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Marti, Hanna; Borel, Nicole; Dean, Deborah; Leonard, Cory A

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Methods

Evaluating the antibiotic susceptibility of *Chlamydia* – new approaches for *in vitro* assays

Hanna Marti¹, ², Nicole Borel¹, Deborah Dean², ³, ⁴, Cory Ann Leonard¹, *, #

¹ Institute for Veterinary Pathology, Vetsuisse Faculty, University of Zurich, Zurich, Switzerland
² Center for Immunobiology and Vaccine Development, UCSF Benioff Children's Hospital Oakland Research Institute, Oakland, California, USA
³ Joint Graduate Program in Bioengineering, University of California, San Francisco, California, USA, and University of California, Berkeley, Berkeley, California, USA
⁴ School of Medicine, University of California, San Francisco, California, USA

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*Correspondence
Cory Ann Leonard
cory.leonard@ufl.edu

#Current Address: Department of Pathology, Immunology and Laboratory Medicine, University of Florida, Gainesville, FL, USA

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Abstract

Pigs are the natural hosts of *Chlamydia suis*, the only *Chlamydia* species known to spontaneously acquire homotypic resistance conferred by a class C tetracycline resistance gene. Various susceptibility assays have existed for several years, but there is no widely accepted, standardized assay to determine chlamydial antibiotic susceptibility. In this study, we developed new approaches to determine the *in vitro* susceptibility of *Chlamydia* to different antibiotics in view of existing protocols. Specifically, the minimal inhibitory concentration (MIC) is based on a consensus of both inclusion number reduction and alteration of inclusion size and morphology upon antibiotic exposure. In addition to these, we employed a recovery assay, allowing observation of the chlamydial response to drug removal and subsequent recovery, as compared to both continued exposure and to the unexposed control. We propose a simple and fast screening method to detect tetracycline resistant *C. suis* strains within two to three days with minimal use of consumables. For proof of principle, we evaluated the susceptibility of three *C. suis* field strains and the reference strain S45/6 to tetracycline, sulfamethoxazole and penicillin, antibiotics commonly used to prevent respiratory and gastrointestinal diseases on fattening pig farms. We found that tetracycline sensitive strains can easily be distinguished from resistant strains using the evaluation parameters proposed in this study. Moreover, we report that S45/6 is sensitive to sulfamethoxazole while all evaluated *C. suis* field strains showed some degree of sulfamethoxazole resistance. Finally, we confirm that Penicillin G induces the chlamydial stress response in all evaluated *C. suis* strains.

Keywords

*Chlamydia*, antibiotic resistance, new diagnostic approaches, antibiotic susceptibility assay, resistance screen, minimal inhibitory concentration, recovery assay
Introduction

In vitro susceptibility assays are essential to detect antibiotic resistance in Chlamydia. Protocols have existed since the 1980s (Henning and Krauss, 1986), but unlike the standardized tests that are in place for extracellular and facultative intracellular bacteria (CLSI, 2012; Jorgensen and Turnidge, 2015), there is no clear consensus among the research community for antibiotic resistance determination in obligate intracellular bacteria such as Chlamydia. Table 1 summarizes previously reported protocols and/or definitions to determine the antibiotic susceptibility in Chlamydia.

The aim of this study was to investigate new approaches to determine the in vitro susceptibility of Chlamydia to different antibiotics. To achieve this goal, the parameters were defined based on consideration of existing protocols (Table 1). Chlamydia (C.) suis was selected as a model for other Chlamydia species because of its high degree of genetic diversity and because it is the only example of homotypic resistance to antibiotics among the Chlamydia, which is conferred by an acquired resistance gene (Borel et al., 2016; Joseph et al., 2016; Lenart et al., 2001; Sandoz and Rockey, 2010; Schautteet and Vanrompay, 2011; Seth-Smith et al., 2017). This gene is located within a genomic island carrying a class C tetracycline resistance gene and can be transferred to other C. suis but also to and among C. trachomatis upon simultaneous co-infection in vitro (Jeffrey et al., 2013; Marti et al., 2017; Suchland et al., 2009). This could have serious implications for human health considering that the DNA of both C. suis and C. trachomatis has been detected in the eye of trachoma patients in Nepal, and that C. suis could be isolated from samples of pig farmers and slaughterhouse workers originating from various anatomical locations (conjunctiva, nose, pharynx and stool samples) (Dean et al., 2013; De Puyssseley et al., 2014, 2015). Apart from tetracycline resistance in C. suis, induced resistance to antibiotics (e.g., rifamycins, fluoroquinolones) through point mutations has also been found in C. suis and other Chlamydia (e.g., C. trachomatis, C. pneumoniae, C. psittaci, C. muridarum and C. caviae) following propagation at sub-inhibitory concentrations of the drug in vitro (Sandoz and Rockey, 2010). Moreover, heterotypic resistance in various Chlamydia species has been observed upon survival of a small proportion of bacteria exposed to antibiotic concentrations well above the minimal inhibitory concentration (MIC) and might be a cause of treatment failure in patients, a phenomenon also referred to as drug indifference, tolerance or persistence (Sandoz and Rockey, 2010; Suchland et al., 2003). In Chlamydia, persistence, or the chlamydial stress response, is known to occur specifically upon exposure to antibiotics affecting cell wall synthesis, such as penicillin (Borel et al., 2014; Leonard et al., 2017; Lewis, 2007; Sandoz and Rockey, 2010; Schoborg, 2011; Xue et al., 2017; Zheng et al., 2015).

In the present study, the MIC was determined according to the methods described by Donati and colleagues (2010) (MIC = ≥90% inclusion number reduction) and by Suchland and colleagues (2003) (transition point MIC [MICTP] = ≥90% alteration of size/morphology; MIC = 2x MICTP). Following the determination of MIC (Donati) and MIC (Suchland), an MIC consensus was established if they were identical. Otherwise, an MIC range was determined unless the MICs were notably different. In addition, instead of applying the generally used protocols to determine the minimal bactericidal/chlamydicidal/lethal concentration (MBC/MCC/MLC), we incorporated an assay based on protocols used to evaluate recovery from the chlamydial stress response (Kintner et al., 2014; Leonard et al., 2015, 2017). With this recovery assay, it is possible to evaluate the in vitro behavior of chlamydial strains upon recovery and continued exposure to low, moderate and high concentrations of the antibiotic in question instead of evaluating a single value that is identical to or only a few twofold dilutions higher compared to the determined MIC value as often observed for MBC/MCC/MLC (Agacfidan et al., 1993; Donati et al., 2010; Ehret and Judson, 1988; Henning and Krauss, 1986; Japan Society of Chemotherapy, 1992a, 1992b; Suchland et al., 2003; Webberley et al., 1987).
In summary, we propose a two-step protocol evaluating two major parameters, the MIC and the recovery. This approach allows initial assessment after two workdays, and a detailed report after eight workdays (Fig. 1). For proof of principle, we investigated the susceptibility of three *C. suis* field strains and the reference strain S45/6 to tetracycline, sulfamethoxazole and penicillin.

**Material and Methods**

The detailed *in vitro* antibiotic susceptibility assay protocol (tetracycline, sulfamethoxazole and PenG) is available as Supplemental Material (Text S1), and includes suggestions regarding optimization of this assay.

**Host cells and media**

LLC-MK2 cells (Rhesus monkey kidney cell line, provided by IZSLER Brescia, Italy) were grown in antibiotic-free growth medium consisting of 500 ml Eagle’s minimum essential medium (EMEM, Gibco, Thermo Fisher Scientific, Invitrogen, Carlsbad, CA, USA) supplemented with 10% heat-inactivated fetal calf serum (FCS, BioConcept, Allschwil, Switzerland), 5 ml L-glutamine (Gibco, Thermo Fisher Scientific) and 6 ml glucose (0.06 g/ml; Sigma-Aldrich Co., St. Louis, MO, USA). Following infection, growth medium was replaced by *Chlamydia* cultivation medium consisting of 500 ml EMEM supplemented with 20% FCS (BioConcept), 5 ml L-glutamine (Gibco, Thermo Fisher Scientific) and 2 g glucose (Sigma-Aldrich) with or without 0.7 ml cycloheximide (1 mg/ml; Sigma-Aldrich) as described (Donati et al., 2010; Wanninger et al., 2016).

