The Clinical and Molecular Epidemiology of Noncarbapenemase-Producing Carbapenem-Resistant Enterobacteriaceae: A Case-Case-Control Matched Analysis

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Background. Risk factors and outcomes associated with carbapenem-resistant Enterobacteriaceae (CRE) acquisitions are derived primarily from cohorts consisting of carbapenemase-producing (CP) strains. Worldwide epidemiology of non-CP-CRE is evolving, but controlled epidemiological analyses are lacking.

Methods. A matched case-case-control investigation was conducted at Shamir (Assaf Harofeh) Medical Center, Israel, on November 2014–December 2016. Noncarbapenemase-producing CRE (as defined by the US Clinical and Laboratory Standards Institute Standards) carriers were matched to patients with non-CRE Enterobacteriales and to uninfected controls (1:1:1 ratio). Matched and nonmatched multivariable regression models were constructed to analyze predictors for acquisition and the independent impact of carriage on multiple outcomes, respectively. Representative isolates were whole genome sequenced and analyzed for resistome and phylogeny.

Results. Noncarbapenemase-producing CRE carriers (n = 109) were matched to the 2 comparative groups (overall n = 327). Recent exposure to antibiotics (but not specifically to carbapenems), prior intensive care unit admission, and chronic skin ulcers were all independent predictors for non-CP-CRE acquisition. Acquisitions were almost exclusively associated with asymptomatic carriage (n = 104), and despite strong associations per univariable analyses, none were independently associated with worse outcomes. Genomic analyses of 13 representative isolates revealed polyclonality, confirmed the absence of carbapenemases, but confirmed the coexistence of multiple other genes contributing to carbapenem-resistance phenotype (multiple beta-lactamases and efflux pumps).

Conclusions. Noncarbapenemase-producing CRE acquisitions are primarily associated with asymptomatic carriage, specifically among prone populations with extensive recent exposures to antibiotics. The prevalent mode of acquisition is “emergence of resistance” (not “patient-to-patient transmission”), and therefore the role of stewardship interventions in reducing the spread of these therapeutically challenging pathogens should be further explored.

Keywords. antimicrobial resistance; case-case control; CRE; multidrug resistance; nosocomial infection.

Infections caused by resistant pathogens is a well established and frequently encountered global threat [1]. The World Health Organization and the Infectious Diseases Society of America listed groups of pathogens that pose the greatest epidemiological threat [2, 3]. Among these are Enterobacteriales, the most common human offending pathogens in many regions [4], which are resistant to carbapenems (ie, carbapenem-resistant Enterobacteriaceae [CRE]).

Carbapenem-resistant Enterobacteriaceae created a massive global pandemic, resulting in devastating outcomes among millions of patients [5]. The initial emergence of CRE was marked by the clonal spread of a certain strain, a transposon Tn440-mediated bla_KPC-producing sequence type (ST) 258 Klebsiella pneumoniae [5–7]. The epidemiology of CRE then rapidly evolved, with dissemination of the bla_KPC to other Enterobacteriales [8] and emergence of Enterobacteriales harboring other carbapenemases [9]. However, our knowledge pertaining to the epidemiological features of CRE are derived primarily of cohorts consisting of carbapenemase-producing Enterobacteriaceae (CPE), specifically bla_KPC [5, 9].
After the global spread of CRE, both the US Clinical and Laboratory Standards Institute Standards (CLSI) and the European Committee on Antimicrobial Susceptibility Testing (EUCAST) revised their recommended diagnostic criteria for CRE and CPE, with considerable reductions (ie, 2–3 doubling dilutions) in the breakpoints for defining nonsusceptibility to carbapenems and enabling the diagnosis of CPE carriage solely on the presence of a carbapenemase [10–13]. This suddenly resulted in identification of a “new” group of CREs with no carbapenemases detected: noncarbapenemase-producing CRE (non-CP-CRE).

