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Bacterial Cytological Profiling (BCP) as a Rapid and Accurate Antimicrobial Susceptibility Testing Method for *Staphylococcus aureus*

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**A B S T R A C T**

Successful treatment of bacterial infections requires the timely administration of appropriate antimicrobial therapy. The failure to initiate the correct therapy in a timely fashion results in poor clinical outcomes, longer hospital stays, and higher medical costs. Current approaches to antibiotic susceptibility testing of cultured pathogens have key limitations ranging from long run times to dependence on prior knowledge of genetic mechanisms of resistance. We have developed a rapid antimicrobial susceptibility assay for *Staphylococcus aureus* based on bacterial cytological profiling (BCP), which uses quantitative fluorescence microscopy to measure antibiotic induced changes in cellular architecture. BCP discriminated between methicillin-susceptible (MSSA) and -resistant (MRSA) clinical isolates of *S. aureus* (*n* = 71) within 1–2 h with 100% accuracy. Similarly, BCP correctly distinguished daptomycin susceptible (DS) from daptomycin non-susceptible (DNS) *S. aureus* strains (*n* = 20) within 30 min. Among MRSA isolates, BCP further identified two classes of strains that differ in their susceptibility to specific combinations of beta-lactam antibiotics. BCP provides a rapid and flexible alternative to gene-based susceptibility testing methods for *S. aureus*, and should be readily adaptable to different antibiotics and bacterial species as new mechanisms of resistance or multidrug-resistant pathogens evolve and appear in mainstream clinical practice.

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1. Introduction

We are locked in an evolutionary race with bacteria that rapidly become resistant to each new antibiotic, necessitating the continued development of new treatments and diagnostic approaches. The aptly coined “ESKAPE” pathogens (Rice, 2008) — *Enterococcus faecium, Staphylococcus aureus, Klebsiella pneumoniae, Acinetobacter baumannii, Pseudomonas aeruginosa* and *Enterobacter* species can exhibit multiresistance to all first line antibacterial drugs and pose significant clinical threats. Current therapeutic agents are losing effectiveness more rapidly than they are being replaced, and new drug discovery has stagnated due to high development costs coupled with a lower profit margin associated with antibiotics compared to other drug classes (Gwynn et al., 2010). Hence, in the recent decade there has been a push for the implementation of antimicrobial stewardship, a set of strategies to reduce the emergence and spread of resistant pathogens and to use existing antimicrobials more effectively. Rapid susceptibility testing plays a key role in antimicrobial stewardship by decreasing emergence of resistance and by reserving drugs of last resort to those cases in which they are warranted (Goff et al., 2012).

Traditional antimicrobial susceptibility tests include the disk diffusion test, E-test gradient diffusion test and broth dilution susceptibility tests. Although the tests are robust, they each rely on the growth of the bacteria to a dense culture while being exposed to the antibiotic panel. Consequently, these methods require 24–48 h to isolate the bacteria from the patient in pure culture and another 24–72 h to complete susceptibility testing (Goff et al., 2012). Due to the long time required to deliver test results compared to the tempo of infection, patients are often treated “empirically” with broader-spectrum regimens under the assumption that the infection could be drug-resistant.

To improve susceptibility testing times there are currently a few rapid platforms gaining acceptance in the clinical laboratory: polymerase chain reaction (PCR), poly-nucleic acid fluorescence *in situ* hybridization (PNA-FISH), and nanosphere hybridization. PCR method detects coding sequences of known resistance genes while PNA-FISH identifies species and/or resistance by hybridizing synthetic oligo-nucleotide fluorescence-labeled probes to species-specific ribosomal RNA and mRNA. Similarly, nanosphere hybridization uses nanoparticle probes to detect DNA, RNA or protein targets. Rapid susceptibility methods can improve patient outcomes and decrease cost of care. In a study...
comparing PCR to traditional methods, time required for identification of methicillin-susceptible \textit{S. aureus} (MSSA) versus methicillin-resistant \textit{S. aureus} (MRSA) was reduced by 1.7 days, cost per patient decreased from $69,737 to $48,350, and outcomes improved with 6.2 fewer days in the hospital and 20% lower mortality (Bauer et al., 2010). Unfortunately, both these current rapid methods depend on \textit{a priori} knowledge of resistance genes.