**Chlamydial strains**

Four *C. suis* strains were used for this study: three field strains (SWA-14, 86 and 141), all isolated from fecal swabs of asymptomatic fattening pigs (Hoffmann et al., 2015; Wanninger et al., 2016) and the *C. suis* reference strain S45/6, which was originally isolated from feces of an asymptomatic pig (Kaltenboeck et al., 1993; Schautteet and Vanrompay, 2011) (Table 2). Stocks were prepared and stored in sucrose phosphate glutamate (SPG) at -80°C (218 mM sucrose [Sigma-Aldrich], 3.76 mM KH2PO4 [Sigma-Aldrich], 7.1 mM K2HPO4 [Merck Eurolab AG, Dietlikon, Switzerland], and 5 mM GlutaMAX-100 [GIBCO]) as described (Leonard et al., 2016) with minor changes: After infected cells were scraped, mechanical disruption was performed by vortexing with 10-20 glass beads (Ø 5mm) for 1 min followed by pushing the suspension through a 20 Gauge needle with a 20 ml syringe.

**Antibiotic reagents**

Tetracycline hydrochloride powder (Sigma-Aldrich) was dissolved in deionized water to reach a final concentration of 10 mg/ml, filter-sterilized with a 0.22 μm syringe filter, aliquoted and stored at -20°C. Sulfamethoxazole (Santa Cruz Biotechnology, Dallas, TX, USA) stocks were prepared in DMSO (Sigma-Aldrich) with a final concentration of 50 mg/ml, filter-sterilized and stored as described for tetracycline. PenG (sodium salt, Sigma-Aldrich) stocks were prepared as described (Leonard et al., 2016) with a stock concentration of 20,000 U/ml in deionized water.

**Important definitions and parameters**

The Minimal inhibitory concentration (MIC) is defined as the consensus of the MIC defined by Donati and colleagues (2010); MIC Donati ≥ 90% inclusion number reduction) and Suchland and colleagues (2003); MIC Suchland ≥ 90% inclusions altered in size and morphology) unless one parameter could not be employed. In the recovery assay, chlamydial recovery upon drug-removal (rec) at 48 hours (h) is compared to both, continued exposure for
96 h (exp) and an untreated control (mock) following determination of the number of inclusion forming units (IFU)/ml (semi-quantitative analysis) and is expressed through three parameters. 1) The resistance potential represents how well Chlamydia resist continuous exposure to the antibiotic in question (exposure to mock). In detail, it provides the highest antibiotic concentration where a) >25% or b) >10% of continuously infected cultures remain infectious compared to untreated controls. 2) The recovery potential represents how well Chlamydia recover from antibiotic treatment (recovery to mock). It provides the highest antibiotic concentration where a) >1% or b) >10% of cultures remain infectious following recovery compared to the mock-exposed cultures. Finally, 3) Survival after continued exposure directly compares the infectivity of continuously exposed to recovered cultures (exposure to recovery) and provides the highest antibiotic concentration where a) >1% or b) >10% of continuously infected cultures remain infectious compared to the recovery data of the same antibiotic concentration.

Overview of the in vitro antibiotic susceptibility assay

The protocol can be performed within two weeks and consists of a) determination of the initial susceptibility phenotype, b) MIC confirmation and c) the recovery assay. In the first week, the initial susceptibility phenotype and the first part of the recovery assay can be performed within 48 h and 96 h, respectively. In the second week, MIC confirmation and the second part of the recovery assay are performed within two workdays. Because all samples required for the second week are stored frozen by the end of the first week, this provides a convenient potential stopping point in the protocol. Immunoﬂuorescence microscopy analysis can be performed on the third day (Fig. 1). For tetracycline, there is the possibility to initially perform a tetracycline resistance screen (Tet screen) for a large number of strains within a short period of time with minimal use of consumables.

Determination of the initial susceptibility phenotype

A Chlamydia/cell suspension was prepared for each C. suis strain containing 300,000 cells per ml cycloheximide-free Chlamydia cultivation medium and a multiplicity of infection (MOI) of 0.5. 100 µl per well of this suspension was added directly to 100 µl of cycloheximide-free Chlamydia cultivation medium containing a serial dilution of antibiotics in a 96-well plate (30,000 cells per well). For tetracycline, the final antibiotic concentrations ranged from 0.0078 to 8 µg/ml (2-fold dilution); for sulfamethoxazole (or other antibiotics with an unknown susceptibility range), concentrations ranged from 4.9E-4 to 1024 µg/ml (2-fold dilution); for PenG (and presumably appropriate for most or all known inducers of the classic chlamydial stress response), the final concentrations ranged from 0.01 U/ml to 100 U/ml in a 10-fold dilution series. For all antibiotics evaluated, an unexposed control (mock) containing only cycloheximide-free medium was added. Moreover, each strain was assayed in duplicate. Following centrifugation (1 h, 1000g, 25°C), cultures were incubated at 37°C with 5% CO₂ for 34 h before they were fixed in -20°C-chilled methanol for 10 min and immunolabeled according to the protocol described below. MIC determination was performed according to MIC (Donati). If an estimate regarding inclusion number reduction was not possible (e.g., sulfamethoxazole), MIC (Suchland) was employed instead.

Recovery assay protocol

Cultures were performed identically to the initial susceptibility phenotype. After incubation for 48 h instead of 34 h, culture supernatants were removed and monolayers were washed twice with antibiotic- and cycloheximide-free medium before addition of medium (150 µl) either without antibiotics (recovery) or with serially diluted antibiotics (continued exposure). After another incubation period (96 h post infection), samples were scraped into
the supernatant and frozen at -80°C until use. For tetracycline, 0.03, 0.5, 2 µg/ml exposed samples and the unexposed mock control were collected both for recovery and continued exposure groups. Alternatively, 0.125 instead of 0.03 µg/ml could be collected. For sulfamethoxazole, 2, 32, 512 µg/ml exposed samples and the unexposed mock control were similarly collected. For penicillin, 1, 10, 100 U/ml exposed samples and the unexposed mock control were similarly collected. In total, each strain resulted in eight samples for IFU/ml determination per antibiotic agent (including mock). For antibiotics being evaluated for the first time, the general suggestion is that samples should be generated close to the lowest and the highest initially determined MIC, as well as including an intermediate concentration in order to evaluate the entire susceptibility range.

In the second week, samples were inoculated in duplicate onto 24-well plates containing coverslips of cells cultured to confluence. Following centrifugation (1 h, 1000g, 25°C), inocula were replaced by 1 ml cycloheximide-containing *Chlamydia* cultivation medium. Monolayers inoculated with samples from the continuously exposed condition (including mock) were additionally washed twice with medium to remove residual drugs. Inoculation volumes were established for each antibiotic and generally consisted of 1 µl of 1 ml total sample volume for the mock controls (details shown in Text S1). For resistant strains (determined according to the initial susceptibility phenotype) 1 µl, 10 µl and 60 µl were generally inoculated for low, intermediate and high concentrations, respectively. For sensitive strains (according to the initially determined phenotype), 1 or 10 µl inocula were used for samples, which exhibited recovery from low antibiotic concentrations (MIC or below) and 60 µl for all other conditions/samples. Following fixation in methanol and immunofluorescence assay, the IFU/ml for each condition (recovery, or continued exposure for mock and antibiotic concentrations) was determined according to previously published methods (Deka et al., 2006). a) The resistance potential, b) the recovery potential and c) survival after continued exposure was determined as described above (“Important definitions and parameters”).

### MIC confirmation

MIC confirmation was performed in the second week. Prepared host cell monolayers on glass coverslips in 24-well plates (150,000 cells per well) were infected with an MOI of 0.1 in 1 ml cycloheximide-containing *Chlamydia* cultivation medium. Following centrifugation (1 h, 1000g, 25°C), inocula were replaced with 1 ml cycloheximide-containing *Chlamydia* cultivation medium either without (mock) or with antibiotics close to the MIC. For tetracycline, 0.03, 0.125 and 0.5 µg/ml were used; for sulfamethoxazole, 64, 128 and 256 µg/ml were used for the field strains, and for S45/6, in addition to these concentrations, 0.0039, 0.0078 and 0.015 µg/ml were tested; for PenG, 1, 10 and 100 U/ml was used. Following incubation for 34 h (tetracycline) or 48 h (sulfamethoxazole, penicillin), monolayers were fixed in methanol and immunofluorescence assay was performed. MIC determination was performed according to definitions by Donati et al. (2010) and Suchland et al. (2003) as described above.