Resistance mechanisms to carbapenems among non-CP-CRE were attributed to the possible production of extended-spectrum beta lactamases (ESBLs) and/or bla∗CP-CRE were attributed to the possible production of extended-spectrum beta lactamases (ESBLs) and/or $\text{bla}_{\text{ampC}}$-cephalosporinases, coupled with outer membrane modifications such as loss of porin channels or up-regulation of efflux pumps [14]. However, molecular epidemiologically controlled analyses and explorative sequencing data are lacking. In addition, clinical epidemiological analyses of this clinical and microbiological entity are lacking: however, although the mode of acquisition and the transmission dynamics of CPE clearly demonstrated “patient-to-patient” horizontal spread [9, 15], the mode of acquisition for non-CP-CRE is still undetermined [15]. In a recent multicenter, retrospective, noncontrolled analysis from Singapore, recent carbapenem exposure was 3 times more common among patients with new acquisitions of non-CP-CRE compared with patients who acquired CPE [16]. Moreover, a small report suggested that non-CP-CRE new isolations are not commonly clustered in time nor space [17], and these 2 findings suggest that non-CP-CRE acquisition stems from endogenous emergence of carbapenem resistance among susceptible strains rather than “patient-to-patient transmission” [15], but controlled data are lacking. This issue is important, because the mode of acquisition should direct the implementation of appropriate preventive resources [18], ie, enhanced isolation and barrier precautions or stewardship initiatives to curb inappropriate usage of carbapenems or antimicrobials in general [19].

In 2 small epidemiological investigations of non-CP-CRE conducted at Johns Hopkins Hospital [20, 21], the overall 14-day mortality of monomicrobial CPE bloodstream infections ([BSIs] n = 37) was significantly worse compared with non-CP-CRE monomicrobial BSI (n = 46) [20], and non-CP-CRE asymptomatic carriage was significantly less associated with later active infection compared with CPE [21]. These data suggest that the epidemiology of non-CP-CRE differ considerably from that of CPE, and therefore well designed and well controlled epidemiological investigations of non-CP-CRE are warranted, to curb the continued emergence of these therapeutically challenging resistant organisms.

A matched case-case-control design is the preferred method to analyze the epidemiology for acquisitions of multidrug-resistant organisms (MDROs) [22–24]. It enables us to control for multiple confounders and to isolate the parameters that are independently associated with the resistance determinants [22, 23]. Our study aims were to study the clinical and molecular epidemiology of non-CP-CRE acquisitions while implementing matched case-case-control design.

**METHODS**

**Ethics Approval**
The institutional ethics committee followed the Helsinki declaration, approved the study before its initiation, and waved the need to obtain written informed consent from participants, due to the historical retrospective chart-based design.

**Setting and Design**
A retrospective matched case-case-control investigation was conducted among patients from all age groups at Shamir (Assaf Harofeh) Medical Center, central Israel (November 2014 to December 2016). The institutional ethics committee had approved the study before its initiation. The study comprised 3 groups: (1) “resistant case patients” with non-CP-CRE from either a screening or a clinical specimen; (2) “susceptible case patients” with an *Enterobacteriaceae* isolate susceptible to carbapenems; and (3) “uninfected control patients,” ie, patients who were screened and had no documented isolation of *Enterobacteriaceae*. Patients were included in the cohort only once. A susceptible case patient and an uninfected control patient were randomly selected (Excel; Microsoft) and matched to a resistant case in a 1:1:1 ratio. Matching criteria (in established order of importance [22]) included the following: the infection versus asymptomatic colonization status [25]; the species, age group, and “time at risk” and the calendar year. For both resistant and susceptible case patients, the time at risk was defined as the length of stay (LOS) from admission to culture [26]. For uninfected controls, the time at risk was adjusted by multiplying the total LOS by a random number (Excel; Microsoft) between 0 and 1 [26]. Patients who were colonized or infected with a nonenteric Gram-negative carbapenem-resistant organism were excluded (eg, *Pseudomonas aeruginosa, Acinetobacter baumannii, Stenotrophomonas maltophilia*). Data were retrieved from all available records and included demographics, underlying conditions, recent exposures to healthcare and to antimicrobials, acute illness indices, and clinical and microbiological outcomes. Posthospitalization deaths were captured from a national registry governed by the Israeli Ministry of Interior.

