ORAL PRESENTATIONS

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CLINICAL VIGNETTE POSTER PRESENTATIONS

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AUTHOR INDEX
A01
SUSCEPTIBILITY RATES AND SPECIES IDENTIFICATION OF CAMPYLOBACTER STRAINS ISOLATED FROM HUMAN ENTERIC INFECTIONS IN A NEW BRUNSWICK REGIONAL HOSPITAL

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OBJECTIVES: Campylobacter (C) is a common human enteropathogen. Published data on susceptibility rates from human Canadian Campylobacter isolates is scarce. We analyzed the characteristics of a 38 month cohort of Campylobacter strains isolated from stool specimens in a New Brunswick regional laboratory. We also present the susceptibility data of a subset of C jejuni isolates to ciprofloxacin (cipro) and erythromycin (erym). METHODS: We retrieved data from stool cultures collected between August of 2011 and October of 2014 in which Campylobacter was isolated. We used Campylobacter CNA agar as the selective medium. Identification was performed using published methods. Isolates not identified as C jejuni or C coli were identified by sequencing of the cpe60 gene at a reference laboratory. Susceptibility of C jejuni to cip and ery was determined by Kirby-Bauer (KB) since May of 2013, using criteria described by Gaudreau. Travel history was obtained from New Brunswick public health.

RESULTS: Campylobacter was isolated in 2.1% of samples. 82 separate episodes of enteritis were identified. Campylobacter was grown from all samples collected in 85% of enteritis cases. We isolated: C jejuni (72%), C lari (10%), C coli (7%), C upadiensis (5%), 2% of other species and 4% of isolates not further identified. 29 isolates of C jejuni were tested by KB. Resistance to nalidixic acid (na) was seen in 6/7 of cip resistant but in no cip susceptible strains. 76% (22/29) of C jejuni isolates were susceptible to cip and ery (colony colour other than brown or mauve) was detected in 353 (22.7%) specimens. 166 (10.7%) specimens had mauve/brown colonies that needed further workup. Of those, 47.6% identified as an organism other than E. coli (32% M. morganii, 22.7% E. hermanii, 13.3% Proteus sp., 8% Enterobacter sp.). Total of 87 (5.5%) specimens had brown/mauve colonies identified as E. coli from the STEC CM and were subsequently tested using the Shiga toxin EIA. 10 specimens (0.64%) tested positive for Shiga toxin by EIA and 100% of them were confirmed to be positive by PCR. 49 E. coli isolated from the STEC plates which were found to be negative by Shiga toxin EIA were sent to SDCCL for Shiga-toxin PCR testing and 100% were confirmed to be negative.

CONCLUSION: Two step algorithm using the STEC CM and Shiga-toxin EIA in an area of low prevalence is an effective method for the detection of O157 and common non-O157 STEC.

A02
IDENTIFICATION OF SHIGA-TOXIN PRODUCING ESCHERICHIA COLI (STEC) FROM HUMAN STOOL SPECIMENS USING A TWO-STEP ALGORITHM: CHROMOGENIC MEDIA (CM) AND ENZYME IMMUNOASSAY (EIA)

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BACKGROUND: Shiga toxin-producing Escherichia coli (STEC) are responsible for haemorrhagic colitis and potentially more serious complications including, hemolytic uraemic syndrome (HUS) and death. The most common serotype of STEC is O157:H7, however as many as 30-50% of STEC infections are caused by non-O157 E. coli. We evaluated a two-step protocol using the STEC CM and Shiga-toxin EIA to screen for O157 and non-O157 STEC.

METHODS: Clinical stool specimens received at the Regina General Hospital medical microbiology laboratory between June 1st and October 31st, 2014 were prospectively evaluated for presence of STEC by being plated onto routine media as per established protocol (BAP, MAC, XLD, CAMPY and E. coli O157 CM) as well as the new STEC CM. Mauve or Brown colonies on the STEC CM identified as E. coli using MALDI-TOF (Bio Merieux, Vitek MS) were then tested for Shiga toxin production using the EIA. All STEC isolates were sent to the Saskatchewan Disease Control Reference Laboratory (SIXCL) for confirmation of Shiga-toxin genes using established protocols. Samples positive by PCR were sent to the National Microbiology Laboratory (NML) in Winnipeg for serotyping.

RESULTS: A total of 1589 stool specimens were screened for STEC. Thirty-four samples were incompletely worked up and therefore were excluded from the subsequent analysis. 1026 (66.0%) of specimens showed no growth on STEC CM. Clinically insignificant growth (colony colour other than brown or mauve) was detected in 353 (22.7%) specimens. 166 (10.7%) specimens had mauve/brown colonies that needed further workup. Of those, 47.6% identified as an organism other than E. coli (32% M. morganii, 22.7% E. hermanii, 13.3% Proteus sp., 8% Enterobacter sp.). Total of 87 (5.5%) specimens had brown/mauve colonies identified as E. coli from the STEC CM and were subsequently tested using the Shiga toxin EIA. 10 specimens (0.64%) tested positive for Shiga toxin by EIA and 100% of them were confirmed to be positive by PCR. 49 E. coli isolated from the STEC plates which were found to be negative by Shiga toxin EIA were sent to SDCCL for Shiga-toxin PCR testing and 100% were confirmed to be negative.

CONCLUSION: Two step algorithm using the STEC CM and Shiga-toxin EIA in an area of low prevalence is an effective method for the detection of O157 and common non-O157 STEC.
A04 PERFORMANCE AND EFFICIENCY OF TWO ALGORITHMS FOR CLOSTRIDIUM DIFFICILE ASSOCIATED DIARRHEA (CDAD) DIAGNOSIS FOR A PROVINCIAL MICROBIOLOGY NETWORK

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BACKGROUND: Algorithms containing both EIA and PCR offer reduced cost and more point of care application as compared to PCR on all stool specimens. EIA for both GDH and Toxin is applied to all specimens and PCR only used for discrepant results. We compared two algorithms to PCR alone.

METHODS: Algorithm 1 combined GDH/Toxin EIA (QuickChek, Alere, $14, 3 minutes hands-on time) and PCR (Illumigene, Meridian Biosciences, $24, 7 minutes). Algorithm 2 combined GDH/Toxin EIA and PCR (AmpliVue, Quidel, $20, 5 minutes). Reference method was PCR alone (Illumigene). 200 consecutive liquid stool specimens submitted to a regional microbiology lab between October and December 2014 were included.

RESULTS: Mean age was 61.9 years (SD 21.2), 120 (60.0%) were female and 114 (57.2%) were inpatients (mean length of stay 15.7 days (SD 48.9)). In both algorithms, only 33/200 (16.5%) specimens required PCR. Algorithm 1 had sensitivity of 100% (95% CI 100-100%), specificity of 99% (98-100%), total cost of $4007.50. Algorithm 2 had sensitivity of 94% (92-98%), specificity of 98% (96-100%), and total cost of $3842.50. Reference standard cost $5500.00.

CONCLUSIONS: Algorithm diagnosis reduced cost without sacrificing performance. Algorithm 1 offered better performance (although reference method was included in algorithm). Algorithm diagnosis could be applied in rural laboratory locations with referral of discrepant to regional centers.

14:45–15:45 Oral Presentations: Session B Room: Johnson, Delta Prince Edward

B01 EVALUATION OF MALDI-TOF FOR IDENTIFICATION OF CAMPYLOBACTER, ARCOCBACTER, HELICOBACTER, AND RELATED ORGANISMS

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OBJECTIVE: MALDI-TOF MS was evaluated for its effectiveness as an identification tool for Campylobacter and related organisms compared to conventional biochemical and molecular methods.

METHODS: For validation, 399 clinical and reference isolates, representing 39 species, belonging to the genera Campylobacter, Arcobacter, Helicobacter, Wolinella, and Bacteroides were subjected to conventional short or full biochemical plus molecular testing (PCR-RFLP, species-specific PCR, 16S sequencing), as well as MALDI-TOF analysis in parallel. In a second implementation stage MALDI-TOF analysis was used as the initial identification screen followed by selective confirmatory conventional testing.

RESULTS: MALDI-TOF results were in agreement with the conventional results at the species level: 100% (322/322) for Campylobacter sp., 100% (22/22) for Arcobacter sp., 84% (46/55) for Helicobacter sp. 100% (1/1) for Wolinella sp., and 100% (1/1) for Bacteroides sp. However, even for isolates that were not in perfect agreement at the species level, MALDI-TOF correctly identified the genus for every isolate in this study with the exception of 1 H. pullorum and 3 H. pylori isolates which generated no reliable information. The conventional biochemical testing allowed for the determination of Campylobacter biotypes, as well as biochemical variants such as hirappricate-negative C. jejuni, urea-positive C. lari, and aberrant antimicrobial susceptibility profiles. While MALDI-TOF was not able to discriminate among some biotypes or biochemical variants, it was able to detect a C. fetus isolate belonging to a distinct lineage of a recently reported third subspecies. Conventional identification was completed in 5 to 21 consecutive days (short and full biochemicals, respectively) whereas MALDI-TOF analysis was completed in 2 to 3 days.

CONCLUSIONS: The identification of Campylobacter and related organisms by MALDI-TOF was an efficient and effective means of identification offering a substantial reduction in time and resources. Although it was unable to determine biotypes, biochemical variants, or molecular anomalies, due to its speed and accuracy, identification by MALDI-TOF analysis is recommended for use in a clinical or reference laboratory to either supplement or substitute conventional species-level identification methods for Campylobacter, Arcobacter, Helicobacter, Wolinella, or Bacteroides.

B02 COMPARATIVE EVALUATION OF BBL CHROMAGAR ORIENTATION AND CONVENTIONAL CULTURE APPROACH FOR IDENTIFICATION AND SUSCEPTIBILITY OF UROPATHOGENS

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OBJECTIVES: Chromogenic agar offers the potential to identify uropathogens while reducing laboratory workload by decreasing subcultures before being tested on automated identification and susceptibility systems. The purpose of this study is to compare the performance of direct inoculation of colonies from BBL Chromagar Orientation to conventional urine culture methods.

METHODS: One hundred eight urine samples were inoculated simultaneously onto blood agar (BA), MacConkey agar (MAC), and Chromagar and incubated for 18-24 hours aerobically at 35C. Colonies from these media were tested by Microscan (Siemens) for identification and susceptibility. Results from conventional media versus chromogenic agar were compared by calculating the essential and categorical agreement for each antibiotic and the percentage of concordant identification. Prior to validation of this test, cephalothin results were used to predict oral cephalosporin susceptibility for uncomplicated urinary tract infections due to certain Enterobacteriaceae.

RESULTS: Identification results from Microscan were identical for organisms isolated from BA/MAC and Chromagar. Categorical agreement for all antimicrobials tested was greater than 90% except for cephalothin (14% minor errors) and piperacillin (2.6% very major error). Essential agreement was greater than 90% except for cephalothin (85%) and piperacillin (88.3%).

CONCLUSION: Direct inoculation of uropathogens from Chromagar using Microscan is a reliable method of identifying uropathogens. Direct inoculation using Microscan for susceptibility testing is a reliable method except for cephalothin and piperacillin. Cefazolin can replace cephalothin as a predictor for oral cephalosporins for uncomplicated UTI.

B03 SHORT-INCUBATION-MALDI-TOF PATHOGEN IDENTIFICATION REDUCES TIME TO APPROPRIATE ANTIBIOTIC THERAPY

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BACKGROUND: Sepsis accounts for significant morbidity and mortality in hospitalized patients. Approximately 30,002 Canadians develop sepsis each year with an associated mortality rate of 30%. Early institution of appropriate antibiotics is associated with improved patient outcomes.

OBJECTIVES: The time to initiation of appropriate antibiotic therapy in patients with blood stream infections was compared during corresponding six week periods in 2012, 2013 to assess the impact of MALDI-TOF identification and a short incubation MALDI-TOF identification method implemented in 2014.

METHODS: As part of an on-going quality assurance program we reviewed and compared the impact of the implementation of a short incubation MALDI-TOF identification method to identify pathogens from positive blood cultures. Charts of patients with positive blood cultures were reviewed for a 6 week period in 2012 (standard incubation and ID using Vitek), 2013 (MALDI-TOF identification after standard incubation) and a
comparable 6 week period in 2014 (short term incubation and ID using Maldi TOF). The time to initiation of appropriate antibiotic, as directed by pathogen identification was compared.

RESULTS: We identified 13/39 patients in 2012, 8/37 patients in 2013 and 10/27 patients in 2014 where empirical antibiotics needed to be changed. The average time to final identification of the pathogen was respectively 86, 74.7 and 68.6 for the respective periods. In 2014, we adapted a process whereby the interim identification obtained from the SMI was reported, this resulted in a reduction in time to initiation of more appropriate antibiotics from 73.7 hours (2012) and 62 hours (2013) to 53.75 hours in 2014.

CONCLUSIONS: We demonstrated that a rapid detection method of pathogens from positive blood cultures reduces time to initiation of appropriate antibiotics in a patient population with blood stream infections. Further studies are required to see whether this impacts patient outcomes.

B04 CHARACTERIZATION OF CANADIAN ISOLATES OF A RECENTLY-DESCRIBED GENUS AND SPECIES NOVUM, EISENBERGIELLA TAYI, OF THE FAMILY LACHNOSPIRACEAE

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INTRODUCTION: Blood culture isolates of an unidentifiable Gram negative bacillus from patients in BC and MB, referred to the NML since 2005, had been found to cocluster as a single taxon group. In 2014, Amir et al [JJSEM 64:907-914] described Eisenbergiella tayi gen. nov. sp. nov., which appeared to be consistent with these NML referrals. Here we describe features of Canadian isolates of this emerging pathogen.

METHODS: Standard methods were used for 16S rRNA gene sequencing, phylogenetic analyses, phenotypic testing (conventional anaerobic biochemical testing, API 32A and BIOLOG panels) and chemotaxonomic studies (CFA composition) were done. Antimicrobial susceptibility testing (AST) was done using broth microdilution (BMD) with CLSI breakpoints for anaerobes being applied.

RESULTS: the NML analysed 8 strains from 7 patients, all from blood cultures. All were described as either Gram-negative or variable curvy bacilli, in contrast to Amir (2014) where these bacteria were described as Gram-negative-staining but Gram-positive structurally. All were strict anaerobes. Interestingly, some grew in the presence of bile in broth or on BBE plates, making blackened colonies (like Bifidobacteria spp). All were reactive to a variety of PY sugars / substrates; reactions for esculin, gelatin and catalase were variable. All were negative for nitrate, indole and urea. All had similar CFAs, the majority being of the saturated and mono unsaturated types. AST by the BMD method suggested that strains were susceptible to most antibiotics, including metronidazole. Phylogenetically, NML strains best fit with E. tayi in the family Lachnospiraceae, part of Clostridiales Cluster XIVa.

DISCUSSION: Characterization of unusual pathogens such as these should include use of a molecular identification approach, in order to analyze novel or newly-described taxa. Pathogenicity remains unknown, but clinicians should be aware of the possibility of Eisenbergiella tayi as a cause of bacteremia.
The majority of Foreign-born cases belong to cluster 12 (24%). The remaining 9 clusters each account for 1.2-3.6% of cases, with fairly consistent yearly rates. Geographic analysis of clusters among regional health authorities revealed more localized cluster variations.

CONCLUSIONS: Tuberculosis in MB is consistently dominated by three MIRU-VNTR types (clusters 1, 3 and 12). Further investigations of strain and populations genetics are required to guide public health efforts to control TB in Manitoba.

C03 EVALUATION OF A LATERAL FLOW IMMUNOASSAY FOR CULTURE CONFIRMATION OF MYCOBACTERIUM TUBERCULOSIS COMPLEX
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BACKGROUND: Potential isolates of M. tuberculosis complex (MTBC) must be identified rapidly for TB patient care and control. The traditional method for identification of MTBC isolates is a non-amplified DNA probe test which requires a significant amount of growth and sometimes fails to detect cultures at early stages of growth. SD Bioline (South Korea) has developed a rapid immunochromatographic test (ICT) kit that claims to have excellent sensitivity and specificity, and is not expensive. This kit has been demonstrated to have a detection limit of $10^5$ CFU/mL and can be used as soon as tiny colonies on solid media are observed or when the auto-detection of positivity is detected in a liquid culture system. We tested this ICT against a standard DNA probe test (Gen-Probe AccuProbe MTBC).

METHODS: Positive cultures from Mycobacterium Growth Indicator Tubes (MGITs, BD Biosciences) or Lowenstein-Jensen pyruvate agar or Myco/F Lytic blood culture vials (BD Biosciences) were used. After smearing each culture, an aliquot was tested with the ICT and the DNA probe test.

RESULTS: Out of 86 isolated tested, 52 were MTBC and the rest were NTMs. The ICT specificity was 99.4%, and the sensitivity was 94% when compared to the DNA probe test. The reduced specificity was due to the DNA probe missing two MTBC-positive samples that were detected by the ICT one due to not enough growth and the other due to the presence of blood. The reduced sensitivity was due to the ICT missing 3 DNA probe positives. The first discrepant isolate was M. bovis BCG which is typically negative by the ICT since it detects Mpt64 antigen. The second discrepant isolate was suspected to have a mutation in mpt64; the gene was detectable by PCR but not by ICT even upon subculturing in different media. The third discrepant isolate was from a lymph node sample that was submitted in an unidentified pink fluid. When an aliquot of the culture media was heat killed in preparation for a confirmatory TB PCR assay, it produced a white precipitate. Subculturing the isolate allowed the ICT to become positive.

CONCLUSIONS: Our results are in agreement with the literature for this ICT. The ICT was simple to use. The rapid identification of MTBC in fifteen minutes without any equipment costs (versus three hours using the Gen-Probe AccuProbe) will not only reduce the culture identification workload but also will be cost efficient while improving turn-around times.

C04 UNDERSTANDING PSEUDOMONAS AERUGINOSA TREATMENT RESISTANCE IN CHRONIC LUNG INFECTION USING DEEP POPULATION SEQUENCING
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Objective: To characterize P. aeruginosa isolates from sick individuals were collected from the lung by deep population sequencing to understand the genetic basis of drug resistance.

Methods: All P. aeruginosa isolates from sick individuals were collected from the lung by deep population sequencing to understand the genetic basis of drug resistance. This analysis revealed the gene encoding penicillin-binding protein 3 (PBP3). These mutations were confirmed by 16S ribosomal RNA sequencing; serotyping was performed by 16S rRNA gene sequencing) we show that P. aeruginosa is the dominant bacterial constituent of the lung, despite numerous courses of multiple anti-pseudomonal antibiotics. We then whole-genome sequenced a total of 235 P. aeruginosa isolates twelve sputum specimens obtained over a one-year period. This revealed multiple intraspecies ‘selective sweeps’, demonstrating that within species population dynamics result in periodic complete or nearly complete strain replacement, producing multiple distinct populations: an ancestral population characterized by higher diversity and recombination, and a sweep population characterized by mutations in the gene encoding penicillin-binding protein 3 (PBP3). These mutations correlate with resistance to anti-pseudomonal antibiotics. These intraspecies dynamics are not apparent with lower resolution analyses, and demonstrate an adaptive mechanism allowing P. aeruginosa to persist in the CF lung during therapy.
D03
ANTIBIOTIC SUSCEPTIBILITY OF INVASIVE HAEMOPHILUS INFLUENZAE STRAINS IN CANADA, 2007 TO JUNE 30, 2014
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OBJECTIVE: Susceptibility of invasive Haemophilus influenzae (Hi) to commonly prescribed antibiotics was determined. Canadian isolates from 2007-2014 were included and they were also characterized based on their serotype and sequence type, to determine any possible correlation with susceptibility profiles.

METHODS: Standard methods were used to determine serotype and MLST. β-lactamase was determined by chromogenic method and amplification of TEM-1 and ROB-1 genes. Disk diffusion and MIC determination were conducted according to CLSI guidelines.

RESULTS: Of the 849 invasive isolates examined, resistance to one or more antibiotics was seen in 48.0% of non-typeable (229/477), 36.4% of serotype b (24/66), 33.3% of serotype e (7/21), 16.7% of serotype c (1/6), 14.0% of serotype f (13/93) and 3.2% of serotype a (6/186) isolates. β-lactamase mediated ampicillin resistance was most commonly observed among all serotypes and non-typeable isolates, with rates as high as 30.3% of the serotype b isolates (20/66) and 21.6% of the non-typeable isolates (103/477). Resistance to trimethoprim-sulfamethoxazole was observed with rates as high as 20.8% of non-typeable (99/477) and 13.6% of serotype b (9/66) isolates. Resistance or reduced susceptibility to a number of other antibiotics was also observed. Multidrug resistance was observed in serotype a, b, c and non-typeable isolates.

CONCLUSIONS: Overall, higher incidence of resistance was observed with non-typeable isolates compared to serotypeable isolates. Resistance rates of non-typeable isolates were also higher for specific antibiotics, with the exception of ampicillin resistance in serotype b isolates. Non-typeable isolates show resistance to more antibiotics, and higher incidence of multidrug resistance compared to serotypeable isolates.

E01
NURSING ADVANCED DIRECTIVE TO REMOVE URINARY CATHETERS AMONG GENERAL INTERNAL MEDICINE PATIENTS: A CONTROLLED BEFORE-AFTER STUDY
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BACKGROUND: Overuse of urinary catheters among general medical inpatients contributes to increased incidence of healthcare-associated urinary tract infection and catheter-associated asymptomatic bacteriuria. Prior studies
suggest physicians are often unaware of whether patients are catheterized, and nurse-led directives for catheter removal can reduce unnecessary use.

METHODS: An advanced directive was implemented allowing nurses to remove urinary catheters for patients admitted to two general medical units who lacked the accepted indication: pre-admission permanent indwelling catheter, bladder outlet obstruction, need for continuous bladder irrigation, stage 3/4 sacral ulcers among incontinent patients, patient wishes for comfort care in end of life, or serum sodium <120 mmol/L with a physician order for strict fluid monitoring. Catheter-days per patient-days was compared monthly to two control general medical units before (1 month) and following (3 months) implementation using a task-oriented nurse acuity tool validated to measure urinary catheter use. Active surveillance for catheter re-insertion within 48-hours was performed with chart reviews to assess for re-insertion due to inappropriate urinary catheter removal.

RESULTS: At baseline, urinary catheter-days per patient-days were 252/1332 (18.9%, 95% CI, 16.9-21.1%) and 264/1410 (18.7%, 95% CI, 16.8-20.8%), on intervention and control units, respectively (P=0.89). Following implementation of nursing advanced directive, catheter-days per patient-days on intervention units decreased to 107/1351 (7.9%, 95% CI: 6.6-9.5%), significantly below control units (194/1539, 12.6%; 95% CI: 11.0-14.4%, P=0.0001). Three-months after implementation, catheter use on intervention units remained significantly lower (96/1564, 6.1%; 95% CI: 5.1-7.4%) compared to control units (254/1669, 15.2%; 95% CI: 13.6-17.1%, P=0.002). Eight catheter re-insertions occurred over 3-months, and none were related to inappropriate catheter removal.

CONCLUSIONS: A nursing advanced directive to remove unnecessary urinary catheters on general medical units led to a significant reduction in urinary catheter use without inappropriate removal. Long-term follow-up is needed to confirm sustainability of this model of care.

E02 IMPACT OF URINARY TRACT INFECTION PROVINCIAL EMPRIC TREATMENT GUIDELINES ON PRESCRIBING OF NITROFURANTOIN IN PRIMARY CARE AND LONG TERM CARE SETTINGS IN THE PROVINCE OF PRINCE EDWARD ISLAND
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OBJECTIVES: The Health PEI Urinary Tract Infection Empiric Treatment Guidelines were distributed to all physicians, nurse practitioners, and pharmacists in Prince Edward Island on May 24, 2013. In these provincial guidelines, nitrofurantoin is recommended as the first-line option for early cystitis in females (if no contraindication for its use) and a twice daily dosing formulation (MacroBID®) is suggested over four times daily dosing formulations to improve patient adherence. The objective of this audit is to examine the potential impact the guidelines have on nitrofurantoin prescribing practices in primary care and long term care settings.

METHODS: Information on nitrofurantoin prescriptions dispensed at community pharmacies and Provincial Pharmacy from January 1, 2012 to October 28, 2014 was obtained from the provincial Drug Information System. Prescriber claims using ICD-9 diagnosis codes for UTI-related diagnoses were retrieved for females from the Medicare Claims System. Prescriber claims using ICD-9 diagnosis codes for UTI-related diagnoses were retrieved for females from the Medicare Claims System. The pre-intervention period was defined as January 1, 2012 to November 2012 and the post-intervention period was June 2013 to October 2014.

RESULTS: When comparing the pre-intervention period to the post-intervention period, the average number of monthly nitrofurantoin prescriptions dispensed for female patients increased by 30.7% (P<0.001). As a potential confounder, the number of diagnoses of UTIs pre- and post-intervention was examined. The average number of monthly female UTI claims increased post-intervention, although not significantly (+7.6%, P=0.06). When controlled for differences in UTI diagnosis claims, the percent of nitrofurantoin prescriptions increased by 21.6% (P<0.001). The percentage of nitrofurantoin prescriptions for MacroBID® increased by 25.7% in the post-intervention period (P<0.001).

CONCLUSIONS: Primary care clinicians are more likely to prescribe nitrofurantoin for female UTIs after the release of the guidelines and this increase was sustained over the post-intervention period. The new guidelines appear to have a positive and lasting impact on the prescribing behaviours of PEI prescribers.

E03 EARLY ONSET INVASIVE CANDIDIASIS IN EXTREMELY LOW BIRTH WEIGHT INFANTS: RISKS AND OUTCOMES IN A MULTICENTRE CANADIAN PROSPECTIVE STUDY
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BACKGROUND: Data on neonatal invasive candidiasis (IC) presenting in the first week of life are limited to case reports. This entity is less well described than later onset IC. We sought to compare the clinical features and outcome in early onset disease (EOD, <= 7d) with those in late onset disease (LOD, >7d) in extremely low birth weight (ELBW, <1000g) infants.

METHODS: All ELBW cases and their controls (2 controls: 1 case) from a Canadian multicentre prospective study of neonatal candidiasis in very low birth weight infants were selected for secondary analysis. Factors associated with occurrence and outcome of EOD in ELBW infants were identified using logistic regression.

RESULTS: EOD occurred in 14 of 45 (31%) ELBW infants with IC compared to 2/24 (8%) of infants ≥1000g (P=0.039). Birth weight <750g, gestation <25 weeks, chorioamnionitis and vaginal delivery were all associated with EOD (P<0.01) versus LOD. Infection with Candida albicans, disseminated disease, pulmonary involvement and cardiovascular disease were significantly more common in EOD than in LOD (P<0.05). The EOD case fatality rate (71%) was higher than in LOD (32%) or controls (15%); P=0.001. The rate of neurodevelopmental impairment and mortality combined was similar in EOD (86%) and LOD (72%) but higher than in controls (32%; P=0.007).

CONCLUSION: Outcomes of EOD in ELBW infants are poor. The association of EOD with chorioamnionitis, vaginal delivery and pneumonia support perinatal acquisition. Dissemination is common and affected infants often die. Empiric antifungal treatment should be considered for ELBW infants with pneumonia, delivered vaginally, whose mothers have chorioamnionitis or an intraterine foreign body.

E04 SUCCESSFUL TREATMENT OF CANDIDA ALBICANS MENINGITIS WITH INTRATHECAL AMPHOTERICIN AND INTRAVENTRINAL FLUCONAZOLE: A CASE REPORT AND REVIEW OF LITERATURE
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OBJECTIVE: To present a case of Candida albicans meningitis post cranio-pharyngioma resection successfully treated with a combination of intrathecal (IT) amphotericin and intravenous (IV) fluconazole.

METHODS: A case report and literature review (1958-2014) on the treatment of Candida albicans meningitis complicating neurological procedures was performed via PubMed and Ovid using search terms: “Candida”, “meningitis”, “shunt-infection”, “neurosurgical”, and “treatment”.

RESULTS: A 59-year-old man underwent resection of a cranio-pharyngioma complicated by a CSF leak necessitating lumboperitoneal (LP) shunt insertion. Postoperatively he developed persistent fever and a reduced level of consciousness. CSF analysis revealed elevated leukocytosis and protein with...
normal glucose. He continued to be febrile while on broad-spectrum antibiotics. Although initial blood and CSF cultures were negative, following removal of LP shunt both CSF and shunt tip grew Candida albicans. After the initiation of IV amphotericin, the patient quickly defervesced with a decrease in CSF leucocytosis. His course was then complicated by acute renal failure requiring a switch to IV fluconazole. After 7 days of therapy his level of consciousness worsened with increased CSF leucocytosis. However, cultures remained negative. IV amphotericin was added and the patient improved clinically and biochemically. Review of the literature revealed 85 cases of Candida infection complicating neurosurgery for which the majority were secondary to Candida albicans. Over 85% of cases received IV amphotericin alone or in combination with IT amphotericin, fluconazole and/or fluvoxazole. No cases reported use of IT amphotericin with IV fluconazole. Cure rates with IV amphotericin with or without IT amphotericin were highest and ranged from 80 to 90%, while fluconazole monotherapy resulted in a cure rate of 60%.

CONCLUSION: Candida infections of the central nervous system are an increasingly recognized post-neurosurgical complication associated with significant mortality. Patients that do not tolerate IV amphotericin may benefit from the addition of IT amphotericin to fluconazole.

E05  
LINEZOLID-RESISTANT MRSA STRAINS SEQUENTIALLY ISOLATED FROM AN ELDERLY PATIENT WITH PNEUMONIA DURING PROLONGED THERAPY WITH LINEZOLID  
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OBJECTIVE: Linezolid is a crucial drug for methicillin-resistant Staphylococcus aureus (MRSA) infection. Here we describe the emergence of linezolid-resistant MRSA strains with a 23S rRNA mutation sequentially isolated from an elderly patient with pneumonia, during long-term linezolid treatment.

METHODS: A retrospective analysis was conducted to calculate the cumulative amount of linezolid administered to an elderly patient with pneumonia during prolonged therapy with linezolid. The sequential isolates were identified and their minimum inhibitory concentration (MIC), homology and drug resistant related genes (cfr, 23S rRNA) were investigated.