In detail, the average number of inclusions per 20x field was determined and compared to the mock for MIC (Donati). The lowest antibiotic concentration, where 10% or less inclusions were present compared to the mock, was defined as the MIC.

For MIC (Suchland), the MIC was based on two different criteria: inclusion size and morphology, neither of which can be as easily quantified as the inclusion number due to possible variability. Despite this drawback, MIC determination was possible as the change from normal to altered inclusions in 90% of the inclusions was abrupt rather than gradual. In order to quantify this change, we semi-quantitatively determined the mean inclusion size. For that, 50 randomly selected inclusions were evaluated in at least 10 fields (400x magnification) per condition and the area (in µm²) was calculated using BonTec measuring and archiving.
software (BonTec, Bonn, Germany). Representative microscopic images were captured using BonTec software (BonTec) and a UI-2250SE-C-HQ camera (uEye, IDS Imaging Development Systems GmbH, Obersulm, Germany) as described previously (Leonard et al., 2015). For conditions with only few or very small inclusions, up to 20 inclusions were analyzed for size and morphology if possible. In parallel, we qualitatively evaluated inclusion morphology. The following criteria were used to classify the inclusion morphology as altered compared to control inclusion morphology: size and/or the presence of aberrant bodies (ABs, aberrant inclusion bodies; diameter ≥ 2 µm, (Matsumoto and Manire, 1970)). Micro-inclusions were defined as inclusions with an area of less than 15 µm². In the case of discrepancies between these parameters, we reported susceptibility ranges (e.g., 64-128 µg/ml).

Tetracycline resistance screen (Tet Screen)

With this method, up to ten strains per 24-well plate can be tested for tetracycline susceptibility. A tetracycline sensitive and a resistant control should be included. Two wells of a 24-well plate with confluent monolayers (150,000 cells/well) on glass coverslips are infected with an MOI of approximately 0.5 for each strain. Following centrifugation (1 h, 1000g, 25°C), inocula are replaced with 1 ml of cycloheximide-containing medium either with or without 0.125 or 0.5 µg/ml tetracycline. After 34 h of incubation, monolayers are fixed in methanol, immunolabeled and tested for the presence or absence of inclusions in tetracycline-containing conditions. If inclusions in tetracycline-exposed cultures are comparable to the corresponding unexposed-control, in terms of inclusion number and morphology, the strain is considered tetracycline resistant. Complete absence of inclusions upon tetracycline exposure indicates that the strain is tetracycline sensitive. Strains falling in between these categories (“intermediate”) should be further analyzed with the in vitro susceptibility assay protocol described above.

Immunofluorescence analysis (IFA)

Inclusions and cell nuclei were visualized as described previously (Leonard et al., 2016). Briefly, primary Chlamydiaceae family-specific mouse monoclonal antibody directed against the chlamydial lipopolysaccharide (LPS, Clone ACI-P; Progen, Heidelberg, Germany; 1:200) and secondary Alexa Fluor 488-conjugated secondary goat anti-mouse antibody (Molecular Probes, Eugene, OR, USA; 1:500) were used to label inclusions. Host and chlamydial DNA were labeled with 1 µg/ml 4′,6-diamidino-2′-phenylindole dihydrochloride (DAPI, Molecular Probes). If applicable (for MIC confirmation and recovery assay), coverslips were mounted with FluoreGuard mounting medium (Hard Set; ScyTek Laboratories Inc., Logan, UT, USA) on glass slides. All coverslips were analyzed with a Leica DMLB fluorescence microscope (Leica Microsystems, Wetzlar, Germany), while 96-well plates were analyzed with a Nikon Eclipse Ti microscope (Nikon, Tokyo, Japan).

Transmission electron microscopy (TEM)

TEM images were produced according to previously published methods (Leonard et al., 2016). Ultrathin (80 nm) sections were mounted on gold grids (Merck) and contrasted with uranyl acetate dehydrate (Fluka; Sigma-Aldrich) and lead citrate (Merck). Sections were subsequently evaluated using a Philips CM10 electron microscope (Software release version 5.1; FEI Company, Hillsboro, OR, USA) and imaged using a Gatan Orius SC 1000 CCD Camera with software version Digital Micrograph 2.30 (Gatan Inc., Warrendale, PA, USA).

Statistical analysis

Unless stated otherwise, results were displayed as means ± standard deviation, of the results from 2 or 3 independent experiments. Statistical significance of the difference of
means was determined by Student’s t-test and Welch t-test (t-test unequal variance) using GraphPad QuickCales Web site: http://www.graphpad.com/quickcalcs/test1/ (accessed January 2018) and the Excel software. p-values of <0.05 were considered significant.

Results

Tetracycline susceptibility can be determined using multiple evaluation parameters

Initial susceptibility phenotype and MIC determination

The initial tetracycline susceptibility phenotype of four C. suis strains (field strains SWA-14, SWA-86 and SWA-141, and reference strain S45/6), all originally isolated from farmed pigs, was determined after two to three workdays. This was accomplished by simultaneously seeding and infecting LLC-MK2 cells with Chlamydia in cycloheximide-free medium with two-fold tetracycline dilutions (Week 1, Table 3) and determining the MIC as described by Donati et al. (2010) (Table 3). In confirmation of previously published results (Dugan et al., 2004), and in accord with tetracycline susceptibility definitions described (Wanninger et al., 2016), S45/6 was sensitive to tetracycline (MIC < 2 µg/ml). Amongst the field strains, SWA-141 was resistant (MIC ≥ 4 µg/ml), and SWA-14 and SWA-86 were sensitive, and no strain showed intermediate tetracycline susceptibility (2 µg/ml ≤ MIC < 4 µg/ml) (Dugan et al., 2004; Wanninger et al., 2016). The MIC was confirmed by infecting confluent host cell monolayers in the presence of tetracycline concentrations several two-fold dilutions above and below the initially determined MIC in cycloheximide-containing medium (Week 2, Table 3). MIC determination was performed both according to Donati et al. (2010) and Suchland et al. (2003) to establish the MIC consensus. Cycloheximide was not added in the first week to ensure cellular growth upon simultaneous seed and infection, while it was added in the second week to a) analyze whether cycloheximide potentially influences the susceptibility of Chlamydia to tetracycline and other antibiotics and b) to avoid overgrowth of the cell monolayers.

Inclusion size analysis

Considering data provided by Suchland et al. (2003), a reduction in inclusion numbers can be expected to be preceded or accompanied by altered inclusion size and/or morphology. In order to quantify and further confirm the MIC (Suchland), we analyzed the average inclusion size for each strain at 0.03 µg/ml, 0.125 µg/ml and 0.5 µg/ml tetracycline. We clearly observed that the average inclusion size of all three tetracycline sensitive strains (SWA-14, SWA-86, S45/6) was significantly reduced at tetracycline concentrations as low as 0.03 µg/ml, while the average inclusion size of tetracycline resistant SWA-141 exposed to ≤ 0.5 µg/ml tetracycline was similar to the unexposed control (mock, Fig. 2).

Tetracycline resistance screen (Tet Screen)

The observed variation in inclusion size in tetracycline resistant versus sensitive strains led us to hypothesize that it might be possible to easily distinguish resistant from sensitive strains by comparing the unexposed controls to cultures exposed to 0.125 µg/ml or 0.5 µg/ml tetracycline, which would allow screening of a large number of strains in less than 48 h post infection (hpi). Fig. 3 shows that the inclusion number and size/morphology of the tetracycline resistant strain SWA-141 was similar regardless of tetracycline concentration, while no or only micro-inclusions (area less than 15 µm²) were visible in tetracycline-exposed cultures infected with tetracycline sensitive SWA-14, 86 or S45/6. To further confirm our hypothesis, we screened additional previously reported tetracycline resistant (5-27b, 1-28b, 5-22b) and sensitive (10-26b, 1-28a) C. suis field strains from previous studies (Seth-Smith et
al., 2017; Wanninger et al., 2016). As expected, we also observed that inclusion number and morphology of the three tetracycline resistant strains was not affected by tetracycline exposure while the two sensitive strains showed no or only micro-inclusions at 0.125 µg/ml and 0.5 µg/ml tetracycline (Fig. S1). In addition, we screened two human chlamydial strains, C. trachomatis serovar E and C. pneumoniae Kajaani 6, and confirmed their susceptibility to tetracycline, as indicated by the described inclusion size criteria, as expected (Fig. S2).