**Microbiological Analysis**
Carbapenem-resistant *Enterobacteriaceae* screening policy at Shamir (Assaf Harofeh) Medical Center, as in every Israeli hospital, is mandated and tightly regulated by the Israeli Ministry of Health [27, 28]. Patients who meet certain criteria are screened upon admission and weekly thereafter [27]. Processing is conducted in accordance with established criteria [11–13] while...
Using selective media and conducting thereafter confirmatory genotypic testing (Xpert Carba-R test; Cepheid) and full identification and susceptibility determination (VITEK 2; bioMérieux, Mercy l’Etoile, France). Non-CP-CRE were either *Klebsiella* species (*K. pneumoniae* or *Klebsiella oxytoca*), *Enterobacter* species, or *Escherichia coli*, demonstrating a meropenem minimum inhibitory concentration ≥2 µg/dL, followed by confirmatory disk diffusion tests [13]. Isolates were tested by assays to determine the presence of *blaKPC*, *blaVIM*, *blaOXA-48*, *blaNDM*, and *blaIMP* (Xpert Carba-R test; Cepheid), and *blaIMI* [29].

**Molecular Analyses**

A sample of 13 *K. pneumoniae* non-CP-CRE strains were sequenced using Illumina MiSeq (Illumina, San Diego, CA) after Nextera Flex library preparation. Bioinformatics analysis was performed as following: FASTQ files underwent quality control, filtering, trimming, and de novo assembly using fastQC (v0.11.8), Kraken (v1, with the DustMasked MiniKraken db 4GB – 2017) [30], and shovill (v1.0.0; with spades, v3.11; using the parameters “--trim” and “--opts '--sc'”) (https://github.com/tseemann/shovill). All assemblies were multilocus sequence typed (MLST) in silico using the tool MLST (v 2.10) (https://github.com/tseemann/mlst) based on the *K. pneumoniae* pubMLST schema (updated April 2018) (https://bigd-db.pasteur.fr/cgi-bin/bigsdb/bigsdb.pl?db = pubmlst_klebsiella_seqdef and pubMLST website [https://pubmlst.org/]) [31]. An ad hoc core genome (cg)MLST schema was generated using chewBBACA (v2.0.9) [32], producing a schema of 3635 loci (at 95% loci presence). Results were visualized using GrapeTree (v1.5) [33]. The assemblies were searched for antimicrobial resistance genes using c-SSTAR (v1.2c, with the ResGANNOT_srst2.fasta db—07082018) (https://github.com/tomdeman-bio/Sequence-Search-Tool-for-Antimicrobial-Resistance-SSTAR/blob/master/ARG-ANNOT.srst2_July-12-2016.fasta) [34, 35] and ABRicate (v0.8.13, with the CARD database from July 2019) [https://github.com/tseemann/abricate] [36].

**Statistical Analyses**

Analyses were all executed with IBM SPSS 25.0 (2018). Univariable matched analyses compared the characteristics of non-CP-CRE and susceptible *Enterobacterales* to the uninfected control group. Variables with *P* < .1 were incorporated into the stepwise backward selection to create 2 separated logistic regression models. In each step, the proposed model was tested for confounding. Subgroup analyses were performed when any interaction effect was identified. Time at risk was entered to both models [26]. The 2 final models of predictors associated with either non-CP-CRE or susceptible *Enterobacterales* were then contrasted to identify those predictors solely associated with non-CP-CRE [22, 23].