\textit{S. aureus} is a prevalent Gram-positive human pathogen that is a leading cause of bacteremia, pneumonia, skin/soft tissue infections and endocarditis worldwide (Tong et al., 2015). Asymptomatic colonization of up to 30% of healthy individuals allows for the continual transmission and proliferation of this pathogen (Wertheim et al., 2005). \textit{S. aureus} is responsible for community-acquired (CA-) and hospital-acquired (HA-) infections in both healthy and immune-compromised individuals, and numerous lineages of MRSA including the USA300 clone have spread throughout the US and internationally (Mediavilla et al., 2012). Currently, rPCR is used by many clinical laboratories to detect the \textit{mecA} gene, which confers beta-lactam resistance in \textit{S. aureus} by encoding the low-affinity penicillin-binding protein, PBP2a (Pinho et al., 2001). Since MSSA lack \textit{mecA}, detection of this gene is considered a gold standard for molecular identification of MRSA. However, occurrence of oxacillin-susceptible \textit{mecA}+ MRSA strains (Hososaka et al., 2007) and emergence of a \textit{mecA} variant, \textit{mecC}, which encodes a protein with <63% AA identity to PBP2a (Laurent et al., 2012; García-Álvarez et al., 2011), highlight limitations in tests that detect only the presence of this gene.

In recent years, rapid phenotypic susceptibility assays for \textit{S. aureus} have been proposed, but many of these assays are based solely on cell lysis and/or growth (Price et al., 2014; Kalashnikov et al., 2012; Choi et al., 2011), highlight limitations in tests that detect only the presence of this gene.

2. Materials and Methods

2.1. Strains and Culture Condition

The strains used in this study are listed in Table S3. MLST, spa type, and agr group were provided from their corresponding sources. MLST typing for \textit{cb} strains were done by following published protocols (Ji, 2007). All cultures were grown in Muller-Hinton Broth (MHB) (HiMedia, Lot #140338) supplemented with 2% NaCl, 12 mg L$^{-1}$ Mg$^{2+}$ and 25 mg L$^{-1}$ Ca$^{2+}$ to ensure that the growth conditions and therefore the cytological profiles of the cells remained consistent for all antibiotics. The only exception was daptomycin, since this antibiotic requires 50 mg/L CaCl$_2$ for activity. Antibiotics were prepared using the following concentrations and solvents: 10 mg mL$^{-1}$ cefoxitin (H$_2$O, Fluka Analytical), 10 mg mL$^{-1}$ oxacillin (H$_2$O, Fluka Analytical), 10 mg mL$^{-1}$ cephalixin (H$_2$O, Sigma), 10 mg mL$^{-1}$ cefotaxime (H$_2$O, MP Biomedical LLC), 10 mg mL$^{-1}$ meropenem (H$_2$O, Sigma), 5 mg mL$^{-1}$ daptomycin (DMSO, Cubist).

2.2. Minimal Inhibitory Concentration Determination and Synergy Assays

Minimum inhibitory concentration (MIC) data shown in Table S1 and Figs. 2–3 were determined using microdilution method while synergy assays (Fig. 5b–c) were performed using checkerboard assays (CLSI, 2007, 2012). In both cases, cells were grown in a culture tube to an OD$_{600}$ of 0.35–0.6 and diluted to $5 \times 10^7$ CFU/mL (or $5 \times 10^6$ CFU/mL for daptomycin MIC) in cation-adjusted MHB. Cell solution was diluted 1:10 into a 96-well plate containing different concentrations of each antibiotic. Clinical isolates with an MIC for oxacillin higher than 4 μg/mL were categorized as MRSA, and used as the training set for the initial LDA experiments. MIC and synergy results were obtained after 24 h incubation at 30°C. MIC values reported and synergy curve shown were calculated from the mean of triplicates ($n=3$).

