Regional spread of blaNDM-1-containing Klebsiella pneumoniae ST147 in post-acute care facilities

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**Summary:**

Whole-genome sequencing of carbapenem-resistant *Enterobacterales* in Chicago area healthcare facilities revealed clonal dissemination of bla<sub>NDM-1</sub> *Klebsiella pneumoniae* sequence type 147 within and between post-acute care facilities.
Abstract

Background

Carbapenem-resistant Enterobacterales (CRE) harboring bla\textsubscript{KPC} have been endemic in Chicago-area healthcare networks for more than a decade. During 2016-2019, a series of regional point prevalence surveys identified increasing prevalence of bla\textsubscript{NDM}-containing CRE in multiple long-term acute care hospitals (LTACHs) and ventilator-capable skilled nursing facilities (vSNFs). We performed a genomic epidemiology investigation of bla\textsubscript{NDM}-producing CRE to understand their regional emergence and spread.

Methods

We performed whole-genome sequencing on NDM+ CRE isolates from four point-prevalence surveys across 35 facilities (LTACHs, vSNFs, and acute care hospital medical intensive care units) in the Chicago area and investigated the genomic relatedness and transmission dynamics of these isolates over time.

Results

Genomic analyses revealed that the rise of NDM+ CRE was due to the clonal dissemination of an ST147 Klebsiella pneumoniae strain harboring bla\textsubscript{NDM-1} on an IncF plasmid. Dated phylogenetic reconstructions indicated that ST147 was introduced into the region around 2013 and likely acquired NDM around 2015. Analyzing the relatedness of strains within and between facilities supported initial increases in prevalence due to intra-facility transmission in certain vSNFs, with evidence of subsequent inter-facility spread among LTACHs and vSNFs connected by patient transfer.
Conclusions

We identified a regional outbreak of blaNDM-1 ST147 that began in and disseminated across Chicago area post-acute care facilities. Our findings highlight the importance of performing genomic surveillance at post-acute care facilities to identify emerging threats.

Keywords:

NDM, Klebsiella pneumoniae, ST147, genomic epidemiology, carbapenem resistance
Introduction

Carbapenem-resistant *Enterobacterales* (CRE) represent an urgent antibiotic resistance threat due to their resistance to first-line antibiotics and transmissibility in healthcare settings [1,2]. The emergence of epidemic lineages of CRE that are resistant to nearly all antibiotics and that cause infections with high mortality rates, such as *Klebsiella pneumoniae* carbapenemase (KPC) containing *Klebsiella pneumoniae* (KPC-Kp) sequence type (ST) 258 [3], has further escalated the need for more effective strategies to interrupt CRE transmission. Most interventions to prevent the spread of CRE and other healthcare-associated antibiotic resistance threats have been implemented at the level of individual healthcare facilities. However, there is now a multitude of evidence that the frequent movement of colonized and infected patients among regional healthcare facilities necessitates regional surveillance and infection prevention strategies [4].

Long-term acute care hospitals (LTACHs) and ventilator-capable skilled nursing facilities (vSNFs) are potentially high-impact settings for implementation of regional CRE surveillance and infection prevention interventions [5,6]. Patients in these facilities have been shown to be colonized with CRE at high rates, likely due to a combination of their chronic severe illness, long lengths of stay, and high rates of prior or on-going antibiotic exposure. Modeling and epidemiologic studies have suggested that the high CRE prevalence in LTACHs in particular has a significant impact on connected healthcare facilities with which they share patients [7,8]. Currently, less is known about the regional influence of vSNFs, although the even longer lengths of patient stay and more limited resources for infection prevention indicate that they might also be important settings in regional amplification of antibiotic resistance.
A bundled infection prevention intervention [9] (Chicago PROTECT [10]) was initiated in July 2017 to control CRE in Chicago-area healthcare facilities, including in vSNFs and LTACHs. Serial point prevalence surveys conducted to monitor the impact of the intervention demonstrated that KPC-Kp levels remained stable across regional facilities. However, during the intervention period, New Delhi metallo-beta-lactamase (NDM) containing CRE prevalence unexpectedly increased in a subset of surveyed vSNFs. Here, we applied whole-genome sequencing to investigate the underlying transmission dynamics related to the increase in NDM prevalence in the region.