To determine whether non-CP-CRE impact hospitalization outcomes, we conducted univariable analyses pertaining to 8 different outcome parameters: in-hospital mortality, 30-days mortality, and 90-days mortality for the entire cohort and among survivors only; LOS from date of culture to discharge (or total LOS for uninfected controls); functional status deterioration [37]; additional hospitalizations (in 6 months); invasive procedures (in 3 months); and discharge to a long-term care facility (LTCF) after being admitted to the index hospitalization from home. Unmatched multivariable regression models for outcomes were constructed while enforcing the non-CP-CRE determinant into all models.

**RESULTS**

Initially, 111 unique patients with non-CP-CRE were enrolled. For 2 patients, no uninfected control patients could be matched due to extreme prolonged time at risk. The final cohort included 327 patients (3 groups of 109 patients). Of the non-CP-CRE, 89 were *K. pneumoniae*, 17 were *E. coli*, and 3 were *Enterobacter* spp. One-hundred four patients were asymptomatic carriers, 4 had active infection (intra-abdominal, skin and soft tissue, catheter-related bloodstream infection, and bloodstream infection with no determined focus), and for 1 patient, no sufficient data were available to determine whether she had had a catheter-associated urinary tract infection or asymptomatic bacteriuria. Among the 104 asymptomatic carriers, 4 patients (3.8%) subsequently developed a non-CP-CRE clinical infection. In the majority of susceptible case patients that were matched to the asymptomatic non-CP-CRE carriers, the susceptible *Enterobacterales* was isolated from patients with asymptomatic bacteriuria (n = 70), or from surveillance sputum (n = 28) or rectal (n = 6) cultures, obtained for infection control purposes. The median age of the cohort was 76 years (interquartile range, 64–85), 240 (74%) were elderly, 218 (68%) were already functionally impaired, and 191 (58%) had stayed in a healthcare facility in the preceding 6 months.

Table 1 summarizes the bivariable analyses conducted between the 3 groups of patients. There were no demographic differences between the various groups. Patients with *Enterobacterales* isolation (both non-CP-CRE or susceptible *Enterobacterales*) were more likely to be functionally dependent, with dementia, chronic renal failure, and recent MDRO isolation. Non-CP-CRE carriers had significant elevated Charlson’s scores [38], prior recent hospitalization(s), and/or intensive care unit (ICU) stay and were more often mechanically ventilated. The exposures to antimicrobials were more common among the non-CP-CRE carriers: namely, exposure overall (to any antibiotic) and specifically to cephalosporins, carbapenems, metronidazole, fluoroquinolones, and vancomycin. Because the cohort primarily consisted of asymptomatic carriers, and the infection versus colonization state was the primary matching
criteria, acute illness indices were indifferent between groups. However, the McCabe scores were lower among non-CP-CRE carriers [39].

In multivariable analyses (Table 2), the independent predictors associated with non-CP-CRE acquisition, but not with susceptible Enterobacteriaceae, were as follows: (1) recent hospitalization in an ICU (or mechanically ventilated in an advanced-care room), (2) chronic skin ulcers (eg, decubitus ulcers, diabetic foot infection), and (3) recent exposure to (any) antibiotics (not specifically to carbapenems). Recent MDRO acquisition was an independent predictor for Enterobacterales acquisition in general: it was independently associated with both non-CP-CRE isolation and with susceptible Enterobacterales isolation.