2.3. Growth Curves

For growth curve determination, cells were grown to OD$_{600}$ of 0.2. Cells were then either incubated with 10 μg mL$^{-1}$ oxacillin or with no antibiotic at 37°C. OD$_{600}$ readings were made every 30 min for 3 h. The exponential slope of the growth curve was used to determine doubling rate see in Table S3.

2.4. Fluorescence Microscopy

Exponential-phase cell cultures (OD$_{600}$ ~0.2) were treated with antibiotics (10 μg mL$^{-1}$ oxacillin; 20 μg mL$^{-1}$ cephalexin; 5 μg mL$^{-1}$ daptomycin) and grown at 37°C in a roller. Samples were collected for imaging every hour for 3 h (and at 30 min for daptomycin). Eight micro-liters of cells was used to 2 μL of dye mix consisting of 2.5 μM SYTOX Green, 10 μg mL$^{-1}$ DAPI and 20 μg mL$^{-1}$ of WGA-647 in 1×-Tbase and transferred to an agarose pad (10% MHB, 1% agarose, 1 oxacillin). The same setup was done after 3 h and transferred onto an agarose pad (10% MHB, 1% agarose, 1 oxacillin). The same field was imaged at 5 min intervals for 4 h.

2.5. Blind-test

In the first blind test, which consisted of MRSA and MSSA isolates, the test administrator reassigned thirty of the 71 isolates to BT 1–30. Similarly, in the second blind test which contained DNS and DS isolates, 12 of the 20 isolates were renamed BT 1–12. The tester and administrator were both unaware of what class (MSSA, MRSA$^{18}$, MRSA$^{13}$; DNS, DS) each isolate belonged to.

2.6. Cytological Profiling

The threshold for the deconvolved images was adjusted on FIJI (Image 2.0.0) in order to remove background and obtain shape
edges. All SYTOX and DAPI intensity measurements were performed on nondeconvolved images. Cell measurements were made using CellProfiler 2.0 (Kamentsky et al., 2011). Each parameter is taken as the weighted average of three images. For each strain, three sets of profiles (each represented by three images), were obtained in three experiments on different days (n = 3).

2.7. Statistical Analysis

LDA analysis was completed using built-in MATLAB (R2013) function. LDA analysis was done on the triplicate data set (n = 3) of measured parameters (22 parameters per antibiotic) for each strain while only the average data set was graph on the LDA plots. Categorization of unknown isolates was determined based on a decision boundary of the LDA analysis. The decision boundary was defined as the mean distance between of the centroid of the two groups. Unweighted Pair Group Method with Arithmetic Mean (UPGMA) dendrogram based on the MLST of each strain was made using saureus.mlst.net online tree creation program. The UPGMA relationship is calculated based on pair-wise differences in the allelic profiles of the seven housekeeping genes.

3. Results

3.1. Rapid Antimicrobial Susceptibility Determination by BCP

To determine if BCP can distinguish between MSSA and MRSA, we created cytological profiles for 71 clinical S. aureus isolates (37 MRSA and 34 MSSA strains, with MRSA strains defined as having an MIC for oxacillin > 4 μg/mL). Isolates were cultured in MHB liquid media and treated with one of two beta-lactam antibiotics: oxacillin or cephalexin. After 1 or 2 h, cells were stained with fluorescent dyes and imaged (Figs. 1 and 2a, b). For each bacterial cell, 22 different parameters were measured to create cytological profiles for each strain in the presence or absence of antibiotic treatment. We used linear discriminate analysis (LDA) to generate a linear classifier that could separate the profiles from this training set into two classes susceptible (S) and resistant (R) (Fig. 2c–f, diamond). Linear discriminant analysis is a supervised pattern recognition and machine learning statistical method used to find combinations of features which separates classes of objects. We input all the parameters measured into the LDA, and the algorithm identifies linear combinations of the parameters which maximize distances between groups and minimize distances within groups. The first LDA projection (LDA1) retains the most between-class variance in comparison to the second (LDA2). This analysis showed that better separation between S and R control strains was generated with cephalexin treatment (Fig. 2d, left) than oxacillin treatment (Fig. 2c, left). Aggregation of the profile data from cephalexin and oxacillin treatments into a single 44 parameter combined profile also generated excellent separation between S and R strains (Fig. 2e, left).

