LCDS1 and LCEA1) are shown relative to the entrance and depth. tN represents true North on the plan, while the profile has an exaggerated vertical profile of 1.5×. doi:10.1371/journal.pone.0034953.g001
Results

Sampling of the Lechuguilla Cave Culturable Microbiome

Since its discovery in 1986, Lechuguilla Cave has been closed to human access without a permit. The sample areas were chosen off the designated trail through the cave, in areas that have experienced very little human impact (National Park records suggest that a maximum of 4–6 people have been in the vicinity of each sample site) (Figure 1). Specific sample locations were chosen due to the unlikelihood that these areas were actually exposed to human visitation: absence of footprints and scuff marks that usually indicate human activity. Three sample sites were chosen, based on location and geology: LCDS1 and LCEAE (deep and within the Capitan Formation) and LCEA1 (mid-depth in the Yates Formation). None of the sites received any direct source of water with any moisture likely to come percolation from the surface or from condensation events within the cave atmosphere. LCDS1 was the deepest sample site (at ~400 m) within a region known as Deep Secrets and LCEAE was in the proximity of the LCDS1 sample site at the same depth. LCEA1 was chosen as an area that has been closed to human access for the past 20 years, with the only human activity in the area being carried out by scientists. The LCEA1 sample is at the bottom of the Yates Formation, which is high in iron and manganese and is the rock unit in which much of the oil and gas is extracted from the Delaware Basin. As a result, this rock unit is usually associated with organic material from ancient sediments [14].

A total of 20 different culture media were inoculated at each sample site, resulting in the isolation of over 500 unique isolates (Text S1) [20]. Of these, 93 that were able to grow in 50% TSB were chosen at random to screen for antibiotic resistance in this fraction of the cave microbiome (Table S1, Figure S1). These bacteria were phylogenetically classified by sequencing of the 16 S rRNA gene sequence (Table S2) and spanned several Gram- and Gram-negative genera known to be associated with cave environments [21,22,23,24,25,26,27].

Antibiotic Resistance Screen of the Lechuguilla Microbiome

The Lechuguilla bacterial collection (93 strains, 33% Gram-positive and 63% Gram-negative) was screened against 26 different antimicrobial agents representing a broad spectrum of natural products, their semi-synthetic derivatives, and completely synthetic molecules (Figure 2). In the primary screen, the level of resistance was assessed by monitoring the growth at 20 μg/ml drug concentration (a high level of antibiotic to select for robust resistance) and resistance in the primary screen was defined as >50% growth in the presence of antibiotic, consistent with other surveys of resistance [11,28]. Resistant and sensitive strains obtained from the primary screen were further quantitatively analyzed by determination of the minimal inhibitory concentration (MIC) of the antibiotics (Figure 3).

Resistance spanned most of the major drug families in both Gram-positive and Gram-negative strains. In the collection of 33 Gram-positive strains, on average, 70% of the strains were resistant to 3–4 different antibiotic classes. Three strains were resistant to 14 antibiotics, all of which were Streptomyces spp. These values are consistent with those found in surface Streptomyces [11]; however, unlike surface strains, we saw no Gram-positive resistance to the synthetic drugs ciprofloxacin and linezolid, the semi synthetic compounds rifampin and minocycline, and the natural product vancomycin (Figures 2–3).

Gram-negative bacteria are intrinsically resistant to many classes of antibiotics due to the presence of the relatively impermeant outer membrane along with the presence of chemo-selective porins and highly efficient small molecule efflux pumps. Therefore, only antibiotics known to have activity against Gram-negatives were included in the data analysis. On average, approximately 65% of the Gram-negative strains showed resistance to 3–4 antibiotic classes. No tetracycline resistance was observed, though it is common in surface bacteria [29,30,31]. On the other hand, resistance to sulfamethoxazole, trimethoprim, and fosfomycin was common; a feature shared with surface bacteria (Figures 2–3).