**Recovery assay**

Finally, we analyzed the infectivity of strains SWA-14, 86, 141 and S45/6, in terms of recovery, upon exposure to tetracycline followed by discontinuation of exposure to tetracycline versus uninterrupted, continuous exposure to tetracycline. Briefly, in parallel to the initial susceptibility phenotype, cultures were prepared for the recovery assay in cycloheximide-free medium. However, instead of fixation of the infected cells on coverslips, followed by immunofluorescence microscopic analysis to determine the phenotype, infected cells were washed two times with fresh medium, to remove residual drugs, and the medium was replaced with either medium alone (recovery, rec) or the same concentrations of tetracycline (continued exposure, exp) and incubated for another 48 h prior to scraping collection of infected cells to determine infectivity. In Week 2, in parallel to performing MIC confirmation, collected samples were inoculated onto fresh cells in cycloheximide-containing medium to determine the IFU/ml (see the Material and Methods section). The complete, detailed protocol is available in Text S1.

We found that even tetracycline sensitive strains (SWA-14, 86, S45/6) are able to recover from tetracycline concentrations as high as 2 µg/ml, despite MIC values of 0.125 µg/ml or less (Fig. 4). In contrast, no or only a minimal number of IFU (<0.001% of mock) were detectable following continued exposure to 0.5 and 2 µg/ml tetracycline for SWA-14, SWA-86 and S45/6, while the infectivity of SWA-141 cultures continuously exposed to 0.5 and 2 µg/ml was equivalent to 43.96% and 1.56% of the mock exposed infection, respectively.

In order to concisely summarize the data (here for tetracycline exposure, also useful for analyses of other antibiotics) and provide descriptive characterization of the strains, we developed the following data analyses for the data (see Table 4): 1) “Resistance potential” (compares exposure to mock groups and represents the degree to which Chlamydia resist continuous exposure to the antibiotic in question) indicates the highest tetracycline concentration at which tetracycline exposed cultures exhibit infectivity equivalent to a) >25% or b) >10% of mock-exposed culture infectivity. 2) “Recovery potential” (compares recovery to mock groups and represents the degree to which Chlamydia recover from antibiotic exposure) indicates the highest antibiotic concentration at which cultures initially exposed to tetracycline, but then further cultured in the absence of tetracycline (recovered cultures), exhibit infectivity equivalent to a) >1% or b) >10% of mock-exposed culture infectivity. Lastly, 3) “Survival after continued exposure” (directly compares the infectivity of continuously tetracycline exposed cultures to recovered cultures [exposed to recovered groups]) indicates the highest antibiotic concentration at which continuously exposed cultures exhibit infectivity equivalent to a) >1% or b) >10% of recovered culture infectivity. The result of these analyses allowed clear differentiation between tetracycline resistant (SWA-141) and sensitive (SWA-14, SWA-86, and S45/6) strains (Table 4).

**Reference strain S45/6 is sensitive to sulfamethoxazole, while all tested C. suis field strains are sulfamethoxazole resistant**

*Initial susceptibility phenotype, inclusion size analysis and MIC determination*
As for tetracycline, we aimed to analyze the susceptibility of the same four *C. suis* strains to sulfamethoxazole. However, determination of the initial susceptibility phenotype according to Donati et al. (2010) was not possible due to the fact that we only observed sulfamethoxazole-dependent changes in inclusion size/morphology, but no sulfamethoxazole-dependent changes in the number of inclusions, even at concentrations as high as 512 µg/ml sulfamethoxazole compared to the mock-exposed control (Fig. S3A). As a result, initial susceptibility phenotype determination was performed only according to Suchland et al. (2003), resulting in MIC of 128-256 µg/ml, 64-128 µg/mL, 128-256 µg/ml and 0.0039 µg/ml for SWA-14, 86, 141 and S45/6, respectively.

To confirm these MIC, we evaluated the average inclusion size per strain at sulfamethoxazole concentrations close to the initially determined MIC as performed for tetracycline. Fig. 5A illustrates the inclusion size differences allowing initial determination of MIC (per Suchland) and subsequent confirmation. The entire data set is summarized in Table 5.

Interestingly, upon MIC confirmation, we observed that the three field strains exposed to sulfamethoxazole at their respective MIC (ranging from 64 to 256 µg/ml) universally contained smaller inclusions by immunofluorescence assay (IFA) compared to the mock-exposed infection. Those inclusions appeared to contain large aberrant bodies (AB) indicative of the chlamydial stress response. In contrast, while a significant inclusion size reduction was apparent upon exposure to 0.0039 µg/ml sulfamethoxazole for reference strain S45/6, ABs were only sporadically present in the reference strain (Fig. 5A).

**Transmission electron microscopy (TEM) analysis of sulfamethoxazole-exposed cultures**

To confirm IFA observations regarding the presence of ABs, we performed TEM analysis (Fig. 5B). ABs were defined as pale, large inclusions of round to irregular shape with \( \geq 2 \mu m \) in diameter (Matsumoto and Manire, 1970). While all mock-exposed cultures had inclusions primarily populated with small elementary bodies (EBs; dark, 0.25-0.5 \( \mu m \) in diameter) and reticulate bodies (RBs; pale, 0.5-1 \( \mu m \) in diameter), field strain cultures exposed to 128 µg/ml sulfamethoxazole contained inclusions populated with a large number of ABs or markedly altered RBs. In contrast, S45/6 cultures exposed to 0.015 µg/ml sulfamethoxazole contained a decreased number of EBs and RBs but no obvious increase in the number of ABs, confirming IFA results.

**Recovery assay**

Finally, we analyzed the infectivity of strains SWA-14, SWA-86, SWA-141 and S45/6 in terms of recovery upon exposure to sulfamethoxazole followed by discontinuation of exposure to sulfamethoxazole versus continuous exposure to sulfamethoxazole. The assay was performed identically to the described tetracycline recovery assay.

As expected from the initial susceptibility phenotype, wherein no sulfamethoxazole-dependent inclusion number reduction was detected, all strains showed recovery up to 512 µg/ml sulfamethoxazole though the number of IFU was distinctly reduced to only 0.18%, 0.22%, 0.35% and < 0.001% of the control for SWA-14, 86, 141 and S45/6, respectively. While the general recovery pattern, as well as survival following continuous sulfamethoxazole exposure, was comparable for the three field strains, S45/6 showed a relative reduction of both parameters compared to the field strains (Fig. 6). This sulfamethoxazole susceptibility difference between the field strains and the type strain was even more pronounced when additional sulfamethoxazole concentrations were evaluated (2, 8, 32, 128, 512 µg/ml sulfamethoxazole for field strains and 0.0039, 0.03, 0.25, 2, 8, 32, 128, 512 µg/ml sulfamethoxazole for S45/6, Fig. S3B).

Again, as for tetracycline analysis above, we compiled a table containing the following data of sulfamethoxazole analyses: 1) “resistance potential” (compares exposure to mock
groups), 2) “recovery potential” (compares recovery to mock groups) and 3) “survival after continued exposure” (compares exposure to recovery groups). The results of these analyses allowed clear differentiation between sulfamethoxazole resistant field strains (SWA-14, 86, 141) and the sulfamethoxazole sensitive reference strain S45/6 (Table 6).