Materials & Methods

Study isolates and metadata

Starting from October 2016 until July 2019, 20 medical ICUs in 20 short-term acute care hospitals, 7 LTACHs, and 8 vSNFs in the Chicago region were invited to participate in serial one day point prevalence surveys of residents. In vSNFs, surveys were performed in ventilator wards. Medical ICUs were surveyed once in 2016-2017; vSNFs and LTACHs were surveyed every 6-12 months. Patients who were present in their room at the time of survey were considered eligible for participation. Written informed consent was waived for this project, and patients who were competent were provided a standardized verbal explanation of the rationale for surveillance and were asked for verbal assent. Local staff obtained a rectal swab sample from each participating patient and collected de-identified patient information assessed at time of survey (age up to 90 years, sex, respiratory support status, length of stay, contact precautions status, facility awareness of resident CRE status). Swabs were processed at a central lab within 6 hours of collection. Overnight growth from MacConkey agar plates was screened for 5 carbapenemase gene families (KPC, NDM, VIM, IMP & OXA-48) using multiplex PCR assays (Acuitas® MDRO gene test and Acuitas® Resistome test,
For all genomic analyses, only the first isolate of a given ST (for *K. pneumoniae* isolates) or species (for all other species) was used for each patient. We used a Fisher’s exact test to test for the statistical significance of the difference in NDM or KPC prevalence between the first and last surveys. Fisher’s exact p-values were corrected using the Benjamini-Hochberg method.

**Whole-genome sequencing**

Genomic DNA was extracted from cultures derived from single sub-cultured colonies. Genomic libraries were prepared with NEBNext Ultra DNA library prep kit and sequenced at the University of Michigan Advanced Genomics Core on an Illumina NovaSeq 6000. All sequenced isolates have been deposited under BioProject PRJNA686897.

**Genomic analysis**

We processed whole-genome sequences [11,12] and identified in-silico multi-locus sequence types [13,14], generated and annotated assemblies [12,15–20], called single-nucleotide variants (SNVs) [21–28], identified phylogenetic clustering of facilities [8], and calculated pairwise SNV distances between isolates [29]. Reference-based whole-genome alignments of study and public ST147 isolates [30–40] were used to generate a phylogenetic tree using IQ-TREE v1.6.12 [41,42]. We inferred ancestral dates of the phylogeny with the R package BactDating v1.0.12 [43–45]. NDM-containing plasmids were identified from publicly available complete plasmids [29,29,46–48,21,49,50]. See supplemental methods for details of the genomic analysis.
Determining patient flow between facilities

We constructed a patient transfer matrix of the Chicago metropolitan region using the Centers for Medicare and Medicaid Services’ minimum data set, Medicare Provider Analysis and Review [MEDPAR] limited data set, Medicaid Analytic eXtract Data from 2010-2012. Using the patient transfer matrix, we constructed a directed weighted patient transfer network of healthcare facilities in the Chicago area using R igraph v1.2.6 [51], including all healthcare facilities in the study, and patient flow was determined as in [8]. See supplemental methods for details about calculating patient flow. Comparison of patient flow for inter-facility isolate pairs ≤12 SNVs vs. >12 SNVs was performed using Wilcoxon tests.

Data analysis and visualization

Data analysis and visualization was performed in R v4.0.2 [52]. Data visualization used the following packages: tidyverse v1.3.0 [53], pheatmap v1.0.12, lubridate v1.7.9.2, tidytree v0.3.3, treeio v1.12.0, ggtree v2.2.4 [54,55], ggplotify v0.0.5, ggreples v0.4.4, and cowplot v1.1.0. Code for analysis and visualization can be found here: https://github.com/Snitkin-Lab-Umich/ndm-st147-chicago-ms.

Ethical Review

Bacterial isolates and de-identified clinical metadata were collected under a prior surveillance project that underwent ethical review at the CDC and was determined to be a nonresearch activity (public health surveillance). The project was also evaluated independently at each participating healthcare facility and either deemed a public health assessment or human subjects research and approved by local review boards where applicable.
Results

Prevalence of NDM, but not KPC, increased over time in certain vSNFs that are not closely connected by patient transfer.