Antibiotic Inactivation by Gram-positive isolates

Enzymes that modify antibiotics are of particular interest as they most likely evolved in direct response to the emergence of specific antibiotics to block their activity in contrast to mechanisms such as efflux that often can target several classes of bioactive compounds of diverse chemical structure [32]. Within the isolates, we detected no inactivation of aminoglycosides, the lincomamide clindamycin, or chloramphenicol; antibiotics where enzymatic inactivation is a prevalent resistance mechanism in surface bacteria (Table 1) [33]. On the other hand, substantial inactivation (22–62% of strains) was seen for β-lactams (the penicillins ampicillin, piperacillin and the cephalosporin cephalexin), which is primarily caused by the hydrolysis of β-lactam ring by β-lactamases (Table 1) [34].

Enzymatic inactivation was detected for macrolide antibiotics in four strains: three isolates of Streptomyces spp. and one of Bacillus paraconglomeratus (Table S3). These strains inactivated both the natural product erythromycin and its 3rd generation semi-synthetic derivative telithromycin. Inactivation by the Streptomyces isolates yielded a product with a mass increase of 162, consistent with mono-glycosylation of the antibiotic [11,35]. Inactive macrolide antibiotics from B. paraconglomeratus extracts on the other hand revealed a mass increase of 80, indicative of inactivation by phosphorylation (Figure 4) [36].

The lipopeptide daptomycin is the newest class of antibiotic approved for clinical use. Three Streptomyces strains were found to be highly resistant to daptomycin (MIC≥256 μg/ml) and inactivated the antibiotic through hydrolysis as assessed by LC/MS (m/z increase of 18 g/mol). This result is consistent with the high level of daptomycin inactivation in surface Streptomyces where this is a constitutively expressed activity [11,37]. Additionally, four isolates of Paenibacillus lautus completely inactivated daptomycin when grown at sub-MIC concentration of the antibiotic. Daptomycin inactivation has not been previously reported in low G+C bacteria (Firmicutes bacteria). Purification of the inactive product followed by MS/MS analysis revealed hydrolytic cleavage of the ester bond between the threonine and kynurenine residues resulting in ring-opening inactivation (Table S4). The inactivation of daptomycin was highly sensitive to inhibition by EDTA, but not significantly by Ser esterase/protease inhibitors (Figure S2), consistent with the possible involvement of a metallo-esterase. This contrasts with our recent analysis of Streptomyces daptomycin esterases, which appear to use canonical Ser catalytic triad chemistry to inactivate the antibiotic [37]. Furthermore, unlike inactivation by Streptomyces, which appears to be constitutively expressed, the production of the inactivating activity in P. lautus was inducible by exposure to daptomycin (Figure 5). To explore if this activity was unique to the cave isolate, we obtained a surface strain of P. lautus (ATCC 43898) and observed similar results in enzymatic inactivation and antibiotic-associated induction of activity.
Antibiotic Inactivation by Gram-negative isolates

We observed little enzyme-mediated antibiotic inactivation in Gram-negative isolates (Table 2), suggesting that there are other molecular mechanisms of resistance at play such as efflux, target modification, or barriers to entry. Three strains belonging to the genera *Agrobacterium* and *Ochrobactrum* displayed chloramphenicol inactivation, which we determined by LC/MS to be the modification by acetylation (Table S5); a well-established resistance mechanism both in Gram-positive and Gram-negative bacteria [33].

Characterization of a *B. paraconglomeratum* macrolide kinase

Phosphorylation of macrolides in pathogenic bacteria is a growing clinical problem [38] catalyzed by a family of macrolide phosphotransferases (MPHs) [36,39]. In order to probe for the presence of possible mph genes in macrolide inactivating *B. paraconglomeratum*, we prepared a draft genome sequence using Roche 454 and Illumina platforms. A gene encoding a predicted macrolide kinase, *mphE*, was identified (Figure 6). We expressed the gene product in *E. coli*, purified the enzyme and determined its activity and specificity using steady state kinetics (Table 3). The enzyme efficiently modified 14-, 15-, and 16-membered macrolide antibiotics aligning this enzyme with known type II MPHs (based on previous characterization of these enzymes in *E. coli* isolates) [40,41]. The regiospecificity of phosphorylation of telithromycin was determined by multidimensional and multinuclear magnetic resonance analysis to be at the 2'-hydroxyl group of the desosamine sugar (Figures S3 and S4, Tables S6 and S7).

