Short communication

Genome sequencing reveals strain dynamics of methicillin-resistant *Staphylococcus aureus* in the same household in the context of clinical disease in a person and a dog

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ABSTRACT

The strain dynamics of methicillin-resistant *Staphylococcus aureus* (MRSA) isolates from people and the household dog were investigated. The isolates were identified in the context of a randomized controlled trial that tested household-wide decolonization of people. Genotypic comparison of MRSA isolates obtained from two household members, the dog, and home surfaces over a three-month period failed to implicate the pet or the home environment in recurrent colonization of the household members. However, it did implicate the pet's bed in exposure of the dog prior to the dog's infection. Whole genome sequencing was performed to differentiate the isolates. This report also describes introduction of diverse strains of MRSA into the household within six weeks of cessation of harmonized decolonization treatment of people and treatment for infection in the dog. These findings suggest that community sources outside the home may be important for recurrent MRSA colonization or infection.

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1. Introduction

Community-associated methicillin-resistant *Staphylococcus aureus* (CA-MRSA), including USA300 strains, plays a major part in the epidemic of MRSA in people in the United States (Klein et al., 2009). Of particular concern is the propensity for cases or their household members to develop recurrent colonization or episodes of MRSA skin or soft tissue infection (SSTI) (Lautenbach et al., 2010; Fritz et al., 2012). Interrupting human-to-human transmission has been tested using household-wide decolonization treatment approaches, but with mixed success (Fritz et al., 2012), suggesting that other sources inside or outside the home may influence recurrent colonization. Within the home, surfaces and companion animals have been identified as potential reservoirs of MRSA, highlighting the importance of a one health approach to investigate drivers of household transmission of MRSA (Davis et al., 2012b).

Previous reports have strongly implicated companion animals in potential maintenance or recurrence of human MRSA colonization or infection (Faires et al., 2009; Bramble et al., 2011; Ferreira et al., 2011), with multiple case reports documenting that treatment of pets was required to clear human MRSA (Manian, 2003; van Duijkeren et al., 2004, 2005; Sing et al., 2008). Hence, as part of a randomized controlled trial that tested household-wide decolonization treatment of people, patients were enrolled with CA-MRSA skin or soft tissue infection (SSTI). In addition to these index cases, their household members and pet(s) also were enrolled. MRSA was isolated from a SSTI in a patient (a dog owner), and also from a subsequent surgical site infection (SSI) in the patient’s dog. MRSA colonization was observed in the people up to three-months later. This report describes the strain dynamics of these isolates using multiple genetic typing methods.

2. Methods

The household, which consisted of the index patient and a white female household member in her 30s, enrolled in the

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Epidemiology and Prevention of MRSA Transmission in the Community trial (NCT00966446) (Cluzet et al., 2015a,b). Four weeks after the index patient’s MRSA SSTI diagnosis, the two people in the household were randomized at the baseline study visit to receive one week of twice-daily nasal mupirocin treatment and two body washes with 4% chlorhexidine gluconate (Hibiclens®, MÖlnlycke Health Care, Norcross, Georgia). Mupirocin and Hibiclens® were received one week prior to development of SSI in the dog. This treatment involved daily reminders via text message during the week of treatment and a daily log of treatment adherence. The household members reported full compliance with the protocol. At the baseline visit and at subsequent biweekly intervals, household members also provided Eswabs® (Cowan Diagnostics, Murrieta, CA) obtained from nares, axilla/groin, and (for the index patient) the healed lesion site to test for S. aureus (MSSA and MRSA) by culture. Household members provided swabs at baseline (visit one), two weeks later (visit two), eight weeks later (visit four), and 12 weeks later (visit seven) (Fig. 1).

The household concurrently enrolled in the Pets and Environmental Transmission of Staphylococci (PETS) study, which involved sampling of the dog and the home environment at visits one and seven. Electrostatic cloths (Swiffer®, Proctor & Gamble) were used as previously described for surface sampling (Davis et al., 2012a). Samples were preserved in sterile, sealed stomacher bags until culture. Sites sampled were (1) the top of the refrigerator, (2) the handle of the refrigerator, (3) the top of the television, (4) the television remote, (5) a kitchen towel, (6) the bathroom faucet handle, (7) the index patient’s pillow, and (8) the dusty surface of the headboard of the bed. Sterile BBL™ culturettes (BD, Franklin Lakes, NJ) were used to collect samples from the dog’s nares, mouth, inguinal skin, and perineum. Electrostatic cloths were used to collect a sample from the “petting zone” on the dorsum of the dog and from the dog’s bed.