We first detected the presence of NDM+ CRE isolates in vSNFs and LTACHs during a regional point prevalence survey conducted in 2017, and subsequently performed three follow-up surveys (Fig 1A). A summary of the patient population for each facility type across the four surveys can be found in Table 1. We found that, while the prevalence of KPC+ CRE generally remained stable over time, the prevalence of NDM+ CRE increased in three vSNFs not closely connected by patient transfer (Fisher’s exact p < 0.05 for vSNFs J, K, and L; Fig 1B, S1, S2; Table S1).

The majority of NDM+ isolates are *K. pneumoniae* ST147 and carry bla\textsubscript{NDM-1} on an IncF plasmid.

To understand the molecular basis for the increase in NDM+ CRE we performed whole-genome sequencing on all CRE isolates from survey 1 and NDM+ isolates from the subsequent three follow-up surveys. We found that the presence of NDM was highly correlated with the presence of a suite of genes present on an NDM+ IncF plasmid isolated from *K. pneumoniae* [56] (Fig S3). Most isolates containing the IncF plasmid were bla\textsubscript{NDM-1} *K. pneumoniae* ST147; however, one bla\textsubscript{NDM-1} *Escherichia coli* ST354 isolate also contained the plasmid (Fig 2). While short-read sequencing data alone is insufficient to provide structural data associating NDM with the plasmid backbone, the high degree of correlation between NDM and the IncF-associated plasmid genes in concert with the phylogenetic relationships between isolates strongly suggests that these genes are co-inherited (Fig S3). In addition to NDM, the IncF plasmid harbored a number of antibiotic resistance genes from several
different resistance classes and the qacE gene, which may confer reduced susceptibility to common biocides [57] (Fig S3).

**Regional ST147 isolates are phylogenetically distinct from all public isolates.**

We investigated the phylogeography of ST147 in the Chicago area to determine whether circulating ST147 could be attributed to one or multiple importation events into the region. To this end we constructed a whole-genome phylogeny that included publicly available ST147 genomes from across the globe. Examination of the phylogenetic reconstruction revealed that all the ST147 isolates from the study form a monophyletic cluster, consistent with a single regional introduction (Fig 3). We also noted that while none of the NDM+ ST147 public isolates harbor the IncF plasmid, the majority of ST147s in the current study contain the plasmid. Moreover, one of the ST147 isolates in the KPC+/NDM- outgroup (from survey 3) contains the IncF plasmid but lacks NDM (Fig S4), suggesting that the plasmid may have been acquired in a locally circulating ST147 strain, followed by integration of a mobile element harboring NDM. A dated phylogenetic analysis of circulating ST147 yielded an estimate of August 2015 for when the NDM+ clade of ST147 first arose in the region (95% credible interval [CI]: February 2015 - March 2016; Fig 4, S5; Table S2), compared to an estimate of July 2013 for when ST147 first entered the region (CI: October 2011 – October 2014; Table S2).

**Genomic evidence indicates that intra-facility transmission is driving prevalence at high-prevalence vSNFs.**

After determining that the increase in NDM prevalence corresponds to a clonal outbreak of bla$_{\text{NDM-1}}$ ST147, we investigated the potential transmission dynamics of this clone within and between healthcare facilities. We observed a substantial clustering of isolates from certain vSNFs on the phylogeny (Fig 5A, S6), which suggests potential intra-facility transmission. To further investigate
whether these clusters may represent within-facility transmission, we calculated pairwise SNV distances among all pairs of isolates and compared these distances for isolates from the same facility (intra-facility pairs) to isolates from different facilities (inter-facility pairs) across surveys (Fig 5B). Indeed, starting in survey 3 we observed a disproportionate representation of small SNV distances (≤12 SNVs; see methods for threshold selection) consistent with intra-facility transmission in vSNFs. Of note, in survey 4 we observed spikes in small SNV distances for both intra- and inter-facility pairs, with closely related intra-facility pairs being primarily from vSNFs and closely related inter-facility pairs being from both vSNF-LTACH and vSNF-vSNF pairs (Fig S7). Putting these closely related inter-facility pairs in the context of the regional patient transfer network supports a potential role of patient transfer in regional blaNDM-1 ST147 spread in survey 4, with vSNF-LTACH and vSNF-vSNF isolate pairs less than 12 SNVs apart being from facilities with higher patient flow between them than isolate pairs with 12 or more SNVs (Wilcox p < 0.001; Fig S8; higher patient flow indicates more movement of patients from source to destination facility, see supplementary methods for details).