**Penicillin G (PenG) induces the chlamydial stress response in C. suis strains**

*Initial susceptibility phenotype, inclusion size analysis and MIC determination*

PenG is known to induce the chlamydial stress response or persistence in chlamydial species such as *C. trachomatis* and *C. muridarum*, the closest phylogenetic relatives of *C. suis* (Joseph et al., 2016; Kintner et al., 2014; Phillips Campbell et al., 2012; Seth-Smith et al., 2017). In our study, we performed a PenG 10-fold serial dilution ranging from 0.001 to 100 units (U)/ml PenG and observed that almost 100% of *C. suis* inclusions appeared to be altered in terms of morphology with the presence of ABs at almost all concentrations tested (0.1-100 U/ml for the field strains and 0.01-100 U/ml for S45/6, data not shown). At 0.001 U/ml PenG, less than 90% of inclusions were aberrant in all evaluated strains. We subsequently performed MIC determinations while evaluating inclusion size and morphology at PenG concentrations of 1, 10 and 100 U/ml. Interestingly, while the inclusion morphology clearly showed signs of persistence in all four strains (SWA-14, SWA-86, SWA-141, S45/6) even at the lowest PenG concentration of 1 U/ml, the average inclusion size remained similar to that of the mock control (Fig. 7A). The observation of PenG-induced ABs in *C. suis* was confirmed by TEM comparing inclusions of S45/6 exposed to 1 U/ml and 100 U/ml PenG with the mock-exposed control. PenG-exposed *C. suis* inclusions consisted of mostly empty inclusions containing few ABs (Fig. S4A) as described recently for *C. trachomatis* and *C. pecorum* (Kintner et al., 2014; Leonard et al., 2016).

**Recovery assay**

As a final step, we analyzed the infectivity of strains SWA-14, SWA-86, SWA-141 and S45/6 in terms of recovery upon exposure to PenG followed by discontinuation of exposure to PenG versus continuous exposure to PenG. Recovery assays were performed identically to the described tetracycline and sulfamethoxazole recovery assays. As expected, all the *C. suis* strains that were evaluated exhibited increased infectivity upon discontinuation of PenG exposure for 48 h, after 48 h of culture in the presence of PenG, compared to the continuous PenG-exposure group. Though the recovery group infectivity never exceeded 0.001% of the infectivity of the unexposed control (which was inoculated with *Chlamydia* and cultured for 48 h in the absence of PenG exposure), discontinuation of PenG exposure for 48 h resulted in infectivity levels 10-20 fold greater than those observed for cultures subjected to continuous PenG exposure for the same duration, indicative of recovery from PenG-induced infectivity reduction. Very little infectivity (<150 IFU/ml)/single inclusions were detected in the recovery assay for continuously exposed cultures for all PenG concentrations evaluated (Fig. 7B). Again, TEM analysis was performed for S45/6 to further demonstrate recovery after removal of PenG. When PenG exposure was discontinued after 48 h of exposure and culture was continued in the absence of PenG for 48 h, inclusions contained normal EBs and RBs, while continuously exposed cultures continued to contain only few inclusion bodies primarily consisting of ABs (Fig. S4B).

**Discussion**

While there are numerous standardized methods to evaluate extracellular and facultative intracellular bacteria regarding susceptibility to antibiotic agents, there are no standardized antibiotic susceptibility assays for *Chlamydia*. Instead, several research groups...
have, over time, developed multiple protocols to determine the antibiotic susceptibility of 
*Chlamydia in vitro* with microscopic methods as summarized in Table 1 (Agacfidan et al.,
1993; Donati et al., 2010; Ehret and Judson, 1988; Henning and Krauss, 1986; Japan Society
of Chemotherapy, 1992a, 1992b; Suchland et al., 2003; Webberley et al., 1987). In addition to
these classic *in vitro* susceptibility protocols, there are other systems such as antibiotic
susceptibility in continuous-infection models instead of regular infection *in vitro* (Kutlin et
al., 1999), *in vitro* susceptibility testing using flow cytometry instead of classic determination
via microscopic reading (Dessus-Babus et al., 1998), and the use of a reverse transcriptase
PCR (RT-PCR)-based method instead of inclusion number determination (Cross et al., 1999).

Regardless of the applied method, a clear definition of the MIC and other evaluation
parameters is needed to implement new approaches to determine the *in vitro* antibiotic
susceptibility of *Chlamydia*. Although there has been no gold standard or consensus in the
field for a standardized protocol for determination of antibiotic susceptibility and resistance,
all research groups prior to 2000 defined the MIC as the lowest concentration where no
inclusions were found (Agacfidan et al., 1993; Ehret and Judson, 1988; Hammerschlag, 1982;
Henning and Krauss, 1986; Japan Society of Chemotherapy, 1992a, 1992b; Webberley et al.,
1987). More recent protocols from the 2000s included the principle of inclusion number
reduction by 90% or more (Donati et al., 2010) or alterations in inclusion size and
morphology (Suchland et al., 2003). Suchland and colleagues (2003) specifically noted that it
might be problematic to define the MIC according to few survivors at high antibiotic
concentrations (heterotypic survival), since micro-inclusions may not be visible depending on
the staining/labeling method and magnification used. In view of these publications, we
developed a consensus MIC considering previous MIC definitions according to Donati et al.
(2010) and Suchland et al. (2003).

Unlike already published protocols to determine the MIC, which are analyzed
differently but processed analogously, the method to determine MBC/MCC/MLC strongly
depends on the research group (Table 1). In detail, while one half of all research groups
defined the MBC/MCC/MLC as the lowest concentration where there were no inclusions after
passaging the strain once in drug-free medium (Agacfidan et al., 1993; Ehret and Judson,
1988; Suchland et al., 2003; Webberley et al., 1987), the rest defined the MBC/MCC/MLC as
the lowest antibiotic concentration where there were no or 90% fewer inclusions following re-
incubation in drug-free medium compared to the mock-exposed control (Donati et al., 2010;
Henning and Krauss, 1986; Japan Society of Chemotherapy, 1992a, 1992b) Regardless of the
protocol, all MBC/MCC/MLC protocols only determine one value, which is usually identical
or only a few twofold dilutions higher compared to its MIC. In this study, in order to
complement the MIC and to further characterize the chlamydial response to antibiotic
exposure *in vitro*, we decided to employ the recovery assay, which has so far been described
in studies investigating the chlamydial stress response (Kintner et al., 2014; Leonard et al.,
2015, 2017). With this recovery assay, instead of determining one single value that gives little
additional information to the MIC, we evaluated low, intermediate, and high antibiotic
concentrations for effects upon chlamydial infectivity after discontinuation of antibiotic
exposure 48 hpi (recovery) or continued exposure for 96 h. Not only does it allow us to
confirm the impact of inclusion size reduction on subsequent infectivity, it further implements
the MBC/MCC/MLC (recovery) and expands previous protocols to observing infectivity upon
continued exposure.

In the present study, we evaluated the antibiotic susceptibility of three porcine field
strains in comparison to the *C. suis* reference strain S45/6 (also of porcine origin) to
tetracycline, sulfamethoxazole and penicillin. These antibiotic agents were chosen because of
their extensive use in the pig farming industry (Hoffmann et al., 2015). Moreover, the
presence of the tetracycline resistance gene *tetA(C)* has been reported in *C. suis* worldwide
(USA, several European countries, Israel, China) (Borel et al., 2012; Dugan et al., 2004; Di
Francesco et al., 2008; Joseph et al., 2016; Lenart et al., 2001; Li et al., 2017; Schautteet et al., 2013; Seth-Smith et al., 2017; Wanninger et al., 2016). Therefore, tetracycline, a bacteriostatic protein synthesis inhibitor preventing the binding of bacterial aminoacyl-t-RNA to the mRNA-ribosome complex, is well suited as proof of concept for susceptibility assay development and evaluation, as there are clear, well-described features separating resistant from sensitive strains (Donati et al., 2010; Suchland et al., 2003). We further found that even sensitive strains are able to recover from tetracycline concentrations well above the MIC, which is indicative of heterotypic survival as reported by Suchland et al. (2003). Nonetheless, the observed infectivity pattern following recovery and continued exposure is markedly different between sensitive and resistant strains, which is expected based on the MIC data.

With application of the initial susceptibility phenotype analysis that can be performed within two to three workdays and subsequent MIC confirmation that can completed within two to three additional workdays, we showed that a) simultaneous host cell seeding and chlamydial infection into culture medium with appropriate antibiotic dilutions yields similar results regarding susceptibility to more the time-consuming infection of confluent monolayers and that b) cycloheximide treatment does not appear to influence MIC determination. With data from this study and those of other research groups (Donati et al., 2016; Suchland et al., 2003), we established a fast and simple screening method to detect tetracycline resistant strains. Our Tet screen allows the evaluation of ten strains per 24-well plate within three days with minimal use of consumables where cultures are treated with 0.5 µg/ml tetracycline and compared to mock-exposed cultures. If inclusion size/morphology and number are similar to that of the corresponding tetracycline-unexposed mock control, the strain is considered to be tetracycline resistant. If there are no, or only small, aberrant inclusions, the strain is considered to be tetracycline sensitive. All “intermediate” stages, such as few but regular sized inclusions, should be processed further by a) determining the initial susceptibility phenotype, b) performing subsequent MIC confirmation and c) performing the recovery assays. So far, no “intermediate” stages were detected, but a larger sample size must be evaluated to further validate this method.