RESULTS: One linezolid-resistant MRSA strain was isolated from sputum of the patient at Xijing Hospital. The MIC of linezolid had increased from 1.5 μg/mL of initial strain 083 in March 2012 to subsequent 4.0 μg/mL of strain 158 in November 2012 and 16.0 μg/mL of strain 231 in March 2013 respectively. The increasing MIC values to linezolid were directly correlated with the cumulative amount of linezolid 118.8 g administered over 99 days. Strain 158 had heterogeneous resistance to linezolid. Molecular typing showed that three MRSA strains belonged to MLST ST-239 (spa t-030) with 98.8% homology. Cfr mutation was not detected but 23S rRNA mutation G2603T was confirmed in strains 158 and 231.

CONCLUSIONS: This is the first case of linezolid-resistant MRSA in China. The long-term use of linezolid for the treatment of MRSA infection was associated with a 23S rRNA G2603T mutation, which mediates heterogeneous resistance to linezolid. And the strain with heterogeneous resistance might progress to a high-level linezolid-resistant strain when re-exposed to linezolid.
**F02**

**GENETIC DIVERSITY AND MOLECULAR EVOLUTION OF RESPIRATORY SYNCYTIAL VIRUS A ON1 GENOTYPE, 2010-2013: A COMPARATIVE ANALYSIS**

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In 2010, a novel genotype of human respiratory syncytial virus (RSV), RSV A ON1, was first detected in Ontario, Canada, and subsequently isolated from various countries. In order to investigate its persistence and dynamics, we tested 406 randomly selected RSV positive samples submitted to Public Health Ontario from August 2011 to August 2012 by real-time RT-PCR. We identified RSV NA1 (n=181), NA2 (n=135), GA5 (n=3), and ON1 (n=51), and RSV-B (n=30). In addition to the 51 ON1 samples identified, we also collected 346 ON1 G-gene sequences from 18 countries from GenBank to evaluate ON1 genetic diversity, lineage distribution, rate of evolution, selection pressure, and the time of most recent common ancestor (tMRCA). Phylogenetic analysis determined that globally there are three ON1 lineages and we re-confirmed a newly defined genotype, ON2, recently reported in Italy. In Ontario there are two ON1 lineages circulating concurrently with a new cluster that has agreement with lineage identification criteria. ON1 is evolving at a rate of 3.29×10\(^{-3}\) substitutions/site/year. Path-O-Gen analysis determined the possible date of tMRCA as 2002. Selection pressure analysis identified 3 positively substituted sites: V225, L274, and Y297 that are heavily selected (dN/dS=3.5) of tMRCA.

**Conclusions:** RSV A-ON1 is rapidly evolving and spreading globally.

**F03**

**SHEDDING OF LIVE-ATTENUATED INFLUENZA VACCINE (LAIV) VIRUSES IN CANADIAN CHILDREN AND ADOLESCENTS WITH AND WITHOUT CYSTIC FIBROSIS**

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**Objective:** To quantify and describe trends in shedding of vaccine-strain influenza viruses following administration of LAIV in Canadian children and adolescents aged 2-19 years with and without cystic fibrosis (CF).

**Methods:** During the 2013-14 influenza season, 76 participants (57 patients with CF and 19 healthy siblings) were followed-up for 7 days post-LAIV administration. Reverse-transcription polymerase chain reaction (RT-PCR) and viral cultures were performed on flocked nasal swabs obtained from each participant immediately before vaccination (day 0) and on days 1, 2, 4 and 7 post-LAIV. Trends in viral shedding were analyzed using a random effects logistic regression model to determine the effect of demographic variables on PCR-detected viral shedding.

**Results:** Using viral culture, 5 participants (all with CF) shed influenza A virus (1 participant each on days 1, 4 and 7; 2 participants on day 2); 3 of these had received LAIV in the prior influenza season. No culture was positive for influenza B. All 5 participants with positive influenza A cultures also had PCR-detected influenza A virus shedding throughout follow-up. The number of participants shedding PCR-detected viruses was highest on day 1 post-LAIV administration for both influenza A and B viruses. Overall, 40 (70%) participants with CF and 11 (58%) healthy participants shed either PCR-detected influenza A and/or B viruses throughout follow-up. Specifically, 21 participants (28%) shed both influenza A and B at least once, 7 (9%) shed only influenza A virus at least once, 23 (30%) shed only influenza B virus at least once, and 25 (33%) did not shed any virus. Finally, there were no statistically significant differences in the odds of PCR-detected viral shedding by sex, CF status and LAIV vaccination in the previous year (P>0.05). However, the odds of shedding either PCR-detected influenza A and/or B virus decreased as age increased (OR 0.87, 95% CI 0.81-0.94, P=0.001).

**Conclusions:** Shedding of live vaccine-strain LAIV viruses does not appear to be frequent in children with CF. Shedding of PCR-detected LAIV viruses was most frequent the day after vaccine administration. The odds of shedding PCR-detected influenza A and/or B viruses are lower in older compared to younger children.
F05 DETECTION OF ENTEROVIRUS D68 IN CANADIAN LABORATORIES
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1Division of Microbiology, Department of Pathology and Laboratory Medicine, Capital District Health Authority, and Department of Pathology; 2Dalhousie University, Halifax, NS; 3Provincial Laboratory for Public Health, and Department of Laboratory Medicine and Pathology; 4University of Alberta, Edmonton, AB; 5Enterovirus and Enteric Virus Laboratory, National Microbiology Laboratory, Winnipeg, MB; 6Laboratoire de Santé Publique du Québec/Institut National de Santé Publique du Québec, Québec, QC; 7Calgary Laboratory Services Centre, Toronto, ON; 8University of Calgary, Calgary, AB; 9Public Health Laboratory & Pathobiology, University of Toronto; 10Sunnybrook Health Sciences Centre, Toronto, ON, ON; 11University of Ottawa, Ottawa, ON; 12Public Health Microbiology and Reference Laboratory, British Columbia Centre for Disease Control, Vancouver, BC; 13Saskatchewan Disease Control Lab, Ministry of Health, Regina, SK; 14Department of Microbiology, Mount Sinai Hospital and University Health Network; 15Department of Laboratory Medicine and Pathobiology, University of Toronto; 16University of Western Ontario, London; 17Sunnybrook Health Sciences Centre, Toronto, ON; 18Public Health Laboratory & Microbiology, St John's, NL; 19The Hospital for Sick Children, Toronto; 20St Joseph's Healthcare, Hamilton, ON; 21University of Calgary, Calgary; 22Medical Microbiology and Immunology, University of Alberta, Edmonton, AB

BACKGROUND: In late summer and early fall of 2014, widespread enterovirus type D68 (EV-D68) activity was described across North America with severe cases described in the US. Of particular concern, EV-D68 infection has been associated with acute flaccid paralysis, and fatal cases have been documented. While nucleic acid amplification tests (NAAT) have been documented. While nucleic acid amplification tests (NAAT) have become standard practice in most clinical virology laboratories, implementing such assays is essential to mitigate inappropriate antibiotic use, initiate appropriate anti-viral prophylaxis or treatment and implement infection control procedures to limit transmission. As such, multiplexed molecular assays have become standard practice in most clinical virology laboratories. The goal of this study was to evaluate and compare the Seegene® Seeplex RV15 detection kit (done on Caliper Life Sciences gel based detection platform) with two next generation assays, the Seegene® Anyplex II RV16 (done on BioRad CFX-96 Real Time PCR platform) and the Luminex® xTAG RVP fast2 (done on MagPix platform) respiratory virus panels.

METHODS: One hundred and sixty-seven retrospective and 36 prospective respiratory samples were tested on all three assays. Samples were deemed to be positive if they tested positive for a virus by 2/3 assays. Negative samples had to test negative by 2/3 assays, while inconclusive samples were defined as those with band signal intensity between 0-100 on the RV15.

RESULTS: There was no statistically significant difference in overall sensitivity and specificity (Table 1). Discordant rates were 47% and 21% for positive and negative samples, respectively.

CONCLUSION: Overall sensitivity and specificity of all 3 assays were similar though higher than expected discordant rates may reflect primer or chemistry differences amongst the three multiplex assays. Hence, factors such as turn-around time, the need to repeat testing and price become important variables for consideration in the decision making process for implementing such assays.

TABLE 1

| Assay Sensitivity and Specificity for RV15, xTAG and RV16 by Virus Type |
|-----------------|-----------------|-----------------|
| Viral Target     | RV15            | xTAG            | RV16            |
| Sens            | Spec            | Sens            | Spec            | Sens            | Spec            |
| Rhino/ Enterovirus | 93              | 95              | 88              | 100             | 92              | 98              |
| RSV A/B         | 83              | 100             | 88              | 100             | 92              | 99              |
| Influenza A     | 83              | 100             | 95              | 100             | 95              | 100             |
| Influenza B     | 85              | 100             | 64              | 99              | 92              | 100             |
| Parainfluenzavirus | 92             | 100             | 94              | 99              | 90              | 100             |
| Metapneumovirus | 82              | 100             | 88              | 100             | 99              | 100             |
| Coronavirus     | 88              | 100             | 92              | 100             | 96              | 100             |
| Adenovirus      | 100             | 100             | 100             | 99              | 70              | 100             |
| Bocavirus       | 50              | 99              | 44              | 100             | 63              | 100             |
| Average         | 84              | 99              | 84              | 100             | 87              | 100             |

* Sens= Sensitivity; Spec= Specificity

GO1 COMPARATIVE EVALUATION OF TWO PHENOTYPIC TESTS FOR DETECTION OF PENICILLIN BINDING PROTEIN 2A (PBPA2) IN DIVERSE RETROSPECTIVE CLINICAL STAPHYLOCOCCUS AUREUS (SA) ISOLATES
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OBJECTIVES: PBPA2 detection is used to distinguish SA resistant to methicillin (MRSA) from those that are susceptible (MSSA). The Denka Seiken PBPA2 monoclonal agglutination has been the most commonly used test in clinical laboratories. But as it does not detect the emergent mecC-protein, and as it is laborious, other quicker tests are now under development.
consideration. This study compared the Denka to the Alere PBP2a Colony Culture Test (CCT) a line assay that claims to detect mec MRSA.

METHODS: 156 highly-characterized SA including 126 genotypically distinct MRSA (116 mecA, 10 mecC) and 30 unrelated MSSA (13 with SCCmec remnants) were randomized to 5% sheep blood agar (BA) and Chromogenic Staphylococcus/Denim Blue agars (CA) for blinded parallel testing by Denka and CCT. As specified by Alere for CCT inoculation, 3 SA colonies were touched but not picked, and even faint blue lines were positive (pos). CCT discrepancies were retested by picking 3 colonies in parallel to 3 touched colonies.

RESULTS: Of 79 SA tested from BA, CCT was pos in 61/61 mecA (18 weak-pos) but 0/4 mecC MRSA, and negative (neg) in 14/14 MSSA, while Denka was pos in 61/61 mecA and 1/4 mecC MRSA, and neg in 13/14 MSSA (1 weak-pos). Of 77 SA tested from CA, CCT was pos in 45/55 mecA (22 weak-pos) but 0/6 mecC MRSA, and neg in 15/16 MSSA (1 weak-pos remnant), while Denka was pos in 55/55 mecA and 0/6 mecC MRSA, and neg in 13/16 MSSA (3 weak-pos remnants). Overall % sensitivities (Sn)/specificities (Sp) from BA and CA combined for CCT and Denka, respectively, were 91 (95% CI: 85-95)/100 (95% CI: 86-100) were repeated. Using CCT data from 3 picked colony repeats, revised %Sn/Sp ties (Sp) from BA and CA combined for CCT and Denka, respectively, were 83 (95% CI: 76-89)/97 (95% CI: 82-99.9) and 93 (95% CI: 87-96)/87 (95% CI: 70-95). 29 SA from failed tests and 19 from non-tested (as controls) were repeated. Using CCT data from 3 picked colony repeats, revised %Sn/Sp from BA and CA combined for CCT and Denka, respectively, were 83 (95% CI: 76-89)/97 (95% CI: 82-99.9) and 93 (95% CI: 87-96)/87 (95% CI: 70-95). 29 SA from failed tests and 19 from non-tested (as controls) were repeated. Using CCT data from 3 picked colony repeats, revised %Sn/Sp from BA and CA combined for CCT and Denka, respectively, were 83 (95% CI: 76-89)/97 (95% CI: 82-99.9) and 93 (95% CI: 87-96)/87 (95% CI: 70-95).

CONCLUSIONS: Denka detected all mecA MRSA from BA and CA, and CCT detected all mecA from BA but only 80% from CA (P=0.0008) using manufacturers’ protocols. A modified protocol using 3 picked instead of touched colonies increased the overall CCT Sn for detecting mecA MRSA from 90.5% to 99.1% (P=0.0052). Denka detected (as weak-pos) 1 of 4 mecC tested on BA; but otherwise Denka and CCT missed all mecC MRSA.

G02

COMPARISON OF TWO BROTH ENRICHMENT AND TWO AGARS FOR THE DETECTION OF GROUP B STREPTOCOCCUS (GBS) FROM VAGINAL-RECTAL SCREENING SPECIMENS

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BACKGROUND: GBS infection can cause both early- and late-onset neonatal disease. Widespread adoption of GBS screening and intrapartum prophylaxis has reduced the incidence of early-onset disease significantly. Further optimization of screening may result in additional benefits. As a result, there has been renewed interest to enhance GBS recovery, using new broths, new agar plates, chromogenic media as well as molecular detection. The purpose of this study was to assess the sensitivity and specificity of novel culture methods to enhance the recovery of GBS.

METHODS: Vaginal-rectal swabs were collected in duplicate from women attending routine pre-natal appointments between 34 and 36 weeks gestation. Swabs were placed in either Todd-Hewitt (TH) broth (Oxoid™) or RambaQuick (RQ) broth (CHROMagar™), which is designed to reduce overgrowth of enterococci isolated in vaginal-rectal samples. Broths were incubated overnight at 35°C. Broths were subcultured to sheep blood agar (SBA) and Colorex™ StrepB (CSB, CHROMagar™). GBS were identified using colony morphology and Lancefield typing from SBA and by chromogenic reaction and Lancefield typing from CSB.

RESULTS: 45 of 204 samples were positive for GBS. There was no significant difference between testing methods. Sensitivity was 97.8% for TH broth subculture to SBA, and 97.8% for TH broth subculture to CSB. Sensitivity was 95% for RQ subculture to SBA and 97.7% for RQ subculture to CSB. Specificity was 100% for all media. CSB allowed easy visual detection of GBS. RQ broth provided short-24 hours at 35 °C. Plates were examined and beta hemolytic colonies were transfered to culture.

CONCLUSION:Detection of GBS by LAMP is faster and more cost-effective than culture with comparable performance. This assay provides promising results for rapid detection of GBS compared to culture.

G04

COMPREHENSIVE EVALUATION OF CEPHEID’S XPERT NASAL COMPLETE G3 PCR (XNC) ON THEIR GENEXPERT FOR DISTINGUISHING METHICILLIN-RESISTANT (MR) FROM METHICILLIN-SUSCEPTIBLE (MS) STAPHYLOCOCCUS AUReUS (SA) AND COAGULASE-NEGATIVE STAPHYLOCOCCUS (CNS) FROM ISOLATES

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OBJECTIVES: Rapid PCR detection of MRSA directly from clinical specimens is problematic due to 1) genetic similarities between mecA in SA and CNS, 2) prevalence of MSSA with remnant (rem-SA) Staphylococcus Cassette Chromosome mec (SCCmec) sequences that no longer carry mecA, and 3) continued evolution of SCCmec and mec determinants. PCR assays are notoriously prone to false-positives (FP) due to 1 and 2) and false-negatives (FN) due to 3. We evaluated the XNC with the aim to speed up institution of appropriate therapy in cases of short-incubation (<6h) blood culture SA identified by MALDI-TOF.
METHODS: 181 highly-characterized genotypically-unique clinical isolates studied included 126 MRSA (116 mecA, 10 mecC), 30 MSSA (13/30 rem-SA) and 25 CNS. On recovery from ~80°C, strains were plated twice: 50%/50% to 5% sheep blood/Chromogenic Staph or Demin Blue agars (Oxoid). Bacterial suspensions prepared in saline equivalent to ~0.1 MacFarland STD were inoculated by swab to XNC buffers, after which testing was conducted as per Cepheid. XNC results including crossing thresholds (Ct) for SPA (detects SA), mec (detects mecA) and SCC (detects SCCmec) were correlated with strain characteristics.

RESULTS: The XNC SPA target correctly identified 156/156 SA (Ct: 16.7-25.7; 100%; 95% CI: 97-100) and excluded SA (and MRSA) in 25/25 CNS (Ct: 0; 100%; 95% CI: 84-100) regardless of agar used. mec and SCC were amplified in 116/116 mecA MRSA (100%; 95% CI: 96.2-100) with Ct ranges 16.7-26 and 17.6-32.7, respectively. Not surprisingly, these targets were detected in 0/10 mecC MRSA leading to an overall MRSA detection sensitivity (Sn) of 92.1% (95% CI: 85.9-95.8). Although 0/30 MSSA was reported as MRSA (100% specificity; 95% CI: 86.5-100), amplification occurred for SCC alone in 11/13 rem-SA (Ct: 19.6-22.8) and 3/17 SA with no remnants (Ct: 27.2-37.6), for mec alone in 1 rem-SA (Ct: 37.9), and SCC and mec were both amplified in 1/17 MSSA (Ct: 36.4, 36.7).

CONCLUSIONS: With the exception of mecC MRSA, this retrospective study data supports the use of the XNC assay directly from purified clinical isolates. However, prior to its clinical implementation for testing short incubation blood culture SA, additional prospective evaluations should be undertaken to ensure 1) all currently circulating MRSA are indeed detected, and 2) that mixed MSSA/MR-CNS cultures do not lead to false-reports of MRSA.

G05
DIRECT REPORTING CEFAZOLIN FROM VITEK 2 AST NR208 FOR E. COLI, K. PNEUMONIAE AND P. MIRABILIS ISOLATED FROM URINE CULTURES USING 2014 CLSI INTERPRETATION
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BACKGROUND: There have been amendments in cefazolin interpretations through the years (see Table). During 2010-3, our laboratory performed a Kirby-Bauer (K-B) confirmation test for cefazolin before reporting Vitek 2 AST results for E. coli, K. pneumoniae and P. mirabilis isolated from urinary specimens to accommodate the lowered breakpoints.

Cefazolin interpretation Susceptible Intermediate Resistant (UTIs, mg/L) (S) (I) (R)
2014 CLSI M100-S24 <= 16 – >=32

This study is to verify if the lab can report cefazolin directly from Vitek 2 testing without an extra K-B test based on the new 2014 breakpoints.

METHODS: Cefazolin susceptibilities of 129 urinary clinical isolates of E. coli, K. pneumonia and P. mirabilis in 2013 were compared, between their Vitek 2 and K-B (reference method) testing results, using the new UTI 2014 CLSI interpretations for cefazolin (30 µg/disc) S: ≥15 mm and R: ≤14 mm. Category agreement was measured and described as minor, major and very major errors.

RESULTS: Based on the 2014 new K-B interpretation, direct reporting cefazolin from Vitek 2 AST against E. coli, K. pneumonia, and P. mirabilis gave a category agreement of 94%.

TABLE 1
Category Agreement between K-B vs Vitek 2 for Cefazolin of 129 isolates

<table>
<thead>
<tr>
<th>Error Type</th>
<th>K-B Method</th>
<th>Vitek 2 Method</th>
<th>No. of Errors (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minor Errors</td>
<td>S or I</td>
<td>I or S</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>R or I</td>
<td>I or R</td>
<td>0</td>
</tr>
<tr>
<td>Major</td>
<td>S</td>
<td>R</td>
<td>4 (3%)</td>
</tr>
<tr>
<td>Very Major</td>
<td>R</td>
<td>S</td>
<td>4 (3%)</td>
</tr>
</tbody>
</table>

CONCLUSION: It is generally acceptable to report cefazolin directly from Vitek 2 for the three species of Enterobacteriaceae isolated from urine cultures using the new 2014 CLSI interpretation. It saves cost >$3000/yr. *Vitek 2 interpretation based on (FDA Cleared Device).

G06
RAPID DIAGNOSTIC TESTING AND LACTOBACILLUS REUTERI THERAPY FOR CHILDREN WITH SEVERE ACUTE GASTROENTERITIS IN BOTSWANA: A PILOT, FACTORIAL, RANDOMIZED, CONTROLLED, CLINICAL TRIAL
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OBJECTIVES: To demonstrate the feasibility of an RCT designed to measure the benefit of rapid enteric diagnostic testing and probiotic therapy for children admitted to hospital in Botswana with severe acute gastroenteritis. METHODS: Children with acute diarrhoeal disease aged 2-60 months admitted to any of three Botswana hospitals were eligible to participate if they did not have bloody stools or signs of sepsis. Participants received standard fluid rehydration, zinc treatment, and were randomized to one of four arms: 1) rapid diagnostic testing (plus targeted antimicrobial therapy if indicated) plus Lactobacillus reuteri therapy × 60 days; 2) rapid diagnostic testing (plus targeted antimicrobial therapy if indicated) plus placebo therapy × 60 days; 3) standard care (delayed diagnostic testing) plus L. reuteri therapy × 60 days; or 4) delayed diagnostic testing plus placebo therapy × 60 days. Rapid testing consisted of multiplex PCR assays to detect Shigella, Campylobacter, enterotoxigenic E. coli LT/ST toxin, Salmonella, and Cryptosporidium. For this pilot study, achievement of feasibility outcomes was primary; these included validation of the diagnostic testing protocols, ensuring that testing and appropriate treatment could be integrated into clinical care, and verifying adequate recruitment rates. Clinical outcomes included height at 60 days, weight at 60 days, 60-day mortality, and recurrence of diarrhoea in the 60-day followup period.

RESULTS: In a 6-month study period, 76 participants were enrolled. The median participant age was 10.8 months, 49% had been exposed to HIV in utero, and 6 had severe acute malnutrition. 34 of 37 children randomized to rapid testing had results reported within 24 hours, and 19 of 20 with a treatable pathogen detected via rapid testing had antimicrobials begun within 24 h. Mean 60-day height was 2.56 cm greater in the rapid diagnostics arms compared to 2.07 cm greater in the standard treatment arms, and only 6 of 34 children (18%) randomized to rapid testing developed repeated diarrhoea in the followup period, compared to 12 of 34 (35%) in the control arms. Probiotic/placebo groups are still blinded.

CONCLUSIONS: We have demonstrated feasibility of the trial protocol. An adequately-powered multicentre RCT to precisely quantify the benefit of rapid diagnostics and L. reuteri probiotic therapy in this population is indicated.
H01
IS LYME DISEASE BEING MISSED IN MULTIPLE SCLEROSIS PATIENTS IN NEW BRUNSWICK?
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INTRODUCTION: Lyme disease is an emerging infection in Atlantic Canada. The concern has been raised that late Lyme disease is being misdiagnosed as other chronic diseases such as multiple sclerosis (MS). We tested a cohort of MS patients in New Brunswick to determine whether any patients within this cohort had laboratory test results consistent with Lyme disease.

METHODS: 90 patients attending the MS clinic in Saint John, New Brunswick were tested for the presence of antibodies for Borrelia burgdorferi using the two-step algorithm in accordance with the IDSA guidelines. Each patient’s serum was first tested with the C6 B. burgdorferi (Lyme) ELISA (Innmetics, Immunoassays, Inc., Boston, Massachusetts, USA). Samples that produced positive or equivocal results on the C6 ELISA were subsequently tested for IgG antibodies using a Western blot (B. burgdorferi US (IgG), Euroimmun, Luebeck, Germany). A positive Western blot required the presence of 5 of 10 significant bands. Although, 6 of 90 (6.7%) MS patients had serum that was reactive on the C6 EIA, none of these reactive EIA specimens could be confirmed using the Western blot.

CONCLUSION: The seroprevalence of antibodies against B. burgdorferi in this cohort of NB MS patients was zero by the IDSA two-step algorithm.

H02
PREVALENCE AND EPIDEMIOLOGY OF METHICILLIN-RESISTANT STAPHYLOCOCCUS AUREUS COLONIZATION IN AN ABORIGINAL COMMUNITY
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BACKGROUND: Canadian Aboriginal populations are at an increased risk for community-acquired MRSA infection for unknown reasons.

OBJECTIVES: To describe the prevalence and molecular epidemiology of MRSA colonization among humans and animals in an Aboriginal community.

METHODS: Adult and child volunteers were sought from clinic visits, house visits, school and work locations, between October 6 and November 27, 2014. A single nasal swab was collected. Dogs undergoing spay/neuter surgery were swabbed in nasal, oral, anal, groin and wound sites. Univariate regression (SPSS 22.0) and spa typing were performed.

RESULTS: 461 residents were approached, 13 refused consent, 4 were missing data, leaving 444 eligible participants. 242/444 (54.5%) were women, mean age was 25.7 yrs (SD 19.8), 169/444 (38.1%) had received antimicrobials in the last 12 months. MRSA prevalence was 111/444 (25.0%). Significant predictors of MRSA colonization included number of rooms in the house (RR=0.73, P=0.01), antibiotic use in the last 12 months (RR=1.18, P=0.04), owning a dog (RR=0.11, P=0.01), feeding a dog regularly (RR=16.14, P<0.01), and using a dog for recreation (RR=0.02, P=0.01). Age, gender, ethnicity, number of people in house, number of sinks in house, admission to hospital, surgery, indwelling medical device, dialysis, prior MRSA infection, history of skin infection, MRSA among contacts, intravenous drug abuse, new tattoos or piercing, chronic skin condition, exposure to healthcare or corrections or homeless shelter or daycare or veterinary care did not predict MRSA colonization. 89/111 (80.1%) of human staphylococci were CMRSA10, 82/89 (92.1%) of which were Spa type t028. 55/157 (35.0%) of dogs were colonized with coagulase-positive staphylococci.

CONCLUSIONS: MRSA colonization is not associated with healthcare contact, hygiene factors, skin disease or contact with MRSA, but may be associated with crowding, antibiotic use and dog exposure. Human MRSA is mostly clonal. Further genotyping may demonstrate human-animal inter-transmission.

H03
EVALUATION OF THREE CHROMOGENIC MEDIA FOR SCREENING SURVEILLANCE SPECIMENS FOR THIRD GENERATION CEPHALOSPORIN RESISTANT ENTEROBACTERIACEAE
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OBJECTIVE: This study evaluated three media for the detection of Enterobacteriaceae that posses genes for extended spectrum beta lactamases, ampC or carbapenemases.

MATERIALS AND METHODS: Chromogenic bi-plates of Colorex™ KPC and Colorex™ CIGR (CIGR) were obtained from Alere Canada. chromID™ CARBA (CIC) plates were obtained from bioMérieux Canada. Rectal swabs from 339 patients were vortexed in sterile saline and 50 μL was applied to each medium. A challenge set of known multi-resistant Enterobacteriaceae (n=31) was also tested. Plates were incubated as recommended and examined for degree of growth and colony colour. Isolates were identified using Vitek® MS (bioMérieux) MALDI-TOF Resistance mechanisms were elucidated using the CLSI ESBL disk test, Neo-Sensitabs™ Carbapenemase Confirmation tablets (Rosco Diagnostica) and PCR.

RESULTS: Both CRE specific agars were very selective. Escherichia coli colonies were pink on all plates. Citrobacter spp. were blue with a pink halo on CIGR, blue on KPC and dark pink on CIC. Klebsiellae and Enterobacter spp. were blue, green or blue-green. From the challenge set, CIGR medium detected 29 CRE (100%), KPC detected 23/29 (79.3%) and CIC detected 26/29 (89.7%). A CTX-M isolate was detected on all media and a plasmid ampC isolate was only detected on CIGR as expected. From routine specimens, CIGR detected 31 ESBL producing organisms, 1 of which grew on KPC agar and 10 AmpC producers, none of which grew on the other media. Seven isolates with an ESBL and plasmid ampC grew on CIGR, 2 of which grew on KPC and 1 on CIC.

CONCLUSIONS: Colour specificity for Escherichia coli was excellent on all media. Differentiation of Enterobacteriaceae from Klebsiellae was not possible and Citrobacters had various presentations. Colorex™ CIGR agar provides the greatest yield of multi-resistant Enterobacteriaceae. Combining CIGR agar with Colorex™ KPC and MALDI-TOF identification provides an efficient and cost effective solution for nosocomial screening.

H04
COMPARISON OF THE NEW MRSASELECT II, BRILLIANCE MRSA 2 MEDIA WITH CURRENT VERSIONS OF MRSASELECT AND DENIM BLUE FOR THE DETECTION OF MRSA
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OBJECTIVES: Two commercial media manufacturers have recently modified their respective formulations of chromogenic MRSA plates. A limited run of 500 of Thermo Fisher Oxoid’s Brilliance MRSA 2 (DB) and Bio-Rad’s MRSASelect II (MS2) were available for evaluation. Each claim to offer more robust growth at reduced incubation times of 18 hrs. To assess, we compared their previous versions MRSASelect (MS) and Denim Blue (DB) with new versions of each at various incubation times in 3 phases 1) retrospectively: 100 characterized MRSA strains split at dilutions of 10³, 10⁴ and 10 CFU/plate 2) 350 prospective nasal, rectal/perineal, wound swabs and 3) 50 recent retrospective low colony count positive swabs.

METHODS: 1) 100 known positive MRSA were freshly subcultured and
Abstracts

split into thirds diluted to each of approximately $10^6$, $10^7$ and $10^{10}$ CFU/plate. General colony size, count and clarity was compared at 12, 16, 18 hrs incubation 2) 350 new incoming samples were set up on each of 4 plates in random order and growth/no growth/breakthrough growth assessed at 12, 16, 18 hrs and 3) 50 plates of each reset up with patient swabs of known low CFU ($+/-$ growth) read at 18 and 24 hrs.

**RESULTS:** 1) Overall Sensitivity at 12 hrs: 61% MS; 85% DB; 88%DB2 and 90% MS2. At 18 hrs 61% all 100%. Growth ratio overall at 18 hrs ($+/+$: $+/2+: +/3+$) : MS 81:19:0; DB 45:45:10; DB2 33:45:22 ; MS2 16:49:36 2) 350 samples identified 34 positives (10% positivity rate) through at least one positive plate. Sensitivity at 18 hrs: MS:92%; DB: 94%; MS2 and DB2:97%. Specificity at 18 hrs: MS2: 92%; DB2: 96% and MS: 99%. 3) Detectability of low CFU DB2>MS2>DB>MS. Bacillus species were the noteworthy false positive growths on DB2 and MS2. Colonial morphology helped to differentiate from true MRSA.