These results suggested that BCP can be applied to rapidly (within 2 h) determine S. aureus antibiotic susceptibility. We therefore performed a blinded experiment to test if BCP can accurately distinguish between MSSA and MRSA clinical isolates by plotting the profiles for 30 blinded isolates (filled circles) on the LDA graph using the linear classifier obtained from the LDA training set (Fig. 2c–f, left). For profiles derived from oxacillin or cephalexin treatment alone or in aggregate, we correctly categorized MSSA and MRSA strains with 100% (n = 71) accuracy after 2 h of antibiotic exposure. When the analysis was performed after only 1 h of antibiotic treatment, the separation between the two groups was somewhat reduced, but the accuracy of discrimination remained at 100% (Fig. 2f, left). LDA values strongly correlated with the MICs for cephalexin and oxacillin and correctly predicted susceptibility (Fig. 2c–f, right).

With high rates of vancomycin treatment failure (Levine, 2006; Charles et al., 2004; Stevens, 2006), daptomycin has emerged as an important treatment option for serious MRSA infections including bacteremia (Murray et al., 2013; Moore et al., 2012). Accordingly, we applied our method to daptomycin non-susceptible (DNS) and susceptible (DS) strains. Performing the same techniques described above but with daptomycin treatment, we observed distinct differences in bacterial cytological profiles between DS and DNS strains (n = 20) (Fig. 3a). LDA graphs showed that we could accurately discriminate between DS and DNS strains as early as 30 min post treatment in a double blinded test (n = 12) (Fig. 3b).

3.2. BCP Profiles can be Mined to Extract Additional Information and Detect Patterns

While cephalexin treatment alone was sufficient to distinguish between MSSA and MRSA strains, oxacillin profiles revealed strikingly different responses to antibiotic treatment among MRSA strains. Principal component analysis (PCA) of the combined profile data (aggregated profile of oxacillin and cephalexin treatment) showed two major clusters corresponding to the MSSA and MRSA groups (Fig. 4a), but large variance within principal component 2 (PC2) hinted at the existence of a third group.
of subgroups within the MRSA cohort. Further inspection of MRSA strains at opposing ends of this range revealed two distinctive phenotypes. LDA of the cytological profiles of oxacillin-treated MRSA strains in the top and bottom quartiles of PC2 was used to define the LDA space onto which all MRSA strains were plotted. This LDA parameter space separated MRSA into two groups with distinctive profiles (MRSALL, low lysis; MRSAHL, high lysis; Fig. 4b). When exposed to oxacillin, strains in the MRSAHL (purple) group have an elliptical cell shape and low levels of cell lysis (average 3%, n = 24 strains), whereas MRSAHL strains (green) are circular with a high degree of lysis (average 20%, n = 12 strains). This result was unanticipated, as previous studies suggested that lysis alone is indicative of susceptibility (Matsuda et al., 1995; Best et al., 1974; Memmi et al., 2012; Kalashnikov et al., 2012); however, these clinical strains with high lysis possessed oxacillin MICs >50 μg/mL.