| Table 1. Selected Bivariable Analyses Comparing Risk Factors and Outcomes of Patients With Non-CP-CRE, Shamir (Assaf Harofeh) Medical Center (November 2014–December 2016), (n = 109 in Each Group) |
|----------|----------------|----------------|-----------------|----------------|----------------|
| Parameter | Non-CP-CRE | Susceptible Enterobacteriaceae | Uninfected Controls | Non-CP-CRE vs Uninfected | Susceptible Enterobacteriaceae vs Uninfected | CRE vs Susceptible Enterobacteriaceae |
|          | No. (%)* | No. (%)* | No. (%)* | Value OR (95% CI) | Value OR (95% CI) | Value OR (95% CI) |
| Age (years), median (IQR) | 76 (64–84) | 75 (63–84) | 76 (64–91) | 9 | .6 | .6 |
| Age group | 2 (1.8) | 2 (1.8) | 2 (1.8) | - | - | - |
|          | 80 (73.4) | 80 (73.4) | 80 (73.4) | - | - | - |
| Female gender | 49 (45.0) | 58 (53.2) | 53 (48.6) | 0.9 (0.5–1.5) | .6 | 1.2 (0.7–2.0) |
| In multivariable analyses (Table 2), the independent pre-| acquisition in general: it was independently associated with both | non-CP-CRE isolation and with susceptible Enterobacterales isolation. |
There were also multiple worse outcomes associated with non-CP-CRE isolation as per bivariable analyses (bottom of Table 1). For the outcomes that were captured among survivors of the index hospitalization (ie, LOS from isolation to discharge, functional deterioration [37], discharge to LTCF after being admitted from home), the non-CP-CRE determinant did not

| Parameter | Non-CP-CRE vs Uninfected | Susceptible Enterobacteriaceae vs Uninfected | CRE vs Susceptible Enterobacteriaceae |
|-----------|---------------------------|---------------------------------------------|-------------------------------------|
| ICU stay in the prior 3 months | 3.3 (1.4–7.7) | <.001 | <.001 |
| Received antibiotics in preceding 3 months before event | 3.1 (1.0–10.3) | .05 | <.001 |
| Chronic skin ulcers | 11.6 (4.3–30.8) | .002 | <.001 |
| Impaired functional status upon admission to hospital | 3.9 (1.7–9.2) | .02 | <.001 |
| Past MDRO in preceding 3 months before event | 52.3 (11.7–233.9) | <.001 | <.001 |
| Time at risk | 0.96 (0.93–1.0) | .04 | 0.06 |
| McCabe score | 2.3 (1.2–4.4) | .006 | <.001 |
| Received cephalosporins in preceding 3 months before event | 0.2 (0.1–0.5) | <.001 | <.001 |

Abbreviations: CI, confidence interval; CP, carbapenemase-producing; CRE, carbapenem-resistant Enterobacteriaceae; ICU, intensive care unit; MDRO, multidrug-resistant organism; OR, odds ratio; SD, standard deviation.

NOTE: Significant associations are highlighted in bold.

Valid percentage: count divided by the total number of valid (ie, nonmissing) observations.

Serum creatinine over 1.5 mg/dL or if patient was regularly undergoing hemodialysis or peritoneal dialysis.

Lower limb diabetic foot wounds, decubitus ulcers, dwelling wound surrounding PEG insertion, surgical-site wounds, surrounding catheters.

Immunosuppression includes any of the following: neutropenia at culture date (<500 neutrophils/mm³), exposure to glucocorticoids in the previous month, chemotherapy in the previous 3 months, radiotherapy, posttransplantation of any kind, or any immunomodulator.

Includes methicillin-resistant Staphylococcus aureus, vancomycin-resistant Enterococcus, extended-spectrum beta lactamase-producing Enterobacteriaceae, Acinetobacter baumannii, and Pseudomonas aeruginosa.

Event was defined as bacterial isolation for the patients who acquired non-CP-CRE or susceptible Enterobacteriaceae and as patients’ discharge date for uninfected patients.

Any type of invasive procedure, including endoscopies, any percutaneous intervention, biopsies, and any type of surgery.

Includes central venous lines, permanent urinary catheters, orthopedic external fixators, gastrostomies, drains. Not included: internal stents, prosthetic heart valve, and prosthetic joints.

For uninfected patients, we considered patients to have had a permeant device if it was placed for more than 48 hours during their entire hospitalization.

Patient received an antibiotic course of at least 48 hours.