**CONCLUSIONS:** Clarity of positive samples: MS2>DB2>DB>MS; Sensitivity MS2>DB2>MS=DB; Specificity: DB=MS>DB2>MS2; relative amount of growth MS2>DB2>DB>MS. Both new formulations are desirable over predecessor. In particular MS2 showed most improvement in sensitivity and clarity but not specificity. DB2 improvements were slight in sensitivity and clarity. Both new versions captured all positive growths within 18 hrs and in most cases could detect MRSA presence at 12 hrs as a possibility for pre-screening earlier.

H05

**A DIRECT COMPARISON OF TOXIGENIC CLOSTRIDIUM DIFFICILE BY DUAL RAPID TEST CASSETTE VERSUS TRADITIONAL ENZYME IMMUNE ASSAY / PCR ALGORITHMIC METHODS**

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**BACKGROUND:** The Prince County Hospital, PEI microbiology laboratory receives about 27 specimens per month for C. difficile with a 15.3% positivity rate. There is only one specimen run per day to the offsite laboratory that does 5 days a week GDH antigen and three times a week PCR testing. Various delays lead to over 40% of positives taking greater than 60 hours to result. A rapid method was studied.

**METHODS:** All samples were evaluated using the Bristol Score for stool consistency and only those having a score of 5 – 7 were tested. 100 screened samples were tested with the C. DIFF QUIK CHECK COMPLETE (a dual rapid test cassette for GDH antigen and toxin A&B) and then subsequently sent on for GDH EIA (Cdiff CHECK-60) and PCR toxin testing (BD GeneOhm Cdiff for tcdA).

**RESULTS:** 86% of samples tested exactly the same by all methods. Another 10% tested antigen positive, toxin negative by the rapid test and antigen positive, toxin positive by the algorithmic method. This was determined to be a direct result of the increased sensitivity of the PCR method. 4% did not show some correlation and were sent out of province for additional investigation. Cost analysis between the 2 methods was performed and ranges between a saving of 7.9% to an increase of 9.6% of non-labour costs were obtained depending on the approach to testing.

**CONCLUSIONS:** 98% of samples were essentially equivalent. 86% of samples tested rapidly so a significant reduction in turnaround time was evident. The one false positive Toxin EIA with a very faint line that in the future will be considered indeterminate. There was one false negative. The overall test performance using the algorithmic method as the gold standard is a sensitivity of 98.9% specificity of 88.9%. The rapid testing method when added to modified algorithm is accurate and at least cost neutral.

H06

**SUPERIORITY OF REAL-TIME PCR OVER OTHER ANTIGEN TESTING FOR SYPHILIS DETECTION**

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**BACKGROUND:** The clinical diagnosis of syphilis is complicated due to its diverse clinical manifestations, complex testing algorithms and variable antigen detection methods. In this study, we focus on comparing our real-time PCR assay to other antigen detection methods, namely, darkfield (DF) microscopy, direct fluorescent antibody (DFA) assay, as well as a first-generation conventional PCR. We intend to highlight trends and infer clinical relevance as we advance our testing methodologies.

**METHOD:** We compared the results of DF microscopy, DFA microscopy, and a conventional PCR which targets poA gene to our real-time PCR which targets both poA and tp47 protein genes. Two by two contingency tables were used for comparative analysis.

**RESULTS:** DF(100%), DFA (100%) and conventional PCR (100%) had comparable specificity to real-time PCR but each test had a much lower sensitivity. DF had the poorest sensitivity at 38.5%, DFA at 57.6%, and conventional PCR at 81.3%. The increasing sensitivity trend can be attributed to the advancement of test method technology which can provide more reliable and consistent results. Real-time PCR has the advantage of detecting T. pallidum down to a single spirochete, whereas microscopy is reliant on sample integrity, technologist’s expertise, and homogeneous sample distribution on the sample slide for detection. Another advantage of the real-time PCR is that more stringent control and quality control settings can be used for added sensitivity and specificity. **CONCLUSION:** Real-time PCR provided greater sensitivity and specificity than traditional microscopic and molecular methods. The semi-quantitative data generated from the real-time assay can also provide a means of using statistical rules such as the Levey-Jennings method to better manage the consistency and quality of the entire analytical process.


I01

**FULL GENOME ANALYSIS OF ENTEROVIRUS D-68 STRAINS CIRCULATING IN ALBERTA, CANADA**

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**INTRODUCTION:** Apart from respiratory tract infections and exacerbation of asthma EV-D68 has also been implicated in several other clinical presentations. Recently acute neurologic illness with focal limb weakness in children and acute flaccid paralysis with anterior myelitis has been linked to EV-D68.

**METHODS:** Four different EV-D68 positive samples from patients in Alberta, Canada were used to determine near full-length genome sequences. Primers designed based on the full genome sequence of the prototype human EV-D68 Fermon strain were used to amplify overlapping fragments spanning the entire genome. Full-length genome sequences were assembled using Seqscape v3.0 and Sequencing Analysis v6.0 (ABI). Nucleic acid and protein sequence alignments were performed using Clustal W. Neighbour joining trees were constructed using the Jukes-Cantor model and 1000 bootstrap replicates in MEGA v6.

**RESULTS:** The currently circulating EV-D68 strains are closely related to the contemporary strains circulating world-wide based on VP1 phylogenetic analysis, but have diverged considerably from the prototype Fermon strain. Phylogenetic analysis shows that the EV-D68 strains sequenced from all four patients in Alberta belong to lineage 2 based on Meijer et al and variant 1.2.1 and 1.2.2 based on the classification scheme by Lauinger. Consistent with these lineage designations for our sequences, the deletion at amino acid 141 in VP1 was absent and two deletion blocks were observed in the 5’UTR. Some of the Alberta strains showed amino acids changes at 142 and 143 that could affect the tertiary structure and protein function, however, no changes were noted in the antigenic epitope regions of the BC and DE loops.

**CONCLUSIONS:** EV-D68 was not a commonly reported enterovirus serotype between 1970 and 2005, however since then reports in the literature have suggested a greater incidence worldwide, independent of
increased surveillance and availability of sensitive detection methodologies. It has been suggested that mutations in the VP1 gene causing altered antigenicity or lack of immunity in the younger population due to the lack of circulation in the recent years may be responsible for the increased detection of EV-D68 worldwide.

IO2
GENERATION OF A CAMPYLOBACTER JEJUNI SURVEILLANCE DATABASE IN ALBERTA UTILIZING THE SUBTYPING METHOD COMPARATIVE GENOMIC FINGERPRINTING
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OBJECTIVES: Campylobacter jejuni infection due to ingestion of contaminated food or water is a leading cause of gastroenteritis. Subtyping methods for C. jejuni surveillance such as pulsed-field gel electrophoresis (PFGE), which is utilized by PulseNet Canada, have been developed. However, these methods are unable to overcome the difficulties of the high genetic diversity and chromosomal rearrangements in C. jejuni. An alternative method known as comparative genomic fingerprinting (CGF) was developed in the Laboratory for Foodborne Zoonoses by the Public Health Agency of Canada in Lethbridge, Alberta. CGF was adopted to build a C. jejuni surveillance database in Alberta as PFGE was not routinely used.
METHODS: A total of 304 known clinical C. jejuni isolates across Alberta health zones from 2006 to 2009 were genotyped for the presence or absence of 40 target accessory genes using capillary electrophoresis following multiplex polymerase chain reaction (PCR). Presence or absence of a gene was determined by analysis of generated electropherograms and gels by the computer software and user verification. Genomic fingerprints for each isolate comprising genotyping results were used to determine genetic relatedness amongst strains and generate clusters sharing 100% homology. RESULTS: There was high concordance between results generated with CGF, PFGE and epidemiological data pertaining to outbreak cases. Furthermore, new clusters between sporadic isolates were discovered, suggesting identification of outbreak cases that were previously not detected due to the lack of evidence or typing method for correlation. Subtyping with CGF is simple, robust, and has a fast turnaround time without sacrificing discriminatory power.
CONCLUSIONS: CGF is an alternative subtyping method for C. jejuni surveillance. Genomic fingerprints generated by CGF showed high concordance with available PFGE results and potentially revealed case associations previously not detected. The Alberta database will continue to expand with the inclusion of additional genomic fingerprints.

IO3
WHOLE GENOME SEQUENCING TO DIFFERENTIATE HIGHLY CLONAL SALMONELLA ENTERICA SEROTYPE HEIDELBERG ISOLATES INVOLVED IN OUTBREAKS IN THE PROVINCE OF QUEBEC
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BACKGROUND: Salmonella enterica is a common cause of food borne outbreaks. Salmonella enterica serotype Heidelberg is the most encountered serovar causing human salmonellosis in Canada and the second one in the province of Quebec. This serotype is highly clonal and over 50% of strains isolated in Quebec exhibit the same PFGE pattern SHEXAI.0001/SHEBNI.0001, leading often to unsuccessful outbreak investigations.
OBJECTIVE: This study aimed to evaluate the core genome SNP phylogeny method for discrimination of highly clonal S. Heidelberg strains involved in different outbreaks.
METHODS: Forty-six strains involved in 3 epidemiologically documented outbreaks in Quebec (2012, 2013 and 2014) were subjected to whole genome sequencing and core genome SNP analysis using SNVPhyl pipeline at the National microbiology laboratory (NML). A custom script to filter out all repeat regions and also remove any known mobile elements such as phage, islands was done. These strains exhibit the same pulsed-field gel electrophoresis pattern (SHEXAI.0001/SHEBNI.0001) and associated to phage types 19 or 26. Additionally, 15 sporadic strains with different PFGE patterns including the common one were included in this study.
RESULTS: Phylogenetic tree based on core genome SNP analysis allowed to clustering outbreak strains into 3 distinct clusters with concordance with epidemiological data. The number of SNP difference within strains from the same outbreak was comprised between 0 and 2 SNP. Interestingly, strains from outbreak 1 (n=11, 2012) and outbreak 3 (n=12, 2014) exhibiting the same PFGE and phage type patterns (SHEXAI.0001/SHEBNI.0001, PT 19) showed a SNP variation of 73. Strains from outbreaks 2 (n=8, 2013) exhibited a distinct phage type (PT 26) and showed a SNP difference of 82 and 147 compared to outbreak 1 and 3, respectively.
CONCLUSIONS: This study demonstrates that despite the high clonality of this serotype, core genome SNP was able to differentiate strains belonging to the same outbreak from others. This tool offers a powerful alternative for strain typing during outbreak investigations.
IO5 DETECTION OF LISTERIAモノCytogenes USING REAL-TIME AMPLIFICATION ASSAYS
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BACKGROUND: Listeria monocytogenes (Lm) is a major foodborne pathogen of concern for human health and food safety in Canada. There have been several different outbreaks in Canada in the past years involving a variety of foods. Current Lm detection methods are costly and time-consuming, therefore highlighting the need to improve detection strategies.

OBJECTIVE: This study aimed to develop and validate new molecular assays for rapid identification of Lm based on the presence or absence of specific biomarkers.

METHODS: Whole genome sequencing was performed on 207 Lm isolates and data was analysed using a newly developed NML-bioinformatics pipeline to identify new and unique biomarkers for incorporation into rapid real-time detection assays. Three additional biomarkers were identified from pre-existing Lm genome sequences. Loop-mediated isothermal amplification (LAMP) and probe based real-time PCR (qPCR) for each target were developed and tested for specificity and analytical sensitivity. The qPCR assays were also tested using different real-time platforms. Validation was performed on two types of surface areas that were artificially inoculated to determine the ability of the assays to detect Lm on surfaces as compared to the standard culture method.

RESULTS: LAMP and qPCR assays targeting three Lm-specific biomarkers, inlA, lmo0733 and mlbA and a single Listeria specific biomarker, pjo were developed. The assays were capable of detecting Lm with high-sensitivity and specificity from clinical, environmental and food isolates. Non-Listeria strains were not detected based on exclusivity testing. The LAMP results on all utilized real-time platforms (ABI 7500, BioRad CFX384, Qiagen RotorGene and Roche Lightcycler 96) demonstrated equivalent detection sensitivity. Both assays were able to detect Lm on artificially contaminated stainless steel and plastic surfaces with 100% concordance with culture results.

CONCLUSIONS: The newly-developed LAMP and qPCR assays accurately detect Lm in a rapid and cost effective manner. Incorporation into routine testing will improve detection strategies for Lm, while decreasing the result turnaround time to hours from days using the current standard Lm detection methods currently applied within food processing facilities.

J02 THE SPECIAL IMMUNIZATION CLINIC NETWORK TO INVESTIGATE PATIENTS WITH ADVERSE EVENTS FOLLOWING IMMUNIZATION AND POTENTIAL CONTRAINDICATIONS TO VACCINATION
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BACKGROUND: Vaccination is a safe and highly effective public health intervention. However, for patients who experienced significant adverse events following immunization (AEFI) and for those with specific underlying medical conditions, the evidence as to how best to approach vaccination or revaccination is lacking. The Special Immunization Clinic (SIC) network was established to standardize patient management, assess outcomes after vaccination or revaccination and develop a research platform. We aim to describe the first 18 months of the network’s implementation.

METHODS: A panel sample containing the tailored educational material. A panel of experts assessed the content for clarity and accuracy. The content was then refined and developed for the concept chosen, and we engaged and interacted were developed and tested in 2 focus groups. Content was developed for the concept chosen, and we engaged and interacted were developed and tested in 2 focus groups. Content was developed for the concept chosen, and we engaged and interacted were developed and tested in 2 focus groups.

RESULTS: Of 124 patients referred for the inclusion criteria, of which 81 (91%) were <18 years of age. Most were referred for prior AEFI (78/89, 88%) followed by vaccine-hesitant (47/89, 53%) and vaccine-accepting group (VHG or VAG). Focusing on differences between VAG and VHG patients, we analysed data for statistically significant differences (P<0.05) between groups for four domains: knowledge of immunization; public perception of immunization; motivators and barriers; and immunization sources and influencers for immunization decisions. Based on these results, two alternative approaches to web-based engagement and interaction were developed and tested in 2 focus groups.

CONCLUSION: ImmunizeAlberta.ca is a tailored and market-researched health-education initiative for immunization education. Our approach to its development highlights a new way to develop immunization education platforms, by harnessing appropriate web-based tools and exploiting social marketing practices to construct a targeted public health intervention.
allergic-like reactions, 8 (9%) for prior neurological symptoms and 19 (21%) for other AEFI. Eleven patients (12%) were seen for underlying conditions that complicated vaccination. To date, 44/58 (76%) patients offered revaccination have been immunized and followed up. Nine patients (20%) experienced adverse events; none were serious (resulted in hospitalization >24 h, permanent disability or death).

CONCLUSIONS: The most frequent reason for referral to a SIC are large local reactions and allergic-like events after immunization. Revaccination was safe in most patients. Through comprehensive expert assessment of patients with AEFI and contraindications to vaccination, the SIC network is helping to strengthen Canada's immunization programs.

**J03**

**USE OF WIRELESS HAND RUB DISPENSER MONITORING (eMONITORING) TO IMPROVE HAND HYGIENE ADHERENCE: A PILOT STUDY**


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**BACKGROUND:** Appropriate hand hygiene among healthcare workers is necessary for reducing healthcare associated infection. Systems providing electronic monitoring (eMonitoring) and feedback of hand hygiene adherence have the potential to support improvements in practice.

**METHODS:** We performed a time-series study of the introduction of eMonitoring (DebMed®) with near real-time feedback on three medical/surgical wards at an academic hospital. Rates of hand hygiene adherence were monitored electronically and by observational audit during the pre-intervention period (December 1 2013 to February 14 2014), intervention period (February 15 to May 31 2014), early post-intervention period (June 1 to August 31 2014), and late post-intervention period (September 1 to November 30 2014). During the intervention, units selected the method of feedback from the eMonitoring system. Hand hygiene adherence was measured with eMonitoring as a hand hygiene compliance index (HHCI), representing the number of dispenses of handrub divided by the number of expected hygiene opportunities per patient hour (percentage).

**RESULTS:** Observational audit adherence and eMonitoring HHCI across the three units for Pre/During/Early Post/Late Post-Intervention periods were: Unit 1 (85%/88%/89%/88% by audit, 24%/30%/33%/28% by eMonitor), Unit 2 (94%/98%/96%/95% by audit, 24%/36%/31%/33% by eMonitor), and Unit 3 (92%/86%/92%/94% by audit, 39%/49%/44%/44% by eMonitor). The overall eMonitoring HHCI increased from pre to during intervention periods (29% to 38%, P<0.001), decreased from during to early post-intervention periods (38% to 36%, P<0.001), and decreased from early to late post-intervention periods (36% to 35%, P<0.001). Observed audit adherence rates did not appear to significantly change from pre to during intervention periods (90% to 87%, P=0.21), from during to early post-intervention periods (87% to 89%, P=0.12), or from early to late post-intervention periods (89% to 92%, P=0.24).

**CONCLUSIONS:** eMonitoring of hand hygiene with feedback improved hand hygiene adherence and may be less susceptible to biases inherent in observational audits. Continued feedback may be necessary to sustain performance.

**J04**

**#INSTAGRAM+ID+ME: INCREASING STUDENT ENGAGEMENT WITH AN INFECTIOUS DISEASES LEARNING TOOL ON PHOTOGRAPHY BASED SOCIAL MEDIA**

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**OBJECTIVE:** We created a virtual photo album of infectious diseases (ID) memory hangers (visual mnemonics with short explanations) on Instagram©, an online, photography based social media platform, to accompany an ID course in 2013. Students who used it found the album was useful for review and the flexible format was helpful. However, student engagement was limited to less than 1/3 of the class. This year we recreated the album and tried to promote student engagement by also using relevant news stories, asking questions, and by challenging students to create their own memory hangers.

**METHODS:** Ethics approval was obtained and a private Instagram© account was created. Each memory hanger was uploaded synchronously after the corresponding face-to-face sessions. Questions and links to current news stories about the memory hangers were also uploaded over time. Instagram© allowed students to access the album from their mobile phones, receive push notifications about new postings, add comments, ‘like’ them or link out to their accounts and display their own memory hangers. The tool was then evaluated with usage analysis.

**RESULTS:** Seventy-six ID memory hangers were uploaded during the course (vs. 35 in 2013) and 8 more about ID issues in the news were uploaded after the course. 170 students (vs. 58 in 2013) followed the account. There were 157 likes on 68 different memory hangers. Based on the number of likes, “Things I am paranoid about after the course”, H. pilori, Harry Houdini and the Enteric Jazz Band and the Penicillin Pyramid were the most popular. Nine postings had multiple student comments related to questions (vs. 2 in 2013), 17 student groups created memory hangers (vs. 0 in 2013), and 2 memory hangers were created by students after the course (C. difficile and a rap about Hip Hop Haemophilus and Rhymovirus). There were 81 likes of the 8 memory hangers posted after the course and 2 had >1 comment.

**CONCLUSIONS:** Student engagement with the Instagram ID memory hangers album increased this year with our interventions. Many students have remained engaged after the completion of the course through creation of their own memory hangers or through commenting on items we have posted related to current ID events in the news.

**J05**

**COMMUNITY-BASED INFECTIOUS DISEASE CLINICS: A TOOL OF ENGAGEMENT FOR VULNERABLE POPULATIONS**

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**BACKGROUND:** As we seek to implement more effective treatment modalities for HIV and HCV infection in all target populations, novel approaches to engagement in care must be considered to ensure to maximize individual success and impact the spread of disease.

**METHODS:** We have designed and evaluated the Community Pop-Up Clinic (CPC) model on the Downtown East Side of Vancouver. The clinic is held at a community-based site on a weekly basis and accepts up to 30 patients/event. It is staffed by 6 individuals, including an infectious disease specialist. Point-of-care testing for HIV and HCV is offered, as well as review of existing infection status, and design of a plan of engagement in care. Follow-up is offered at the established clinic, as a walk-in or by appointment. Patients are also invited to fill out a questionnaire since January 2014 to record demographic information and knowledge of disease state.

**RESULTS:** Since 2013, 1025 patients have been tested during CPCs, 305 HCV positive (29.8%), 240 Male (78.7%), 22 co-infected with HIV (2.1%), 6 HIV positive (0.6%). Out of 311, 80 people showed up at least once, 7 HIV positive patients are now on antiretroviral therapy at VIDC, with 57% having an undetectable viral load. We have treated 14 patients for HCV infection, 7 with all-oral regimens. The SVR rate is 86%, with 3 more patients awaiting the initiation of HCV therapy in the coming 3 months.

**CONCLUSIONS:** The CPC model has proven highly effective for the identification and engagement in care of vulnerable individuals infected with HIV and HCV, many of whom were not engaged despite multiple previous attempts. CPC will be a powerful tool for addressing the dual HIV and HCV epidemics in Canada, with particular impact on treating those who are most likely to be core transmitters of HCV and HIV.
K01  
**PCR DETECTION OF CARBAPENEMASE-PRODUCING ORGANISMS (CPO) USING CEPHEID'S XPERT CARBA-R (XCR) VERSION 1 AND 2 ASSAYS**  
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1Mount Sinai Hospital/University Health Network, Toronto, ON; 2National Microbiology Laboratory, Winnipeg, MB; 3William Osler Health Sciences Centre, Brampton; 4University of Toronto; 5Michener Institute, Toronto; 6University of Western Ontario, London, ON  
**OBJECTIVES:** Timely detection and confirmation of CPO is integral to effective infection prevention and dissemination control critical to patient care. Most laboratories are not able to perform conventional PCR on suspected CPO. Using a comprehensive collection of retrospective clinical isolates, this study verified version 1 (v1) of the low-complexity XCR assay which claims to detect IMP1, KPC, NDM, OXA48-like and VIM genes. Additional tests were performed on a subset of CPO and non-CPO following the initial study using XCR version 2 (v2) after modifications were made to improve OXA48-like gene detection.  
**METHODS:** 228 test Gram-negative bacilli (mostly Enterobacteriaceae) included 194 CPO and 34 non-CPO isolates. Only 187 CPO were expected to be detected by XCR (100 KPC, 53 NDM, 13 OXA48, 6 VIM, 6 OXA181, 3 OXA232, 3 OXA181, 2 OXA232, 1 OXA244) as genes in 7 CPO (3 SME, 2 NMC, 1 IMP7, 1 OXA24) were not claimed, and the reduced-susceptibility to carbapenems in 32 non-CPO was due to other mechanisms [ESBL, ampicillin-resistance, ptoin mutations]. It was unclear how XCR would perform for a Shewanella (chromosomal OXA252) and an NDM-revertant. XCR was run as per Cepheid except that buffers were inoculated from lightly touched isolates and not specimens.  
**RESULTS:** XCR v1 detected 100% of KPC (95% CI: 96-100), NDM (95% CI: 93-100), VIM (95% CI: 96-100), OXA48-specific (95% CI: 73-100), OXA244 (95% CI: 22-100) CPO and the OXA232 progenitor gene. 0/14 OXA48-like genes were detected (0/9 OXA181 and 0/5 OXA232). XCR was negative for SME, NMC, OXA24 and all non-CPO and the NDM-revertant. Overall sensitivity (Sn)/specificity (Sp) for claimed genes was 92% (95% CI: 88-96)/98% (95% CI: 86->99.9). Testing on XCR v2 detected all OXA48 and OXA48-like genotypes, resulting in an overall Sn of 100% (95% CI: 98-100) for claimed CPO genes.  
**CONCLUSIONS:** The XCR v1 was simple to perform, providing highly accurate (100%) CPO results in 1h for claimed KPC, NDM, VIM and OXA48-specific genes. Modifications made in v2 enabled XCR to detect the increasingly common OXA48-like genotypes (OXA181 and OXA232) that vary from OXA48 by only minor sequence mutations. The XCR v2, currently a research-use-only assay, has been submitted to FDA for release as an IVD assay to enable use from clinical specimens for detecting the listed CPO genotypes.

K02  
**COMPARISON OF THE ABACUS DIAGNOSTICA GENOMERA CDX™ (CDX) PCR AND THE FOCUS DIAGNOSTIC SIMPLEXATM UNIVERSAL DIRECT C. DIFFICILE (CD) PCR FOR THE DIAGNOSIS OF C. DIFFICILE INFECTION (CDI)**  
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**OBJECTIVE:** The CDX direct PCR for the diagnosis of CDI is a rapid and new amplification system that detects the *C. difficile* (CD) tcdB gene. The purpose of this study was to evaluate the performance of the CDX assay compared to the Simplexa FCD assay for the direct detection of *C. difficile* in stool samples.  
**METHOD:** Glutamate Dehydrogenase (GDH) positive and negative samples were tested by PCR using the CDX and FCM assays according to manufacturer’s instructions. All samples were also cultured using the CHROMagar™ CD fluorogenic chromogenic agar (CHROM). Colonies suspicious for CD were identified using MALDI-TOF (Bruker). Cultures positive for CD with corresponding negative PCRs were confirmed to be toxigenic by testing isolates for the tcdB gene using the FCD PCR. A CD positive result was defined as a sample positive by toxigenic culture or positive by GDH and both the FCD and CDX PCRs.  
**RESULTS:** Of the 148 samples tested, 63 and 85 met the definition of a CD positive and CD negative sample respectively. Of the 63 CD positive samples, 43 (68%) were positive by culture and both the CDX and FDX assays. There were 101 samples negative by culture, 16 (16%) of these were positive by GDH, FDC and CDX PCR. We tested 34 GDH negative samples and all were negative by culture and both PCRs. The sensitivity, specificity, negative and positive predictive values of the CDX and FDX assays were 94%, 96%, 95%, 95% and 95%, 98%, 97%, 96%, respectively.  
**CONCLUSION:** The performance of both amplification methods was very similar. Toxigenic cultures using the fluorogenic CHROM agar failed to identify CD in 16% of samples positive by both PCRs and many fluoro reactive colonies did not confirm as CD. Although this selective media can be used for epidemiological purposes, the high rate of false negatives and low specificity may not be suitable for diagnostic purposes. Overall the Abacus CDX assay provides an accurate, rapid and simple to use alternative amplification method for the diagnosis of CDI.

K03  
**DETECTION OF GROUP B STREPTOCOCCI (GBS) DIRECTLY FROM E-SWABS BY ISOTHERMAL AMPLIFICATION: AN AFFORDABLE, AND RAPID POINT-OF-CARE ALTERNATIVE TO EXPENSIVE COMMERCIAL MOLECULAR DIAGNOSTIC TESTS**  
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**OBJECTIVES:** GBS is a leading cause of neonatal morbidity and mortality. It is the most common life threatening sepsis and meningitis in neonates especially the first week of life. It can be prevented if the mother diagnosed during antenatal period and treated at labor. Current guidelines recommend that all pregnant women be screened for vaginal/rectal colonization at 35-37 weeks of gestation. Rapid commercial molecular tests are too expensive to be used in GBS screening. The objective of this study was to develop an accurate method that is affordable in performance to culture but faster and cheaper. We evaluated the performance of LAMP assay to detect GBS colonization in the anogenital tract of pregnant women using E-swab specimens and compared to culture.  
**METHOD:** Randomly selected E-swab specimens from 275 women from the period January 2014 to September 2014 sent for routine screening at Hamilton Health science was used. For LAMP assay E-swab fluid was concentrated and added to a lysis solution prior to boiling for 15 min to obtain a crude extract of nucleic acids. For LAMP 210bp fragment of *cbb* gene of GBS was amplified using primers designed by Primer Explorer V4 software (Eiken Chemical Co). LAMP was carried out at 65 °C for 30 min using a Genie® II standard reaction mixture containing *Bst* 2.0 WS DNA polymerase (New England BioLabs). LAMP amplification was detected using Genie® II (OptiGene, UK). For culture 10 μL of E-swab fluid was inoculated into LIM broth and incubated overnight at 35 °C followed by plating on Chromagar for 24 hour before examining LAMP. The cost for LAMP is about Cdn$3.00 per test.

**RESULTS:** Sixty four specimens were GBS positive by both culture and LAMP. There were 4 discordant results of which 2 were culture positive and LAMP negative and two were LAMP positive and culture negative. The performance characteristics of LAMP as compared to culture were as follows: 97% sensitivity, 99% specificity, 97% PPV and 99% NPV. The turn-around-time for LAMP was 60 min as compared to 24h hours for culture. The cost for LAMP is about Cdn$3.00 per test.

**CONCLUSION:** Detection of GBS by LAMP using the amplification of *cbb* gene is rapid and cost-effective than culture with comparable performance to commercial molecular tests. This assay provides promising results for rapid detection of GBS during screening.
K04 DETECTION OF GROUP B STREPTOCOCCUS (GBS) FROM VAGINAL-RECTAL SCREENING SPECIMENS BY LOOP MEDIATED AMPLIFICATION (ILLUMIGENE® GBS) ASSAY FROM TWO BROTHS

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BACKGROUND: GBS is a common cause of early-onset neonatal sepsis. Widespread use of GBS screening and antibiotic prophylaxis has reduced the incidence of disease significantly. Technical improvements in screening may result in further reductions in disease prevalence. One method of reducing turn-around time for detection of GBS without compromising sensitivity is direct detection of GBS from enrichments broths using convenient, affordable and random-access molecular platforms. One such platform is the illumigene® GBS assay, which uses loop-mediated amplification technology to detect GBS in selective broths after overnight broth enrichment. The purpose of this study was to assess the sensitivity and specificity of GBS detection from two types of enrichment broths after overnight incubation of pre-natal vaginal-rectal swabs.

METHODS: Vaginal-rectal swabs were collected in duplicate from women attending routine pre-natal appointments between 34 and 36 weeks gestation. Swabs were placed in either Todd-Hewitt (TH) broth (Oxoid™) or RAMBAQuick(RQ) broth (CHROMagar™), which is designed to reduce overgrowth of enterococci isolated in vaginal-rectal samples. Broths were incubated overnight at 35°C. GBS was detected using the illumigene® GBS assay following manufacturer's instructions (off-label use for RQ broth). was Broths were sub-cultured to sheep blood agar (SBA) and Colorex™ StrepB (CSB, CHROMagar™). GBS were identified using colony morphology and Lancefield typing from SBA and by chromogenic reaction and Lancefield typing from CSB. GBS growth from the broth on either SBA or CSB was considered the gold standard.