Human Eswab™ (Cowan Diagnostics, Murrieta CA) samples were streaked onto CHROMagar MRSA agar plates (BD, Sparks MD) and incubated at 37°C for 24–48 h. Environmental cloths and animal swabs and cloths were cultured for methicillin-susceptible and methicillin-resistant coagulase-positive staphylococci using parallel broth enrichment protocols. Presumptive staphylococcal isolates were identified on Columbia CNA blood agar (BD, Sparks MD) and individual isolates were sub-cultured to Baird Parker agar (BD, Sparks MD) and incubated at 37°C for 48 h as previously described (Davis et al., 2012a). Isolates were frozen at –80°C in Microbank™ tubes (Pro-Lab Diagnostics, Canada) until further testing.

The staphylococcal species was confirmed using a multiplex PCR assay that amplifies species-specific segments of the nuc gene (nuc) (Hirotaki et al., 2011). MRSA isolates were confirmed to carry mecA by presence of a universal mecA/C sequence and absence of the mecC gene using sequential PCR, with ATCC43300 as mecA-positive and LGA251 as mecC-positive controls (Garcia-Alvarez et al., 2011). Isolates were tested for antimicrobial susceptibility to amikacin (AMK), cefoxitin (FOX), chloramphenicol (C), ciprofloxacin (CIP), clindamycin (CLI), erythromycin (ERY), gentamicin, linezolid (LZD), quinupristin-dalfopristin (SYN), tetracycline (TE), and trimethoprim-sulfamethoxazole (SXT) prior to cryopreservation using Kirby-Bauer disc diffusion testing according to CLSI standards (CLSI, 2013). Mupirocin and vancomycin susceptibilities were evaluated via E-test® analysis (Biomérieux, Marcy l’Etoile, France).

A priori, isolates were compared using PCR methods to identify Panton-Valentine leukocidin (PVL) genes, pulses-field gel electrophoresis (PFGE), S. aureus protein A (spa) typing, and staphylococal chromosomal cassette (SCCmec) typing (Shopsin et al., 1999; McDougal et al., 2003; Milheiro et al., 2007; Zhang et al., 2008). Isolates then were subjected to pyrosequencing via 454 chemistry using the Genome Sequencer FLX System (Roche, Branford, CT, USA). Single-nucleotide polymorphisms (SNPs) were compared to assess overall relatedness, using a cut-off of ≤60 SNPs for cluster assignment (Tong et al., 2015). Details of these methods and additional results are provided in the online supplement.

3. Results

The human index patient was a white male in his 30s who presented as an outpatient to the emergency department (ED) after a two-week history of soft tissue swelling of the neck. The patient had a temperature of 37.2°C and reported no co-morbidities present. The attending physician diagnosed a non-draining, indurated abscess on the neck; incised and drained the abscess; administered one dose of intravenous clindamycin in the ED; and prescribed oral, twice daily trimethoprim-sulfamethoxazole and Hibiclens® use for ten days. Culture of pus from the abscess confirmed MRSA. The isolate was discarded by the hospital laboratory before study staff could request it for additional testing.

A 23-month old female spayed 32 kg Italian Mastiff presented for outpatient Tibial Tuberosity advancement surgery to repair a ruptured cranial cruciate ligament on the right stifle two days after the study baseline visit (visit one) and four weeks after the index patient’s ED visit. This surgery involved implantation of a 6-prong plate, 12 mm cage titanium implant, and cancellous bone graft, as well as additional C-laser treatments. Four weeks later, SSI was diagnosed by the veterinary surgeon. A commercial veterinary diagnostic laboratory identified the infecting organism as MRSA (Isolate B). An additional swab was submitted to study staff. A three-week course of oral, twice daily clindamycin was prescribed and the dog’s bed was laundered.