The genome of another *Brachybacterium* species has been reported, *Brachybacterium faecium* DSM 4810, a terrestrial soil isolate...
The *B. faecium* genome includes a putatively annotated aminoglycoside kinase, that is 72% identical at the amino acid level to the *B. paraconglomeratum* MPH. Expression, purification and analysis of the *B. faecium* enzyme revealed similar antibiotic substrate profiles and activity as the MPH encoded by the *mphE* gene in *B. paraconglomeratum*, with no aminoglycoside modification activity (Table 4).

**Table 1. Summary of Antibiotic Inactivation Studies for Gram-positive Isolates.**

| Antibiotic       | Resistant Strains | Number of antibiotic Inactivation Strains | Mechanism of Inactivation       |
|------------------|-------------------|------------------------------------------|----------------------------------|
| Apramycin        | 7                 | 0                                        | ------                           |
| Gentamicin       | 5                 | 0                                        | ------                           |
| Neomycin         | 8                 | 0                                        | ------                           |
| Streptomycin     | 8                 | 0                                        | ------                           |
| Tetracycline     | 4                 | 0                                        | ------                           |
| Minocycline      | 1                 | 0                                        | ------                           |
| Clindamycin      | 6                 | 0                                        | ------                           |
| Chloramphenicol  | 12                | 0                                        | ------                           |
| Synercid         | 2                 | 0                                        | ------                           |
| Erythromycin     | 7                 | 1                                        | Phosphorylation                  |
| Telithromycin    | 4                 | 4                                        | Phosphorylation Glycosylation    |
| Linezolid        | 2                 | 0                                        | ------                           |
| Novobiocin       | 5                 | 0                                        | ------                           |
| Cephalexin       | 17                | 5                                        | Hydrolysis                       |
| Ampicillin       | 8                 | 5                                        | Hydrolysis                       |
| Pipercillin      | 9                 | 2                                        | Hydrolysis                       |
| Daptomycin       | 24                | 7                                        | Hydrolysis                       |

Strains were grown in 50% TSB for 5 days in presence of 20 μg/ml antibiotic. Conditional media was used for setting up disk diffusion assays and LC-MS analyses. Inactivation was defined as the absence of a zone of clearance around the disk. Hydrolytic mechanism of β-lactam resistance is inferred.

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Figure 3. MIC of antibiotics determined in this study. Heat Plot for (A) Gram-positive strains (B) Gram-negative strains against various antibiotics. Antibiotics are grouped according to their mode of action and the gradient from light blue to dark blue represents the range from lowest MIC value (0.3 μg/ml) to highest MIC value (256 μg/ml) as shown in the legend. White means no MIC was determined.

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The mph genes in both *Brachybacterium* species are organized on comparable regions of the chromosome and flanked by similar genes and synteny (Figure 6); a survey 10 kb upstream and downstream of mph revealed eleven common and five different genes (Figure 6) and the gene products were 75–80% similar. Phylogenetic analysis shows that MPHs from *Brachybacterium* strains cluster together as a separate group among the known and putative members of the MPH family (Figure S5).

**Discussion**

Antibiotic resistance is manifested through a number of different mechanisms including target alteration, control of drug influx and efflux, and through highly efficient enzyme-mediated inactivation. Resistance can emerge relatively quickly in the case of some mutations in target genes and there is evidence that antibiotics themselves can promote such mutations [43,44,45,46]; however, resistance to most antibiotics occurs through the aegis of extremely efficient enzymes, efflux proteins and other transport systems that often are highly specialized towards specific antibiotic molecules. Such elements are the result of evolution through natural selection; this therefore implies that antibiotic resistance has a long evolutionary past. A growing body of evidence suggests that non-pathogenic environmental organisms are a reservoir of resistance genes that have the potential to be transferred to pathogens [31,47,48]. The problem of antibiotic resistance in clinical settings therefore likely has its origins in the environment.

One of the challenges in studying the evolution and prevalence of resistance is the massive use of antibiotics in the clinic and in agriculture over the past seven decades that makes identifying environments that have not been impacted by anthropogenic antibiotics difficult. Studying resistance in pristine environments that have not been exposed to human antibiotic use provides a critical measure of the genetic diversity of resistance that is essential to our understanding of resistance gene prevalence and evolution. Lehguilla Cave provides an outstanding ecosystem that has been isolated for over 4 million years. The cave’s geologic features, including the impermeable siltstone caprock which prevents rapid influx of surface water, great depth, and long isolation from the surface, rules out the possibility of exposure to...
anthropogenic use of antibiotics as well as antibiotic contamination through water bodies. As a result, Lechuguilla Cave is an ideal ecosystem for investigating microbes that have not been exposed to anthropogenic antibiotics.