Time-lapse microscopy was performed to follow the progression of cytological profiles over time during treatment with oxacillin (Fig. 4c). The expected response to beta-lactam treatment is that MSSA strains halt growth and lyse while MRSA strains continue to grow with relatively little cell lysis. As expected upon oxacillin treatment, MSSA halted growth, became enlarged and began to lyse. In contrast, MRSAHL strains exhibited an initial burst of lysis but then quickly recovered, proceeding to actively grow and divide. Interestingly, MRSAHL strains continued to undergo cycles of growth, division and lysis for the duration of the time-lapse experiment. Comparison of the growth rates of these three groups (MSSA, MRSALL, MRSAHL) on agarose pads without antibiotics revealed that MRSAHL and MSSA have comparable growth rates (generation time = 27 min) while MRSALL grew slower (generation time = 32 min) (Fig. 4d). With the addition of oxacillin, MSSA lysed, resulting in a negative growth rate while MRSALL grew faster than MRSAHL (Fig. 4e). In addition to a faster growth rate in the presence of oxacillin, MRSAHL recovered after an initial burst of lysis, whereas MRSAHL continued to lyse throughout the assay (Fig. 4f).

These results suggest that MRSA strains are diverse in their phenotypic response to beta-lactam antibiotics. High lysis strains are better adapted for rapid growth in the absence of antibiotics, while low lysis strains are better adapted for growth in the presence of beta-lactams. The MRSAHL fitness advantage over MRSAHL in the presence of antibiotics is reflected in higher MICs of low lysis strains compared to high lysis strains (Table S1). This result parallels observations that repeated selection for high level resistance to beta-lactams comes at a fitness cost to growth in the absence of antibiotic (Ender et al., 2004;
Andersson and Levin, 1999). Although the majority of MRSAll strains have lower growth rates than MRSAHl when grown without antibiotic and higher growth rates than MRSAHl under antibiotic treatment, this trend did not apply to all the strains in our collection (Table S2). Such heterogeneity among MRSA strains explains why growth rate alone is insufficient to distinguish between MRSAll and MRSAHl. In contrast to growth rate measurements, BCP evaluates a multitude of parameters allowing rapid differentiation of MSSA and MRSA strains and the identification of subgroups within MRSA isolates.

Next we performed a genetic analysis of the S. aureus clinical isolates to determine if their cytological profiles correlated with specific genotypes. Specifically, we questioned if the two groups of MRSA identified as displaying distinct responses to beta-lactam antibiotics were genetically different. We classified our clinical isolates of S. aureus using a number of genetic markers including the accessory gene regulator agr locus, protein A, and a collection of seven housekeeping genes (Multi Locus Sequence Type, MLST) (Table S3). From the MLST, we constructed a dendrogram illustrating the relationship between all of the MRSA strains in our study (Fig. 5a). We found that all of the MRSAHl strains are closely related to each other (Fig. 5a, green) and to the community-acquired strain USA300. A majority of the MRSAHl belong to multilocus sequence type (MLST) ST8 (Table S3), a marker for USA300 lineage community-acquired MRSA in the U.S. (Nimmo, 2012). However, genotyping alone could not accurately predict whether strains would be low or high lysis. Three of the MRSAll strains were closely related to the community-acquired strain USA300, while the 9 MRSAll strains were more closely related to the hospital-acquired strain USA100 (Fig. 5a, purple). MRSAll isolates belonged to ST8 as well as ST5, a sequence type associated with hospital acquired (HA) MRSA (Gordon and Lowy, 2008; Takano et al., 2013). The three MRSAll strains with genetic identities close to USA300 have high MICs and most likely represent strains that have evolved through continued antibiotic selection to become similar to low lysis, highly antibiotic-resistant strains.

BCP profiles for USA300 (TCH1516) and USA100 (N315) showed that the strains correctly cluster with MRSA isolates (Fig. 2e), but they do not cluster into the same MRSA subgroups in the LDA analysis as they do on the dendrogram (Fig. 4b). N315, a pre-MRSA, exhibits a heterogeneous response to oxacillin exposure (Aiba et al., 2013; Ito...
et al., 1999). The fact that N315 shows genetic similarity to MRSAHL and phenotypic similarity to MRSALL demonstrates that genetic testing alone is insufficient for determining phenotypic response to antibiotic treatment. Because of the constant evolutionary pressures driving strain evolution, phenotypic assays such as BCP paired with molecular genetic tests can be used as a tool to track the evolution of antibiotic susceptibility patterns.