RESULTS: 45 of 204 samples were positive for GBS in TH broth by culture and 44 were positive in RQ broth by culture. Sensitivity of illumigene® was 97.7% (43/44) from the RQ broth and 100% (45/45) from the TH broth. Specificity of the illumigene® assay was 100% for both broths and an invalid illumigene® result occurred in 1 (0.2%) test.

CONCLUSION: illumigene® performed equally well from both broths and had very high sensitivity and specificity. The illumigene® assay was easy to perform with minimal training or expertise and provided rapid results without the need for colony picking or additional testing (e.g. Lancefield typing). Turnaround time was reduced by one day compared to conventional broth-enrichment culture.

K05 COMPARATIVE EVALUATIONS OF COMMERCIAL AVAILABLE RAPID PHENOTYPIC TESTS FOR DETECTING CARBAPENEMASE-PRODUCING ORGANISMS (CPO)

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OBJECTIVES: While molecular tests cannot target every CPO, the phenotypic Carb-NC (CPN) claims to rapidly distinguish all CPO from non-CPO. But many studies have found the CPN insensitive for OXA48-like CPO. This study compared capabilities of similar CPO detection tests from Rosco [Carba Blue (CBL); Rapid Carba Screen (RCS); modified RCS (miRCS)] and bioMérieux [Rapidec Carba-NC (RCN)] using characterized clinical strains.

METHODS: 206 Gram-negative bacilli (mostly enterobacteriaceae) were randomized for testing. These included 175 CPO [91 KPC, 49 NDM, 10 OXA48, 6 OXA181, 6 VIM, 3 OXA232, 2 NDM+OXA181, 2 NMC, 2 SME, 1 IMP, 1 NDM+OXA232, 1 OXA24, 1 OXA244] and 31 non-CPO (variously carbapenem-insusceptible ESBL, AmpC or OMP-mutants). RCN, RCL and RSC were tested and interpreted as per manufacturer while miRCS used 0.8% saline (pH 8.0) in lieu of extraction buffer. Compared to controls, positive endpoints at 15, 30, 60 and 120 min of yellow or orange (Y/O), or Y/O/red-orange (Y/O/RO) for RCN, RCS, and miRCS and Y/green (Y/G) or Y/G/blue-green (Y/G/BG) for CBL, respectively, were noted.

RESULTS: The RCN detected 100% non-OXA CPO but only with criteria of Y/O/RO at 120 min. Using these criteria, 77% OXA CPO went undetected and 12.9% of non-CPO were erroneously pos. If an RCN endpoint of Y/O at 120 min was applied, detection of non-OXA CPO was reduced to 97.4% and OXA CPO to 8% but 0% false-poss were incurred. Using an endpoint of Y/O at 120 min, the miRCS outperformed RCS, detecting 99.4% vs 79.9% non-OXA CPO and 42.9% vs 10.5% OXA, respectively, with no false-pos calls for non-CPO strains. For CBL, if an endpoint of Y/G at 120 min was used, non-OXA CPO detection was 98.7% and OXA-CPO detection was 42.9%, but 9.7% non-CPO were also incorrectly called pos. Including Y/G/BG endpoints at 30, 60 or 120 min increased OXA-CPO detection to 71.4%, 66.6% and 66.6%, respectively, but also substantially increased the false-pos rates to 51.6%, 48.4% and 35.5%, respectively.

CONCLUSIONS: No assay detected all CPO despite modifications or consideration of various pos colour endpoints. While these assays are valuable when definite colour changes are obtained, laboratories must be aware that none work alone to detect all CPO and most have associated false-poss, especially when weaker colour-changes are considered pos.

L01 ANTIBIOTIC RESISTANCE SURVEILLANCE FOR THE COMMUNITY: CLOSING THE GAP THROUGH PUBLIC-PRIVATE COLLABORATION IN BC

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OBJECTIVES: A review of antibiotic resistant organisms (ARO) surveillance in Canada revealed a gap outside of the hospital setting. We hypothesized that, where a large proportion of susceptibility testing is concentrated in a single laboratory network, collaboration between the providing laboratory and surveillance units could fill the void.

METHODS: LifeLabs collects specimens from 80 community-based patient service centres located throughout British Columbia (BC) representing approximately 80% of community-based testing and is also well represented in other provinces. Steps in collaboration with the BC Do Bugs Need Drugs program at BCCDC included proposal review, datasharing agreement, extension of a three-party non-disclosure agreement to include work with the Public Health Agency of Canada (PHAC), data transfer to a secure server, analysis and collaborative report production involving surveillance staff and LifeLabs microbiologists.

RESULTS: The entire process took four months, with about 80% of the time being required to finalize agreements. Data from 2007-2013 were provided and included 744,155 records from two information systems, 121,732 records for 2013 alone. Data structure included age-group, sex, local health area, date of collection, anatomical site, organism and variables for each antibiotic of interest. Extraction from a reporting database meant some limitation in organism-drug combinations and a need to assure denominators were accurately accrued. The process allowed reporting for 14 organisms of interest, detailed analysis of trends by age-group, gender and region within the province and provided very large sample sizes for robust confidence intervals. The process appears sustainable through twice-yearly data extraction.

CONCLUSIONS: The public private collaboration model makes use of existing data, avoids the costs of de novo surveillance and is easily achieved when supported by laboratory leadership. Its application to other regions
will be explored with PHAC. In-depth analysis (e.g. by age, region and over time) can provide professionals with new tools to combat AROs.

L02
ANTIFUNGAL SUSCEPTIBILITY OF RESPIRATORY ASPERGILLUS ISOLATES FROM CANADIAN HOSPITALS: RESULTS OF THE CANWARD 2014 STUDY
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OBJECTIVES: CANWARD is an ongoing national surveillance study that assesses pathogens causing infections in patients attending Canadian hospitals, as well as determines the prevalence of antimicrobial resistance in these isolates. Here we present the distribution of species and in vitro antifungal susceptibility of Aspergillus isolates collected in 2014 from patients visiting Canadian hospitals.

METHODS: Clinical Aspergillus isolates recovered from respiratory specimens at 13 participating medical centres during the 2014 study period were tested against amphotericin B (AMB), itraconazole (ITRA), posaconazole (POSA), voriconazole (VORI) and caspofungin (CASP) by broth microdilution using the CLSI M38-A2 method. Growth endpoints were measured as per CLSI M38-A2 and values above the published epidemiological cutoff values (mg/L) were scored as non-wildtype (non-WT). Clinical breakpoints are not available for Aspergillus susceptibility interpretation.

RESULTS: Of the 713 Aspergillus isolates recovered, A. fumigatus, A. niger, A. flavus, and A. terreus represented 71.9%, 11.2%, 6.5%, and 3.9% of the population, respectively. A. fumigatus isolates were recovered primarily from Clinic outpatients (68%) and Medicine inpatients (19%) from sputa (62%) and bronchoscopy (30%) specimens. Very few Aspergillus isolates exhibited non-WT MIC and MEC values. Twenty isolates (4%) of A. fumigatus had non-WT VORI MICs (2 to 4 mg/L) and 5% of A. niger had non-WT MICs to VORI, ITRA, or POSA. There was very little evidence of microbiological resistance to echinocandins across the species. Six A. caldoidus isolates from six centres were recovered, all exhibiting non-WT MICs to the azoles and variably high MECs to caspofungin.

CONCLUSION: Epidemiological cutoff values have been published to aid the detection of acquired microbiological resistance in Aspergillus. In this study, WT isolates of A. fumigatus were most prevalent and the rate of non-WT isolates was very low and comparable to rates from the previous two years of surveillance. Intrinsically resistant species like A. caldoidus remain very uncommon in respiratory cultures.

L03
ANTIMICROBIAL RESISTANCE IN PATHOGENS ISOLATED FROM CANADIAN HOSPITALS: CANWARD 2014
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OBJECTIVES: The CANWARD study assesses the pathogens causing infections in patients affiliated with Canadian hospitals, and evaluates the prevalence of antimicrobial resistance in these isolates.

METHODS: Thirty tertiary-care centres across Canada submitted pathogens causing infections from patients attending clinics (C), emergency rooms (ER), medical and surgical wards (W) and intensive care units (ICU) in 2014. Susceptibility testing was performed by CLSI broth microdilution methods.

RESULTS: A total of 3,196 isolates were collected: 40.0%, 40.2%, 10.1%, and 9.7% from blood, respiratory, urine and wound/IV site specimens, respectively. Patient demographics were as follows: 56.4/43.6% male/female, 13.8% ≤17 years, 42.3% 18-64 years and 43.9% ≥65 years. Isolates were from patients on W 39.5%, ER 23.3%, ICU 18.8%, and C 18.4%. The most common pathogens were: E. coli 19.6%, S. aureus (MSSA) 19.5%, P. aeruginosa 11.0%, K. pneumoniae 5.8%, S. aureus (MRSA) 5.0%, and S. pneumoniae 4.9%. Resistance rates for E. coli were: 0% for meropenem and ticarcillin, 0.3% ertapenem, 1.6% piperacillin/tazobactam, 9.6% gentamicin, 12.0% ceftriaxone, 24.4% ciprofloxacin and 28.8% trimethoprim/sulfamethoxazole (SXT). For P. aeruginosa, resistance rates were 1.1% colistin, 7.1% gentamicin, 8.6% piperacillin-tazobactam, 16.6% meropenem, and 17.7% ciprofloxacin. Resistance rates for MRSA were: 0% vancomycin, daptomycin and linezolid, 5.0% tigecycline, 5.0% SXT, 38.7% clindamycin, 80.0% ciprofloxacin, and 82.5% clarithromycin. Overall, the prevalence of MRSA and vancomycin-resistant enterococci (VRE) was 20.5% and 7.6%, respectively.

CONCLUSIONS: In Canada, resistance rates for E. coli remain lowest for meropenem, tigecycline and piperacillin-tazobactam, while for P. aeruginosa, rates are lowest with colistin, gentamicin, and piperacillin-tazobactam. No resistance was observed in MRSA with vancomycin, linezolid, or daptomycin.

L04
ANTIMICROBIAL RESISTANT ORGANISMS IN CANADA: JANUARY 1, 2009 TO DECEMBER 31, 2013
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BACKGROUND: Surveillance of antimicrobial resistant organisms (AROs) in hospitalized patients is vital to informing strategies to reduce the risks and impacts of antimicrobial resistance.

METHODS: The Public Health Agency of Canada collects national data on AROs in hospitalized patients through the Canadian Nosocomial Infection Surveillance Program (CNISP). As of December 2013, CNISP conducted surveillance in 59 major acute-care hospitals in Canada.

RESULTS: Canadian healthcare-associated Clostridium difficile infection (HA-CDI) rates remained stable from 2009 (5.81 per 10,000 patient days) to 2013 (5.36 per 10,000 patient-days). HA-CDI attributable mortality rates peaked in 2011 (6.4%) and fell by 52% in 2013 (3.1%). National methicillin-resistant Staphylococcus aureus (MRSA) infection rates have decreased since 2009, with the most dramatic reduction seen in healthcare-associated rates (2.56 to 1.77 per 10,000 patient days). Vancomycin-resistant enterococci (VRE) infection rates in Canada remain low but have been slowly increasing since 2009 (0.32 to 0.54 per 10,000 patient days). Regional variations exist with the highest rates in the western region (0.79 per 10,000 patient days) and lowest in the east (0.08 per 10,000 patient days). Rates of carbapenemase-producing organisms (CPO) and carbapenem-resistant enterobacteriaceae (CRE) in CNISP participating hospitals remain low. The central region of Canada reported the highest number of CRE and CPO cases, the majority of which are likely due to an outbreak at one hospital.

CONCLUSIONS: In Canada, ARO rates among hospitalized patients have been relatively stable since 2009. Regional variations have been reported and may be partially explained by differences in the virulence of circulating strains, infection control practices, antimicrobial stewardship programs and surveillance and screening activities. These data highlight the importance of continuing to monitor trends to ensure the detection of new and re-emerging threats associated with antimicrobial resistance.

L05
ANTIFUNGAL SUSCEPTIBILITY OF INVASIVE CANDIDA ISOLATES FROM CANADIAN HOSPITALS: RESULTS OF THE CANWARD 2014 STUDY
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OBJECTIVE: CANWARD is an ongoing national surveillance study that assesses pathogens causing infections in patients admitted to Canadian hospitals, as well as determines the prevalence of antimicrobial resistance (R) in these isolates. Here we present the antifungal susceptibility data for candidemia isolates collected in 2014.

METHODS: Candida species isolated from bloodstream infections were collected from 12 participating medical centres during the 2014 study

Abstracts
period. Antifungal susceptibility testing and interpretation was performed as per CLSI M27-S4 broth microdilution method and interpretation guidelines for fluconazole (FLUC), caspofungin (CASP), and micafungin (MICF). Epidemiological cut-off values of \( \leq 2\) mg/L for amphotericin B (AMB) and 0.5 mg/L for voriconazole (VORI) against C. glabrata (CG) were used in the absence of M27 breakpoints.

**RESULTS:** Of 328 Candida spp. collected, C. albicans (CA) was predominant (48.8%), followed by C. glabrata (CG, 22.8%), C. parapsilosis (CP, 9.8%), and C. tropicalis (6.1%). The majority of cases were identified in ICU (31.4%), medicine (36.3%), and surgical wards (15.6%). Susceptibility (S) values are in Table 1 below and limited acquired resistance was detected. High level azole R was detected in one CA and two CP isolates. Echinocandin R was detected in four CG isolates (5.6%), all of which contained \( \beta \) mutations.

**CONCLUSION:** CANWARD surveillance of invasive Candida isolates since 2011 shows stable distribution of species and antifungal MICs in hospitals patients, and limited evidence of resistance. C. glabrata is a common cause of candidemia and echinocandin resistance of ~5% in Canada is associated with glucon synthase mutations.

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**POSTER PRESENTATIONS**

**P01**

**ANTIPROTOZOAL POTENTIAL OF PHENOLIC COMPOUNDS FROM KLEINIA ODORA**

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Today, over one billion people worldwide are at risk of tropical diseases caused by parasitic protozoa. Several antiparasitic drugs have been derived directly from natural sources, such as quinine, artemisinin and amphoterin B. In this study phenolic constituents of *Kleinia odora* growing in Saudi Arabia were isolated and assessed for antiprotozoal activity. They were screened in vitro against erythrocytic schizonts of *Plasmodium falciparum*, intracellular amastigotes of *Leishmania infantum*, *Trypanosoma cruzi* and free trypomastigotes of *T. brucei*. Cytotoxic activity was determined against MRC-5 cells to assess selectivity. The compounds were identified as dihydrodehydrodiconiferyl alcohol (1), (7\(^,8\))-1-(3,4-dimethoxyphenyl)-2-(2-methoxy-4-omegahydroxypropylphenyl) propane-1,3-diol (2), (7\(^,8\))-(2-methoxy-4-O-methyl-3,4-dimethoxyphenyl)-1-(2- methoxy-4-omegahydroxypropylphenyl) propane-1,3-diol (3) caffeic acid (4), (S)-1-(4-hydroxy-3-methoxyphenyl)-2-(2-methoxy-4-omegahydroxypropylphenyl) propane-1,3-diol (5) and (3\(^,\))-(2-methoxy-4-O-methyl-3,4-dimethoxyphenyl)-1-(2- methoxy-4-omegahydroxypropylphenyl) propane-1,3-diol (6).

**P02**

**IN-VITRO ANTIVIRAL ACTION OF AMYLMETACRESOL (AMC) AND 2,4-DICHLOROBENZYL ALCOHOL (BCDA) AGAINST THE RESPIRATORY PATHOGEN PARAINFLUENZA TYPE 3**

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**OBJECTIVES:** Antibiotic resistance continues to be a global threat and is in part fuelled by their inappropriate use for primarily viral respiratory tract infections. With viral infections accounting for 80-90% of all sore throats and a wide variety of potential virus types involved, there is a need for better treatment options for this common condition.

**METHODS:** Parainfluenza Type 3 is a common respiratory pathogen which can cause sore throat. The ASTN international standard method E152-11 for testing microbicides against viruses in suspension was used to evaluate the effect of AMC and DCBA in combination and a dissolved lozenge containing both AMC and DCBA against parainfluenza type 3 virus under in-vitro conditions.

**RESULTS:** 0.6 mg AMC and 1.2 mg DCBA in 5 ml of artificial saliva exhibited a 4.18 – 4.56 log\( \text{reduction in viral titre at 1 minute. The lozone containing 6.6 mg of AMC and 12.4 mg of DCBA dissolved in 5 ml of artificial saliva exhibited a 2.43 – 2.68 log\( \text{reduction in viral titre at 1 minute. A placebo lozenge without AMC and DCBA exhibited a 0 – 0.43 log\( \text{reduction in viral titre. CONCLUSIONS:** AMC and DCBA as free active substances and when incorporated into a lozenge delivery format are able to rapidly inactivate parainfluenza virus type 3 under in-vitro conditions. AMC and DCBA have already been demonstrated to inactivate other enveloped respiratory viruses including influenza, respiratory syncytial virus and coronavirus making them a potential candidate for further clinical investigation as an alternative to antibiotics in controlling viral respiratory tract infections.**
might be an ultra metabolizer of voriconazole. His respiratory status worsened and chest CT showed worsening opacities. He improved and was switched to oral voriconazole. A week later, a 48-year-old male, allogeneic hematopoietic stem cell transplant recipient with extensive graft versus host disease was admitted to medical center, Chicago, IL, USA. He was a 48-year-old male, allogeneic hematopoietic stem cell transplant recipient with extensive graft versus host disease who was admitted to medical center, Chicago, IL, USA. He was a 48-year-old male, allogeneic hematopoietic stem cell transplant recipient with extensive graft versus host disease. He was discharged home on micafungin with plans to switch to liposomal amphotericin B if there was any worsening.

RESULTS: Genetic polymorphism of CYP2C19 modulates enzyme activity and significantly different plasma concentrations are observed despite identical dosing schedules. Studies reveal polymorphisms to contribute to pharmacokinetic variability. Individuals may be categorized as ultra rapid metabolizers (UM) (CYP2C19*1/*17 CYP2C19*17*17), extensive metabolizers (EM) (CYP2C19*1/*1); intermediate metabolizer (IM) (CYP2C19*1/*2 and CYP2C19*1/*3) and poor metabolizers (PM) (CYP2C19*2/*2, CYP2C19*2/*3 and CYP2C19*3/*3)(5). This study highlighted that voriconazole is influenced by CYP2C19 polymorphism and gene based adjusted dosing is important to achieve therapeutic drug levels.

CONCLUSIONS: Clinicians should keep CYP2C19 polymorphisms in the differential when evaluating very low voriconazole levels for treatment of invasive fungal infections.

P06 CYP2C19*17 GENETIC POLYMORPHISM AND THERAPEUTIC CHALLENGES IN AN ALLOGENEIC STEM CELL TRANSPLANT RECIPIENT WITH INVASIVE PULMONARY ASPERGILLOSIS

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BACKGROUND: A 48-year-old male, allogeneic hematopoietic stem cell transplant recipient with extensive graft versus host disease was admitted with pulmonary aspergillosis 6 months after transplant. Chest computerized tomography (CT) scan showed bilateral lung opacities and he required intubation. Antimicrobials included intravenous cefepime, azithromycin and voriconazole. Bronchoalveolar lavage (BAL) fluid cultures grew 10,000 cfu/mL Pseudomonas aeruginosa and 1,200 cfu/mL Aspergillus fumigatus. Nucleic acid amplification testing was positive for Mycoplasma pneumoniae. He improved and was switched to oral voriconazole. A week later, his respiratory status worsened and chest CT showed worsening opacities. Voriconazole serum levels were undetectable twice. We suspected that he might be an ultra metabolizer of voriconazole.

METHODS: Cytochrome assay was requested. The methodology employed was DNA analysis of the cytochrome P450 2C19 gene including the alleles for poor metabolizers *1, *2, *3, *4, *5, *6, *7 and *8 as well as the ultra-metabolizer alleles *17. We added micafungin and one week later imaging showed improvement. CYP2C19 genotype testing was positive for the *17 translocation indicating he was an ultra metabolizer for voriconazole. He was discharged home on micafungin with plans to switch to liposomal amphotericin B if there was any worsening.

RESULTS: Genetic polymorphism of CYP2C19 modulates enzyme activity and significantly different plasma concentrations are observed despite identical dosing schedules. Studies reveal polymorphisms to contribute to pharmacokinetic variability. Individuals may be categorized as ultra rapid metabolizers (UM) (CYP2C19*1/*17 CYP2C19*17*17), extensive metabolizers (EM) (CYP2C19*1/*1); intermediate metabolizer (IM) (CYP2C19*1/*2 and CYP2C19*1/*3) and poor metabolizers (PM) (CYP2C19*2/*2, CYP2C19*2/*3 and CYP2C19*3/*3)(5). This study highlighted that voriconazole is influenced by CYP2C19 polymorphism and gene based adjusted dosing is important to achieve therapeutic drug levels.

CONCLUSIONS: Clinicians should keep CYP2C19 polymorphisms in the differential when evaluating very low voriconazole levels for treatment of invasive fungal infections.

P07 NANO PARTICLES TO TARGET HIV RESERVOIRS

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BACKGROUND: Although the current antiretroviral drugs (ARVs) can effectively inhibit replication of HIV-1 in infected patients, the virus is able to persist in cellular and anatomical viral reservoirs leading to latent re-infection and development of drug resistance. Macrophages constitute one of the most important viral reservoirs and are able to transport HIV into other reservoir sites such as the lymphoid organs and the central nervous system (CNS). ARVs hardly penetrate into the CNS and macrophages therefore, it is highly critical to improve drug delivery in these compartments.

OBJECTIVE: Here we evaluated the lipid-based nanosystem Neutraplex (Nx) for the transport of therapeutic molecules into macrophages and in the brain to target HIV viral reservoirs.

METHODS: We have determined the cytotoxicity profile of the Neutraplex nanosystem using different cell models (TZM-bl, THP-1, Be(2)-M17) and various cytotoxicity assays. Cellular uptake was evaluated by confocal microscopy. In addition, Nx capability to cross the blood-brain barrier (BBB) was investigated using the immortalized adult rat brain microvascular endothelial cell model (SV-ARBEC) and effect on inflammatory response was evaluated using an in vitro neutrophil apoptosis assay.

RESULTS: Nx nanoparticles (NP) were not found cytotoxic up to the highest dose tested, 35 µg/mL. Confocal microscopy studies showed that Nx NPs are rapidly and efficiently taken up by THP-1 macrophages and were still abundantly found in the cells 48 h following exposure, at doses as low as 4 µg/mL. In addition, Nx NPs were able to transmigrate the endothelial cell monolayer and were not found cytotoxic for neuronal cells, suggesting that they have the capability to deliver drugs through the BBB without affecting neuronal viability. Finally, Nx NPs did not induce apoptosis of polymorphonuclear neutrophils indicating their low interference with inflammatory response.

CONCLUSION: Altogether these results indicate that the Neutraplex nanocarrier shows potential as a delivery strategy aiming to target HIV in cellular and anatomical viral reservoirs.

P05 NOVEL ISO STEVIOIL ISOLATED FROM PITTOSPORUM TETRASPERMUM EXHIBITED ANTIMICROBIAL, ANTIBIOFILM AND ANTICANCER ACTIVITIES

VA Mariadhas, NA Al-Dhabi

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Drugs derived from medicinal plants as green medicine in recent years has attracted many researchers because of its widespread applications in curing infectious diseases, compared with costly synthetic drugs that have adverse effects. Aim of this study was to investigate the in vitro antimicrobial, antibiofilm and anticancer properties of isosteviol isolated from endangered medicinal plant Pitto sporum tetraspermum. Pure compound was obtained and characterized by column chromatography followed by 1H NMR, 13C NMR, IR and Mass spectral analysis. The antibacterial and antifungal activities of the compound were assessed by the broth micro dilution method.

Anticancer study was evaluated by following MTT assay. Primary screening revealed that the organic extract of the compound inhibited the growth of microbial pathogens. Further, column purification and spectroscopical analysis guided to identify novel isosteviol from the crude ethyl acetate extract. The compound exhibited significant activity against bacteria such as Staphylococcus epidermidis (125 µg/mL), Staphylococcus aureus (125 µg/mL) and Klebsiella pneumoniae (62.5 µg/mL). The MIC of the compound against Candida albicans, Aspergillus niger and Trichophyton mentagrophytes were 62.5, 125 and 500 µg/mL respectively. The compound showed comparatively better antibiofilm activity against the uropathogens such as E. coli, S. typhi and P. aeruginosa. Furthermore, it exhibited good anticancer properties against Vero and MCF7 cell lines. Novel isosteviol would be useful to reduce the infectious diseases caused by pathogenic bacteria and fungi or slowing the progress of various oxidative stress-related diseases.
INCREASED RECOVERY AND DETECTION OF PATHOGENS USING COPAN’S WASPLAB™ TOTAL LABORATORY AUTOMATION (TLA)

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BACKGROUND: The objective of this study was to observe the effects of using Copan’s WASPLab™ Total Laboratory Automation (TLA) on the recovery and detection of pathogens from clinical specimens.

METHODS: A wide range of specimens were processed on TLA using a customized automation platform (inoculation, incubation in Smart Incubators, and digital image reporting). Consecutive specimens were processed in real-time simultaneously and blinded on TLA and manually according to standard operating procedures. Analytical and post-analytical processes followed incubation of plates in the Smart Incubators and manually in off-line incubators. Identification of pathogens was achieved using a combination of mass spectrometry, Vitek2, chromogenic media, and standard biochemical tests.

RESULTS: A total of 874 specimens were evaluated in parallel comparing processes followed incubation of plates in the Smart Incubators and manually with CHROMagar, 16% (n=43) of specimens had positive cultures (≥A total of 874 specimens were evaluated in parallel comparing processes followed incubation of plates in the Smart Incubators and manually with CHROMagar, 16% (n=43) of specimens had positive cultures (≥

P09 USEFULNESS OF PREVIOUS METHICILLIN-RESISTANT STAPHYLOCOCCUS AUREUS SCREENING RESULTS IN GUIDING EMPIRIC THERAPY FOR S. AUREUS BACTEREMIA

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1University of Ottawa, Ottawa; 2Mount Sinai Hospital; 3Leslie Dan Faculty of Pharmacy; 4University of Toronto, Toronto; 5Lakeridge Health, Oshawa; 6Queen’s University, Kingston; 7Trillium Health Partners, Mississauga; 8North York General Hospital; 9University Health Network; 10Institute for Clinical Evaluative Sciences, Toronto, ON

OBJECTIVES: Staphylococcus aureus bacteremia (SAB) is an important infection. Methicillin-resistant Staphylococcus aureus (MRSA) screening is performed in hospitalized patients to guide infectious disease control practices. We conducted a retrospective cohort study to assess the usefulness of past MRSA screening in guiding empiric antibiotic therapy for SAB.

METHODS: A retrospective cohort study examined consecutive patients with confirmed SAB and prior MRSA swab screening from 6 academic and community hospitals between 2007 and 2010. Diagnostic test properties were calculated for MRSA swab screening in predicting methicillin susceptibility of SAB.

RESULTS: In the study, 799 patients had MRSA screening swabs prior to SAB. Of the 799 patients, 95 (12%) patients had a positive and 704 (88%) had a negative prior MRSA screening swab. There were 150 (19%) patients with MRSA bacteremia. Overall, prior MRSA screening swab had a positive likelihood ratio of 33 (CI: 18-60) and negative likelihood ratio of 0.45 (CI: 0.37-0.54). Diagnostic accuracy differed depending on mode of acquisition (i.e. community-acquired, nosocomial, or healthcare-associated infection) (P<0.0001) and hospital (P=0.0002). Mathematical modeling of this data demonstrates that the post-test probability of MRSA for a negative MRSA screening result is close to or below 10% when MRSA prevalence in SAB is less than 25%.

CONCLUSIONS: MRSA screening swabs can be helpful in guiding empiric antibiotic therapy for SAB in areas with moderate MRSA prevalence.

P10 INADEQUATE SYMPTOM DOCUMENTATION OF URINARY TRACT INFECTION IN THE MEDICAL CHART

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OBJECTIVES: Small studies suggest clinicians inconsistently document symptoms in the medical chart for patients suspected of having urinary tract infection (UTI). Our goal was to prospectively evaluate the completeness of clinician documentation, and to compare it with data obtained from a prospective interview with the medical team caring for the patient.

METHODS: Consecutive inpatients at two academic, tertiary care hospitals with positive urine cultures from January 30 to April 17, 2012 were included. Urinary symptoms were documented prospectively by interviewing physicians and nurses caring for the patient and compared to an independent retrospective review of the patient’s chart upon discharge.

RESULTS: During the study period, 341 patients were eligible and 312 were included for analysis (29 were excluded due to inability to interview the medical team caring for the patient). Amongst these patients, 174/312 (56%) had symptomatic UTI when combining information from the interviews and charts, which we considered our gold standard (Table).

CONCLUSIONS: Due to insufficient documentation of symptoms, a significant number of patients with UTI would be misclassified as asymptomatic bacteriuria with retrospective chart review alone. Infection surveillance or audits for asymptomatic bacteriuria should incorporate prospective data collection from health care providers in addition to chart review to avoid underestimating the number of symptomatic patients.
**Abstracts**

**P11**
**ALL-ORAL 12-WEEK COMBINATION TREATMENT WITH DACLATASVIR (DCV) AND SOFOSBUVIR (SOF) IN PATIENTS INFECTED WITH HCV GENOTYPE (GT) 3: ALLY-3 PHASE 3 STUDY**

DR Nelson1, JN Cooper2, JP Lalezari3, E Lawitz4, PJ Pockros5, N Gitlin6, BF Freilich7, ZH Younes8, W Harlan9, R Ghalib10, G Oguchi11, M Bennett12, T Hawkins17, N Ravendhran18, AM Sheikh19, church, vA; 3quest Clinical Research, San Francisco, cA; 4Texas Clinical Research Institute, Arlington, TX; 10Midland Florida Clinical Research Center, DeLand, FL; 15Hofstra North Shore-Long Island Jewish School of Medicine, Manhasset, NY; 16medical Associates Research Group, San Diego, CA; 17Southwest CARE Center, Santa Fe, NM; 19Digestive Disease Associates, Atlanta, GA; 18Digestive Disease Associates Research Center, DeLand, FL; 21Swedish Medical Center, Seattle, WA; 22bristol-Myers Squibb, Princeton, NJ, USA

**BACKGROUND:** Options for treating HCV GT 3 infection are limited in both treatment-naive and -experienced patients; the only currently available all-oral regimen requires 24-week treatment that includes ribavirin (RBV). Newer combinations without RBV are being studied to shorten treatment duration. The efficacy and safety of the combination of DCV (potent, pangenotypic NS5A inhibitor) and SOF (NS5B polymerase inhibitor) for 12 weeks were evaluated in patients chronically infected with GT 3.