A MRSA isolate (Isolate A) was cultured from the dog’s bed at visit one, prior to the surgery. No other environmental isolates at either visit were found to be S. aureus. No MRSA isolates were cultured from the index patient and household member at baseline, at visit two, or at visit four. No S. aureus isolates were identified from the dog at visit one. At visit seven, MRSA was isolated from both household members (Isolates C-F). At visit seven, a methicillin-susceptible S. aureus (MSSA) was cultured from the dog’s perineum (Isolate G).

MRSA isolates from the dog bed at visit one, prior to the surgery, and from the dog’s surgical site infection at visit three were 100% identical by PFGE and had an identical spa type (t121, a deletion variant of spa type t098—USA300) (Fig. 2). Both isolates were positive for the PVL gene. At visit seven, both household members were positive for MRSA from multiple sites. PFGE indicated that the isolates from the index patient’s axillae/groin, the index patient’s healed lesion and the household member nares were related
(Dice coefficient of similarity = 0.968), had an identical spa type (t008—USA300), and were positive for the PVL gene. In addition, one MRSA isolate was obtained from the nares of the index patient. This isolate was spa type t002 (USA 100) and was negative for the PVL gene. While the dog infection and three of the human colonizing isolates had the same pulsed-field profile, they had minor differences in spa type (t121 versus t008). Whole genome sequencing and SNP analysis revealed that the dog bed and dog SSI MRSA isolates clustered together, but did not cluster with the subsequent human MRSA isolates (Fig. 3).

4. Discussion

This report is notable in that (1) it describes laboratory-confirmed MRSA infections in a person and his dog in temporal and spatial proximity; (2) it implicates an environmental surface (the pet’s bed) in exposure of the dog prior to onset of the dog’s infection, which to our knowledge is the first report of a potential linkage between MRSA contaminated pet bedding and subsequent pet disease: and (3) it fails to implicate the pet in human MRSA outcomes, which also is distinct in the case literature. This study further demonstrates the critical importance of using multiple typing methods to investigate household transmission. In the absence of spa typing data, we would have concluded that the human and animal strains were related; whole genome sequencing was required to confirm that the dog bed and dog SSI MRSA isolate cluster was distinct from related strains found subsequently in people. In the prior literature, less than half (5 of 11 case reports or case series) were documented using more than one molecular typing technique, typically the same combination (PFGE+spa typing) that we initially employed here (Sing et al., 2008; Rutland et al., 2009; Faires et al., 2009; Ferreira et al., 2011). This example highlights the value of using whole genome sequencing methods for strain comparison to assess the dynamics and epidemiology of household MRSA contamination.

While it can be speculated that the MRSA-infected owner contaminated the dog’s bed or another fomite responsible for exposure, and that this led to the dog’s SSI, this route of indirect transmission cannot be confirmed because the clinical isolate from the index patient could not be obtained. Similarly, both dog and owner may have been exposed to a common source outside the home. A third explanation is that the dog was exposed to MRSA through veterinary contact prior to the surgery. However, during 2012 and 2013, the veterinary surgeon performed 300 surgeries, with three staphylococcal surgical site infections (1% SSI rate), including this case. Only one case (this report) was MRSA; the other two dogs were infected with methicillin-resistant S. pseudintermedius, an animal-associated pathogen. While veterinarians and their staff may become MRSA colonized, which could be a risk factor for transmission to their animal patients, Hanselman et al. found MRSA positivity rates of 4.4% among small-animal veterinarians attending a U.S. conference (Hanselman et al., 2006).

The finding of diverse MRSA strains in this household within six weeks of cessation of harmonized decolonization treatment in both people and treatment for infection in the dog is of great concern as it demonstrates rapid re-introduction of MRSA to the household. The scenario described in this report represents one explanation for observed failure of household-wide decolonization protocols to eradicate S. aureus carriage from index cases (Fritz et al., 2012; Cluzet et al., 2015a) and is unique in that it neither implicates the home environment nor the pet dog in re-colonization of the index patient following successful treatment. While future case assessments and research studies should include multiple evaluations of home environmental contamination and companion animal carriage over time to capture the potentially dynamic nature of intra-household transmission, attention to community sources outside the home may be necessary to understand drivers of recurrent colonization or infection.

Data access

Genome data from this study have been deposited in the Short Reads Archive under accession numbers A: SAMN03755562, B: SAMN03755563, C: SAMN03755566, D: SAMN03755565; and E: SAMN03755564.
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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.jvetmic.2015.09.007.

References