Information regarding the susceptibility of *C. suis* to sulfonamides such as sulfamethoxazole, a bacteriostatic inhibitor of folate synthesis by competition with the substrate para-aminobenzoic acid (PABA) (Marwaha et al., 2014), is very limited. Reports generally state that *C. suis* appears to be sensitive to sulfonamides with the exception of a few tetracycline resistant strains (Andersen and Rogers, 1998; Sandoz and Rockey, 2010). These reports are contrary to our findings wherein all three field strains, of which two were sensitive to tetracycline, had sulfamethoxazole MICs of 64 µg/ml or higher. Moreover, MIC determination was only possible according to the MIC determination method by Suchland et al. (2003), because the inclusion number was not significantly influenced by sulfamethoxazole. Here, the recovery assay was crucial to confirm that the alteration in terms of size and morphology strongly impacted the infectivity upon recovery and continued exposure and therefore further served as a confirmation of the MIC. Interestingly, reporting results from a study on *C. trachomatis*, which defined the MIC as the concentration of sulfamethoxazole for which no inclusions are seen, the authors were able to determine susceptibilities for sulfamethoxazole in the range of 2 to 128 µg/ml (Hammerschlag, 1982).

However, the authors specifically noted that only low inocula (≤ 1000 IFU) could be used to yield these results. Larger inocula did not yield an MIC according to their definition, but they showed a sulfamethoxazole concentration-dependent alteration in inclusion size and morphology. The susceptibility range (2-128 µg/ml) to sulfamethoxazole reported for *C. trachomatis* appears to be similar to that found for *C. suis* although the MIC of S45/6 was below 0.01 µg/ml. From the small number of strains investigated in our study, it appears that field strains are mostly resistant to sulfamethoxazole independent of resistance to tetracycline, while the *C. suis* type strain S45/6 is sulfamethoxazole sensitive. However, a larger sample
size is necessary to confirm this finding. Moreover, these and other *C. suis* field strains should also be investigated for resistance to N-acylated sulfonamide derivatives because, despite sharing the structural core with sulfamethoxazole and sulfafurazole, they operate via a distinct working mechanism (Marwaha et al., 2014; Mojica et al., 2017): These antibiotics do not affect folate synthesis but instead bind directly to the 3-oxoacyl-(acyl carrier protein [ACP] synthase II [FabF]) thus inhibiting the essential type II fatty acid synthesis (FASII) pathway.

Given that almost every chlamydial species evaluated to date responds with the chlamydial stress response upon treatment with β-lactam antibiotics (Beatty et al., 1994; Galasso and Manire, 1961; Hogan et al., 2004; Matsumoto and Manire, 1970; Schoborg, 2011; Tamura and Manire, 1968; Wyrick, 2010) with few exceptions (Dumoux et al., 2013), it is not surprising that all four investigated *C. suis* strains develop persistence from PenG treatment. However, while this in vitro assay allowed us to detect and describe persistence in our cultures, more appropriate protocols are available to investigate the chlamydial stress response (Kintner et al., 2014; Leonard et al., 2015, 2016, 2017). Nevertheless, we showed that *C. suis* could recover from PenG exposure in a similar manner to *C. trachomatis*, though infectivity was reduced by more than 99% upon exposure to 1 U/ml in both chlamydial species (Kintner et al., 2014). Interestingly, while TEM analysis in the *C. trachomatis* study revealed large ABs that filled the inclusion after exposure with penicillin, we found few grossly enlarged ABs that did not fill the entire inclusion. Ultrastructural differences in the chlamydial stress response are primarily caused by different persistence inducers, host cells and the *Chlamydia* species (Goellner et al., 2006; Hogan et al., 2004; Mukhopadhyay et al., 2006; Schoborg, 2011). Therefore, host cells (HeLa vs. LLC-MK2) or the chlamydial species (*C. trachomatis* vs. *C. suis*) may have caused the ultrastructural differences in this study compared to Kintner et al. (Kintner et al., 2014). In contrast, HeLa cells infected with the porcine *P. pecorum* strain 1710S and PenG-exposed (Leonard et al., 2016, 2017) revealed similar inclusions to those present in PenG-exposed *C. suis* strains described in this study.

One of the limitations of our study is that we did not test our protocols using different host cells, unlike previous reports from other authors (Ehret and Judson, 1988; Suchland et al., 2003). Suchland et al. (2003) found that there are notable differences regarding the MIC for macrolides (azithromycin, erythromycin) for *Chlamydia* spp. depending on the cell line used, for example McCoy, HeLa, BGMK, HEp-2, HL, or Vero cells. But the same was not shown for tetracycline, ofloxacin and doxycycline. The authors proposed that McCoy cells should be consistently used for *C. trachomatis* and HEp-2 for *C. pneumoniae* antibiotic susceptibility assays. So far, there is no specific recommendation for *C. suis* but McCoy, BGM and LLC-MK2 cells have all been successfully used for isolation and antibiotic susceptibility assays (Donati et al., 2010; Dugan et al., 2004; Schauette et al., 2013). For tetracyclines, cell-specific differences likely play a minor role as observed for *C. trachomatis* (Suchland et al., 2003), but no comparable study similar to that of Suchland et al. (2003) has been conducted for *C. suis* so far. Other potentially relevant factors not assessed in this study include medium constituents, incubation conditions and centrifugation protocols as well as timing/duration of antibiotic exposure and immunolabeling of inclusions (Giemsa, iodine, immunofluorescence) (Ehret and Judson, 1988). Additionally, the inoculum size is generally considered irrelevant for MIC determination but could influence the recovery assay if the inoculum size is below 5000 IFU/well (Suchland et al., 2003).

Furthermore, our study has primarily focused on the classic microscopic reading methods. In the future, it is vital consider alternative methods that were first published in the late 1990s such as the RT-PCR method (Cross et al., 1999). They found that smaller and/or aberrant inclusions still produce detectable levels of mRNA and are therefore potentially viable further supporting the results of the recovery assay applied in this study and the high MBC/MCC/MLC values from other protocols (Agacfidan et al., 1993; Donati et al., 2010; Ehret and Judson, 1988; Henning and Krauss, 1986; Japan Society of Chemotherapy, 1992a,
Another method is the use of flow cytometry (Dessus-Babus et al., 1998), which was not considered to be as sensitive as the direct microscopic assessment method but still needs to be considered for its reproducibility and objective interpretation. Additionally, flow cytometry must be considered as a potential titration method (Käser et al., 2016) for the recovery assay as it has shown to be highly reproducible, faster with lower material cost than traditional titration methods.

In summary, we propose new approaches to evaluate the antibiotic susceptibility of *C. suis* and other *Chlamydia* spp. by creating a consensus MIC based on inclusion number reduction and size/morphology alteration. This approach allows the determination of a susceptibility range for antibiotic agents and chlamydial species that have not been tested so far. Finally, we propose a simple and fast screening method to detect tetracycline resistant *C. suis* strains.

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Author Contributions Statement

Substantial contribution to the conception and design of the work: HM, CL, NB, DD. Acquisition, analysis, interpretation of data: HM, CL, NB. Draft and/or critical revision of the manuscript: HM, CL, NB, DD. Final approval of the version to be published: HM, CL, NB, DD. All authors agree to be accountable for all aspects of the work.

Conflict of Interest Statement

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Figure Legends

Figure 1. Overview of the in vitro antibiotic susceptibility assay protocol. Shown is an overview of the in vitro assay protocol grouped into the first and second week of the assay and divided into initial susceptibility phenotype/MIC confirmation and the recovery assay.