**METHODS:** Two cohorts consisting of treatment-naive or -experienced patients (prior treatment failures, including prior SOF- or asunaprevir-treated) patients received open-label DCV 60 mg + SOF 400 mg QD for 12 weeks. Efficacy (sustained virologic response at posttreatment Week 4 [SVR4]) and safety outcomes are reported. SVR12, the primary endpoint, will be available for presentation.

**RESULTS:** 152 patients were treated: 101 (66%) treatment-naive and 51 (34%) treatment-experienced; 21% were cirrhotic, 61% non-CC IL28B genotype, 71% HCV RNA ≥800K IU/mL. Baseline characteristics were comparable between the 2 cohorts except for a higher proportion of cirrhotic patients in the treatment-experienced cohort. Overall, 91% and 86% of treatment-naive and -experienced patients, respectively, achieved SVR4; SVR4 rates were 94% for patients with a META VIR category of F0-F3 and 70% for those with F4. One patient had detectable HCV RNA at the end of treatment and 15 had relapse posttreatment (mostly cirrhotic patients). One serious adverse event was reported on-treatment: a gastrointestinal bleed considered not related to study treatment. There were 7 serious adverse events, all considered unrelated to study treatment, and 3 (<1%) adverse events leading to treatment discontinuation. The most common adverse events (in >10% of patients) were headache, fatigue, diarrhea, and nausea.

**CONCLUSIONS:** In this Phase 3 study of 415 patients, 12 weeks of all-oral treatment with DCV/ASV/BMS-791325 FDC achieved high SVR12 rates in patients with chronic HCV GT 1 infection and was well tolerated. These findings demonstrate the potent antiviral activity, safety, and tolerability of the DCV 3DAA regimen in treatment-naive and treatment-experienced patients without cirrhosis.

<table>
<thead>
<tr>
<th>Virologic outcomes, n (%)</th>
<th>Treatment-naive (N=312)</th>
<th>Treatment-experienced (N=103)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SVR12</td>
<td>287 (92)</td>
<td>92 (89)</td>
</tr>
<tr>
<td>Posttreatment relapse</td>
<td>15 (5)</td>
<td>6 (6)</td>
</tr>
<tr>
<td>On-treatment failure</td>
<td>9 (3)</td>
<td>4 (4)</td>
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<tr>
<td>Missing data</td>
<td>1 (&lt;1)</td>
<td>1 (&lt;1)</td>
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</table>

**P12**
**ALL-ORAL, FIXED-DOSE COMBINATION THERAPY WITH DACLATASVIR/ASUNAPREVIR/BMS-791325 FOR NON-CIRRHOTIC PATIENTS WITH CHRONIC HCV GENOTYPE 1 INFECTION: UNITY-1 PHASE 3 SVR12 RESULTS**

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**INTRODUCTION:** The all-oral combination of daclatasvir (DCV; pangenotypic NS5A inhibitor), asunaprevir (ASV; NS3 protease inhibitor), and BMS-791325 (non-nucleoside NS5B inhibitor) – DCV 3DAA regimen – was evaluated without ribavirin in HCV genotype (GT) 1-infected treatment-naive and -experienced patients without cirrhosis in a Phase 3, open-label, international clinical trial.

**METHODS:** Patients received a fixed-dose combination (FDC) of DCV 30 mg, ASV 200 mg, and BMS-791325 75 mg twice daily for 12 weeks. SVR12 rates in the treatment-naive and -experienced cohorts were evaluated separately as key efficacy outcomes.

**RESULTS:** SVR12 was achieved by 92% of treatment-naive patients (Table). Among treatment-experienced patients, 89% achieved SVR12. Virologic failure occurred in 34 (8%) patients overall. Baseline characteristics were comparable between the treatment-naive (N=312) and treatment-experienced (N=103) cohorts. Overall, patients were 58% male and 26% IL28B (rs1297980) CC genotype; 73% were infected with GT 1a and 27% with GT 1b. One death reported posttreatment was considered not related to study treatment. There were 7 serious adverse events, all considered unrelated to study treatment, and 3 (<1%) adverse events leading to treatment discontinuation. The most common adverse events (in >10% of patients) were headache, fatigue, diarrhea, and nausea.

**CONCLUSIONS:** In this Phase 3 study of 415 patients, 12 weeks of all-oral treatment with DCV/ASV/BMS-791325 FDC achieved high SVR12 rates in patients with chronic HCV GT 1 infection and was well tolerated. These findings demonstrate the potent antiviral activity, safety, and tolerability of the DCV 3DAA regimen in treatment-naive and treatment-experienced patients without cirrhosis.
P13
ALL-ORAL FIXED-DOSE COMBINATION THERAPY WITH DACLATASVIR/ASUNAPREVIR/BMS-791325, + RIBAVIRIN, FOR PATIENTS WITH CHRONIC HCV GENOTYPE 1 INFECTION AND COMPENSATED CIRRHOSIS: UNITY-2 PHASE 3 SVR12 RESULTS

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INTRODUCTION: The all-oral combination of daclatasvir (DCV; pangenotypic NS5A inhibitor), asunaprevir (ASV; NS3 protease inhibitor), and BMS-791325 (325; non-nucleoside NS5B inhibitor) – DCV 3DAA regimen – was studied with and without ribavirin (RBV) in treatment-naïve and treatment-experienced patients with HCV genotype (GT) 1 infection and compensated cirrhosis in a Phase 3, international clinical trial.

METHODS: Patients were randomly assigned to receive a fixed-dose combination (FDC) of DCV 30 mg, ASV 200 mg, and ‘325 75 mg, with blinded RBV or placebo, twice-daily for 12 weeks. SVR12 rates in the treatment-naïve and experienced cohorts were evaluated separately as key efficacy outcomes.

RESULTS: SVR12 results in treatment-naïve - and -experienced cirrhotic patients are in the table below. Virologic failure was observed in 13 (6%) patients. Baseline characteristics were comparable between treatment-naïve (N=112) and treatment-experienced (N=90) groups. Overall, patients were 66% male and 27% IL28B CC genotype; 74% of patients had GT1a infection and 26% had GT1b. There were 3 serious adverse events (SAEs) considered related to treatment, 1 AE leading to discontinuation, and no deaths. The most frequent AEs (>10% of patients) included fatigue, headache, anemia, diarrhea and pruritis. Hemoglobin <9 g/dL on treatment was observed in 5% of patients in the DCV 3DAA regimen – was studied with and without ribavirin (RBV) in treatment-naïve and treatment-experienced patients with HCV genotype (GT) 1 infection and compensated cirrhosis in a Phase 3, international clinical trial.

CONCLUSIONS: Twelve weeks of all-oral treatment with DCV/ASV/BMS-791325 FDC, with or without ribavirin, achieved high rates of SVR12 in 202 cirrhotic patients with GT1 infection. These results demonstrate the potent antiviral activity, tolerability and safety of the DCV 3DAA regimen in patients with compensated cirrhosis.

- Treatment-naïve cirrhotics* - Treatment-experienced cirrhotics*
  - Virologic outcomes, n (%) 0 0
    - DCV 3DAA (N=57) 53 (93) 54 (98)
    - DCV 3DAA + RBV (N=55) 39 (77) 42 (93)
    - DCV 3DAA + RBV (N=45) 0

- Posttreatment relapse 4 (7) 0 0
  - 0 (7) 0
  - 1 (2) 1 (2)

- On-treatment failure 0 0 0

- Missing data 0 0 0

Cirrhosis was determined by one of the following: (1) liver biopsy with Metavir F4 or equivalent, (2) Fibroscan >14.6 kPa within 12 months prior to screening, or (3) Fibrotest ≥0.75 and AST/platelet ratio index >2 at screening.

P14
RECURRENT MICAFUNGIN-RESISTANT Candida glabrata CANDIDEMIA WITH PROLONGED MICAFUNGIN TREATMENT FOR MULTIPLE INTRA-ABDOMINAL ABSCESSES: A CASE REPORT

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BACKGROUND: Candida glabrata (CG) is the second leading cause of candidemia in North America. The reports of invasive infections caused by echinocandin-resistant CG are increasing, usually in immunocompromised patients receiving prolonged echinocandin treatment.

CASE: A 45-year-old female patient with complicated medical and surgical history was admitted for multiple intra-abdominal abscesses. The medical history included Addison’s disease on long-term glucocorticoid replacement therapy, poorly controlled type II diabetes mellitus, severe idiopathic pancreatitis complicated by pseudocyst requiring prolonged drainage, and persistent enterocutaneous fistula that failed to close despite multiple surgical interventions. Micafungin (MICA)-susceptible (S) CG was isolated from a retroperitoneal abscess and the patient was treated with MICA for 42 days. Two days after MICA was discontinued, CG was isolated from blood cultures and MICA therapy was used for another 13 days until susceptibility results detected MICA-resistant (R) and the patient was switched to fluconazole for 14 days. Six months later, CG was cultured from Jackson-Pratt drainage of re-progressing intra-abdominal abscesses, which was treated with MICA for 14 days. On day 10 of therapy, MICA R CG was again isolated from blood culture. DNA sequencing of the FSK1 and FSK2 glucan synthase loci showed mutations associated with echinocandin resistance. Patient was deceased before the susceptibility result was available.

CONCLUSION: We describe a case of recurrent MICA-R CG candidemia with prolonged MICA treatment for multiple intra-abdominal abscesses. Echinocandin resistance in Candida is uncommon but most often associated with CG and recent echinocandin exposure. Glucan synthase mutations confer high MICs, likely due to the haploid CG chromosome, and strongly correlate with clinical failure.

P15
ACTIVE ENVIRONMENTAL SURVEILLANCE OF LYME DISEASE IN BRITISH COLUMBIA (2013 AND 2014)

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OBJECTIVES: Lyme disease cases are increasing in many parts of Canada. However, it remains low in British Columbia. In response to public concerns regarding the low positivity rate, we conducted an active field surveillance for two years (2013-2014) to look at the prevalence of Borrelia burgdorferi in the vector (Ixodes pacificus ticks) and host (Peromyscus maniculatus) populations.

METHODS: Mice trapped from 11 different sites were tested by Immunofluorescence Antibody Test (IFA) and Western Blot (WB) to determine the presence of B. burgdorferi antibodies. Ticks were pooled from trapped mice (up to 5 ticks from each mouse per pool) for DNA extraction and real-time PCR, which targeted the 23S rRNA gene of Borrelia spp. and ospA gene of B. burgdorferi. Two organs (brain and bladder) from each mouse were subjected to the same molecular tests. A subset of tick DNA extracts (n=96) were also sent to CDC Fort Collins where multiplex PCR targeting flaD and gB31 genes for B. burgdorferi was performed to check for concordance between results from the two labs.

RESULTS: A total of 885 ticks at different developmental stages were retrieved from 483 mice. Serologically, 51 mice were confirmed to have antibodies against B. burgdorferi. For molecular testing, 5 tick pools were positive for B. burgdorferi but all mouse organs tested were negative. Of the
positive tick pools, the corresponding mice were also serologically positive. The subset of ticks tested by the two laboratories had identical results.

CONCLUSIONS: The positivity rate of B. burgdorferi in the British Columbia foxes tick population remained low and was found to be 0.6%. The exposure rate in the mouse population to B. burgdorferi was determined to be 11%. The low incidence in vector and low infection rate in host population may explain the continued low incidence of Lyme disease in British Columbia.

P16
INVESTIGATING THE POTENTIAL ROLE OF BACTERIAL INFECTION IN GRANULOMATOSIS WITH POLYANGIITIS (GPA)
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BACKGROUND AND OBJECTIVE: Granulomatosis with polyangiitis (GPA; formerly Wegener's disease) is a rare chronic inflammatory disorder that is characterized by granuloma formation in the lung and necrotizing-small vessel vasculitis. The available evidence indicates that GPA develops as a result of a dysregulated host immune response directed against host neutrophil-derived antigens; however, the precise etiology of this and related diseases remain largely uncertain. Importantly, in addition to genetic influences, it is thought that environmental factors also contribute to susceptibility to GPA. Recent studies have identified a number of human pathogens including Staphylococcus aureus and Burkholderia spp., among others, that have been associated with related vasculitides. Whether these or other pathogens may contribute to GPA pathogenesis is not known and was investigated here.

METHODS: Whole blood was collected and plasma isolated from 24 GPA patients and 7 healthy controls. Plasma endotoxin levels were measured using a commercial ELISA. DNA was extracted from plasma and PCR was used to detect the presence of bacteria using Burkholderia-specific and 16S rDNA primers.

RESULTS: No difference was observed in the plasma endotoxin concentration of GPA patients compared to healthy controls. In addition, analysis of the correlation between the inverse ANCA titres and plasma endotoxin level showed no significant difference among GPA patients. Burkholderia DNA (specifically B. pseudomallei, B. thailandensis, and B. cenocepacia complex) were not detected, nor were any 16S rDNA sequences detected in any of the GPA or control samples.

CONCLUSION: In these preliminary studies, we did not detect the presence of bacteria in plasma samples from GPA patients. Additional studies are required to further evaluate the role of bacteria, and possibly other pathogens, in the pathogenesis of GPA.

P17
EVALUATION OF ANTIGEN DETECTION FOR CONFIRMATORY TESTING OF HEPATITIS C VIRUS INFECTION
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OBJECTIVES: Diagnosis of hepatitis C virus (HCV) infection by serological testing requires multiple tests. Many patients with HCV antibodies do not have active infection. We investigated the use of an antigen detection assay for use in the confirmation of HCV infection.

METHODS: During a five month period, 23,096 serum specimens were tested for HCV antibodies using an automated chemiluminescent immunomassay (Siemens). 1024 specimens that were reactive in this assay were further tested for HCV antibodies using a recombinant line immunomassay (INNO-LIA) and for HCV antigen using an automated chemiluminescent immunomassay (Abbott).

RESULTS: 600 specimens (58.5%) were positive for HCV antigen, of which 390 were also positive by INNO-LIA. 8 specimens were indeterminate and 2 were negative by INNO-LIA. Seven of these samples were from patients who subsequently seroconverted, two were not re-tested and one was determined to be a false-positive HCV antigen result. 424 specimens were HCV antigen-negative, of which 384 were positive by INNO-LIA or were from previously confirmed patients. Nine specimens were INNO-LIA indeterminate, of which one patient subsequently seroconverted, and 31 specimens were negative by INNO-LIA.

CONCLUSIONS: Of 1024 serum specimens reactive in the screening assay, 96% (984) were confirmed positive by either HCV antigen test or immunoblot. Almost 60% of positive specimens were correctly identified by HCV antigen testing, allowing faster confirmation of HCV infection at lower cost than a conventional two step antibody detection algorithm. In addition, positive results for hepatitis C antigen are consistent with active infection and can be used to identify patients for further investigation.

P18
RECURRENT INVASIVE HAEMOPHILUS INFLUENZAE TYPE A INFECTION IN AN INFANT
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1Vaccine Preventable Bacterial Diseases, National Microbiology Laboratory, Winnipeg, MB; 2Saskatchewan Disease Control Laboratory, Regina; 3Prince Albert Parkland Health Region, Prince Albert, SK

BACKGROUND: Before introduction of conjugated Haemophilus influenzae serotype b (Hib) vaccines into the routine childhood immunisation programs, Hib was a major cause of meningitis in infants and children under the age of 5. In the post-Hib vaccine era, the epidemiology of invasive H. influenzae has changed substantially with most invasive diseases now caused by non-Hib strains, including H. influenzae serotype a (Hia) and serotype f, as well as non-encapsulated or non-typeable strains. This presentation describes the microbiological characterization of a H. influenzae serotype a (Hia) strain involved in causing recurrent invasive infections in an infant. The current knowledge of Hia, including methods of protection will be discussed.

METHODS: Identification and serotyping of H. influenzae were done by standard biochemical tests, slide agglutination with serotyping antisera as well as PCR methods. Multi-locus sequence typing (MLST) and pulsed-field gel electrophoresis (PFGE) were used to analyse the molecular genetics of Hia.

RESULTS: Three H. influenzae isolates were recovered from blood and CSF cultures obtained 3 months apart from the infant who succumbed to the second episode of the infection. All 3 isolates belonged to serotype a, sequence type-23, shared identical PFGE profiles, and did not possess the partial deletion involving the capsular IS1016-bexA genes.

CONCLUSION: Hia has emerged as a significant invasive pathogen in the post Hib vaccine era. MLST and PFGE serve as useful techniques for typing of Hia. Many attributes of Hia and the disease it causes bear resemblance to Hib and Hib disease, including the ability to cause recurrent infections. This raises the potential for protection by vaccination and chemoprophylaxis.

P19
OPTIMIZING THE DIAGNOSTIC YIELD OF JOINT FLUID/TISSUE CULTURE FOR THE HAMILTON REGIONAL LABORATORY MEDICINE PROGRAM
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OBJECTIVE: The aim of the study was to look at our current joint and tissue culture practices for the HRLMP to determine if these were optimal when compared to expected culture positivity rates published in the literature. Additional questions we hoped to answer were whether or not sonication and/or extending culture incubation duration could potentially increase culture yield without increasing the false positivity rate.

METHODS: In the first phase, we did a literature review to determine published culture positivity rates for joint tissues and fluids and also explored the evidence supporting prolonged culture incubation times and sonication of prostatic joints. In the next phase, we retrospectively analyzed our hospital database for joint fluid culture and in the final phase, tissue culture, both over a one year period. A total of 426 joint fluids and 56 tissue samples were assessed from culture during this time period.
RESULTS: Joint fluid culture was positive in 68 samples (16.7% of total) and tissue culture was positive in 15 samples (26.7% of total). This is lower than would be expected from the literature. Additionally Propionibacterium acnes, a slowing growing bacteria which is a relatively common cause of joint infection especially in shoulder, was not isolated in any of the cases. According to literature, prolonged incubation of joint fluid/tissue of up to 2 weeks increases the culture yield, especially for slow growing organisms, without increasing the false positivity rate.

CONCLUSION: Based on our low culture positivity rate and the available evidence we would recommend extending the incubation period from 5 days to 7-14 days to increase the culture yield for the slow growing bacteria. We are also exploring the option of sonication of prosthetic joints to increase the yield of culture and help our clinical colleagues in the diagnosis and management of joint infections.

P20

PUSHING THE LIMITS OF CHEMISTRY POINT-OF-CARE TESTING FOR THE MANAGEMENT OF PATIENTS UNDER INVESTIGATION FOR EBOLA VIRUS DISEASE

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BACKGROUND: With the recent outbreak in West Africa, hospitals worldwide have been developing protocols for suspect of cases of Ebola virus disease (EVD). Patients with EVD present with a severe gastroenteritis leading to dehydration and electrolyte abnormalities and as such, routine chemistry analysis is essential for patient management. While point of care (POC) testing can be used for rapid chemistry analyses in a containment level 2 laboratories with enhanced precautions, significant delays could ensue before specimens arrive to the laboratory due to additional precautions required during collection and transport. This study evaluated the stability of eight chemistry analytes up to four hours post-collection.

METHODS: Blood was collected by venipuncture from 20 healthy volunteers and tested at times 0, 30, 60, 90, 120, and 240 hours. Approximately 100 μL of blood was dispensed into a CHEM 8+ Cartridge and processed on a model 300 i-STAT 1 Analyzer (Abbott Point of Care Inc.).

RESULTS: While the manufacturer recommends testing within 30 minutes of collection, no significant variation in sodium, potassium, chloride, and creatinine was observed with time points extending up to four hours. For urea, ionized calcium, and bicarbonate, significant differences were observed but the concentration differences were no more than 5% from the initial time point and would not be considered clinically significant. In contrast, glucose concentrations decreased significantly (P<0.0001) over time at an average rate of 0.003 mmol/L per minute (R² =0.9989). After 90, 120, and 240 min, the decreases would be considered clinically significant at 5.7%, 7.8%, and 15.1%, respectively.

CONCLUSIONS: This study provides supporting data suggesting that delays up to four hours can be tolerated, giving ample time for collection and transport of specimens to the clinical laboratory. For glucose, POC testing could still be used, taking into account the collection time and the average rate of decrease.

P21

INCREASE IN METHICILLIN-SUSCEPTIBLE STAPHYLOCOCCUS AUREUS WITH REDUCED SUSCEPTIBILITY TO TRICLOSAN OVER THE LAST 20 YEARS

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OBJECTIVE: Triclosan as an antibacterial agent in consumer products, such as soaps, deodorants, toothpastes, plastics, and fabrics, has been used widely over the last 20 years. Triclosan targets the enoyl-acyl carrier protein reductase (FabI) which is involved in fatty acid biosynthesis. Triclosan is effective in inhibiting growth of Gram-positive and Gram-negative organisms including S. aureus and Escherichia coli. Widespread use of triclosan has the potential to lead to alterations in FabI resulting in increased triclosan MICs. We previously showed a significant increase in the proportion of clinical methicillin-susceptible S. aureus (MSSA) and methicillin-resistant S. aureus (MRSA) with elevated triclosan MICs (MICs ≥1 mg/L) isolated from 2004-2005 compared to 1993-1997. This study was completed to determine if triclosan MICs have continued to rise.

METHODS: 308 MSSA and 342 MRSA from 2012-2014 MSSA and MRSA isolates were randomly selected from a Toronto-area clinical laboratory’s archived sterile and non-sterile site isolates. Triclosan MICs were determined by agar dilution (AD) following Clinical and Laboratory Standards Institute guidelines. Results were compared to AD MIC results of the sterile and non-sterile 397 MSSA and 315 MRSA from 1993-1997 which had been randomly selected from archived isolates collected from Ontario laboratories as part of surveillance programs; and to AD MIC results of the sterile and non-sterile 436 MSSA and 362 MRSA from 2004-2005 which were randomly selected from clinical isolates.

RESULTS: The proportion of MSSA with triclosan MICs ≥1 mg/L rose from 0.2% to 4.6% to 9.1% in 1993-1997, 2005-2006, and 2012-2014, respectively (P=0.0001, χ² test for trend). There was no significant trend in the proportion of MRSA with triclosan MICs ≥1 mg/L [1.9%, 5.2%, and 0.2% in 1993-1997, 2005-2006, and 2012-2014, respectively (P=0.1590, χ² test for trend)]. Results were comparable for sterile and non-sterile isolates.

CONCLUSIONS: The proportion of MSSA with triclosan MICs ≥1 mg/L rose significantly from 0.2% to 9.1% over the last 20 years from sterile and non-sterile sites. The lack of a similar trend in MRSA, the associated mechanisms of triclosan resistance, and the possible impact on MICs to novel drugs targeting fabI should be further investigated.
C. albicans and significantly enhances the activity of FLC and ITC. Iron chelation by DIBI can inhibit the growth of C. albicans continued to be suppressed up to 72-96 h when the azole was combined with DIBI. We failed to detect a synergistic relationship using checkboard alone. Recovery of C. albicans in vitro was inhibited following 24 h exposure to 0.5 μg/mL FLC or 32 μg/mL DIBI; or 48 h exposure 1-4 μg/mL and 64-250 μg/mL respectively. We failed to detect a synergistic relationship using checkboard alone. Recovery of C. albicans in vitro was inhibited following 24 h exposure to 0.5 μg/mL FLC or 32 μg/mL DIBI; or 48 h exposure 1-4 μg/mL and 64-250 μg/mL respectively. We failed to detect a synergistic relationship using checkboard alone. Recovery of C. albicans in vitro was inhibited following 24 h exposure to 0.5 μg/mL FLC or 32 μg/mL DIBI; or 48 h exposure 1-4 μg/mL and 64-250 μg/mL respectively. We found an increase in the activity of fluconazole (FLC) or itraconazole (ITC). RESULTS: All 3 C. albicans were sensitive to both azoles (FLC 0.12-0.5 μg/mL, ITC 0.25-0.5 μg/mL). MICs to DIBI and deferiprone were 1-4 μg/mL and 64-250 μg/mL respectively. We failed to detect a synergistic relationship using checkboard alone. Recovery of C. albicans was inhibited following 24 h exposure to 0.5 μg/mL FLC or 32 μg/mL DIBI; or 48 h exposure to 8 μg/mL FLC or 64 μg/mL DIBI. However, C. albicans was inhibited at lower concentrations of 0.25 μg/mL FLC and 4 μg/mL DIBI in combination. Growth curves demonstrated that C. albicans continued to be suppressed up to 72-96 h when the azole was combined with DIBI. CONCLUSIONS: Iron chelation by DIBI can inhibit the growth of C. albicans and significantly enhances the activity of FLC and ITC in vitro.

P25
SNAP—SEPSIS NOW A PRIORITY; DEVELOPMENT AND IMPLEMENTATION OF A SEPSIS ALGORITHM IN THE EMERGENCY DEPARTMENT OF AN ACADEMIC HOSPITAL
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OBJECTIVES: We developed a sepsis algorithm based on the results of a chart audit of 364 patients presenting to the ER at Mount Sinai Hospital (MOSH) in Toronto, Canada, with sepsis in 2010-11. Our algorithm was aimed at improving sepsis management and outcomes by identifying patients early, and providing them with rapid, protocolized care in the MOSH emergency department (ED).
METHODS: The prior chart audit was the basis for the development of the Sepsis Now A Priority (SNAP) Recognition and Management algorithm, identifying gaps or deficiencies in care to be addressed. The practice changes implemented were: 1) employment of the SNAP algorithm into the ED with aggressive timelines for clinicians; 2) pre-printed order sets for initial and ongoing management of sepsis; 3) linking patient-tracking board data with collection reports; 4) revised electronic order set for sepsis symptoms; and 5) revised nursing medical directives that align with the algorithm.
RESULTS: The SNAP algorithm was implemented on July 21, 2014. As of January 1, 2015, there were 378 patients entered into the algorithm at triage. An initial chart audit of 30 patients entered into the algorithm demonstrated that 53% (16/30) of patients had a diagnosis of sepsis. Of those, 19% (3/16) had severe sepsis and 6% (1/16) had septic shock. Data collected to date has demonstrated that the timelines of the algorithm are being met and that septic patients are being flagged and treated swiftly. Ongoing post-implementation review will evaluate patient outcomes including mortality, morbidity, length of stay and process measures (i.e. time to diagnosis, time to appropriate fluid resuscitation, antibiotics, etc.).
CONCLUSION: The ability to recognize sepsis early is essential to improving outcomes. The intent of the SNAP algorithm is to allow for early recognition of septic patients, with rapid, protocolized care. By implementing this concise quality improvement tool to optimize diagnosis and treatment, we hope to improve outcomes for patients with sepsis. Our algorithm prioritizes identifying patients with possible sepsis over having a specific tool, which would identify fewer patients who do not end up having sepsis.
RESULTS AND CONCLUSION: As a result of the presentation and discussion, the relationships, roles, and knowledge needs of the varied disciplines and stakeholders will be better understood. This should be a useful step forward for better public health knowledge translation to address the three pillars of the federal framework.

P28
OVERSIGHT OF HUMAN AND ANIMAL PATHOGENS AND TOXINS
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BACKGROUND: In 2009, the Human Pathogens and Toxins Act (HPTA) received Royal Assent with the intent to establish a national safety and security regime to protect the health and safety of the public against the risks posed by human pathogens and toxins. With the passage of the HPTA, select provisions came into force to provide interim measures until a complete program and regulatory framework could be developed.

METHODS: Regulated and interested parties were consulted over 4 years to develop the proposed Human Pathogens and Toxins Regulations (HPTR) and the Canadian Biosafety Standard (CBS) to support the full implementation of the HPTA scheduled for December 2015.

RESULTS: The proposed HPTR would set out a risk-based licensing scheme for facilities conducting controlled activities with human pathogens and toxins. Furthermore, a biological safety officer who meets certain knowledge requirements set out in the proposed regulations would be designated before a licence can be issued by the Agency. The updated CBS sets out the physical containment, operational practice, and performance and verification testing requirements for the safe handling and storing of human and terrestrial animal pathogens and toxins. The Agency will also use other policy tools to strengthen pathogen accountability and oversight in Canada.

CONCLUSION: The Agency is developing a risk-based program and regulatory framework based on requirements set out in the HPTA, input from stakeholders, policy decisions and other considerations such as domestic and international best practices. This framework facilitates the best and most innovative science at Canadian university and science and technology facilities, enables Canadian public health labs to respond to disease outbreaks and threats of unknown pathogens as efficiently as possible and finally enables Canadian companies to maintain a competitive edge in the economy, all while ensuring activities are conducted in a manner that is safe and secure as possible.

P29
WITHDRAWN

P30
THE RETROSPECTIVE STUDY OF THE PREVALENCE AND CLINICAL SIGNIFICANCE OF HEPATITIS B VIRUS PRECORE AND BASAL CORE PROMOTER VARIANTS
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OBJECTIVES: Hepatitis B Virus (HBV) Precore (PC) and Basal Core Promoter (BCP) variants are recognized for so long but extrapulmonary manifestations also occur. <5 years of age. Pneumonia and asthma exacerbation are most common, but extrapulmonary manifestations were included.

RESULTS: 96 children, aged <5 years, were identified; 19 were excluded on the basis of an alternative diagnosis. Median age was 36 months (2 weeks to 59 months); 7 were ≤1 year of age. Male to female ratio was 1:3.1. MP was detected in 76 respiratory samples (throat swabs [n=52], nasopharyngeal swabs [n=12], bronchoalveolar lavage [n=9]), aural suction [n=2], pleural fluid [n=10]) and 4 CSF samples. Sixty-seven (87%) had symptoms restricted to the respiratory tract, 56 with pneumonia and 4 with asthma exacerbation. Multilobar infiltrates were seen in 13 of 70 (19%) patients who had chest radiographs. There was no difference in the proportion of those with radiographic evidence of multilobar involvement between patients <24 months of age and those who were older. Of the 10 (13%) who had extrapulmonary manifestations, 5 had neurologic disease (2 encephalitis, 2 ataxia; 1 motor axonal neuropathy), 4 had dermatologic manifestations (3 erythema multiforme, 1 serum sickness rash), and 1 had autoimmune hemolytic anemia. Four (40%) of those with extrapulmonary manifestations did not have documented concurrent or preceding respiratory symptoms.