For uninfected patients, we calculated the total length of stay.
remain significantly associated with any of those adverse outcomes. For the mortality outcomes (ie, in-hospital, 30-days, and 90-days), the non-CP-CRE determinant that was enforced into the models (as depicted under Methods) became a “protecting predictor” to the outcome of interest, implying a strong confounding effect called the “Simpson’s paradox” [40].

Phylogenetic analysis of the 13 isolates subjected to whole-genome sequencing is depicted in Figure 1. Six of the isolates belonged to ST-395, and others belonged to ST-101, ST-37, ST-147, ST-54, and ST-1. It is notable that the 13 K pneumoniae non-CP-CRE isolates displayed a wide range in the number of differing alleles (7–3234). Several isolates (n = 6) appeared to be clustered to some extent at a range of 7–40 alleles, which could coincide with theoretical epidemiological relatedness, at least for few of those isolates across time, but the majority of isolates appeared polyclonal and unrelated. None of the genomes harbored a known carbapenemase gene. All isolates had >1 mechanism of resistance. It is notable that each of the 13 strains harbored 4 distinct beta-lactamase genes including blaOXA-1, blaTEM-1, blaSHV (including ESBLs), and blaCTX-M-15. Typical RND-family efflux pump genes associated with beta-lactamase resistance such as acrB were found in all of the samples, in addition to efflux pumps of the oqxAB family, which mainly contribute to quinolone resistance.

DISCUSSION

The CRE healthcare-associated pandemic is one of the biggest challenges and threats in modern medicine [2]. Its epidemiology in recent years rapidly evolved, and along with (or partially due to) the CLSI and EUCAST breakpoints modifications [11–13], a new “epidemiological entity” of non-CP-CRE has emerged and has been reported in studies from around the world [15]. Despite the early reports displaying wide differences between the epidemiology of CPE and non-CP-CRE [20], controlled analyses pertaining to the predictors, outcomes, mode of acquisition, and transmission dynamics of non-CP-CRE are lacking. As far as we know, this is the first matched case-case-control investigation in this research field.

There are currently several definitions and recommendations pertaining to the diagnosis, treatment, and prevention of non-CP-CRE [10, 11]. Some have not even addressed the issue and use a broad definition of “CRE” to guide the management of both CPE and non-CP-CRE. Even the 2015 guidelines by the
Centers for Disease Control and Prevention do not differentiate between the 2 entities in terms of infection control-suggested measures (although it does acknowledge the issue) [10]. As depicted in this study and by others [15, 16, 20, 21], the epidemiology of non-CP-CRE differ considerably from that of CPE, and therefore measures should be tailored based of controlled analyses.

The study included 327 patients (consisting of 3 groups of 109 patients), but only 8 had acute active infection (ie, 4 with non-CP-CRE and 4 with susceptible Enterobacterales). This might imply, as suggested by others, that current non-CP-CRE isolates display low virulence properties and reduced endemic profiles, compared with CPE isolates, but this necessitates directed controlled investigations [15, 16, 20, 21]. However, theoretically, the therapeutic challenges associated with non-CP-CRE active infection resemble those of CPE [41], and therefore the exploration of the clinical and molecular epidemiological features of non-CP-CRE, and the mode of their acquisition and spread in hospitals, is still of paramount importance. Moreover, as displayed in our cohort as well [21], some asymptomatic carriers (4%), eventually develop active infection, including BSIs.

In this matched case-case-control investigation, we were able to provide some genuine insights into the clinical and the molecular epidemiology of non-CP-CRE. The fact that the majority of our cohort consisted of asymptomatic carriers, and not patients with active infection (as studied in other cohorts [20]), is one of the study's advantages, because this is the most common mode in which CRE is first diagnosed in most regions [42], and early prevention interventions could be applied. We discovered that the independent predictors for non-CP-CRE acquisitions were recent admission to an ICU, chronic skin ulcers (mostly decubitus ulcers, and/or chronic diabetic foot infections), and recent exposures to (any) antibiotics. Recent ICU admissions and chronic skin ulcers are somewhat nonmodifiable predictors that might serve as markers for exposures to (1) high selective pressures and/or colonization pressures in units [43], (2) severe burdens of active and/or chronic background conditions, and (3) deteriorated functional status in baseline. However, exposure to (any) antibiotics is a modifiable predictor and should therefore direct our future allocation of non-CP-CRE prevention resources.

