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1231. Patient-Level Factors Associated with Vancomycin-Resistant Enterococci Transmission to Healthcare Workers Gowns or Gloves

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Session: 137. Healthcare Epidemiology: MSSA, MRSA and Other Gram Positive Infections

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Background. Vancomycin-resistant *Enterococcus* (VRE) is transmitted from person-to-person, most commonly by healthcare workers (HCW) whose hands or attire have become contaminated while interacting with an infected or colonized patient. Our group recently found that VRE colonized patients transmitted this pathogen to HCW gowns or gloves 15% of the time. This study aims to describe patient-level factors associated with higher risk of transmission of VRE to HCW gowns or gloves and thus likely to subsequent patients.

Methods. We analyzed a prospective cohort that included 43 VRE-colonized patients and 215 HCW-patient interactions in medical or surgical intensive care units at the University of Maryland Medical Center. HCWs' gowns and gloves were cultured for VRE after performing patient care and before doffing. Univariate and multivariable logistic regression models, using generalized estimating equations to account for patient clustering, were used to estimate the odds ratios associated with specific patient-level factors (i.e., age, race, Elixhauser comorbidity score components obtained by ICD-10 codes, diarrhea, and devices). Multivariable models with and without stool VRE burden were created.

Results. In the initial multivariable model, having a nasogastric tube, diarrhea, complicated diabetes, rheumatoid arthritis/collagen vascular diseases, neurological disorders or psychoses doubled (OR greater than 2) the patient's risk of VRE transmission. After adjusting for VRE stool burden (OR 2.1 (95% CI 1.5–3.0)), having a nasogastric tube (OR 3.6 (95% CI 1.3–9.8)), diarrhea (OR 3.3 (95% CI 1.4–8.1)), or rheumatoid arthritis/collagen vascular diseases (OR 4.8 (95% CI 1.6–14.7)) remained significant in the model.

Conclusion. Patient-level factors associated with higher risk of VRE transmission to HCW gowns or gloves were identified even after adjusting for VRE stool burden, highlighting the importance of patient characteristics in VRE transmission. These patient-level factors may facilitate transmission by either increasing VRE stool shedding to the environment or the need for direct HCW-patient contact. These factors could be used to target more aggressive infection control interventions for these patients.

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1232. Phylogenomics of *Enterococcus faecium* From South America: Revisiting Worldwide VRE Population Structure

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Background. Previous studies have suggested that the population structure of *E. faecium* is composed of two main clades; a commensal clade (designated clade B) and a hospital-associated clade (Clade A) that encompass most of the clinical and animal isolates. The phylogenetic analyses leading to these results have been accomplished with the notable absence of isolates from diverse geographical regions (including South America). We aimed to refine the worldwide population structure of *E. faecium* by including 55 representative genomes from isolates obtained from five Latin American countries recovered between 1998 and 2014.

Methods. We sequenced our 55 representative isolates and selected other 285 genomes, from public databases, obtained across different regions (36 countries), different sources (animal, commensal, and clinical strains) and a wide range of dates of isolation (1946–2017). We characterized the genomes by presence/absence of resistance, virulence and mobile elements, and of CRISPR-*cas* systems. We analyzed the phylogeny of the entire population, selected the genomes belonging to clade A to examine recombination patterns and performed Bayesian molecular clock analysis excluding recombinant regions.

Results. Two major clades were identified, as previously reported. However, a higher degree of variation in clade A was found. Indeed, we identified a subclade (subclade I) that diverged ~894 years ago, and clearly distinguished clinical isolates from those of animal origin (distributed among a number of smaller early-branching subclades). A further split within the clinical subclade (subclade II) that diverged around ~371 years ago was also evident. Latin American isolates were distributed within subclades I (48%) and II (42%). Isolates in "animal" branches exhibited an average recombination of 34 Kbp, where it was 5 Kbp and 21 Kbp for subclades I and II, respectively. More resistance determinants were found in subclade III (62%), followed by I (54%) and absence of *cas* was the norm in the clinical subclades.

Conclusion. Inclusion of *E. faecium* isolates from diverse geographical region supports a continuous evolution of these organisms causing human infections. Important evolutionary events seem to favor emergence of novel subclades capable to cause important morbidity and mortality.

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1233. An Automated E-mail Notification Systemic to Infectious Disease Specialists and Effect on the Management of *Staphylococcal aureus* Bacteremia in a Community Hospital setting

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Background. Staphylococcus aureus is the leading cause of community and healthcare-associated bacteremia and carries a high burden with a substantial mortality, ranging from 20 to 40 %. Evidence suggests infectious disease (ID) consultation improves mortality and adherence to the Infectious Diseases Society of America (IDSA) guidelines. Due to complications from a lack of ID consultation, a notification system consisting of automated e-mails to ID providers was implemented. The objective of this study was to review the impact of the automatic notification to ID consultants with positive blood culture results in a community hospital system.

Methods. Cases of staphylococcus aureus bacteremia were identified from the microbiology database by at least one positive blood culture. The automated e-mail notification system was implemented in December 2014. ID providers were encouraged to verbally contact primary providers for positive results. Cases of bacteremia prior to implementation of the automated notification system were compared with those post-intervention. Patients under age 18 were excluded. Data gathered included mortality, re-admission rates, and compliance with IDSA guidelines.

Results. There were no significant differences in inpatient mortality (9 vs. 18%, P = 0.180). 30-day mortality between the two groups (18 vs. 20%, P = 0.815). The 30-day readmission rate among surviving patients was reduced by 50% (40% vs. 19%, P = 0.014). Compliance with antibiotic duration in complicated bacteremia increased post-intervention (57% vs. 85%, P = 0.04).

Conclusion. An automatic notification to ID specialists reporting patients with *Staphylococcus aureus* bacteremia led to improved compliance with IDSA guidelines regarding antibiotic duration and reduced re-admission rates. There was no effect on overall mortality.

Table 1: Patient Demographics

	Pre Intervention $(N = 57)$	Post Intervention $(N = 60)$	<i>P</i> -value
Average patient age (years)	64.4	62.2	0.448
Male	63%	63%	1
Immunosuppressed	16%	13%	0.80
Complicated bacteremia	70%	69%	1

Table 2: Patient Outcomes

	Preintervention $(N = 57)$	Postintervention $(N = 60)$	<i>P</i> -value
Inpatient mortality	9%	18%	0.180
30-day mortality (%)	18%	20%	0.815
Readmitted within 30 days	40%	19%	0.014
Bedside ID consult	75%	78%	0.888
Appropriate antibiotic duration -complicated bacteremia (>28 days)	57%	85%	0.04

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