CONCLUSIONS: MP infection can cause significant disease in children <5 years of age. Pneumonia and asthma exacerbation are most common, but extrapulmonary manifestations also occur.

P32
EMERGENCY DEPARTMENT UTILIZATION OF RESPIRATORY VIRUS TESTING FOR FEBRILE YOUNG INFANTS UNDER SIX WEEKS OF AGE
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BACKGROUND: Well appearing febrile infants with viral illnesses cannot be distinguished from those with occult life-threatening infections by history and physical exam alone. However, infants with confirmed respiratory viruses are at lower risk of serious bacterial infection. This study sought to assess the utilization of respiratory virus testing (RVT) among pediatric ED patients.
physicians in the evaluation of febrile infants under 6 weeks of age.

METHODS: A scenario-based survey describing 2 hypothetical cases of febrile infants without a focus aged 3 and 5 weeks was sent to ED physicians at 15 pediatric tertiary centers across Canada. Participants were asked multiple choice questions to evaluate the use of RVT. In parallel, we performed a single center retrospective analysis of RVT among all infants under 6 weeks evaluated in the ED for sepsis from 2009 to 2013.

RESULTS: Survey response rate was 81% (n=171). Based on the hypothetical scenarios, RVT was performed more frequently in 3-week infants (40% vs. 29%, P<0.05). Conversely, more ED physicians felt that RVT would have an impact on the management of 5-week infants (35% vs. 15%, P<0.001). RVT was performed more among physicians reporting test results available under 6 hours compared to those reporting test results available only after 24 hours, for both 3-week (62% vs. 32%, P<0.01) and 5-week-old infants (41% vs. 14%, P<0.01). There were significant variations in rates of RVT between centers, ranging from 0%-75% for 3-week-old infants, and from 0%-63% for 5-week-old infants. In clinical practice, among 1292 infants evaluated in a single ED for sepsis, RVT was performed more frequently for infants 4-6 weeks of age than for infants under 4 weeks (73% vs. 58%, P<0.001).

CONCLUSIONS: Utilization of RVT for the evaluation of febrile young infants differs by infant age and among centers. More widespread access to rapid RVT may increase the utilization of this tool in the pediatric ED.

SP01

COPD SYMPTOM AND DISEASE SCREENING IN AN HIV POPULATION

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BACKGROUND: HIV-positive individuals represent a population that appears to be at a higher risk of developing chronic obstructive pulmonary disease (COPD). In this study, we sought to determine the effects of smoking on respiratory symptoms and smoking related diseases among HIV-positive patients and to determine if symptomatic patients are being appropriately screened for COPD.

METHODS: HIV-positive individuals were asked to complete a self-administered questionnaire regarding respiratory symptoms and diseases. The effects of smoking on respiratory symptoms and diseases were reported as estimates of odds ratio. The screening criteria used to determine at risk patients for COPD were adapted from the Canadian Thoracic Society (CTS) guidelines.

RESULTS: A total of 247 patients were recruited. The median age was 49 years; 75% were male and 92% were currently on combination antiretroviral therapy (cART). Current and former smokers represented 66% of the population. Smoking had a statistically significant effect on respiratory symptoms including wheeze (OR 4.8 [95% CI 1.6-14.2]), phlegm production (OR 4.9 [95% CI 2.2-10.5]), current cough (OR 7.0 [95% CI 3.0-16.2]), chronic cough ≥3 months (OR 5.2 [95% CI 2.3-11.8]) and dyspnea (OR 7.2 [95% CI 1.7-31.2]). Smoking also had a statistically significant effect on respiratory diseases including COPD (OR 4.9 [95% CI 1.1-21.9]), bronchitis (OR 3.8 [95% CI 1.9-7.7]), asthma (OR 6.0 [95% CI 2.0-17.7]) and pneumonia (OR 2.1 [95% CI 1.2-4.0]). Among HIV-positive smokers, 40% met the CTS criteria for COPD screening, while only 12% of smokers self-reported a diagnosis of COPD and 9% reported use of inhaled puffers. The burden of smoking in the HIV population is significant. HIV-positive smokers are more likely to report both respiratory symptoms and diseases than HIV-positive non-smokers. A discrepancy exists between patients who meet the CTS COPD screening guidelines and those who have been diagnosed with COPD, raising the concern for under-recognition and under-diagnosis of COPD in the HIV-positive population. Our study is limited by self-reporting of symptoms and disease states.
RESULTS: Of the cohort, 91% of patients received empiric antibiotics, whereas only 53% had urine cultures that were positive for an organism (an over-estimate of ‘true’ UTI). Significant risk factors for receiving empiric treatment were a history of fevers or chills and a positive urine culture (RR 12.0 and 1.9, respectively). Significant risks for receiving IV antibiotics were incontinence, confusion or delirium, fever, positive nitrite on urinalysis, and history of diabetes or neurological disorders (RR 1.6 to 3.8). However, classic symptoms such as dysuria, frequency, and urgency were not significant predictors for receiving empiric antibiotic therapy.

CONCLUSIONS: There was a high prevalence of antibiotic use in this elderly cohort presenting with suspected UTI. Multiple features were associated with increased risk of receiving antibiotics, but were not classic features of UTI. Further studies are required in order to assess the efficacy of these clinical features in accurate diagnosis of UTI in this age group in order to minimize inappropriate antibiotic use.

SP05
THE INFECTIONOUSNESS AND/OR VIRULENCE OF ISONIAZID-RESISTANT VS. SUSCEPTIBLE PULMONARY TUBERCULOSIS: A CASE-CONTROL STUDY IN ALBERTA
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OBJECTIVE: In Canada, isoniazid (INH) is the most common first-line anti-tuberculosis (TB) drug to which isolates of Mycobacterium tuberculosis are resistant. The aim of this retrospective case control study is to determine whether isoniazid (INH)-resistant strains are as transmissible/virulent as INH-susceptible strains.

METHODS: Over the 20 years, 1991-2010, all adults (age ≥14 years) with culture-positive pulmonary TB in the Province of Alberta, Canada were identified. From within this cohort, those with phenotypically INH-resistant isolates were identified and defined as “cases”. Phenotypic resistance was confirmed and INH- conferring mutations sought in each isolate. From among these that were INH-susceptible patients, two age (±5 years), sex, population group, smear status and cavitation status “controls” were randomly selected. Conventional and molecular epidemiological tools were then used to identify secondary cases from among contacts. The proportion of contacts that were secondary cases were compared in cases and controls.

RESULTS: A total of 96 pulmonary TB cases were infected with INH-resistant TB strains; 76 with high-level and 20 with low-level phenotypic resistance. katG-only mutations were present in 61 strains (all high-level), inhA-only mutations in 16 strains (14 low-level, 2 high-level), both katG and inhA mutations in 4 strains (3 high-level, 1 low-level) and no mutation or an aphC-only mutation in 15 strains (10 high-level, 5 low-level). Contacts of cases were significantly less likely than contacts of controls to be secondary cases (1 of 2203 vs. 20 of 4331), P <0.05.

CONCLUSION: katG mutations were associated with high-level INH resistance. INH-resistant strains were less transmissible/virulent than INH-susceptible strains.

SP06
MEROPENEM MIC PROFILE OF CARBAPENEMASE-PRODUCING ENTEROBACTERIACEAE IN A LOW PREVALENCE REGION
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BACKGROUND AND OBJECTIVE: Over the past decade, transmission of carbapenemases has become a major problem and carbapenemase producing Enterobacteriaceae (CPE) have been identified in countries worldwide. Of significance, infection with CPE has been associated with substantial morbidity and mortality. Thus, there is a critical need to curtail their dissemination. In an effort to better inform CPE screening practices we analyzed the meropenem MIC distribution of a large set of clinical Enterobacteriaceae isolates.

METHODS: A total of 1022 clinical/screening isolates submitted to the Public Health Ontario Laboratories (PHOL), from January 2011 to March 2014, were analyzed. Only isolates displaying a meropenem or ertapenem MIC of ≥0.25 or 1 μg/mL (determined by agar dilution) were included. Carbapenemase-positive isolates were identified by multiplex PCR.

RESULTS: We identified 189 carbapenemase-positive isolates (90 KPC, 63 NDM, 30 OXA-48-like, 4 KdpA, 1 IMI, 1 BLABE, 1 BLAAOM, 1 BLACQ-OM, 1 OXA-48-like) among the data set, largely within the Klebsiella spp, Escherichia coli and Enterobacter spp. Interestingly, 15 or 20% of CPE displayed meropenem MICs within the susceptible range based on current CLSI (≤1 μg/mL) or EUCAST (≤2 μg/mL) breakpoint tables, respectively. The majority of meropenem-susceptible CPE were observed among E. coli isolates; however, this phenomenon was not exclusive to any one genus. Among the most identified carbapenemase types, a substantial proportion of the OXA-48-positive (40%/50% based on CLSI/EUCAST breakpoints) and KPC-positive (22%/33%) isolates were found to be meropenem susceptible, whereas few, if any, of the NDM-positive isolates (0%/2%) were meropenem susceptible. Notably, application of CLSI carbapenemase screening recommendations captured only 85% of carbapenemase-producing isolates, whereas application of EUCAST recommendations captured >98% of CPE.

CONCLUSION: Our findings demonstrate that in a region with low carbapenemase prevalence, meropenem-based screening approaches will require a cut-off MIC near the epidemiological wild-type cut-off in order to achieve near optimal CPE identification.

SP07
CREATION OF A NOVEL PEDIATRIC ANTIMICROBIAL STEWARDSHIP ADVANCED FELLOWSHIP TRAINING PROGRAM
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OBJECTIVE: To describe the development and structure of a Pediatric Antimicrobial Stewardship (AS) Advanced Fellowship program in Canada.

BACKGROUND: Though some Canadian training programs with a focus on AS in adult medicine exist, there are currently no programs with a focus on pediatrics.

METHODS: A Pediatric AS Advanced Fellowship program was developed for the 2014-2015 academic year. Recognition by the Post-graduate Medical Education (PGME) office was obtained following submission of a formal application for an advanced fellowship program detailing the structure, objectives and evaluation strategy for the fellowship along with specific CanMeds objectives.

RESULTS: The intent of the fellowship is to prepare a physician with training in Pediatric Infectious Diseases for a career trajectory in pediatric AS by learning the core knowledge required to perform AS related program development, research and clinical work. The program is comprised of multidisciplinary collaboration with the adult/pediatric pharmacy teams and Infection Prevention and Control teams at tertiary care sites in the city. The trainee is also involved in the local and provincial AS committees and other projects including review of local clinical guidelines, education and collaborative projects. Foundational theory in the principles of AS is gained through a longitudinal research component, working with experienced physicians/pharmacists, and independent learning through accredited antimicrobial stewardship courses. Successful completion of the fellowship will be acknowledged with a certificate awarded by the PGME office. Application for Royal College recognition as an “Area of Focused Competence” is in progress.

CONCLUSIONS: This novel training program provides a unique opportunity to develop skills in pediatric antimicrobial stewardship and could serve as a prototype for other pediatric centres to develop similar programs.
SP08 EMERGING ANTIMICROBIAL RESISTANCE AMONG STAPHYLOCOCCUS PSEUDINTERMEDII, A POTENTIAL ZOOONES OF CANINE ORIGIN
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BACKGROUND: Staphylococcus pseudintermedii is a ubiquitous colonizer and opportunistic pathogen of dogs. S. pseudintermedii is coagulase positive, biochemically similar to and indistinguishable from S. aureus on CHROMagar S aureus and Denim Blue Agars. Although rarely recognized backgROund, S. pseudintermedii infections are typically zoonotic, stemming from contact with dogs. The emergence of antimicrobial resistance, including methicillin resistance (MRSP), is therefore of interest to physicians and veterinarians.

METHODS: Pharyngeal and rectal swabs were taken from 100 healthy dogs presenting for vaccination to the Western College of Veterinary Medicine in Saskatoon, SK. Samples were cultured for staphylococci and S. pseudintermedii was identified biochemically including tests for acetoin and hyaluronidase production which differentiate it from S. aureus. Antimicrobial MICs were determined and interpreted using the CLSI guidelines. Methicillin resistance was confirmed by PCR amplification of the mecA gene.

RESULTS: 78% of dogs were colonized with S. pseudintermedii including 7% with MRSP. No isolates resistant to fluoroquinolones, rifampin, nitrofuranto-in, vancomycin, linezolid, daltopycin + dalfoprinid were identified nor was inducible clindamycin resistance. The most common resistance phenotypes were penicillin + ampicillin (31.7% of isolates) and pan-susceptibility (30.3% of isolates). Resistance to tetracycline (20.8% of isolates), erythromycin, clindamycin, trimethoprin + sulfamethoxazole, gentamicin and chloramphenicol (all ≤5% of isolates) was also found. None of the MRSP colonized dogs had received antimicrobials in the previous 6 months, indicating a community reservoir.

CONCLUSIONS: Compared to a similar S. pseudintermedii surveillance study conducted in 2008, resistance (including MRSP) was more frequently identified. The introduction of high throughput highly discriminatory bacterial identification technologies such as MALDI-TOF may increasingly result in the identification of S. pseudintermedii from human infections. Physicians should be aware of this potential zoonoses and consider its infection control implications.

SP09 POSITIVE IMPACT OF INFECTIOUS DISEASE CONSULTATION ON QUALITY OF CARE, MORTALITY AND LENGTH OF STAY IN STAPHYLOCOCCUS AUREUS BACTEREMIA: RESULTS FROM A LARGE MULTICENTER COHORT STUDY
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OBJECTIVES: We assessed the impact of Infectious Disease (ID) consultation on management and outcome in Staphylococcus aureus bacteremia (SAB).

METHODS: A retrospective cohort study examined consecutive SAB patients from 6 academic and community hospitals between 2007 and 2010. Quality measures of management including echocardiogram, repeat blood culture, removal of infectious foci, and antibiotic therapy were compared between ID consultation (IDC) and no ID consultation (NIDC) groups. A competing risk model with propensity score adjustment was used to compare in-hospital mortality and time to discharge.

RESULTS: Of 847 SAB patients, 506 (60%) patients received an ID consultation and 341 (40%) patients did not. During hospital stay, 371 (73%) IDC patients and 191 (56%) NIDC patients received an echocardiogram (P<0.002). Repeat blood cultures in 2-4 days of bacteremia were performed in 207 (41%) IDC patients and 107 (31%) NIDC patients (P=0.0058). There was no statistical difference in removal of infectious foci between the two groups. For patients who were alive when the antibiotic course was completed, 285/422 (68%) IDC patients and 141/262 (54%) NIDC patients received the appropriate duration of antibiotic therapy (P=0.0004).

During hospital stay, 204 (24%) patients died: 104 (21%) IDC patients and 100 (29%) NIDC patients. Matched by propensity score, ID consultation had a sub-distribution hazard ratio (shHR) of 0.72 (95% CI: 0.52-0.99, P=0.0451) for in-hospital mortality and 1.28 (95% CI: 1.06-1.56, P=0.012) for being discharged alive.

CONCLUSIONS: ID consultation is associated with adherence to quality measures, reduced in-hospital mortality and earlier discharge in SAB patients.

SP10 PROSPECTIVE OBSERVATIONAL STUDY OF DIAGNOSIS OF URINARY TRACT INFECTION AND RESPONSE TO THERAPY IN LONG TERM CARE
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BACKGROUND: Prevalence of asymptomatic bacteriuria among residents of long-term care (LTC) facilities is 40%, and is a source of inappropriate antibiotic use.

OBJECTIVES: To determine the signs and symptoms associated with positive urine culture and if antibiotic therapy is associated with functional improvement.

METHODS: LTC residents were prospectively observed following urine culture submission.

RESULTS: 174 consecutive specimens from 6 LTC facilities were considered and 101 specimens from 101 residents were eligible. Mean age was 84.0 years (SD 8.6), 79.2% were female, with a mean of 1.8 comorbidities per patient (SD 1.0). Baseline ADL score was 11.9 (SD 8.7), with zero representing total independence and 28 representing total dependence. 38/101 specimens (37.6%) had significant growth. Reasons for collection ranged from one to seven reasons with dysuria, change in character of urine and change in mental status being the most common. Most episodes did not have vital signs or blood testing performed in order to apply published diagnostic criteria. Using univariate regression, two predictors were associated with culture positivity (male sex, RR 5.58, 95% CI = 1.23-23.43; change in mental status, RR 13.83, 95% CI = 1.81-105.81). Antibiotic therapy was prescribed in 4/101 (47.5%) of episodes, but treatment decisions did not correlate with significant growth (kappa=0.44). 19/48 (40%) of treatments were given to residents without significant growth and 9/53 (17%) of episodes with significant growth were not treated. 4/28 (14.3%) of treatments given were reported as resistant. Baseline, 2 day and 5-7 day antibiotic courses were prescribed in 48/101 (47.5%) of episodes, but treatment decisions did not correlate with significant growth (kappa = 0.44). 19/48 (40%) of treatments were given to residents without significant growth and 9/53 (17%) of episodes with significant growth were not treated. 4/28 (14.3%) of treatments given were reported as resistant. Baseline, 2 day and 5-7 day ADL scores did not significantly change in treated or untreated groups.

CONCLUSIONS: The observed rate of significant growth is approximately the same as the expected rate of asymptomatic bacteriuria. Antibiotic treatment did not lead to functional improvement. Restricting access to urine culture may be a future strategy to improve antibiotic stewardship.

SP11 MOLECULAR EPIDEMIOLOGY OF VANCOMYCIN HETEROENZYMATIC COAGULASE-NEGATIVE STAPHYLOCOCCI IN A NEONATAL INTENSIVE CARE UNIT IN QUEBEC
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OBJECTIVES: Coagulase negative staphylococci (CoNS) have emerged as a leading cause of bloodstream infections (BSIs) in ICUs, in particular in premature neonates. The study aimed to describe the molecular and epidemiological characteristics of heterogeneous vancomycin intermediate coagulase negative staphylococci (hVICoNS) central line associated BSI (CLABSI) in a single tertiary-care NICU in Quebec.
METHODS: Between November 2009 and April 2014, all CoNS causing CLABSI were identified in a single tertiary care NICU in Québec through the laboratory information system and the infection control database. Heterogeneous resistance to vancomycin (hV) was determined by both the Macro E-test and E-test GRD. Antibiotic susceptibility to daptomycin and linezolid was determined by E-test, following the manufacturer's recommendations. Clonal relationships were determined by pulsed-field gel electrophoresis (PFGE). Population analysis profile/area under the curve (PAP/AUC), as the gold standard for hV, is being performed.

RESULTS: Eighty-eight CoNS strains were collected. Decreased vancomycin susceptibility was identified in all isolated strains. The most common species of IVicoNS causing CLABSI were Staphylococcus epidermidis (96.6%), S. warneri (2.3%), and S. capitis (1.1%). Eighty-two IVicoNS strains (93.2%) were susceptible to daptomycin, whereas only 52 (59.1%) of isolated strains were susceptible to linezolid. PFGE revealed 27 distinct patterns (A to AB), with 6 strains untypeable. PFGE pattern E strains spanned from 2009-2014, pattern M strains spanned from 2010-2014, and pattern O strains spanned from 2011-2013. CONCLUSIONS: IVicoNS seems more common than currently realized in the NICU, although PAP/AUC results are still pending. Treatment of IVicoNS CLABSI with linezolid may no longer be appropriate, with 40.9% of strains demonstrating resistance. PFGE pattern E, M, and O strains spanned several years, indicating that certain clones can become endemic to the NICU.

SP12 INCREASING INCIDENCE OF GROUP C AND GROUP G STREPTOCOCCAL BACTEREMIA ATTRIBUTED TO INCREASED BACTERIAL VIRULENCE FACTORS AND POOR OUTCOMES
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BACKGROUND: β-hemolytic streptococci are responsible for several human diseases, including skin and soft tissue infection, bacteremia, and streptococcal toxic shock syndrome. Mortality rates in bacteremia have been reported up to 20%. We have described an increased incidence of group C and G streptococcal (GCS and GGS, respectively) bacteremia from January 2007 to December 2012 in our center. We hypothesize acquired virulence factors accounts for increased pathogenicity and poor clinical outcomes.

OBJECTIVES: To identify streptococcal virulence factors in our GCS and GGS blood isolates compared to controls.

METHODS: The study population included all patients with blood cultures for GCS and GGS in two tertiary care centers from January 2012 to December 2014. Polymerase chain reaction (PCR) was performed on a subset of 15 patients and 15 non-invasive controls obtained from throat culture. 5 superantigens previously described in invasive group A streptococcal bacteremia (speA, speG, speK, smeZ, ssa) were explored. Whole genome sequencing, protein profiling via MALDI-ToF, clinical parameters such as demographics, co-morbidities, disease severity and outcomes via retrospective chart review will be completed in the near future.

RESULTS: 90 blood and 33 throat isolates were retrieved. Of the blood isolates, 57 (63%) and 33 (37%) were Lancefield group G and C, respectively. Of the throat isolates, 17 (52%) and 16 (48%) were Lancefield group G and C, respectively. MALDI-ToF mass spectrometry confirmed all but one blood isolate as Streptococcus dysgalactiae grp species. The other was identified as Streptococcus equi. Superantigen genes speA, speG, speK, smeZ, and ssa were not detected by PCR in our subset.

CONCLUSIONS: No invasive pathogenic factor has been identified in this study to account for the increased incidence. Further studies are underway to analyze whole genome sequencing and protein profiling in our cohort.
was never seen at the MCH). A total of 136 PCR positive samples underwent ELISA testing; 34 were positive (40%) for toxin A or B. The average age of the entire cohort was 8.5 years (SD 6.2). There was no (p<0.05) significant difference in age, gender, CIDI clinical manifestation, previous medical problems, and management between patients positive or negative by ELISA. However, patients with positive ELISA results were more likely to have had a recent exposure to antibiotics (67.9% vs. 50%, P=0.04 for ELISA+ and - , respectively).

CONCLUSION: In our pediatric population, 60% of patients with CIDI diagnosed by PCR had no toxin detectable by ELISA. PCR+/ELISA− patients are less likely to have received an antibiotic recently than PCR+/ELISA+ patients. These results highlight the need to standardize laboratory criteria for the diagnosis of CIDI in children.

SP16 SURVEILLANCE OF HUMAN PARAINFLUENZA TYPES 1-4 AND ASSOCIATION WITH ACUTE LOWER RESPIRATORY TRACT INFECTIONS IN HOSPITALIZED CASES, ALBERTA, CANADA (2010 – 2013)

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OBJECTIVE: To describe the association of the Human Parainfluenza virus types (PIV 1-4) in hospitalized cases, with acute respiratory illnesses such as bronchiolitis, croup, pneumonia, and mixed illnesses in Alberta, Canada.

METHODS: ProvLab Alberta tests all respiratory samples for influenza A/B. Influenza A/B negative specimens from non-community patients undergo the Respiratory Virus Panel classic assay, which detects PIV 1-4. Positive PIV specimens were merged with Alberta Public Health administrative databases to define cases and hospitalization status, using unique patient identification numbers for all Alberta patients. ICD-9 codes were used to determine if cases had one event of croup, bronchiolitis, pneumonia, or mixed events (any combination of the three acute respiratory illnesses). All statistical analysis was done using SPSS (Version 19.0.0, IBM Corp© 2010).

RESULTS: 1204 hospitalized PIV 1-4 cases were identified. PIV 3 was the most common type in Alberta, 44.8% (539/1204) followed by PIV 1, 25% (301/1204). Of all PIV 1-4 cases, pneumonia was the most common acute respiratory illness 33.5%, (463/1204) followed by bronchiolitis 19.5%, (235/1204). The majority of group cases 58.9% (103/175) were due to PIV 1 (Fischer’s Exact=166.242, P<0.001) while pneumonia cases 49.1% (198/403) were most likely to be associated with PIV 3 (Fischer’s Exact=14.937, P=0.002). There was no statistical significance between the pneumonia cases and PIV 1 (Fischer’s Exact=166.242, P<0.001) while pneumonia cases 49.1%, (198/403) were most likely to be associated with PIV 3 (Fischer’s Exact=14.937, P=0.002). There was no statistical significance between the pneumonia cases and PIV 1 (Fischer’s Exact=166.242, P<0.001) while pneumonia cases 49.1%, (198/403) were most likely to be associated with PIV 3 (Fischer’s Exact=14.937, P=0.002). There was no statistical significance between the pneumonia cases and PIV 1 (Fischer’s Exact=166.242, P<0.001) while pneumonia cases 49.1%, (198/403) were most likely to be associated with PIV 3 (Fischer’s Exact=14.937, P=0.002).

CONCLUSIONS: This study shows association of PIV 1-4 types with specific illnesses including croup and pneumonia. Periodic surveillance of PIV types can be used to evaluate the epidemiology and burden of PIV in hospitalized cases with acute respiratory illnesses.
CONCLUSION: LAIV appears to be well tolerated in children and adolescents with CF. We did not detect a signal for oculo-respiratory syndrome post-LAIV in this population, as previously postulated. Study results support a national vaccine recommendation for preferential use of LAIV in post-LAIV in this population, as previously postulated. Study results support a national vaccine recommendation for preferential use of LAIV in this population, as previously postulated. Study results support a national vaccine recommendation for preferential use of LAIV in post-LAIV in this population, as previously postulated. Study results support a national vaccine recommendation for preferential use of LAIV in post-LAIV in this population, as previously postulated. Study results support a national vaccine recommendation for preferential use of LAIV in post-LAIV in this population, as previously postulated.

SP19
SEVERE SKIN LESIONS INDUCED BY A PREDOMINANT CA-MRSA STRAIN USA300 ARE ASSOCIATED WITH A UNIQUE NEUTROPHIL RESPONSE AND INVOLVEMENT OF PROTEINASE ACTIVATED RECEPTORS (PARS)

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OBJECTIVES: The mechanisms of severe skin lesions induced by a predominant CA-MRSA strain USA300 remain unknown. We established a murine intradermal infection model capable of demonstrating dermatopathological differences between USA300 and other MRSA strains, and capable of revealing new potential mechanisms contributing to development of severe skin lesions.

METHODS: BALB/c mice were monitored for skin lesions and systemic infection after intradermal injection with virulent MRSA USA300, USA400 and avirulent M92. Biopsies of the core cutaneous lesion were subjected to detailed histopathological examination (H & E, Gram, and Esterase staining). Neutrophil infiltration and adhesion/emigration were quantified by myeloperoxidase (MPO) assay and spinning disk confocal microscopy, respectively. RB6-8C5 antibody was used to deplete neutrophils 1 day prior to (early) or post (late) infection. PAR1/2 KO and MyD88−/− mice were used to study the role of PARs in USA300 dermatopathology.

RESULTS: Skin lesions induced by USA300 uniformly presented as extensive ulcers with a profound inflammatory cell infiltrate. In contrast, USA400 and M92 only caused localized cutaneous infection. In this model, no systemic infection was observed, although Gram staining indicated deeper bacterial invasion in USA300-infected tissues. Esterase staining and MPO assay confirmed that USA300 induced greater neutrophil infiltration than other strains (P<0.05). Live cell imaging displayed significantly increased neutrophil (but not CD4) adherence and emigration in USA300-infected mice compared to other strains (all P<0.05). Early (but not late) neutrophil-depletion exacerbated skin lesion sizes (P<0.05), and both early and late depletion resulted in extensive bacterial dissemination and systemic infection in USA300-infected mice. Results with PAR-null mice showed that USA300 induced smaller lesion sizes with less neutrophil infiltration in PAR1-KO than WT mice. Similar lesion size but faster healing with decreased neutrophil infiltration was observed in PAR2-KO relative to WT mice.

CONCLUSIONS: We were able to demonstrate the key clinical characteristics between CA-MRSA USA300 and other MRSA strains in a murine intradermal infection model. In part, the mechanism of USA300-induced severe skin lesions is related to over-activation of neutrophils possibly through activation of PARs.

SP20
SURVEILLANCE OF ANTIMICROBIAL USE IN QUÉBEC
ACUTE CARE HOSPITALS: A DREAM OR ALMOST REALITY?
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OBJECTIVES: In 2011, the Québec Ministry of Health (MSSS) issued a directive to acute care hospitals to implement surveillance of antimicrobial (AM) use in admitted patients. We thus aimed to describe 1) available pharmacy data; 2) hospitals’ actual practices in qualitative and quantitative surveillance of AM use; 3) hospitals’ motivation to perform surveillance of AM use.

METHODS: A web-based questionnaire was sent to chief pharmacists in the province of Québec acute care hospitals for them to complete or forward to the pharmacist in charge of AM use surveillance.

RESULTS: Of 110 questionnaires sent, 44 were at least partly filled (40%), representing 49 hospitals. A database describing AM use was available in 43 hospitals (6 had individual-level data only, 30 had aggregated data only and 7 had both). Ninety percent of hospitals (36/40) had a qualitative surveillance program or were about to implement one (4 had implemented it before 2011); most monitored indication (89%), AM choice (89%), dosage (78%), duration of treatment (78%) and route (72%). Eighty percent of hospitals (32/40) had a quantitative AM use surveillance program (QASP) or were about to implement one (5 had implemented it before 2011); most of these hospitals (84%) monitored defined daily doses of AM per patient-days, but non distributed and non administered doses were analyzed differently across hospitals. In 80% of hospitals, the participant was favourable to the implementation of a provincial QASP. Foreseen problems were the lack of human, financial and information technology resources; comparisons between hospitals were viewed as both a methodological challenge and useful information.

CONCLUSION: AM use surveillance has been implemented in most participating hospitals, but databases are not readily available and indicator definitions vary. However, most participants had a positive view regarding an eventual provincial surveillance program.
significantly from CD-PCR- cases with regards to serum albumin, serum creatinine, temperature, presence of abdominal pain, presence of inflammatory bowel disease or concomitant antibiotic therapy. Mean fecal calprotectin values were higher in CD-PCR- patients (P<0.001), as was the mean serum PCT (p=0.021) in those subjects. The area under the curve (AUC) for CPT-HR predicting CD-PCR+ was 0.82 (95% CI 0.70-0.94) with a sensitivity of 87% and a specificity of 75% at a HR-CPT of 135 mcg/L. CPT was less specific with an AUC of 0.8 (CI 0.67-0.92), specificity 57%, specificity 55%, at a CPT of 81.5 mcg/L. The AUC for PCT was not significant.