Figure 2. Average inclusion size analysis upon tetracycline exposure. The bar graphs compare the average inclusion size (µm²) of unexposed controls (mock) with tetracycline-exposed (0.03, 0.125 and 0.5 µg/ml) cultures at 34 hours post infection (mean ± SD). Shown are the results for strains A) SWA-14, B) SWA-86, C) SWA-141 and D) reference strain S45/6. Asterisks indicate a statistically significant difference between the tetracycline-exposed cultures and the mock by both Student’s t-test and the Welch t-test (ns = not significant; * p < 0.05; ** p < 0.01). Three independent experiments were performed (n = 3); 0.125 and 0.5 µg/ml tetracycline were only evaluated in two of the three experiments (n = 2).

Figure 3. Tetracycline resistance screen of strains SWA-14, 86, 141 and S45/6. Shown are immunofluorescence images illustrating the results of the tetracycline resistance screen (TET Screen) wherein inclusion number and morphology of mock-exposed chlamydial strains are compared to cultures exposed to 0.125 or 0.5 µg/ml tetracycline. The white bar represents 50 µm.

Figure 4. Recovery assay following chlamydial infection and tetracycline exposure. The bar graphs compare the average inclusion forming units per ml (IFU/ml) of unexposed controls (mock) with tetracycline-exposed (0.03, 0.5 or 2 µg/ml) chlamydial cultures (mean ± SD). Cultures were either continuously exposed (exp) to tetracycline for 96 hours (h) or exposed to tetracycline for 48 h, then further cultured in tetracycline-free medium for 48 h (recovery, rec). Shown are the results for strains A) SWA-14, B) SWA-86, C) SWA-141 and D) reference strain S45/6. Three independent experiments were performed (n = 3).

Figure 5. The effects of sulfamethoxazole exposure. A) Shown are bar graphs comparing the average inclusion size of sulfamethoxazole-treated cultures to the mock-exposed cultures (left; mean ± SD) as well as representative immunofluorescence images of the inclusion morphology of the untreated control (right, top) and the MIC<sub>TP</sub> (right, bottom) for strains SWA-14, SWA-86, SWA-141 and S45/6 at 48 hours (h) post infection. Asterisks indicate a statistically significant difference between the sulfamethoxazole-exposed cultures and the mock by both Student’s t-test and the Welch t-test (ns = not significant; * p < 0.05; ** p < 0.01). Three independent experiments were performed (n = 3). B) Representative transmission electron microscopy (TEM) images are shown for each strain wherein the mock is presented on the left part of the panel and sulfamethoxazole-exposed cultures are presented on the right. Inclusions are indicated in green (I), the nucleus in dark red (N) and, if applicable, the nucleolus in bright red (Nc). Cultures were fixed at 48 h post infection for processing for TEM analysis.

Figure 6. Recovery assay following chlamydial infection with sulfamethoxazole exposure. The bar graphs compare the average inclusion forming units per ml (IFU/ml) of unexposed controls (mock) with sulfamethoxazole-exposed (2, 32 and 512 µg/ml) cultures (mean ± SD). Cultures were either continuously exposed (exp) to sulfamethoxazole for 96 hours (h) or exposed to sulfamethoxazole for 48 h, then further cultured in sulfamethoxazole-free medium for 48 h (recovery, rec). Shown are the results for strains A) SWA-14, B) SWA-86, C) SWA-141 and D) reference strain S45/6. Three independent experiments were performed (n = 3).
**Figure 7. Penicillin G induces the chlamydial stress response in all C. suis strains. A)** Shown are bar graphs comparing the average chlamydial inclusion size of Penicillin G (PenG)-exposed cultures to the mock-exposed control (left; mean ± SD) as well as representative immunofluorescence microscopy images of inclusion morphology of both the unexposed control (right, top) and cultures exposed to 1 units (U)/ml of PenG (right, bottom) for strains SWA-14, SWA-86, SWA-141 and S45/6 at 48 hours (h) post infection. Two independent experiments were performed (n = 2). Asterisks indicate a statistically significant difference between the PenG-exposed cultures and the mock by both Student’s t-test and the Welch t-test (ns = not significant; * p < 0.05; ** p < 0.01). **B)** The bar graphs compare the average inclusion forming units per ml (IFU/ml) of unexposed controls (mock) with PenG-exposed (1, 10 and 100 U/ml) cultures (mean ± SD). Cultures continuously exposed to PenG for 96 h (exp) were compared with cultures exposed to PenG for 48 h and then further cultured in PenG-free medium for 48 h (recovery, rec). Shown are the results for strains SWA-14 (top left), SWA-86 (top right), 141 (bottom left) and the reference strain S45/6 (bottom right). Two independent experiments were performed (n = 2).

**Supplementary Material**

**Figure S1. Tetracycline resistance screen of additional C. suis field strains.** Shown are immunofluorescence images illustrating the results of the tetracycline resistance screen (TET Screen) where inclusion number and morphology of mock-exposed chlamydial strains are compared to cultures exposed to 0.125 or 0.5 µg/ml tetracycline. Strains used in this experiment were previously analyzed according to the Donati method (Donati et al. 2010. *Antimicrob. Agents Chemother.* doi:10.1128/AAC.00553-10), as reported in (Wanninger et al. 2016. Plos One. doi: 10.1371/journal.pone.0166917). In the previous study, 10-26b was interpreted as sensitive to tetracycline, 5-27b and 5-22b as resistant to tetracycline. Also during the course of the previous study, strains 1-28a and 1-28b were also determined as sensitive and resistant, respectively (unpublished data). The respective absence and presence of the tetracycline resistance gene was confirmed in a follow-up comparative genomics study (Seth-Smith et al. 2017. Genome Biol Evol. doi: 10.1093/gbe/evx043. ENA deposition PRJEB17986).

**Figure S2. Tetracycline resistance screen of human-infecting pathogens, C. trachomatis serovar E and C. pneumoniae Kajaani 6 (K6).** Shown are immunofluorescence images illustrating the results of the tetracycline resistance screen (TET Screen) wherein inclusion number and morphology of mock-exposed chlamydial strains are compared to cultures exposed to 0.125 or 0.5 µg/ml of tetracycline. Strains used in this experiment were two reference strains, a human genital strain *C. trachomatis* (Ci) serovar E, originally obtained from S. P. Wang and C. C. Kuo (University of Washington, Seattle, WA, USA; Holt et al. 1967. Am J Ophthalmol) and the human *C. pneumoniae* (Cp) Kajaani 6 (K6) strain (kindly provided by Dr Claudia Dumrese and Dr Urs Ziegler, Center for Microscopy and Image analysis, University of Zurich, Switzerland). Neither strain is positive for the tetracycline resistance gene (ENA deposition PRJEB17986).

**Figure S3. Additional analyses for sulfamethoxazole exposure. A)** The bar graphs show the average inclusion number reduction upon exposure of cultures to 256 or 512 µg/ml sulfamethoxazole (SULF), expressed as percent of the mock-exposed control (mean ± SD). **B)** The bar graphs compare the mean inclusion forming units per ml (IFU/ml) of unexposed controls (mock) with sulfamethoxazole-exposed *Chlamydia* cultures (2, 8, 32, 128 and 512 µg/ml SULF for the field strains and additionally 0.0039, 0.03 and 0.25 µg/ml SULF for the reference strain S45/6; mean ± SD). Cultures continuously exposed (exp) to sulfamethoxazole ...
for 96 hours (h) were compared with cultures grown for 48 h in SULF-free medium after 48 h of exposure to SULF (recovery, rec). Shown are the results for strains SWA-14 (top left panel), SWA-86 (top right), 141 (bottom left) and reference strain S45/6 (bottom right). Three independent experiments were performed (n = 3).

Figure S4. Transmission electron microscopy (TEM) analysis of Penicillin G exposure in C. suis strain S45/6. A) Shown is the inclusion morphology of the reference strain S45/6 39 hours (h) post infection for mock-exposed cultures (left), cultures treated with 1 U/ml (middle) and 100 U/ml (right) Penicillin G (PenG). The scale bar indicates 5 µm. B) A recovery assay was performed for the mock (no PenG) and 1 U/ml PenG groups. At 39 h post infection, antibiotic-free (mock) or PenG-containing (1 U/ml) supernatant was removed from the monolayer and replaced with PenG-free (mock, recovery) or PenG-containing (continued exposure) medium. At 87 h, cultures were fixed and further processed for TEM analysis. The scale bar indicates 5 µm.

