Infection Control Hospital Epidemiology







Infection Control & Hospital Epidemiology

Volume 46 2025 Number 7

CONTENTS

SHEA White Paper

665 SHEA/APIC/IDSA/PIDS multisociety position paper: Raising the bar: necessary resources and structure for effective healthcare facility infection prevention and control programs

Thomas R. Talbot, Christopher Baliga, Rebecca Crapanzano-Sigafoos, Tania N. Bubb, Mohamad Fakih, Thomas G. Fraser, Ibukunoluwa C. Kalu, Vidya Mony, Anupama Neelakanta, Ann-Christine Nyquist, Catherine O'Neal, Jan E. Patterson, David K. Warren and Sharon B. Wright

Commentary

Infection prevention and control for measles in healthcare settings

Irene Frantzis, Karen P. Acker, EO Benjamin, Rookmin Pato, Stephanie Legall, Harjot K. Singh and Lisa Saiman

Original Articles

- Risk factors for catheter-related bloodstream infections in a high-risk cancer patient population Andrea Haddad, Rita Wilson Dib, Anne-Marie Chaftari, Ying Jiang, Mohamed Moussa, Hiba Dagher, Ann Philip, Ray Hachem and Issam Raad
- Epidemiology and perioperative factors contributing to postoperative pneumonia in patients undergoing isolated coronary artery bypass grafting: a retrospective Chinese study *Yuxiao Zhan, Yang Yang, Jian Zhang, Rui Yang and Guojun Zhang*
- 703 Usage of oral vancomycin for acute Clostridioides difficile infection (CDI) resulting in later acquisitions of vancomycin-resistant enterococci (VRE)
 Evgeny Rogozin, Husam Maree, Majdi Masarwi, Rozan Hasona, Herschel T. Horowitz, Ruth Bouganim and Dror Marchaim
- 710 A discrete choice experiment to evaluate healthcare personnel preferences regarding risk-tailored policies for contact precautions for patients with methicillin-resistant *Staphylococcus aureus*Lyndsay M. O'Hara, David P. Calfee, Graham M. Snyder, Elise M. Martin, Nicholas F. Angelino,
 Nathan N. O'Hara and Anthony D. Harris
- 716 Testing for methicillin-resistant *Staphylococcus aureus* in the anterior nares for antibiotic de-escalation in patients presenting with acute skin and soft tissue infections: systematic review and meta-analysis *Sara Bohjanen, Connor Goldstick and Maria Hordinsky*
- 721 Characterizing presenteeism among healthcare personnel at an academic medical center across eras of the COVID-19 pandemic
 Amanda Brown Marusiak, Emily Sickbert-Bennett, Hilary Babcock, Daniel Westreich, Justin Lessler and David Weber
- 730 COVID-19 prevention training with video-based feedback in nursing homes: impact on staff safety behaviors

 Victoria Ngai, Joshua B. Hsi, Raveena D. Singh, John E. Mitchell, Raheeb Saavedra, Shruti K. Gohil, Emily A. Hsi, Robert Pedroza, Chase Berman, Kristine P Nguyen, Matthew Zahn, Emily Fonda,

Cover image: The Dynamics of Bacterial Evolution, 2020

Susan S. Huang and Gabrielle M. Gussin

- 738 Understanding hand hygiene adherence in neonatology: a qualitative study of behavioral determinants Tamara C. Bopp, Yvonne Strässle, Colette Wyler, Marie-Theres Meier, Lauren Clack, Walter Zingg, Jehudith R. Fontijn and Aline Wolfensberger
- 747 Reducing inappropriate antibiotic use in febrile neutropenia in hematology patients through the implementation of an antibiotic de-escalation protocol

 Jinghao Nicholas Ngiam, Victor Ling, Matthew Chung Yi Koh, Mohamed Nasar Fathima Rofina Farveen,
 Shi Hui Clarice Choong, Li Mei Michelle Poon, Liang Piu Koh, Nares Smitasin and Lionel Hon-Wai Lum
- 753 Sustainability and spread of best practices for peri-operative antimicrobial stewardship following discontinuation of mandatory public reporting: perceptions from the Society for Healthcare Epidemiology in America research network

 Jacquelyn Pendergast, Hillary Mull, Marlena Shin, Ryann Engle, A. Rani Elwy, Judith Strymish,

 Samuel Golenbock, Kierstin Hederstedt, Rory Ostrow, Matthew Bidwell Goetz, Mary Hawn and Westyn Branch-Elliman

Review

759 Outbreaks in hematopoietic stem cell transplant units: a systematic review Loukas Kakoullis, Georges Chedid, Bradley Walker, Vasiliki Xirou, Sufian Abdel Hafez, Sokratis N. Zisis, Shreya Arora, Costas A. Christophi, George Panos, Courtney E. Harris and Robert Colgrove

Letter to the Editor

770 Two-dimensional chart for antimicrobial spectrums and costs: a tool for antimicrobial stewardship *Yohei Manabe and Hideharu Hagiya*