We surveyed the antibiotic susceptibility of 93 bacterial strains isolated from Lechuguilla Cave. This was a genetically diverse collection of oligotrophic organisms (Figure S1), highly adapted to survive in a nutrient limited environment [22]. Like surface organisms [11], the majority of these strains were multidrug resistant indicating that antibiotic resistance is a common and widespread phenotype in pristine, unimpacted environments; however, there are differences in the pattern of resistance. For example, we measured little resistance to the synthetic antibiotics ciprofloxacin and linezolid, while resistance to natural product antibiotics was more prevalent. Unlike surface bacteria, we also detected very little resistance to tetracycline, glycopeptide (vancomycin), rifamycin (rifampicin) and lipopeptide (daptomycin) natural product antibiotics. There are several possible reasons for these differences. First, this survey includes multiple bacterial genera across five phyla, while our original sampling focused on

Table 2. Summary of Antibiotic Inactivation Studies for Gram-Negative Isolates.

| Antibiotic       | Resistant Strains | Number of antibiotic Inactivation Strains | Mechanism of Inactivation |
|------------------|-------------------|------------------------------------------|---------------------------|
| Apramycin        | 10                | 0                                        | ----                      |
| Gentamicin       | 11                | 0                                        | ----                      |
| Neomycin         | 24                | 0                                        | ----                      |
| Streptomycin     | 30                | 0                                        | ----                      |
| Telithromycin    | 17                | 0                                        | ----                      |
| Clindamycin      | 39                | 0                                        | ----                      |
| Chloramphenicol  | 28                | 2                                        | Acetylation               |
| Trimethoprim     | 35                | 0                                        | ----                      |
| Cephalaxin       | 36                | 14                                       | Hydrolysis                |
| Cefotaxime       | 12                | 4                                        | Hydrolysis                |
| Ampicillin       | 30                | 13                                       | Hydrolysis                |
| Piperacillin     | 30                | 16                                       | Hydrolysis                |

Strains were grown in 50% TSB for 5 days in presence of 20 µg/ml antibiotic. Conditional media was used for setting up disk diffusion assays and LC-MS analyses. Inactivation was defined as the absence of a zone of clearance around the disk. Hydrolytic mechanism of ß-lactam resistance is inferred.

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Figure 6. Genetic context of mph genes in Brachybacterium strains. A genetic map was constructed using available genome sequences of Brachybacterium strains and shown above is a schematic of translated protein query based on BLAST analysis. The MPH sequence is shown in red and homologous sequences are marked with identical colors.

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We recently showed that high G\textsuperscript{+} results with more confidence. Metagenomic analysis are therefore necessary to interpret the and cave microbiomes including more extensive cultivation and non-culturable microbiome. Further studies of both surface have likely not examined the full resistome of both the culturable and smaller in this study in comparison to our previous study and we enriched in resistance elements. Second, the isolate sample size is.

**Table 3. Kinetic parameters for MPH (2')-II from B. paraconglomeratum.**

| Substrate     | Lactone Ring | K\text{in} (μM) | k_{cat} (s\textsuperscript{-1}) | K\text{m} (μM) | k_{cat}/K\text{m} (s\textsuperscript{-1} M\textsuperscript{-1}) |
|---------------|--------------|-----------------|-------------------------------|----------------|-------------------------------------------------|
| Erythromycin\textsuperscript{1} | 14           | 43.8±11.3       | 0.101                         | 1.3           | 3.2×10\textsuperscript{7}                         |
| Clarithromycin | 14           | 21.9±5.45       | 0.106                         | 1.3           | 4.8×10\textsuperscript{7}                         |
| Telithromycin  | 14           | 12.2±5          | 0.04                          | 600±285       | 3.5×10\textsuperscript{7}                         |
| Roxithromycin  | 14           | 58.98±10.9      | 0.202                         | 6.2           | 3.4×10\textsuperscript{7}                         |
| Azithromycin   | 15           | 53.97±7.59      | 0.168                         | 2.6           | 3.1×10\textsuperscript{7}                         |
| Spiramycin     | 16           | 45.6±14.9       | 0.173                         | 292±108       | 3.7×10\textsuperscript{7}                         |
| Tylosin        | 16           | 222±29.2        | 0.197                         | 66.4±8.92     | 8.8×10\textsuperscript{7}                         |
| GTP\textsuperscript{2} | -            | 10.2±0.53       | 0.093                         | -             | 9.3×10\textsuperscript{4}                         |