Text S1. Complete in vitro antibiotic susceptibility assay protocol. Shown is the detailed protocol for the complete in vitro assay for tetracycline, sulfamethoxazole and penicillin. In addition, suggestions regarding assay optimization are included.
| Group/Ref | Definition of minimal inhibitory concentration (MIC) and minimal bactericidal/chlamydicidal/lethal concentration (MBC/MCC/MLC) | Cells | Species |
|----------|-------------------------------------------------------------------------------------------------|-------|---------|
| Suchland et al. (2003) | **MIC**: “The transition point MIC (MIC<sub>TP</sub>) was defined as the concentration of drug in which 90% or more of the inclusions were altered in size and morphology. The MIC was defined as the concentration of drug that is one twofold dilution more concentrated than the MIC<sub>TP</sub>.” (p. 637)  
**MCC**: “The lowest concentration of drug that produced no morphologically normal inclusions by one freeze-thaw passage (…) in antimicrobial-free medium.” (p. 637) | HeLa, BGMK, HEp-2, Vero | CT<sup>1</sup>, Cpsi<sup>2</sup>, Cpne<sup>3</sup> |
| Donati et al. (2010) | **MIC**: “The lowest concentration that reduced the number of inclusions more than 90% compared with the level for drug-free controls.” (p. 5380)  
**MBC**: “The MBC concentration was measured by aspirating the antibiotic-containing medium, washing the monolayer twice with PBS, and incubating it in (…) antibiotic-free medium for 48 hours (h) at 35°C. (…). The MBC was the lowest concentration of the drug reducing more than 90% demonstrable inclusions after monolayers were re-incubated in antimicrobial-free medium.” (p. 5380) | LLC-MK2 | CT<sup>1</sup> |
| Japan Society of Chemotherapy (1992a, 1992b) | **MIC**: “The lowest concentration, which completely inhibits inclusion body formation.” (p. 309)  
**MLC**: “The lowest concentration, which completely inhibits the reformation of chlamydial inclusion in infected HeLa 229 cells even after elimination of the drug from the culture medium (after 24 h).” (p. 318) | HeLa, McCoy | CT<sup>1</sup>, Cpsi<sup>2</sup>, Cpne<sup>3</sup> |
| Ehret and Judson (1988), Hammerschlag (1982) | **MIC**: “The MIC should be defined as no inclusions seen.” (Ehret, p. 1298; Hammerschlag, p. 500)  
**MBC**: “(…) the MBC should be defined as no inclusion seen on passaging.” (Ehret, p. 1298) | McCoy | CT<sup>1</sup> |
| Webberley et al. (1987) | **MIC**: “The MIC was defined as the lowest concentration of antibiotic that inhibited inclusion development after 48 h.” (p. 409)  
**MLC**: “(…) the lowest concentration that prevented the reappearance of inclusions after re-incubation in antibiotic free media.” (p. 409) | McCoy | Cpsi<sup>2</sup>, Cpne<sup>3</sup> |
| Schachter et al. (Agacfidan et al., 1993) | **MIC**: “The MIC for *Chlamydia* was defined as the concentration of antibiotic that allowed no inclusions on the first passage.” (p. 1993)  
**MCC**: “The MCC was defined as the lowest concentration that allowed no inclusions in a further passage in the absence of antibiotics.” (p. 1993) | McCoy | CT<sup>1</sup> |
| Henning and Krauss (1986) | **MIC** (MHK, Minimale Hemmkonzentration): “The lowest concentration where there are no inclusions.” (p. 448)  
**MBC** (Washed and reincubated, W & R): “The | BGM, McCoy | Cpsi<sup>2</sup> |
lowest concentration of drug that produced no inclusions after removal of the drug and several washing steps during incubation in drug-free medium.” (p. 448)

1 C. trachomatis, 2 C. psittaci, 3 C. pneumoniae

Table 2: Overview of Chlamydia suis strains used in this study

| Sample ID | Year of isolation, country of origin | Antibiotic treatment of pig | ENA deposition | Ref |
|-----------|-------------------------------------|-----------------------------|----------------|-----|
| SWA-14 (alt. 2-7 a R) | 2013, Switzerland | unknown | PRJEB17986 | This study |
| SWA-86 | 2013, Switzerland | unknown | PRJEB17986 | This study |
| SWA-141 (alt. 4-29 b R) | 2013, Switzerland | tetracycline | PRJEB17986 | (Seth-Smith et al., 2017; Wanninger et al., 2016) |
| S45/6 (ref) | 1960s, Austria | unknown | SRP076849 | (Joseph et al., 2016; Kaltenboeck et al., 1993) |

Table 3: Summary of the initial MIC determination (tetracycline)

| Week | Sample ID | MIC (per Donati) | MIC (per Suchland) | MIC consensus | Interpretation |
|------|-----------|------------------|--------------------|---------------|----------------|
| Week 1 | SWA-14 | 0.03 µg/ml | 0.03 µg/ml | 4 µg/ml | 0.06 µg/ml |
|       | SWA-86 | 0.03 µg/ml | 0.03 µg/ml | 4 µg/ml | 0.06 µg/ml |
|       | SWA-141 | 4 µg/ml | 4 µg/ml | 4 µg/ml | 0.06-0.125 µg/ml |
|       | S45/6 | 0.06 µg/ml | 0.03 µg/ml | 2 µg/ml | 0.06-0.125 µg/ml |

Table 4: Summary of the recovery assay (tetracycline), strain characterization

| Resistance potential | SWA-14 | SWA-86 | SWA-141 | S45/6 |
|----------------------|--------|--------|---------|-------|
| 1) Exposure to mock |        |        |         |       |
| >25%                 | <0.03 µg/ml | <0.03 µg/ml | 0.5 µg/ml | <0.03 µg/ml |
| >10%                 | <0.03 µg/ml | <0.03 µg/ml | 0.5 µg/ml | 0.03 µg/ml |
| 2) Recovery to mock |        |        |         |       |
| >10%                 | <0.03 µg/ml | <0.03 µg/ml | 0.5 µg/ml | 0.03 µg/ml |
| >1%                  | 0.03 µg/ml | <0.03 µg/ml | 2 µg/ml | 0.03 µg/ml |
| 3) Exposure to recovery |        |        |         |       |
| >10%                 | <0.03 µg/ml | <0.03 µg/ml | 2 µg/ml | 0.03 µg/ml |
| >1%                  | <0.03 µg/ml | <0.03 µg/ml | 2 µg/ml | 0.03 µg/ml |
Table 5: Summary of the MIC determination (sulfamethoxazole)

| Initially determined MIC | MIC confirmation | MIC consensus | Interpretation |
|--------------------------|-----------------|---------------|----------------|
| *SWA-14* 128-256 µg/ml | 256 µg/ml       | 128-256 µg/ml| resistant      |
| *SWA-86* 64-128 µg/ml | 64-128 µg/ml    | 64-128 µg/ml | resistant      |
| *SWA-141* 128-256 µg/ml| 128-256 µg/ml   | 128-256 µg/ml| resistant      |
| *S45/6* 0.0039 µg/ml | 0.0078 µg/ml    | 0.0039-0.0078 µg/ml | sensitive |

Table 6: Summary of the recovery assay (sulfamethoxazole), strain characterization

| 1) Exposure to mock | SWA-14 | SWA-86 | SWA-141 | S45/6 |
|---------------------|--------|--------|---------|-------|
| Resistance potential| >25%   | < 2 µg/ml | < 2 µg/ml | 2 µg/ml | < 0.0039 µg/ml |
|                      | >10%   | 2 µg/ml | 2 µg/ml | 2 µg/ml | < 0.0039 µg/ml |
| 2) Recovery to mock | SWA-14 | SWA-86 | SWA-141 | S45/6 |
| Recovery potential   | >10%   | 2 µg/ml | 2 µg/ml | 2 µg/ml | < 0.0039 µg/ml |
|                      | >1%    | 32 µg/ml | 2 µg/ml | 32 µg/ml | 0.0039 µg/ml |
| 3) Exposure to recovery | SWA-14 | SWA-86 | SWA-141 | S45/6 |
| Survival post exposure| >10%   | 2 µg/ml | 2 µg/ml | 2 µg/ml | < 0.0039 µg/ml |
|                      | >1%    | 2 µg/ml | 32 µg/ml | 2 µg/ml | 0.0039 µg/ml |

Interpretation: resistant resistant resistant sensitive