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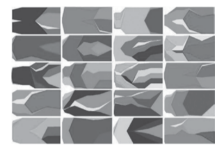
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About the cover:

Beginning with volume 43 (January 2022), the cover of *Infection Control & Hospital Epidemiology* (ICHE) will feature art inspired by or reflective of topics within the scope of the journal and their impact on patients, healthcare personnel and our society. These topics include healthcare-associated infections, antimicrobial resistance, and healthcare epidemiology. The intent is to feature original artwork that has been created by individuals who have a personal connection to one or more of these topics through their clinical work, research, or experience as a patient or an affected patient's family member, friend or advocate. The goal is to provide readers with a visual reminder of the human impact of the topics addressed in the journal and the importance of the work being done by those who read or contribute to ICHE and by all who are trying to make healthcare safer through the elimination of healthcare-associated infections.

For more information about the ICHE cover and how to submit artwork for consideration for a future cover, please visit the ICHE website: https://www.cambridge.org/core/journals/infection-control-and-hospital-epidemiology/front-covers

2025



Title: The Dynamics of Bacterial Evolution, 2020

Artist: Angharad Ellen Green, PhD

Medium: The artwork is made up of individual Muller plots representing *Streptococcus pneumoniae* bacteria lineages that were evolved separately within nasopharynx and lung environments. The command line program muller (v0.6.0 - https://pypi.org/project/muller/), with default parameters applied, was used to produce genotypes and trajectories tables for each of the evolved lineages. These tables were then used as inputs for ggplot2 (v3.3.2) and ggmuller (v0.5.4) in R-Studio (v4.0.2), to produce Muller plots. The individual plots were then assembled to produce the resulting artwork.

Dr. Green spoke to ICHE about her artwork.

What was the inspiration for this artwork? My postdoctoral research used an *in vivo* experimental evolution model to understand how *Streptococcus pneumoniae* (the pneumococcus) adapts to the lung and nasopharynx environments. The pneumococcus was experimentally evolved through a lung infection model and a nasopharynx infection model, producing independently evolved lung and nasopharynx lineages. We sequenced the evolved lineages and compared them to the ancestor to understand how their genomes had changed. This work also enabled us to determine how environmental differences between the upper and lower airways might shape pneumococcal adaptation and evolution. The resulting sequencing dataset was very large and complex with lots of interesting results. I wanted to use an effective method of visualising the data and Muller plots were chosen to display the evolutionary dynamics of mutations found in each evolved lineage over time. In these plots, each mutation is grouped as a genotype, which is represented by a different colour, and the blocks of colour expand when the genetic changes make the bacteria better able to survive in their local conditions. After completing the data analysis and publishing this work, I created this artwork as a memento of my postdoctoral research and I have a canvas of this work hanging in my apartment. Additionally, I wanted to demonstrate how scientific artwork can help visualise the complexities of evolution dynamics and help us to better understand bacterial processes.

What is your personnel connection to the content of ICHE? Throughout my career as a microbiologist, I have carried out research to investigate bacterial pathogenesis and antimicrobial resistance (AMR) of WHO-defined bacterial priority pathogens, such as *Pseudomonas aeruginosa*, methicillin-resistant *Staphylococcus aureus* (MRSA) and *Streptococcus pneumoniae*. I have actively promoted the importance of microbial genomic research to confront current global challenges, such as AMR and healthcare-acquired infections. I have championed microbiology research through my various roles in academia, volunteering on the Microbiology Society's Policy Committee and as a Research Manager at the Healthcare Infection Society. It is an honour for my bacterial evolution artwork to be on the cover of ICHE.

Given the scope of the journal, why is this work appropriate for the cover of *Infection Control & Hospital Epidemiology*? This artwork is made up of a collection of graphs called Muller plots, which are used to visualize how bacteria evolve when grown in diverse environments. The colours represent genetic changes that have taken place in the presence of environmental factors, such

Cover image: The Dynamics of Bacterial Evolution, 2020

as antimicrobials and the host immune system. The dynamics of evolution are complex and being able to visualise this process enables scientists to better understand bacterial processes, including the development of AMR. This artwork is appropriate for the cover of ICHE as it was created as a direct result of scientific research into how bacteria can adapt and evolve in diverse host niches to cause disease. Additionally, this artwork makes it possible for scientists to visualise the complexities of the dynamics of evolution and comprehend how bacteria adapt to different host environments.

Dr. Green is a Senior Research Data Steward in the Advanced Research Computing Centre (ARC) at UCL in London. Her postdoctoral research at the University of Liverpool was supported by a Sir Henry Dale Fellowship, awarded by the Wellcome Trust and the Royal Society (grant number 204457/Z/16/Z) to Dr. Daniel R Neill. The research from which this artwork was derived was published in Molecular Biology and Evolution (Green AE, Howarth D, Chaguza C, et al. Pneumococcal colonization and virulence factors identified via experimental evolution in infection models. Mol Biol Evol 2023; 38: 2209-2226).