\textsuperscript{1} GTP held at 200 μM for antibiotic substrates, 2-erythromycin held at 400 μM. doi:10.1371/journal.pone.0034953.0003

Aminoglycoside antibiotic resistance was more common in Lechuguilla Cave isolates as compared to surface actinomycetes. This may reflect the biosynthetic capacity of antibiotic producing bacteria in Lechuguilla Cave and the production of these antibiotics by species within the cave. A survey of the actinomycetes in our collection using oligonucleotide primers designed to amplify aminoglycoside biosynthetic genes failed to identify potential aminoglycoside producers; however linking resistance to antibiotic production will require an extensive and systematic survey of the cave microbiome and resistome that is beyond the objectives of this work.

The mechanisms of antibiotic modification and inactivation are evidence of highly specific evolutionary adaptations to evade the cytotoxic action of these antibiotics. The high level of β-lactam antibiotic resistance by hydrolysis parallels that of surface bacteria and the result of genetically diverse β-lactamases that are widespread in microbial genomes. Similarly, chloramphenicol acetylation was also detected, an activity that is well established widespread in microbial genomes. Similarly, chloramphenicol lactam antibiotic resistance by hydrolysis parallels that of surface the cytotoxic action of these antibiotics. The high level of β-

Table 4. Kinetic parameters for MPH (2')-II from B. faecium DSM 4810.

| Substrate     | Lactone Ring | K\text{in} (μM) | k_{cat} (s\textsuperscript{-1}) | K\text{m} (μM) | k_{cat}/K\text{m} (s\textsuperscript{-1} M\textsuperscript{-1}) |
|---------------|--------------|-----------------|-------------------------------|----------------|-------------------------------------------------|
| Erythromycin\textsuperscript{1} | 14           | 38.2±6.24       | 0.192                         | -              | 5.0×10\textsuperscript{7}                         |
| Clarithromycin | 14           | 420±6.26        | 0.286                         | -              | 6.8×10\textsuperscript{7}                         |
| Telithromycin  | 14           | 23.9±4.22       | 0.168                         | -              | 7.0×10\textsuperscript{7}                         |
| Roxithromycin  | 14           | 19.3±2.60       | 0.262                         | -              | 1.3×10\textsuperscript{7}                         |
| Azithromycin   | 15           | 60.5±6.86       | 0.312                         | -              | 5.1×10\textsuperscript{7}                         |
| Spiramycin     | 16           | 173±3.18        | 0.157                         | -              | 9.1×10\textsuperscript{7}                         |
| Tylosin        | 16           | 14.5±4.00       | 0.088                         | -              | 6.0×10\textsuperscript{7}                         |
| GTP\textsuperscript{2} | -            | 16.0±1.3        | 0.203                         | -              | 1.2×10\textsuperscript{7}                         |

\textsuperscript{1} GTP held at 200 μM for antibiotic substrates, 2-erythromycin held at 400 μM. doi:10.1371/journal.pone.0034953.0004

Antibiotic Resistance in Cave Bacteria of a specific receptor for daptomycin or a non-specific response to the physiological impact of daptomycin bioactivity.

The observation of two distinct macrolide inactivation mechanisms in the Lechuguilla bacterial isolates was also intriguing. In the resistant Streptomyces strains we determined that antibiotic modification by glycosylation was the primary mechanism of inactivation, a mechanism that is known in surface actinomycetes [11]. On the other hand, we established that the mechanism of macrolide inactivation in B. paraconglomeratum is through phosphorylation at position 2′ catalyzed by a member of the MPH class of antibiotic kinases. Previously identified mph genes are encoded on plasmids found in clinically resistant isolates of the pathogens Escherichia coli, Staphylococcus aureus, Pasteurella multocida and Pseudomonas aeruginosa [36]. This is the first report of mph genes from environmental bacteria as a potential source of the genes currently circulating in pathogens. The presence of a transposase-like gene upstream of mph from a surface strain of B. faecium points to a potential history of horizontal gene transfer (Figure 6). It is possible that mph genes have been circulating among bacterial populations before the cave was sealed off millions of years ago, resulting in an mph gene that is a shared trait between both terrestrial and cave bacteria.