3.3. BCP Detects Groups that Respond to Different Potential Treatment Options

The existence of a MRSA group with high lysis in the presence of a low level (\(< 1/10\)th MIC) of oxacillin led us to explore the potential of eradicating the remaining population with a second antibiotic. Previous work showed that community-acquired strains of MRSA respond to dual beta-lactam therapy in vitro (Memmi et al., 2008). We thus tested each MRSA clinical isolate in synergy assays to determine if they were susceptible to physiologically relevant combinations of beta-lactams. We found that all of the MRSAHL strains responded synergistically to combinations of oxacillin + cefoxitin (Fig. 5b) or cefotaxime + meropenem (Fig. 5c). The MRSAHL strains (green) did not grow when treated with both antibiotics at concentrations below 1/4 Cmax (maximum IV concentration), modeling physiologically achievable antibiotic concentrations (ADD-Vantage®, 2015; Pfizer, 2009; Novation, 2014). Strains that did not grow at or below this cutoff are candidate responders to dual antibiotic combination therapy. Testing of the remaining strains (MRSALL, purple) revealed either no synergy or synergy at concentrations that were too high to be physiologically achievable. Finally, two MRSALL strains (dotted purple) that bordered the cutoff were not reproducibly inhibited below 1/4 Cmax. Together these results show that BCP can detect two distinct subgroups of MRSA and predict which strains will be susceptible to combinations of beta-lactam antibiotics.

4. Discussion

Our study demonstrates that applications of BCP extend beyond determination of antibiotic mechanism of action to determination of antibiotic susceptibility and identification of optimally effective combinatorial therapies. This single cell phenotype-driven approach provides a rapid susceptibility test that can be performed in parallel with traditional MIC testing, but which delivers a result within 1–2 h, rather
than 1–2 days. The test is precise, yielding no major or minor errors in our blinded tests. Rather than relying on a single parameter (such as growth or permeability), BCP as implemented here assesses 22 different parameters that together capture the wide array of affects antibiotics have on bacterial cells, including changes in cell length, width, permeability, and in chromosome number, compactness and shape (Lamsa et al., 2012; Nonejuie et al., 2013). This likely explains why BCP is more accurate (as well as more rapid) than other proposed phenotype-driven tests (Choi et al., 2014; Ligozzi et al., 2002). Indeed, phenotypic methods that rely only on lysis (Price et al., 2014; Kalashnikov et al., 2012; Choi et al., 2013) would have miscategorized the MRSA HL strains as MSSA, whereas methods that rely only on growth would have failed to identify the two classes of MRSA strains that differ in their susceptibility to combinatorial drug therapies.

Our mechanistic studies have demonstrated that BCP works for all antibiotics, natural products and antimicrobial peptides tested so far, and for all species tested (S. aureus, Escherichia coli, Bacillus subtilis, E. faecium, Enterococcus faecalis, Streptococcus pneumoniae and A. baumannii) (Lamsa et al., 2012; Nonejuie et al., 2013; Sakoulas et al., 2015a, 2015b; Lin et al., 2015; Hindler et al., 2015; Werth et al., 2014). A major strength of BCP is that, in contrast to the rapid gene and genome-based susceptibility tests, it does not rely on prior knowledge of the mechanism of antibiotic resistance or prior identification of all possible genes and mutations that confer resistance. This is critical, even for drugs that bind well-defined targets, such as the ribosome, since the number of mutations that confer drug resistance in the clinic continues to expand (Wilson, 2014). It is even more critical for drugs such as daptomycin, for which several different mutations confer various levels of resistance (Diaz et al., 2014; Mishra et al., 2014; Bayer et al., 2015, 2013; Berti et al., 2015). The reliance of BCP on phenotype rather than genotype makes it highly adaptable, allowing future applications to new drug resistant pathogens and new antibiotics. BCP can be readily optimized to provide a more accurate test for different species or antibiotics. For example, the susceptibility test reported here relies on a different combination of fluorescent stains than used in our previous BCP assays.