Discussion

We performed genomic analyses of CRE isolates collected through serial point-prevalence surveys in the Chicago area to investigate an increase in NDM+ CRE prevalence across a regional healthcare network. Our analysis supports the increase in NDM+ CRE being due to the clonal dissemination of a single blaNDM-1 ST147 strain of K. pneumoniae that emerged in 2015. Putting genomic analysis in the context of the regional healthcare network supports this strain first reaching high prevalence in a small number of vSNFs due to intra-facility transmission, followed by inter-facility spread to connected healthcare facilities.
Whole-genome sequencing showed that the majority of bla\textsubscript{NDM-1} ST147 harbored an IncF multidrug resistance plasmid. Incorporating public data into the analysis revealed that these isolates formed a monophyletic clade, suggesting a single introduction into the region, either through importation of a pre-existing NDM+ ST147 strain or acquisition of bla\textsubscript{NDM} by a locally circulating ST147 strain. Furthermore, examination of the global phylogeny indicates that, while NDM+ ST147 has evolved multiple times in different locations and sometimes resulted in clonal outbreaks, none of the global NDM+ ST147 isolates we included in our analysis harbor the IncF plasmid found in our study isolates. The rapid and widespread dissemination of this strain in the region indicates that the NDM-carrying IncF plasmid we identified here can stably associate with an ST147 strain with epidemic potential. Given the potential negative impact of epidemic NDM-carrying \textit{K. pneumoniae}, this possibility warrants close monitoring.

By combining regional surveillance with genomic analysis, we were able to discern that NDM initially spread in three vSNFs, likely via intra-facility transmission, with evidence of subsequent spread to healthcare facilities connected by patient transfer. There are several factors that likely contributed to the spread of this NDM+ ST147 clone. First, vSNF patients are a high-risk population for carriage of CRE as they are chronically ill, are usually admitted from ICUs or LTACHs, and are often exposed to antibiotics [58]. Second, patients in vSNFs generally have long lengths of stay — often much longer than patient stays at LTACHs [59] — meaning that they have a longer period of time to acquire a multi-drug resistant organism. Furthermore, multibed rooms are common and the facilities themselves are often under-resourced from a staffing and infection control perspective [6], both of which could facilitate intra-facility spread. Our findings paired with these observations indicate that vSNFs may be important healthcare facilities to detect emerging threats and potentially contain them before widespread dissemination. In the current study, we note that NDM+ isolates were
uncommon in ICUs, and the outbreak of ST147 might not have been detectable until much later if sampling were restricted to ICUs.

Our study has several strengths. Active surveillance of diverse types of healthcare facilities in the region allowed us to identify and investigate a potential multi-drug resistant organism threat earlier than would have been possible if serial point-prevalence surveys across several facility types were not ongoing. Furthermore, cross-sectional patient sampling within each survey allowed us to obtain a complete snapshot of CRE prevalence at a given facility at a given point in time, and to detect the increase in NDM+ isolates over time. Finally, we were able to leverage information from whole-genome sequencing to investigate the relatedness of isolates, as well as the intra- and inter-facility transmission dynamics of NDM across the healthcare network.

Our study also has several important limitations. First, the cross-sectional study design could have led to potential biases in the number of NDM+ isolates sequenced at facilities given that the patients at these facilities had different average lengths of stay, and precluded a more nuanced examination of NDM-1 intra-facility transmission dynamics and associated patient risk factors. Second, we lacked data from short-term acute care or community settings, particularly in the last three surveys, which limited our ability to examine the relative importance of other regional reservoirs for NDM+ ST147. However, the short-term acute care data that was available did not support their role in blaNDM-1 ST147 expansion. Third, we used facility-level aggregate patient transfer data to infer the likelihood of patient-level exposure to facilities; lack of patient-level facility exposure data precluded us from performing a more detailed exposure network analysis [60]. Lastly, we used short-read sequencing data, which limited our ability to investigate more complex plasmid dynamics. While we plan to perform long-read sequencing on a subset of these isolates in the future, we find it notable that we...
were able to leverage publicly available complete plasmid sequences to determine that NDM was carried on the same plasmid in the majority of isolates.