Exposure to any antibiotics, not specifically to carbapenems, was a predictor for non-CP-CRE in this trial, as opposed to a previous investigation [16]. This again highlights the advantages associated with the case-case-control design, because it enables us to isolate the factors that “truly” impact, independently, the emergence of the resistance determinant(s) among susceptible strains. Moreover, this distinction of recent exposure to any antibiotic versus specific exposure to carbapenems was also illustrated in the past, among cohorts of CPE, that were analyzed by the matched case-case-control design, versus the “case-case design” [24]. In univariable analyses, exposures to many antibiotics were associated with non-CP-CRE acquisition, even recent exposures to vancomycin and metronidazole (Table 1), which have no established activity against Enterobacterales.

Whole-genome sequencing of 13 K pneumoniae non-CP-CRE strains (all obtained from asymptomatic carriers) displayed notable polyclonality (Figure 1). In addition, isolates displayed a coexistence of several mechanisms for beta-lactam resistance but not carbapenemases. These molecular features, along with the clinical epidemiology features as depicted above (ie, the independent association with recent exposure to any antibiotic), should lead to stewardship interventions that focus on reducing usage of antimicrobials consumption in general and not specifically focus solely on restriction of carbapenems.

Despite the fact that the cohort consisted mostly of asymptomatic carriers, non-CP-CRE was a strong marker for severe chronic and background conditions, with elevated Charlson's scores [38]. In univariable analyses, non-CP-CRE was significantly associated with worse morbidity and mortality outcomes (Table 1). However, when we enforced this parameter into the outcomes’ regression models, along with other parameters associated with worse outcomes, it became a protecting factor, illustrating the Simpson's paradox [40]: although non-CP-CRE was diagnosed among asymptomatic carriers, acute illness indices (eg, mechanical ventilation, McCabe score), which are naturally associated with worse outcomes of the acute condition, diluted the potential impact of being a non-CP-CRE asymptomatic carrier.

The study has several limitations. First, it is a retrospective, chart review-based, single-center analysis, with all its inherent limitations. However, the matched case-case-control design could overcome, at least in part, some of these nonmodifiable limitations, particularly regarding the predictors’ (for acquisition) analysis. Second, we sequenced the whole genome of only 13 isolates of non-CP-CRE. This might limit our conclusions pertaining to clonality, but in terms of representing the prevalent mechanisms of resistances to carbapenems and genetic relatedness, the conclusions are probably less affected by the low sample size of sequenced isolates. In addition, we did not perform molecular analysis of gene expression of porins, which could also contribute to carbapenem-resistant phenotypes. Third, there are inconsistencies in many clinical laboratories in non-CP-CRE diagnostic recommendations [11]. Therefore, some isolates that grow on the initial screening selective media and are directly subjected to a genotypic test of already known carbapenemases might not have been enrolled. Finally, because the majority of patients with non-CP-CRE were asymptomatic carriers, the impact of the resistance determinant on patients’ outcomes and efficacy analyses pertaining to the therapeutic management of non-CP-CRE active infection are limited.
CONCLUSIONS
This is the first study to explore, by using case-case-control design, the epidemiology of non-CP-CRE, which has become prevalent in many facilities around the world. Due to lack of controlled data, recommendations pertaining to non-CP-CRE diagnostics, prevention, and management are inconsistent. These analyses clearly illustrate that focusing on stewardship interventions, to limit usages of antibiotics in hospitals, could curb the continued emergence and spread of non-CP-CRE.

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