**CONCLUSIONS:** Elevated fecal CPT-HR is a valid predictor of CD-PCR+ in a hospitalized patient with diarrhoea, however further studies are required to determine its quality as a marker of CDI versus colonization.

**SP22** SUCCESSFUL USE OF INTRATHECAL COLISTIN IN A CASE OF PSEUDOMONAS AERUGINOSA VENTRICULITIS: A CASE REPORT AND BRIEF REVIEW OF THE LITERATURE

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**BACKGROUND:** The use of intrathecal antibiotics is rare, but may represent an additional treatment option in cases of ventriculitis with multi-drug resistant (MDR) organisms or in patients with persistently positive cultures.

**METHODS:** We searched PubMed for English language articles using the terms “intrathecal,” “colistin,” and “Pseudomonas”. We present a case where intrathecal colistin was used successfully in a case of Pseudomonas aeruginosa ventriculitis, as well as a summary of six additional similar cases from the literature.

**RESULTS:** The most common presenting diagnosis was subarachnoid haemorrhage in five of seven patients. In all seven cases, intrathecal therapy was used in conjunction with systemic antibiotic therapy and removal or externalization of the infected catheter. In two cases, the renal side effects of intravenous (IV) antibiotics prompted a switch to intrathecal colistin. Only one study reported side effects with intrathecal colistin in the form of self-limited paresthesias of the arm. In two cases, the renal side effects of intravenous (IV) antibiotics alone had failed to do so. In our case, the CSF was sterilized by day seven of IV colistin alone, but a subsequent rise in CSF leukocytes suggested persistent infection, prompting intrathecal therapy. In three patients where IV and intrathecal colistin were used concomitantly, CSF sterilization occurred by day 2-4 of therapy, and no recurrence occurred.

**CONCLUSIONS:** Intrathecal colistin has been successfully employed in CNS shunt infections with multi-drug resistant Pseudomonas aeruginosa in conjunction with IV antibiotics resulting in rapid CSF sterilization, and physicians should be aware of this potential strategy. Patients should be monitored for neurological side effects during therapy.

**SP23** PREVALENCE OF ANEMIA IN THE SASKATCHEWAN POPULATION WITH HIV INFECTION

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**BACKGROUND:** Anemia is a common hematologic complication in patients with Human Immunodeficiency Virus (HIV) infection. It is an independent risk factor for disease progression and adverse clinical outcomes that include increased morbidity and mortality. Appropriate management of anemia in patients with HIV infection improves functional outcomes and quality of life. The prevalence of anemia, risk factors for developing anemia, and rates of intervention for patients with anemia are currently unknown in the Saskatchewan population with HIV infection.

**METHODS:** A retrospective chart review of 292 adult Saskatchewan patients with HIV infection was performed. The primary endpoint was the presence of anemia as defined by the World Health Organization. Six variables and their impact on the prevalence of anemia were examined. Rates of anemia investigation and intervention by physicians were also assessed.

**RESULTS:** The current prevalence of anemia in patients with HIV infection is 26.3% (95% CI, [21.2% - 31.4%]). Anemia was present in 40% of patients prior to initiating Highly Active Anti-Retroviral Therapy (HAART) and 14% following effective therapy with a 26% net reduction. Multivariate analysis demonstrated that female sex (OR 2.0; 95% CI, 1.08-3.74; P=0.028), First Nations and Métis ethnicity (OR 2.3; 95% CI, 1.07-4.91; P=0.032), and lack of HAART (OR 3.2; 95% CI, 1.75-5.99; P=0.002) were associated with significantly increased risk of developing anemia. HCV co-infection (OR 1.9; 95% CI, 0.96-3.75; P=0.065), other ethnic minorities (OR 3.35; 95% CI, 0.85-13.15; P=0.083), and rural habitation (OR 1.2; 95% CI, 0.62-2.17, P=0.652) were not associated with increased risk of developing anemia. Of patients with anemia, 18.4% (95% CI, [9.5% – 27.3%]) had appropriate investigations and 16% (95% CI, [7.5% – 24.5%]) had a documented anemia management plan.

**CONCLUSION:** Anemia is a negative prognostic marker that is prevalent in the Saskatchewan population with HIV infection. Female sex, First Nations and Métis ethnicity, and lack of HAART are significant risk factors for developing anemia. More aggressive strategies are required to investigate and manage anemia in patients with HIV.

**SP24** RAPID IDENTIFICATION OF BACTERIA IN POSITIVE BLOOD CULTURES USING SMUDGE PLATE PREPARATION AND MALDI-TOF MS SYSTEM

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**OBJECTIVES:** Matrix-assisted laser desorption ionization time of flight mass spectrometry (MALDI-TOF MS) is a novel method for rapid identification of bacteria and fungi in clinical microbiology laboratories. We evaluated the performance of MALDI-TOF MS for the rapid identification of bacteria directly from positive blood cultures using “smudge plate” preparations.

**METHODS:** Blood cultures were incubated in the Bactec 9240 system (Becton Dickinson). When a blood culture flagged as positive, a 3 mL aspirate from the blood culture bottle was obtained, microorganisms were separated from blood cells and concentrated in a serum separator. The bacterial concentrate was then plated as a lawn on a chocolate agar (“smudge”) plate and incubated at 35 °C in CO2. After 1, 2 and 4 hours incubation, organisms were recovered by sweeping the smudge plate, and inoculated on a target slide for identification by Vitek MS (BioMérieux) MALDI-TOF. The identification from MALDI-TOF MS was compared to that from conventional phenotypic identification methods. Discrepancies were resolved using MALDI-TOF MS on the colony isolated from the routine subculture and ultimately, the 16S rRNA sequencing method.

**RESULTS:** We prospectively examined 400 positive clinical blood cultures with 248 Gram positive isolates and 152 Gram negative isolates. The rate of correct identification was 90% overall to both species and genus level, 97% for Gram negative isolates, 85% for Gram positive isolates. It also correctly identified all 6 anaerobes. Among the correctly identified isolates, 72% of them were identified with 1 hour incubation, and 87% with 2 hours incubation, and additional 13% after 4 hours incubation. The rate of discordant ID was 1.8% and no ID 8.5%, mainly from Gram positive isolates.

**CONCLUSIONS:** In conclusion, combination of smudge plate preparation and VITEK MALDI-TOF MS system is a simple and efficient way to provide identification of bacteria directly from positive blood cultures within 1 to 4 hours.
SP25

ASSESSMENT OF THE MANAGEMENT OF SUSPECTED URINARY TRACT INFECTIONS FOR IDENTIFICATION OF TARGETED ANTIMICROBIAL STEWARDSHIP INTERVENTIONS

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BACKGROUND: Inappropriate management of suspected urinary tract infections (UTI), such as treating asymptomatic bacteriuria (AB), leads to emergence of antimicrobial resistance and secondary complications including infection with Clostridium difficile.

OBJECTIVES: This study examined empiric antimicrobial selection for suspected UTI to identify target interventions for stewardship initiatives.

METHODS: Retrospective chart review was completed for 344 consecutive inpatient urine cultures. Data were collected regarding clinical indications and rationale for empiric UTI antimicrobial therapy.

RESULTS: 278 patients met eligibility criteria and underwent chart review. 190 (68.3%) patients were female, and average age was 73.7 years. 224 (80.6%) patients were given antimicrobial therapy. The three leading empiric antimicrobials prescribed were ciprofloxacin (n=71, 31.7%), nitrofurantoin (n=39, 17.4%), and trimethoprim-sulfamethoxazole (n=34, 15.2%). Detailed review of the medical record led to a diagnosis of AB in 170 (61.2%) patients. Of patients diagnosed with AB, 119 (70.0%) were treated with antimicrobials. In 37 (13.3%) of the charts reviewed, the only documented abnormality was a nursing note of abnormal urine color or odor. This led to subsequent AB treatment in 25 (67.6%) of patients. 78 (28.1%) charts reviewed did not have a documented physician order for urine submission. The presence of an MD order was significantly associated with appropriateness to submit urine for culture (46.0% vs 24.4%, P=0.002).

CONCLUSIONS: Our results indicate high levels of fluoroquinolone use and high levels of AB treatment, driven in large part by inappropriate and non-physician directed specimen submission. These results will be used to design and prospectively evaluate an institutional UTI stewardship bundle directed at reducing fluoroquinolone use and AB treatment, which will involve physician and nurse education, as well as pharmacy and lab-based interventions.

SP26

CLINICAL FEATURES AND OUTCOMES OF INFLUENZA INFECTION IN COMMUNITY DWELLING OLDER ADULTS PRESENTING TO SIX URBAN EMERGENCY DEPARTMENTS

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OBJECTIVES: To describe the clinical features, outcomes of influenza infection and health care utilization for individuals 60 years and older with laboratory-confirmed influenza presenting to emergency departments (ED) in Ontario.

METHODS: We identified community-dwelling older adults presenting to six EDs with influenza-compatible symptoms during the 2011/12 and 2012/13 influenza seasons. Clinical characteristics were collected from patient interviews and chart review. Influenza vaccination history was collected; nasopharyngeal swabs were tested for influenza. Follow-up was conducted with cases up to 30 days after symptom onset.

RESULTS: Of 1337 participants, 147 (11%) had flu (94 AH3N2, 12 AH1N1, 4 A (not subtyped), 37 B). The most commonly reported symptoms were cough (93%), fatigue (92%), (84%) and weakness; 44% of patients had a measured temperature ≥37.5°C at time of triage. CDC and PHAC influenza-like illness (ILI) definitions captured 32% and 23% of all cases, respectively. Among cases, 83 (55%) were hospitalized with a median length of stay of 5 days (IQR: 3-7 days); four patients (3%) died. Of the 101 patients at home on day 30 from symptom onset, 28 (28%) were not well enough to return to regular activities. Median time to return to regular activities was 18 days (n=134; range 2 to >30 days).

CONCLUSIONS: Older adults commonly present with influenza with non-specific symptoms (such as cough, weakness & fatigue) and the standardized definitions fail to capture the majority these individuals. The majority of older adults with influenza who present to the ED are hospitalized, and recovery is prolonged.

SP27

ENGAGING INFECTIONS DISEASES AND MICROBIOLOGY RESIDENTS IN IMPROVING THE VALUE OF HEALTHCARE: CHOOSING WISELY

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BACKGROUND: Rising costs present a major threat to the sustainability of healthcare delivery. Resource stewardship is increasingly becoming an expected competency of physicians. We used the Choosing Wisely framework to introduce resource stewardship at a national educational retreat for Infectious Diseases (ID) and Microbiology (M) residents.

METHODS: During the 2014 Annual Canadian ID and M Resident Retreat in Toronto, we engaged ID (n=50) and M (n=17) residents from 11 Canadian universities to participate in a modified Delphi panel. Participants were asked in advance of the retreat to submit up to five practices that ID and M specialists should not routinely perform due to lack of proven benefit and/or potential harms to patients. Submissions were discussed in small and large group forums using iterative, electronic polling until consensus was reached on five practices. A follow-up survey at two-months was performed.

RESULTS: Consensus was reached on the following practices that were considered to be of low-value and may be considered for elimination: (i) repeat diagnostic imaging in all cases of vertebral osteomyelitis; (ii) serial CD4 measurements in virologically-controlled HIV patients; (iii) tuberculin skin testing in the diagnosis of active tuberculosis; (iv) transesophageal echocardiography in all cases of nosophomital Staphylococcus aureus bacteremia; and (v) use of parenteral antibiotics when highly-bioavailable oral antibiotics options are available in appropriate clinical situations. Twenty participants (32%) completed the follow-up survey. The majority of respondents (75%) felt that the session was at least as relevant as other sessions at the retreat, with 95% indicating that at least some of the material discussed was new to them. Since returning to their home institutions, 45% of respondents have incorporated what they learned into their daily practice; 4 (20%) reported that they have considered initiating a project related to the session; and 1 (5%) reported having initiated a project.

CONCLUSIONS: This educational forum demonstrated that trainees can become actively engaged in the identification and discussion of low-value practices and guidelines. Embedding teaching about resource stewardship into residency training programs, as demonstrated at our national retreat, will be necessary to improve the value of care offered by the future members of our profession.

SP28

INCIDENCE AND OUTCOMES OF HOSPITAL-ACQUIRED C. DIFFICILE INFECTION AT A LARGE COMMUNITY HOSPITAL BEFORE AND AFTER A MULTIPRONDED PREVENTION AND MANAGEMENT INITIATIVE

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OBJECTIVES: The aim of this study is to determine the impact of a multifaceted intervention, with an emphasis on antimicrobial stewardship, to prevent and manage hospital-acquired C. difficile infection (HA-CDI).

METHODS: We conducted a retrospective before and after cohort study to evaluate the impact of a multimodal intervention to reduce C. difficile infection at a large, community hospital. The intervention included initiating a facility-wide C. difficile outbreak and utilizing an Incident Management Team to rapidly engage administrators and leaders and to disseminate education to staff. Other elements during the outbreak included: enhanced environmental services; “kitkit” use by nursing staff to help assess and manage potential cases;
and a C. difficile pre-printed order set. Antimicrobial Stewardship (ASP) utilizing audit-feedback was instituted on high-risk medical wards. ASP is the only intervention that remained fully active after the outbreak was declared over. We compared the pre-intervention period of July 2010-September 2012 with the post-intervention period of October 2012-July 2014. HA-CDI was defined according to public reporting definitions in Ontario. Incident data was acquired from an Infection Control database. Detailed chart review was performed to gather demographic, comorbidity, treatment and outcome data.

**RESULTS:** A total of 173 cases of HA-CDI occurred during the study period. A reduction in HA-CDI incident rates was observed after implementation of the intervention (0.29 vs. 0.17 per 1000 patient-days; P<0.001). Overall all-cause (0.62 vs. 0.19 per 10,000 admissions; P<0.001) and attributable (0.36 vs. 0.09 per 10,000 admissions; P<0.05) 30-day mortality in patients with HA-CDI decreased significantly post-intervention. Overall hospital-wide antimicrobial use decreased significantly (183 vs. 148 DOT per 1000 patient-days; P<0.001). No significant differences in demographics, comorbidities, or appropriateness of treatment were noted before and after the intervention.

**CONCLUSIONS:** A multimodal intervention resulted in a significant, sustained decline in incident HA-CDI cases at a large, community hospital. A significant sustained decrease in antimicrobial utilization likely helped to maintain lower HA-CDI rates.

**SP29**

**THE CONTINUED RISE OF EXTENDED-SPECTRUM B-LACTAMASE-PRODUCING ESCHERICHIA COLI IN CANADIAN HOSPITALS: CANWARD 2007-2013**

A Denisuik1, HJ Adam1,2, P Lagacé-Wiens1,2, P Sinmer2, MR Mulvey1,3, M Baxter1, M Gilmour1,3, JA Karlowsky1,2, DJ Hoban1,2, GG Zhanel1

1University of Manitoba; 2Diagnostic Services Manitoba; 3National Microbiology Laboratory, Winnipeg, MB

**OBJECTIVE:** To assess the prevalence, patterns of antibiotic resistance, and molecular characteristics of ESBL-, AmpC-, and carbapenemase-producing EC and KPN isolated from Canadian hospitals.

**METHODS:** 6,606 EC and 2,058 KPN were collected from January 2007 to December 2013 as part of the ongoing CANWARD national surveillance study. Antimicrobial susceptibility testing was performed according to CLSI guidelines and putative ESBL-, AmpC-, and carbapenemase-producers were identified. All putative isolates were characterized by PCR and sequencing to detect resistance genes and by PFGE to assess clonal spread. The EC ST131 clone was identified by an allele-specific PCR for the pabB gene.

**RESULTS:** The prevalence of ESBL-EC [2007: 3.4%, 2013: 9.5%], AmpC-EC [2007: 0.7%, 2013: 3.1%], and ESBL-KPN [2007: 1.5%, 2013: 5.7%] increased significantly during the study period, with all three antibiotic resistant organisms reaching peak incidence in 2013. Antimicrobials demonstrating the greatest activity against ESBL-EC, AmpC-EC, and ESBL-KPN in this study were colistin, amikacin, ertapenem, and meropenem, whereas 78.8%, 34.9%, and 66.7% of ESBL-EC, AmpC-EC, and ESBL-KPN, respectively, were multidrug resistant. The prevalence of the ST131 clone was higher in ESBL-EC (56.9%) compared to AmpC-EC (31.7%; P<0.001). CTX-M-15 was the dominant genotype in both ESBL-EC and ESBL-KPN (66.5% and 48.0%, respectively), whereas the dominant genotype in AmpC-EC was CMY-2 (53.2%). KPC-3 represents the dominant genotype among carbapenemase-producers (n=4). In total, 5 isolates demonstrating a meropenem MIC ≥1 µg/mL were collected in 2013, this in comparison to a total of 5 such isolates for the years 2009 to 2012 combined.

**CONCLUSIONS:** The prevalence of ESBL- and AmpC-producing EC and KPN increased significantly between 2007 and 2013. The prevalence of carbapenem-resistant Enterobacteriaceae remains low in Canada.

**SP30**

**INPATIENT CARE OF COMMUNITY ACQUIRED PNEUMONIA: OPPORTUNITIES FOR IMPROVEMENT**

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**OBJECTIVE:** Community acquired pneumonia (CAP) remains a common reason for hospitalization and guideline concordant care may be associated with improved outcomes. We describe a local population of patients with CAP to identify areas for quality improvement.

**METHODS:** We conducted a retrospective review of 130 consecutive adult patients with CAP admitted between January 2013 and June 2014 to our clinical teaching units (97 beds of 917 total). Patients were identified through our antimicrobial “self-stewardship” program wherein the indication is documented for all antibiotics. Empyema and healthcare-associated pneumonia were excluded.

**RESULTS:** The median age was 75 (IQR 61-83) with a median length of stay 9.5 days (IQR 6-14). The most common comorbidities were chronic obstructive pulmonary disease (33%), diabetes (29%) and coronary artery disease (25%). Hypoxia (36%), tachycardia (11%) and hypotension (5%) were rare on presentation. The chest x-ray was interpreted as lobar consolidation in 68 (52%). The PSI was greater than 3 in 74 (57%). Blood and sputum cultures were rarely positive (7/83, 8% and 17/52, 33%). Streptococcus pneumoniae was the most common organism (20%). Viral PCR was positive in 20 of 81 (62%). Antibiotics were administered within 8 hours in 82% and were guideline-concordant in 34%. A common “transgression” was the use of piperacillin-tazobactam in 25%. Therapy exceeded 7 days in 67%. C. difficile infection occurred in 13 (10%), ICU transfer in 6 (4.6%), and inpatient death in only 4 (3.1%). All deaths occurred in those with a PSI of 4 or greater.

**CONCLUSION:** We describe a cohort of patients admitted with CAP to our institution. Microbiological diagnosis was rare and departures from our institution’s guidelines were common. Outcomes were excellent with few deaths or transfers to ICU but C. difficile was a frequent and potentially avoidable morbidity.

**SP31**

**FEASIBILITY OF UV-C LIGHT DISINFECTION IN A CANADIAN HOSPITAL**

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1McMaster University; 2Hamilton Health Sciences, Hamilton, ON

**BACKGROUND:** Devices using UV light have been shown to be effective in deactivating multiple microorganisms and potentially reducing the transmission of pathogens in hospitals. We conducted a study to evaluate a pulsed xenon ultraviolet (PX-UV) technology disinfection system (Xenex®) to evaluate feasibility of use and its impact on hospital acquired infections (HAI).

**METHODS:** The study took place on two internal medicine units of 35 and 31 beds respectively, including 4 private rooms each. The plan was to deploy the PX-UV machine (provided by Xenex®) following routine environmental cleaning for all isolation discharge and all patient bathroom days for 6 months on each unit while the other ward served as a control, with a cross-over for another 6 months. Bed spaces in multi-bed rooms were isolated using opaque blanket before each disinfection to allow utilization of PX-UV while the room was not vacated. The efficacy was measured by comparing the rate of HAI.

**RESULTS:** Feasibility was hampered by the fact that hanging an opaque blanket before disinfecting non-private rooms was too time consuming. Therefore, the machine was used almost exclusively for private rooms after terminal cleaning and all bathrooms daily. Also, due to staff limitation, the machine could only be used between 8am and 4pm on business days. No difference in the rates of HAI were detected with or without the intervention (Table 1).

**TABLE 1**

<table>
<thead>
<tr>
<th>Mean HAI rate</th>
<th>Intervention group</th>
<th>Control group</th>
<th>P value (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clostridium difficile infection</td>
<td>1.03</td>
<td>0.71</td>
<td>3.67 (−0.59, +1.21)</td>
</tr>
<tr>
<td>Methicillin resistant Staphylococcus aureus</td>
<td>1.28</td>
<td>0.82</td>
<td>1.04 (−1.04, +1.95)</td>
</tr>
<tr>
<td>Vancomycin-resistant enterococcus</td>
<td>3.60</td>
<td>2.71</td>
<td>0.15 (−1.69, +3.48)</td>
</tr>
</tbody>
</table>

**CONCLUSION:** Because of the significant percentage of non-private rooms, deployment of the (PX-UV) disinfection system proved to be impractical and did not affect HAI outcomes in our institution.
SP32
NON-POLO ENTEROVIRUS TYPES ASSOCIATED WITH ACUTE FLACCID PARALYSIS – A SYSTEMATIC REVIEW
S Suresh, J Robinson, S Forgé
University of Alberta, Edmonton, AB

OBJECTIVES: With polio vaccination programs having almost eliminated wild type poliovirus as a cause of acute flaccid paralysis (AFP), there is increased interest in AFP due to other enteroviruses (EV) such as EVD 68. This study aims to characterize the nonpolio enteroviruses (NPEV) associated with acute flaccid paralysis (AFP), transverse myelitis (TM) or Guillain Barre Syndrome (GBS) through a systematic review of all published cases.

METHODS: PubMed, MEDLINE, EMBASE and Web of Science databases were searched from Jan 1960 to Sept 2014 with the search terms “human”, “enterovirus”, “paralysis”, “paralytic”, “transverse myelitis” and “Guillain Barre Syndrome.” Articles were included if they described one or more cases of suspected AFP with concurrent detection of NPEV by any method from any site. Articles were excluded if they studied only poliovirus, viruses other than enteroviruses, or only clinical presentations not consistent with AFP such as meningocencephalitis, hand-foot-mouth disease, and hemorrhagic conjunctivitis.

RESULTS: The search yielded 1618 articles of which 148 met the inclusion criteria. The majority of reported NPEV AFP cases occurred outside North America, particularly in Asia. At least 50 different NPEV subtypes have been associated with AFP, with the most common being EV 71, Echovirus 11, Echovirus 6, Echovirus 9 and Coxsackie B3. EVD68, in the time frame of the study, was not commonly associated with AFP. Of those reports that detailed clinical followup, the majority of patients had residual paralysis.

CONCLUSION: NPEV AFP cases have been predominantly reported in Asia, with EV 71 as the most commonly implicated subtype. EVD 68, though a recent concern, has not historically been associated with AFP. Standardized AFP surveillance programs may help identify the epidemiology and enterovirus subtypes related to AFP in North America.

SP33
EVALUATION OF MATRIX ASSISTED LASER DESORPTION IONIZATION-TIME OF FLIGHT MASS SPECTROMETRY (MALDI-TOF MS) FOR YEAST IDENTIFICATION
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BACKGROUND: For most yeast species, antifungal susceptibility can be predicted based on yeast identification. MALDI-TOF was applied in this study to assess its accuracy when compared to traditional phenotypic methods for rapid yeast species identification.

METHODS: The microbiology database was used to identify yeast isolates recovered from blood and other sterile sites from January 2012 to December 2014. Isolates were sub-cultured twice to BAP (Oxoid, Basingstoke, UK), then to Brilliance Candida (Oxoid- Basingstoke, UK) for presumptive identification. Final identification was confirmed using microscopic and biochemical tests. MALDI-TOF MS (Bruker Daltonics, Basingstoke, UK) was used for presumptive yeast species identification. Identification of yeast species by MALDI-TOF is accurate and reliable. The rapid identification of yeast by MALDI-TOF compared to conventional methods will substantially improve fungal diagnostics and choice of appropriate antifungal therapy.

RESULTS: This study tested 62 Candida and 10 Cryptococcus isolates. Concordant identification between phenotypic methods and MALDI-TOF was 100% to genus and 95.8% to species level (Table 1). Conventional methods failed to identify one isolate and ITS later confirmed MALDI-TOF identification.

CONCLUSIONS: Identification of yeast species by MALDI-TOF is accurate and reliable. The rapid identification of yeast by MALDI-TOF compared to conventional methods will substantially improve fungal diagnostics and choice of appropriate antifungal therapy.

TABLE 1
Summary of Maldi-TOF identification results

<table>
<thead>
<tr>
<th>Yeast Species (n)</th>
<th>C. albicans (29)</th>
<th>C. glabrata (12)</th>
<th>C. parapsilosis-s (12)</th>
<th>C. tropicalis (5)</th>
<th>C. lusitaniae (3)</th>
<th>C. krusei (2)</th>
<th>C. guilliermondii (1)</th>
<th>C. gattii (3)</th>
<th>C. neoformans (7)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phenotypic ID</td>
<td>29</td>
<td>12</td>
<td>12</td>
<td>5</td>
<td>3</td>
<td>2</td>
<td>No reliable id</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td>Maldi-spot id</td>
<td>26</td>
<td>11</td>
<td>8</td>
<td>3</td>
<td>3</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>7</td>
</tr>
<tr>
<td>Maldi-tube id *</td>
<td>3</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>-</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>-</td>
</tr>
<tr>
<td>ITS sequencing</td>
<td>-</td>
<td>-</td>
<td>2</td>
<td>-</td>
<td>-</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

*Tube extraction performed only for isolates failed to achieve reliable id scores with spot extraction.
HOSPITALIZED INFLUENZA PATIENTS DURING 2013-2014: A COMPARISON OF ICU ANDWARD TREATED PATIENTS INCLUDING ANTIMICROBIAL THERAPY, ADVERSE EVENTS, AND OUTCOMES

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OBJECTIVES: To describe the epidemiology of hospitalized patients with laboratory confirmed influenza infection during the 2013-2014 season at a single tertiary care hospital; to compare ICU and non-ICU patients; and to specifically describe antimicrobial therapy, outcomes and adverse events in the ICU cohort.

METHODS: Laboratory and epidemiologic data were collected at the University of Alberta Hospital through the Serious Outcomes Surveillance Network (SOS). Additional detailed diagnostic, clinical and outcome data were collected by retrospective chart review of the ICU cohort and stratified by diagnostic features association with bacterial infections.

RESULTS: Of 96 hospitalized influenza patients, 39 (41%) required ICU care, 4 (4%) required extracorporeal support, and 6 (6%) patients died. H1N1 was the predominant strain in both ICU (79%) and ward (69%) patients. ICU patients were less likely to be vaccinated (2; 5% vs. 14; 25%); however vaccination status was unknown in a large number (49; 51%) of patients. ICU patients were younger, with a higher BMI and comorbidity burden, had longer hospital stays, and higher rates of anti-bacterial use. Fourteen (36%) ICU patients had bacterial infection on presentation. Twenty-one suspected or proven nosocomial infections were treated, including 3 presumed hospital/ventilator-acquired pneumonia (2 with antifungal therapy for Candida spp. on respiratory culture), 3 central venous catheter infections, 4 episodes of C. difficile infection and 3 cases of clinical sepsis with unknown source. In ICU patients the mean oselatamivir therapy duration was 8.8 days (range 0-19), and initial antibiotic course was 9.2 days (range 0-18), with those classified as low likelihood of bacterial infection receiving 6.8 days (range 0-17). Directed antimicrobial treatment duration was 10 days (range 5-15) for C. difficile and 7.8 days (range 2-23) for other nosocomial infections.

CONCLUSION: Patients admitted to the ICU with influenza infection were younger, had higher BMI and comorbidity burden, and lower vaccination rates. Regardless of risk for bacterial infection, antibiotic use was high in the ICU, with documented adverse outcomes such as C. difficile infection. Risk stratification for bacterial co-infection on admission may identify patients unlikely to benefit from antimicrobials thus minimizing unnecessary use.

SP36 CLINICAL SPECTRUM OF RHINOVIRUS INFECTIONS IN HOSPITALIZED CHILDREN IN ISRAEL – A SINGLE CENTER’S EXPERIENCE

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BACKGROUND: Human Rhinoviruses are among the most common causes for acute respiratory illnesses (ARI), involving both the upper and lower respiratory tract (LRT). Clinical spectrum ranges from mild flu-like illness to severe respiratory compromise necessitating intensive care unit (ICU) admission. A known association exists between rhinoviral infections and childhood wheezing illnesses including asthma.

OBJECTIVES: To estimate for the first time the magnitude and characteristics of Rhinoviral infections in hospitalized patients in Israel.

METHODS: All the respiratory samples that were taken from hospitalized patients with ARI between July 2011 and June 2012, were tested for the presence of rhinovirus and other respiratory viruses by real time RT-PCR. Positive rhinovirus samples were further tested by Sequencing to classify the species. Demographic data, past and chronic illnesses, signs and symptoms of current illness, vital signs, laboratory tests, imaging and treatment were all documented from the patients’ records.

RESULTS: 245 children tested positive for rhinovirus, which was perenially detected. In 44% a viral co-infection, most commonly adenovirus was present. 58% had significant co-morbid chronic illness (pulmonary, cardiac and prematurity). Common symptoms were cough (67%), rhinorrhea (59%), shortness of breath (44%), and decreased appetite (39%). Abnormal breathing sounds were found in 55% of the patients. 21% were admitted in ICU. Laboratory data showed monocytosis in 51%, and hypercalcaemia in 61% (findings not previously reported. Chest x-ray was performed in 60% of the patients of them 59% had pathological findings: 63% received antibiotic treatment, 60% needed inhalations, 42% received systemic steroids. LRT involvement was identified in 75% of patients. There weren’t any significant differences between rhinoviral species among these parameters.

CONCLUSION: Rhinovirus infections pose an extensive burden of respiratory illness in hospitalized children. Lower respiratory tract involvement is more common than that reported in the literature. Associated hypercalcaemia is a new finding, as yet, unexplained.