There are two likely explanations for retention of the mph genes with same biochemical properties despite the long isolation of the Brachybacterium strains: (i) these genes could serve a physiologic or metabolic function unrelated to antibiotic resistance (although the genetic context (Figure 6) does not suggest an obvious role); or (ii) these genes are resistance elements for conferring antibiotic resistance. We could not detect any macrolide biosynthetic gene clusters in bacteria collected in the same region as B. paraconglomeratum (as evidenced by a absence of the signature macrolide D-desosamine biosynthesis gene, euyCIT, not shown); however the actinomycete small sample size does not rule out the possibility of the presence of hitherto undetected macrolide producers within the cave.

This work demonstrates that antibiotic resistance is widespread in the environment even in the absence of anthropogenic antibiotic use. Lechuguilla Cave represents a remarkable ecosystem that has been isolated for millions of years, well before the clinical and agricultural use of antibiotics. The presence of multidrug resistant organisms even in this pristine environment reinforces the notion that the antibiotic resistome is an ancient and pervasive component of the microbial pan genome. Given the nutrient-limited nature of the cave environment, it is likely that competition for resources plays a dominant role in species...
Antibiotic Resistance in Cave Bacteria

Materials and Methods

Antibiotic Resistance Screen of Cave Strains

A sample of 93 bacterial strains were isolated from three deep, remote sample sites in Lechuguilla Cave (Figure 1) under a permit for sample collection provided by the US National Park Service (Permit CAVE-2007-SCI-0009 for Study CAVE-00049). These strains were isolated on a variety of culture media that resembled the carbon and energy sources thought to be available to microorganisms within the cave (see Text S1). Following single colony isolation, these isolates were grown in dilute (50%) Tryptic Soy Broth (TSB) at 30°C for 3–5 days and frozen stocks were prepared in 96 well plate format (1:10 dilution of cultures in 80% glycerol). Master plates containing 150 μl 50% TSB were inoculated from the frozen stock plates using a replica head platter and grown at 30°C for 3–5 days. The inoculum plate was prepared from the master plate as a 1:100 dilution in 50% TSB media and supplemented with antibiotic, (ten 2-fold dilution of antibiotic, final concentration ranging from 0.5 μg/ml to 128 μg/ml). Cultures were diluted to OD₆₀₀ of 0.08–0.1 in 0.85% NaCl and MIC plates were inoculated with 1:20 dilution of this suspension. Plates were incubated at 30°C for 5 days. Escherichia coli ATCC 29212, Staphylococcus aureus ATCC 29213 and Enterococcus faecalis ATCC 29212 were used as control strains. MIC testing was performed in duplicate, where MIC was defined as the drug concentration showing no visible growth.

In order to evaluate resistant strains for antibiotic inactivation, cultures containing 5 ml of 50% TSB in the presence and absence of 20 μg/ml of antibiotic were inoculated with 3–5 single isolated colonies of strain of interest and cultures were grown at 30°C for 5 days. Uninoculated controls were also prepared. After growth at 30°C for 5 days, cultures were centrifuged for 20 minutes at 16,800×g and conditioned media was collected. One of the following susceptible organisms/test organisms was used for antimicrobial susceptibility testing (antimicrobial disk diffusion assay): Bacillus subtilis, Micrococcus luteus and Staphylococcus saprophyticus ATCC 15305. Inocula of test organisms were prepared to the 0.5 McFarland standard using the direct colony suspension assay:

B. paraconglomeratum LC44 Genome Sequencing and Assembly

B. paraconglomeratum LC44 was grown in 5 ml 50% TSB at 30°C for 5 days. Genomic DNA was isolated using QIAGEN DNeasy Blood and Tissue Kit 250 (Qiagen, Germany) with one modification of the manufacturer’s protocol for cell lysis: 4 μl of RNase (100 mg/ml in Buffer TE) was added to the reaction mixture and incubated at room temperature for 2 minutes before washing the spin column. Genomic DNA was submitted for shotgun sequencing to Roche 454 Life Sciences Genome Sequencer at Farncombe Metagenomics Facility, McMaster University. Approximately one quarter of a 454 PTP was used for Titanium pyrosequencing on a 454 GS FLX. Additionally, 8059104 Illumina GAIIx 71 bp paired end reads were sequenced by Ambry Genetics (Aliso Viejo, California). After aggressive quality trimming and filtering, the approximately 5.5 million remaining Illumina reads (about 70× coverage) and 262995 454 reads (about 23× coverage) were assembled using MIRA version 3.4 with the ‘-job = denovo,genome,454,accurate,solexa’ switches [52]. The resulting contig sequences of the assembly were deposited in Genbank as a WGS project under accession AGSO000000000. The protein sequence of E. coi MPH (2’-Ia (accession BAA03776) was used to query the assembled sequences using translated blast in order to find a gene responsible for the observed macrolide phototransferase activity. A good candidate was found on contig AGSO000000004 beginning at bp 76533 and a blastx search was performed against the NCBI non-redundant protein database where the top hits corresponded to sequences annotated as putative APH and MPH proteins.
Cloning, Expression and Purification of Macrolide Phosphotransferases

The macrolide phosphotransferase from *B. paraconglomeratum* (mphE) was synthesized by GenScript (USA) with codon optimization for *E. coli* expression and cloned into pET28b with *Nde I* and *Hind III* restriction sites. The plasmid containing MPH was transformed into *E. coli* BL21 (DE3) and the resulting colonies were grown overnight at 37°C in LB media supplemented with 50 μg/ml kanamycin. For overexpression, 1 L LB cultures supplemented with 50 μg/ml kanamycin were subcultured from the overnight culture at 1% (v/v) and grown at 37°C for 16 hours. The cells were harvested and stored at −20°C until further use. For protein purification, the cells were resuspended in lysis buffer (50 mM HEPES, 300 mM NaCl, 10 mM Imidazole, pH 7.5), 1 mM phenylmethanesulfonylfluoride and 1 μg/ml pancreatic-bovine DNase and cells were lysed using T-S series cell disrupter. The crude homogenate was filtered and stored in 10% glycerol at −20°C.

Steady State Kinetic Analysis of MPH(2')

Kinetic parameters of MPHs were determined in triplicate using Pyruvate Kinase/Lactate Dehydrogenase coupled assay [53]. Phosphorylation of macrolides by MPHs was monitored by absorbance of NH2DH (340 nm) in 96 well format using a SpectraMax reader. The reaction was initiated with macrolide or nucleotide (GTP). When monitoring macrolide dependence, 200 μM GTP was used and the macrolide concentration ranged from 3.2 μM and 400 μM. For nucleotide dependence, 200 μM macrolide was used and the final concentrations of GTP ranged from 7.82 μM to 2000 μM. All reactions were performed in triplicate.

The initial rates were fit to equation 1 or 2 (substrate inhibition) using Grafit 4.0 software (Erithacus Software, Staines, UK):

\[ v = \frac{V_{\text{max}}}{K_m + [S]} \]  
\[ v = \frac{V_{\text{max}} * S}{(K_m + S * (1 + S/K_s))} \]

Growth Analysis of *Paenibacillus lautus* LC231 with Daptomycin

Analyses were performed in 96-well flat bottom plates (200 μl volume total) using 50% TSB supplemented with 1.25 mM CaCl2 as a growth medium. Inocula represented 1:200 dilutions of an overnight culture, standardized to an OD600 of 0.1. The following conditions were tested in duplicate: (1) no daptomycin, (2) 4 μg/ml daptomycin and (3) 4 μg/ml daptomycin added after 10.5 hours (early log phase). Plates were incubated while shaking at 30°C and the data was collected as an OD600 every 30 minutes using a Tecan Sunrise plate reader. Similar studies were conducted with a surface strain of *P. lautus* (ATCC 43898) except that the organism was grown in full strength TSB.

Supporting Information

Text S1 Supporting materials and methods.

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