BCP provides a robust and rapid technique that is readily adapted to identify opportunities for antibiotic treatment synergy. Such synergies often derive from the additive effects of multiple genetic factors, making it difficult or impossible to detect them using a gene-based technique. This was found to be the case for the beta-lactam plus daptomycin synergy in E. faecium, which depends on multiple PBPs (Sakoulas et al., 2015a) and for daptomycin resistance in S. aureus and E. faecium for which multiple mutations contribute to resistance (Bayer et al., 2013) by modifying either the cell membrane or the cell wall (Humphries et al., 2015).
et al., 2013). Hence we propose that BCP could be used to guide treatment options including the usage of dual beta-lactam treatment or the substitution or addition of different class antibiotics to lower the chances of treatment failure. Identification of synergistic combina
tions in vitro followed by appropriate clinical correlation studies to determine the efficacy in vivo allows for the assessment of new treatment options using existing antibiotics.

The cytological library we have developed provides a versatile reference data set for the analysis of S. aureus strains that can be expanded to include new data from reference strains as well as new strains as they evolve. Indeed, realizing the clinical potential of BCP as an antimicrobial susceptibility test will require that we expand our current reference data set to include additional clinical isolates to generate a more robust training set that includes strains from more diverse locations. Currently we have tested culture samples, in the future we would like to expand our method to test directly from patient samples for a variety of bacte
erial pathogens and antibiotics. Additionally we expect that BCP could work for determining susceptibility in mixed cultures by categorizing the species according to cytological variation. Rapid, mass spectrometry based methods for species identification might also be applied, which would help identify the appropriate species-specific BCP reference library to determine susceptibility.

5. Conclusions

The objective of this study was to apply BCP as a rapid susceptibility test for S. aureus. We demonstrated that BCP could be used to distin
guish MRSA from MSSA and DN from DS S. aureus for a set of clinical isolates on a timescale competitive with existing rapid susceptibility tests. BCP provided information beyond susceptibility testing and iden
tified two subgroups of MRSA that responded differently to oxacillin and to specific combinations of antibiotics. These subgroups were not predicted based on the presence of commonly assessed genetic markers (mecA), highlighting the potential power of phenotypic cell based assays to rapidly identify the best treatment options. Because cytological profiles consist of tens of parameters, one could retrospectively mine them for parameters that correlate with factors such as new sequence types and groups corresponding to treatment failure. For every new strain tested and added to the library, BCP data can be coupled with sus
cceptibility information, sequence and strain type, and clinical data, such as infection type and treatment outcome. Since sequence typing of dif
derent isolates is routinely performed for phylogenetic and population-based analysis, BCP profiles that are obtained during susceptibility de
termination can complement future epidemiological studies by serving as an early indicator of emerging trends that require more public health attention. Combining sequence and BCP data will also allow us to better understand the cellular consequences of the genetic changes that occur as bacterial pathogens adapt to the clinic and acquire resistance to new antimicrobial agents.

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Conflict of Interest Statement

JP and KP have an equity interest in Linnaeus Bioscience, Inc. (La Jolla, CA). The terms of this arrangement have been reviewed and ap
dproved by the University of California, San Diego in accordance with its conflict of interest policies. US Patent application #13/816,471 has been filed related to this topic. Linnaeus Bioscience, Inc. has licensed the rights to US Patent application #13/816,471 from UCSD. GS has re
cieved consulting and speaking honoraria fees from Merck and Actives Pharmaceuticals and research grant funds from Actives Pharmaceuti
cals. VN has received consulting fees from Cidara Therapeutics and Loxbridge Research, and has laboratory sponsored research agreements with Roche, InhibRx and Alterrnute Technologies.

Author Contributions

GS, JP and KP each contributed to the concept design, data interpre
tation and writing of the paper. VN provided general counsel and contributed to the writing of the paper. DQ contributed to the performance of the experiments, analysis of the data and writing of the paper.

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Appendix A. Supplementary Data

Supplementary data to this article can be found online at http://dx.
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