SP37 PROSTHETIC JOINT INFECTIONS: PATTERNS OF PRACTICE FOR DIAGNOSIS, MANAGEMENT, AND OUTCOME

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OBJECTIVES: The IDSA guidelines in 2013 established a standard of management for prosthesis joint infections (PJI); however, implementation of the guideline tends to be challenging given the variability in clinical practice that has developed in the last centuries. We examine PJI cases at our centre comparing the outcomes of cases managed in accordance with the IDSA guidelines versus the outcomes of cases managed via an alternative approach.

METHODS: A retrospective study of PJIs (NHSN/CDC criteria) from 2005 to 2012 at a tertiary care centre was conducted to determine clinical management variability and outcomes of deep and organ/pace PJIs.

RESULTS: Of 8,505 hip and knee arthroplasty cases completed, 283 (3.4%) were subsequently diagnosed with a PJI and 63 (0.7%) cases were identified as deep or organ space PJI. Of these, 22 were knees (PJI rate of 0.2%) and 41 were hips (PJI rate of 0.5%). Eleven (17%) had medical management only, and were excluded. Of those treated with a surgical approach, 28 (44%) underwent irrigation and debridement (I&D), 17 (27%) had a one stage revision, and 7 (11%) underwent two stage revision. In those treated according to the IDSA guideline, the risk for failure was significant lower (11/29 versus 8/11; odds ratio 0.14, 95% Confidence interval 0.03-0.64). In the subgroup of those treated with I&D, 1/19 and 5/9 failed, respectively (OR 0.04, 0.004-0.49). Of the 17 patients who underwent a one stage revision, 1/7 and 3/10 failed, respectively (OR 0.39, 0.03-4.80). Of the 17 patients who underwent a two stage revision, 6 were treated according to the guidelines with one failure (16% failure rate). The other patient had hardware left in place and was not classified due to the surgical complications.

CONCLUSIONS: We found significant variability in the management PJIs, and that patients that were and could be managed in accordance with the 2013 IDSA guidelines had a significant lower failure rate, especially with respect to candidates for I&D. This review provides evidence to support the use of the IDSA guidelines for treatment of PJIs.

SP38 URINARY BIOMARKERS IN PATIENTS WITH CYSTIC FIBROSIS

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1Queen’s University Belfast, Cystic Fibrosis & Airways Microbiology Research Group, Belfast; 2Mologic Ltd, Bedford, UK

BACKGROUND: Urine biomarkers are potentially useful for early detection of lung inflammation or injury and may be a medium to detect pulmonary exacerbations in Cystic Fibrosis (CF). Optimal urinary biomarkers have not been identified and correlation with clinical outcomes has not been established. The aims of this pilot study were to 1) explore whether 24 biomarkers were detectable in the urine 2) determine if the levels of these biomarkers differ between CF patients and healthy individuals and 3) explore if biomarker levels correlate with disease severity data.

Can J Infect Dis Microbiol Vol 26 No 2 March/April 2015
METHODS: CF patients (n=129, aged 6-68 yr), enrolled at the paediatric or adult CF centres in Belfast, provided a total of 146 urine samples (135 when stable, 11 during exacerbation). Samples from healthy individuals (n=34, aged 24-60 yr) were used as controls. Clinical and microbiology data were recorded. Samples were analysed with a combination of in-house (Mologic Ltd) and commercial assays (ELISA, lateral flow, protease substrate assay,zymography).

RESULTS: Every biomarker tested was detectable in urine and 9 were excrated at statistically different levels between patients and controls (Mann-Whitney, P<0.05). No strong and clinically significant correlation between biomarker levels and clinical data was highlighted. However, CF patients infected with methicillin-resistant Staphylococcus aureus (n=3) excrated higher levels of alpha-1 antitrypsin than those infected with Pseudomonas aeruginosa or Burkholderia cepacia complex (P=0.038 and P=0.026). Although not statistically significant, a positive trend in desmosine, an amino acid related to elastin degradation, was found between stable and exacerbation time points (Wilcoxon signed-rank test, P=0.069).

CONCLUSIONS: These results indicate the potential of urine biomarkers in CF. Further longitudinal studies are needed to determine whether fluctuation in biomarker levels correlates to clinical outcome and can be used to predict exacerbation onset.

This study was supported by the UK NHS NOCRI Translational Research Partnership & US Ireland Partnership Grant. E.V. work is supported by AMMI Canada/Pfizer Post-Residency Fellowship and Cystic Fibrosis Canada.

SP39
CARBON SOURCE MODULATES PARADOXICAL GROWTH EFFECT OF CASPOFUNGIN IN CANDIDA ALBICANS
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1University of Alberta; 2Provincial Laboratory for Public Health, Edmonton, AB

BACKGROUND: In vitro susceptibility testing of caspofungin by broth microdilution has revealed an unexplained paradoxical growth (PG) effect in which noticeable growth occurs at concentrations above the minimum inhibitory concentration (MIC) of susceptible isolates. The frequency of the PG effect is variable in different growth media, which vary most notably in the amount and types of carbon. We evaluated the effect of carbon source on the degree of PG in C. albicans to identify differences that may be applied to in vivo models and understand its clinical significance.

METHODS: C. albicans was grown in RPMI or minimal media containing 2% galactose, 2% lactate, and 3% glycerol and with added caspofungin. Cultures were evaluated by time kill analysis. Samples collected over 72 hr were evaluated for growth by colony forming units (CFU) and optical density at 600 nm (OD600).

RESULTS: C. albicans demonstrated a fungicidal effect at 0.075 and 2 mg/L caspofungin in all media. Growth at 16 mg/L caspofungin (PG) was fungistatic by CFU determination. By OD600 evaluation, C. albicans grown in fermentable sugars (RPMI, galactose) showed a statistically significant increase in PG compared to growth in non-fermentable sugars (lactate, glycerol). Large cell aggregates were seen in fermentable sugars, and un-budding yeast in non-fermentable sugars.

CONCLUSIONS: These data support a correlation between carbon source and the degree of PG in C. albicans, where the effect is significantly increased in fermentable carbon sources. There are significant physiological differences between yeast grown at PG and inhibitory caspofungin concentrations. Our observations suggest that further study of PG in different infection models may reveal what role PG plays in sustaining C. albicans in the different host niches where nutrients vary, despite apparent adequate antifungal therapy.

SP40
COST-EFFECTIVENESS OF MALDI-TOF MS IN CLINICAL ANALYSIS OF URINE SPECIMENS IN NOVA SCOTIA
H Al Sidairi1,2, T Hatchette1,2, D Haldane1,2, R Davidson1,2
1Dalhousie University; 2Queen Elizabeth II Health Sciences Centre, Halifax, NS

BACKGROUND: Matrix Assisted Laser Desorption Ionization Time-of-Flight mass spectrometry (MALDI-TOF) is an automated microbial identification system revolutionizing bacterial identification. It is anticipated that the MALDI-TOF system will result in significant cost savings for the laboratory. We sought to evaluate the impact (financial and recovery) of using MALDI-TOF along with sheep blood agar (SBA) for urine cultures as an alternative to our current protocol utilizing Orientation agar.

METHODS: Prior to implementation, we retrospectively examined 3 years worth of urine culture data to determine our negative and positive culture rates and most commonly isolated pathogens. Prospectively, we continue to compare our isolation rates with historical data.

RESULTS: We process approximately 64,000 urine specimens per year of which 77% are negative. E. coli accounts for 55% of our positive isolates followed by K. pneumoniae, Enterobacter spp. and Enterococcus spp. (12% and 11% respectively). When cost of each agar are considered (Orientation $0.62, SBA $0.27), a savings of over $17,000 was realized for our negative cultures. We examined the additional cost of identification (SBA plus MALDI-TOF vs. Orientation agar plus biochemical) and realized another $1,000 in savings. Our prospective study comparing SBA and MALDI-TOF to Orientation agar has yet to demonstrate a significant difference in the isolation rate or identification of any pathogen with one exception. We have noticed a small increase in “significant” Proteus spp., likely due to its spreading nature on SBA compared to Orientation agar.

CONCLUSION: The introduction of MALDI-TOF and the subsequent return to SBA for urine specimens have produced an annual cost savings of $18,000 for our laboratory with no significant impact on isolation rate or identification of urinary tract pathogens.

SP41
ROLE OF TYPE IV PILI IN THE CLOSTRIDIUM DIFFICILE PATHOGENIC PROGRAM
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1University of Calgary, Faculty of Medicine, Calgary, AB; 2University of Maryland, School of Medicine, Baltimore, MD, USA

Clostridium difficile is an anaerobic Gram-positive microorganism responsible for causing C. difficile infections, or CDI. CDI varies from mild diarrhea to life-threatening inflammation of the colon. CDI has been associated with changes in the normal intestinal microflora mainly due to antibiotic therapy affecting especially hospitalized elderly people. The main virulence factors of Clostridium difficile are TcdA and TcdB toxins. However, our recent discoveries demonstrated that C. difficile can assemble Type 4 pili (T4P). T4P are filamentous structures expressed on the surface of bacteria and have been studied in many Gram-negative bacteria where they play an important role in colonization, adherence, biofilm formation and twitching motility. Assembly of T4P requires structural pilin proteins, extension and retraction ATPases, and pre-pilin peptidases, among others. Objective: To use a mutagenic approach to study the function of T4P in the C. difficile pathogenic program.

METHODS: TEM analysis was employed to investigate T4P expression by group II intron-targeted C. difficile mutant strains. Colonization was investigated using the antibiotic-treated mouse CDI model.

RESULTS: TEM analysis revealed that, unlike the wild-type parental strain, C. difficile pilA1 and pilU (pilin subunits), and pilD1 (pre-pilin peptidase) mutants were unable to express T4P. Moreover, initial results revealed no significant differences in the ability of the pilD1 mutant and wild-type strain to persist or cause illness in mice.

CONCLUSIONS: Our results suggest that, although T4P have been shown to play a role in colonization by Gram-negative bacteria, their role in C. difficile pathogenesis and persistence in mice, at least, remains to be determined.
SP42 DETECTION AND CHARACTERIZATION OF A 2014 ESCHERICHIA COLI O157:H7 OUTBREAK IN ALBERTA BY MULTIPLE MOLECULAR METHODS INCLUDING WHOLE GENOME SEQUENCING

B Berenger1,2, C Berry4, C Nadon4, L Tschetter4, P Fack5, S Delannoy5, V Li7, T Peterson1, L Chui1,3
1Alberta Provincial Laboratory for Public Health; 2University of Alberta Department of Medical Microbiology and Immunology; 3University of Alberta Department of Laboratory Medicine and Pathology, Edmonton, AB; 4Public Health Agency of Canada National Microbiology Laboratory, Winnipeg, MB; 5Food Safety Laboratory, Maisons-Alfort, France

OBJECTIVES: Over a three month period in 2014 a large E. coli O157:H7 outbreak occurred in the province of Alberta, Canada. We sought to characterize this outbreak using different molecular typing techniques including whole genome sequencing. A standardized method of whole genome analysis for determining strain relatedness has yet to be established, therefore in the context of this outbreak, we assessed the ability of core genome single nucleotide polymorphisms (SNPs) and K-mer phylogenies to discriminate between outbreak and sporadic E. coli O157:H7 isolates.

METHODS: We selected isolates received for molecular typing between July 10 and September 22, 2014 in addition to isolates selected from a 2012 beef-associated outbreak. Pulsed-field gel electrophoresis (PFGE), multi-locus variable number tandem repeat analysis (MLVA), and shiga toxin (stx) subtyping were performed as per standardized protocols. Real-time PCR dynamic arrays were used to detect the presence or absence of 49 shiga toxin-producing E. coli virulence genes. Genome assembly and annotation were done using SPAdes and Prokka. Core genome SNP phylogenies were generated using the National Microbiology Laboratory bioinformatics SNVpyl pipeline.

RESULTS: 155 patient isolates were received: 10 from a small outbreak with indistinguishable PFGE/MLVA profiles and 111 from the 2014 outbreak comprising 23 with closely-related PFGE/MLVA profiles. All isolates from the 2014 outbreak were stx1 and stx2. No differences in virulence gene profiles were observed between the outbreak and sporadic isolates. In concordance with PFGE and MLVA, both core genome SNP and K-mer phylogenies clustered the 2014 and 2012 outbreaks as distinct events.

CONCLUSIONS: Core SNP and K-mer phylogenies are appropriate methodologies to detect and investigate outbreaks of E. coli O157. Before whole genome analysis can be implemented for routine public health use, issues surrounding cost, technical expertise, software standardization, and data sharing/comparsions need to be addressed. Despite the unique characteristics of the 2014 E. coli O157 outbreak (ie. source, multitude of PFGE/MLVA profiles and size), no unique virulence gene profile could be identified.

SP43 HEAT RESISTANT SHIGA TOXIN-PRODUCING ESCHERICHIA COLI (STEC) FROM CLINICAL CASES IN ALBERTA

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OBJECTIVES: Foodborne disease due to the consumption of foods contaminated with Shiga toxin-producing E. coli (STEC) continues to be a global public health concern. With mechanical tenderization of meat in food processing plants, there is the potential for inoculation of STEC into deep, sterile tissue. Undercooked meat that does not reach temperatures capable of inactivating STEC further escalates the risk of disease. Environmental strains of E. coli and other Proteobacteria have demonstrated heat resistance but there is no evidence of heat resistant STEC involved in clinical disease.

METHODS: Heat resistance studies were performed on selected Alberta STEC isolates from 2000 to 2014. These isolates were inoculated in Luria-Bertani (LB) broth with sodium chloride (NaCl) concentrations of 1%, 2%, and 4%, reflective of NaCl concentrations used in meat preservation. Recovered cells were exposed to 60°C heat treatment in a water bath for 15, 30, 45, and 60 minutes. Survival was quantified by colony enumeration of heat treated isolates on LB agar and those yielding 50 or more colonies were considered heat resistant. Protein gel electrophoresis was performed on all heat resistant isolates and compared to heat sensitive isolates. Isolates were exposed to 37°C and 60°C heat treatment for 60 minutes prior to protein denaturation to investigate differences in protein expression between isolates and whether expression was inducible or constitutive.

RESULTS: Heat resistance was confirmed in three clinical STEC isolates. Serotypes of the these STEC isolates were ONT:H25, O11:H30, and O157:H3. All positive isolates survived 60°C heat treatment for greater than 60 minutes. Serotype O11:H30 displayed greater heat resistance in 1% NaCl whereas 4% NaCl was favoured for serotypes ONT:H30 and O157:H3. Upon protein gel electrophoresis, evidence suggests that positive isolates express additional protein bands at 35 kda and 110 kda that were visualized both in isolates exposed to 37°C and 60°C heat shock.

CONCLUSIONS: Heat resistant STEC involved in clinical infection in Alberta was confirmed by in vitro studies. Positive isolates displayed the ability to survive 60°C heat treatment for greater than 60 minutes. The presence of heat resistance associated with STEC plays an important role in food safety, especially in determining the duration of cooking temperatures.
THE ROLE OF INTER-HOSPITAL SAMPLE TRANSPORT IN TOTAL ASSAY TIME FOR MICROBIOLOGICAL SCREENING TESTS

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BACKGROUND: McGill University Health Centre uses PCR to screen all admitted medical patients for MRSA and VRE, and if symptomatic, for respiratory viruses. PCR can yield same day results, which helps to optimize isolation precautions and patient flow. However, with planned centralization of laboratories, such as in Quebec, there is concern that inter-hospital transport of samples may retard results, impacting patient flow.

METHODS: The Royal Victoria Hospital sends its screening samples to the Montreal General Hospital, 1.7km away. MRSA, VRE and respiratory virus PCR screenings for admitted medical patients at the Royal Victoria Hospital were analysed to determine the time required for each step of the screening process and their contribution to total assay time.

RESULTS: Data from 100 admitted to internal medicine was recorded and pertinent results are summarized in Table 1. Importantly, the time between sample collection and receipt in lab increased significantly after 15:00 with the time spent “in transport” making up a sizable proportion of the overall analytic time. This led to a substantial delay in result availability and potentially caused prolongation of unnecessary isolation precautions and delays in the transfer of patients from the emergency room to the inpatient units.

CONCLUSIONS: The transportation of specimens between sites contributes significantly to the overall turn-around time and as laboratory technical time decreases, the importance of this contribution increases. It thus represents an important barrier to the timely completion of screening assays as even short inter-hospital transportation may be a rate limiting step in the timely completion of assays. Such delays ought to be considered when planning for a centralized laboratory system such as Quebec’s Optilab project.

| TABLE 1 |
| Test performance characteristics and contribution of transport time to assay time |
| | MRS | VRE | Resp. Virus |
| Positive Rate (%) | 6.7 | 11.0 | 40.4 |
| Mean Total Turn-around Time (hours) | 44.2 (±22.2) | 51.1 (±23.8) | 18.2 (±10.8) |
| Average Proportion of Time spent “in transport” (%) | 28 | 14 | 60 |

INVESTIGATION OF MATRIX ASSISTED LASER DESORPTION/IONISATION TIME-OF-FLIGHT MASS SPECTROMETRY (MALDI-TOF MS) FOR DETECTION OF CLOSTRIDIUM DIFFICILE TOXIN A AND B FROM STOOL SAMPLES

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OBJECTIVE: This study aims to investigate the use of MALDI to directly detect C. difficile toxin A and B from stool samples.

METHODS: Stool samples sent to Royal Inland Hospital that were tested positive for C. difficile by glutamate dehydrogenase and toxin were diluted 10-fold in deionized water or phosphate buffered saline and vortexed to create a relatively homogeneous suspension. Samples were then centrifuged and the pellet removed. Proteins in the supernatant were precipitated with acetonitrile or ammonium sulfate and the solution was centrifuged again. The pellet was then resuspended in deionized water or TA30 and spotted on a MALDI target plate with a sinnapinic acid or SDHB matrix co-crystalized.

RESULTS: MALDI analysis showed no difference between samples diluted in deionized water and those resuspended in phosphate buffered saline. Protein precipitation with acetonitrile produced higher quality spectra than protein precipitation with ammonium sulfate. Sample co-crystalization with sinnapinic acid provided higher quality spectra than SDHB and no difference in spectra quality was seen between samples resuspended in deionized water and those resuspended in TA30. MALDI analysis showed a peak at 56 kDa in four samples. No peaks were seen in the 63 kDa range in any of the samples. It is speculated that this 56 kDa protein is a metabolite of C. difficile toxins. Autocleavage of a commercially known toxin A also failed to show the expected peak at 63 kDa.

CONCLUSION: These findings show that we were unable to use MALDI to directly detect C. difficile toxin A and B from crude stool protein extracts. Further studies would be required to ascertain the possibility of using this technological tool to detect C. difficile toxin as an alternative method of diagnosis to the tests currently available. Although inconclusive, this study is a starting point for the use of MALDI as a diagnostic tool in a clinical setting.
80% of the children were able to keep their lines for >30 days post repair. Series, there were no CVC BSI in the immediate post repair period, and conclusions: It is difficult to draw conclusions about the blood-

tion within 30 days of line repair. Six had a positive cultures- 3 were positive in the pre-

period 5/10 were on broad spectrum antibiotics. 0/10 were on antibiotics 4 received continuous or intermittent TPN. In the pre-repair (30 day) nal atresia and short gut syndrome. Nine were frequently neutropenic, and

diagnosis, antibiotic exposure, total parenteral nutrition (TPN), immune status, organisms isolated and resistance patterns, other complic-

ations and date of CVC removal after repair was collected. Bloodstream infection data was obtained from the Infection Prevention and Control database (prospectively collected data using NHSN Definitions for CVC BSI) and laboratory database. Primary outcome was the rate of CVC BSI post repair and the secondary outcomes were risk factors affecting the inci-

dence of CVC BSI post repair.

RESULTS: Ten patients were included (age range 2-16 years, 7 males): Nine had malignancies (3- AML, 1-ALL, 1- neuroblastoma, 1- epedy-

mona, 2- medulloblastoma and 1- Burkitt’s Lymphoma), one had intesti-

nal atresia and short gut syndrome. Nine were frequently neutropenic, and 4 received continuous or intermittent TPN. In the pre-repair (30 day) period 5/10 were on broad spectrum antibiotics. 0/10 were on antibiotics 30 days post repair. Six had a positive cultures- 3 were positive in the pre-

repair (30 days) period (E. coli, viridans group streptococci and M. catarrhalis), and 3 were positive >30 days after CVC repair (S. maltophilia, Leuconostoc sp. and Diphtheroids). Eight of the repairs were successful (i.e. line retained >30 days). None of the patients had clinical deteriora-

tion within 30 days of line repair.

CONCLUSIONS: It is difficult to draw conclusions about the blood-

stream infection risk after CVC repair in children. However, for this case series, there were no CVC BSI in the immediate post repair period, and 80% of the children were able to keep their lines for >30 days post repair. A prospective study with larger enrolment would be more helpful in pre-

dicting the short and long-term complications of CVC repair.

SP49
ACUTE ODONTOGENIC INFECTION MANAGEMENT IN THE OPAT CLINIC SETTING: A QUALITY IMPROVEMENT STUDY IN CALGARY, ALBERTA

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BACKGROUND: Acute odontogenic infections (AOI) account for an estimated 60% of all non-traumatic dental emergencies and over six hun-

dered thousand visits to Canadian emergency departments each year. The objectives of this study were to quantify and characterize AOI amongst adults presenting to acute medical care and managed through a city-wide out-patient antibiotic therapy program (OPAT). Secondary objectives were to estimate the healthcare costs of AOI management and develop an integrated care pathway.

METHODS: Adult acute medical care and OPAT AOI referrals were quant-

ified using diagnoses codes in a regional reporting system. OPAT referrals consenting to participation in the study completed a questionnaire and had their OPAT medical records reviewed upon completion of care. Through collaboration with local medical and dental services a standardized medical template and referral pathway was developed. The study was conducted in Calgary, Alberta, between February and June, 2014.

RESULTS: Of 704 adult individuals presenting to Calgary area acute care facilities with AOI during the study period 343 [49%] were referred to OPAT [5.5% of all OPAT referrals]. A total of 110 [32%] individuals were included in the study (women 60 [54%], mean age 44 years [range 18-96]). The major-

ity of individuals with AOI were smokers (53 [53%]), had dental insurance (70 [65%]), and reported prior dental infections (84 [77%]). Thirty-nine [46%] reported current AOI as a recurrence. Median monthly pre-tax household income was $4800 with 16% (19) of those responding falling below the Canadian Low-Income Cut-off. Median length of parenteral antibiotic therapy was 3 days (range 1-13, IQR 2) resulting in an average 15-day total course of antibiotics (range 0-35, IQR 4). Estimated total cost saving of OPAT care compared with hospitalization for intravenous portion of antibiotics was $597,434.

CONCLUSIONS: AOI represent a major preventable cause of recurrent morbidity resulting in extended antibiotic exposures and accounting for substantial healthcare costs. OPAT appears to provide a cost-effective model for initial AOI management. Integrated care pathways for AOI may facili-
tate collaboration with dental services allowing timely provision of definitive management and prevent reoccurrence.
METHODS: The multiplex assay runs in two tubes. The first tube detects influenza A and B, RSV and the internal control. The second tube tests for parainfluenza 1, 2 and 3, metapneumovirus and adenovirus. The assay runs on the Rotorgene (Qiagen) real time thermal cyclers which have five detection channels. The crimson channel in tube one was available for rhinovirus/enterovirus detection.

Primers and a probe were designed using sequences from GENBANK and aligned using Clustal Omega. Specificity was checked using NCBI BLAST. The final design has two forward rhinovirus primers and one forward enterovirus primer with one reverse primer and a probe that cover both rhinovirus and enterovirus. Detection of rhinovirus, A, B and C was tested with previously sequenced patient samples. Detection of enterovirus was tested using stock strains of echovirus, enterovirus and coxsackie A and B viruses.

Reaction efficiency and sensitivity were compared by running the multiplex assay at the same time as uniplex assays for each target in the tube. Assay sensitivity was considered acceptable if the crossing threshold (Ct) of the multiplex was within 2 Cts of the uniplex Ct. The LOD of the assay for rhinovirus/enterovirus using enterovirus-D68 as the target is 306 copies/mL. Detection levels for Influenza A and B and RSV were not compromised. External performance assessment challenges are obtained from IQM4, NML and CAP. Data on positivity rates for each virus are tabulated weekly. The expanded assay has been performed 1-3 times daily by the clinical laboratory since September 2013.

OUTCOMES: Since September 2013 we have tested 11,930 respiratory specimens. Of these 1777 (14.8%) were positive for rhinovirus/enterovirus. The most commonly detected pathogen in our population. From August to October 2014 a spike in rhinovirus/enterovirus detection (661 positives) was noted concurrent with reports of enterovirus-D68 (EV-D68) in the USA. An EV-D68 specific assay was quickly developed in September and validated against sequencing results. Testing for EV-D68 was performed within 24 hours of obtaining a positive rhinovirus/enterovirus result. Of 647 rhinovirus/enterovirus positive samples tested in the specific assay, 183 (28.3%) were EV-D68 positive. The addition of rhinovirus/enterovirus to the multiplex assay was welcomed by the clients we serve, especially the pediatric infectious disease physicians and allowed us to react to the emergence of enterovirus-D68 in a timely manner. The addition of the EV-D68 specific assay to the testing protocol was very important to Infection Control at the McMaster Children’s Hospital allowing them to cohort infected children and prevent nosocomial spread of enterovirus-D68.

RESULTS: Only 160 out of the 376 consented patients had an evaluable stool sample for PCR testing. We identified over 10% of tested patients as asymptomatic carriers of C. difficile (17 patients tested positive for C. difficile). Results long with explanation of the significance of C. difficile carriage were communicated to the attending physician. Identifying C. difficile carriers among the most vulnerable patients may offer alternatives to avoid iatrogenic CDI.

CONCLUSIONS: In this preliminary study, we tested and identified C. difficile carriers upon admission to the hospital among a selected group of patients at high risk of developing CDI (oncology patients). Our goal is to extend the screening to all high risk hospital admission for C. difficile to identify carriers at risk of severe disease. Identification of these high-risk patients, combined with hand washing, a strict antimicrobial stewardship program and careful environmental decontamination may help in reducing the rates of CDI significantly.

CV01

ADENOVIRUS INTERSTITIAL NEPHRITIS IN A PATIENT WITH RENAL TRANSPLANT

S Ala

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We present the case of a 65-year-old woman with a history of diabetes mellitus, a CABG in 1996 and end-stage renal failure leading to renal transplant in June 2005. She was well until November 2, 2013 when she developed cough and a decrease in oral intake. She was prescribed Doxycycline for PCP with no improvement. November 8, 2013, she developed weakness and fever and fell. She was admitted for dehydration and hyponatremia, but despite hydration, her creatinine did not return to baseline. All the investigations were unremarkable except for her kidney biopsy, which revealed acute adenoviral (ADV) nephritis (interstitial nephritis). Her ADV PCR was positive at 64,500 copies (blood) and 10,000,000 copies/mL (urine). We reviewed the case management and provide a brief review of the literature.

CV02

A CASE OF CAPNOCYTOPHAGA CANIMORSUS MENINGITIS

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The case is that of a 57-year-old man, presenting with clinical signs and symptoms of bacterial meningitis and treated empirically with ceftriaxone and vancomycin. Though his CSF profile was consistent with bacterial meningitis (WBC of 1329×106/L (90% neutrophils), glucose of 2.6 mmol/L, protein of 2.06 g/L), and the CSF Gram stain demonstrated rare Gram-negative rods, the organism failed to grow in culture, leaving us puzzled as to the etiology of his illness.

After his eighth day of hospitalization, he was feeling well enough to return home for recovery and ongoing therapy with ceftriaxone. After discharge, the result of a 16S ribosomal RNA sequencing of his CSF revealed the etiology of his meningitis as Capnocytophaga canimorsus. Only on subsequent direct questioning after the diagnosis did the patient endorse both a bite from his household dog 1-2 days prior to his illness, and excessive alcohol intake (of approximately 24 bottles of beer per week), both risk factors for systemic disease due to C. canimorsus.

The patient was reassessed in the Infectious Diseases ambulatory clinic two weeks after discharge, and had fully recovered, except for mild hearing loss, after 21 days of ceftriaxone. Further investigation of his liver and spleen health revealed no apparent impairments. It was recommended that he undergo vaccination as would a hyposplenic patient.
ATYPICAL CASE OF ATYPICAL PNEUMONIA: DISSEMINATED MYCOPLASMA HOMINIS INFECTION IN A DOUBLE LUNG TRANSPLANT RECIPIENT

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We present the case of a 67-year-old man who underwent a double lung transplant for end stage interstitial lung disease. On post-operative day (POD) 6, while still intubated, the patient developed a fever and increasing oxygen requirements. There were no other infectious symptoms noted. His past medical history was otherwise significant for coronary artery disease with a prior NSTEMI in 2001. As well, the patient had a history of gastro-esophageal reflux disease and hepatic steatosis. Pertinent medications included tacrolimus 2.5 mg daily, MMF 1500 mg BID and prednisone 60 mg daily. He had been on meropenem perioperatively and continued postoperatively. He was also on fluconazole for a BAL growing scant C. albicans as well as septrin and acyclovir prophylaxis. Investigations were significant for an elevated WBC count 14.6 and mildly elevated LDH 352. Serial chest x-rays showed fluctuating areas of bibasilar atelectasis/consolidation with bilateral pleural effusions. Multiple microbiologic specimens were sent including BAL and chest tube site swabs as well as blood cultures all of which were negative. On POD 6, a BAL specimen submitted on POD 3 showed growth on blood agar plates. The original Gram stain from the specimen was 2+ PMN with no organisms seen. Attempts to perform a Gram stain on the isolate growing on BAP did not reveal any organisms. Given the suspicion of an atypical respiratory pathogen, the isolate was sent to the reference laboratory for identification and azithromycin and moxifloxacin were added to the patient's treatment regimen. The original isolate was identified as Mycoplasma hominis by 16S PCR. This organism was also isolated from several repeat BAL specimens, swabs from the patient's chest tube sites and from a culture of the patient's internal jugular catheter tip, although all blood cultures failed to grow. The patient was diagnosed with disseminated Mycoplasma hominis infection with a respiratory focus occurring post-double lung transplant.
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