In conclusion, our study identified an emerging \textit{bla}_{NDM-1} ST147 clone of \textit{K. pneumoniae} with epidemic potential. The identification of this clone, and characterization of its ability to disseminate within and between healthcare facilities, was made possible through whole-genome sequencing of NDM+ isolates from serial point-prevalence surveys at vSNFs, LTACHs, and ICUs. Our findings highlight the importance of performing surveillance of multidrug-resistant organisms not only in acute care hospital ICUs, but also in post-acute care facilities such as LTACHs and vSNFs. vSNFs in particular appear to be especially important as sentinel sites of active surveillance for rare and emerging resistant pathogens.
NOTES

Authors’ contributions

All authors developed methodology and reviewed and edited the manuscript. ML, ES, MH, AMJ, and ZL conceptualized the research goals and aims. ZL, RC, AP, and ES developed and implemented software and curated the data. ZL, RC, and AP performed formal analysis. ML, MH and ES provided resources. ZL and ES wrote the original draft. ZL and RC visualized the results. ML, MH, AMJ, and ES provided supervision. ML, MH, and ES managed the project. ML, MH, and ES acquired funding.

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Resistome test in kind, during 2016-2017. Any opinions, findings, conclusions, or recommendations expressed in this material are those of the authors and do not necessarily reflect the views of the Centers for Disease Control and Prevention, National Science Foundation, or the National Institutes of Health.

**Conflicts of interest**

M.L. and M.H. have received research support in the form of contributed product from OpGen, LLC and from Sage Products (now part of Stryker Corporation). M.L. has also received an investigator-initiated grant from CareFusion Foundation (now part of BD). M.H. and R.W. have participated in clinical studies where participating healthcare facilities received contributed product from Sage Products Inc., Molnlycke, Clorox, or Medline. Neither M.H., R.W. nor their hospitals received product, funding, payments, or any other form of compensation. M.L. reports honorarium from antibiotic resistance symposium (Medical College of Wisconsin) and participates on CDC’s Healthcare Infection Control Practices Advisory Committee (HICPAC). No other author has any potential conflicts to disclose.
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Table 1: Summary of point prevalence survey results from ICUs, LTACHs, and vSNFs for surveys 1, 2, 3, and 4

|                           | ICU  | LTACH | vSNF |
|---------------------------|------|-------|------|
| Number of surveys         | 1    | 4     | 4    |
| Number of facilities      | 20   | 7     | 8    |
| Number of patients eligible | 238 | 1338  | 1325 |
| Number of patients surveyed, n (% of eligible) | 212 (89) | 1188 (89) | 1154 (87) |
| Age, mean yrs (SD)        | 62 (17) | 62 (15) | 60 (15) |
| Male, n (%)               | 119 (56) | 644 (54) | 627 (54) |
| Length of stay, median days (IQR) | 5 (3-10) | 21 (11-37) | 126 (33-410) |
| Mechanical ventilation, n (%) | 102 (48) | 405 (21) | 477 (33) |
| Tracheostomy collar, n (%) | 0 (0) | 250 (21) | 384 (33) |
| Contact precautions, n (%) | 58 (27) | 736 (62) | 679 (59) |
| Carbapenemase gene:       |      |       |      |
| KPC, n (%)                | 11 (5) | 182 (15) | 360 (31) |
| NDM, n (%)                | 2 (1) | 30 (3) | 146 (13) |
| OXA-48, n (%)             | 1 (0) | 0 (0) | 1 (0) |
| IMP, n (%)                | 0 (0) | 1 (0) | 0 (0) |
| VIM, n (%)                | 0 (0) | 4 (0) | 66 (6) |
| Any carbapenemase gene, n (%) | 14 (7) | 201 (17) | 479 (42) |
| Of carbapenemase-positive:|      |       |      |
| Carbapenemase-positive with contact precautions, n/N (%) | 9/14 (64) | 169/201 (84) | 364/479 (76) |
| Carbapenemase-positive known to facility, n/N (%) | 6/14 (43) | 107/201 (53) | 303/479 (63) |
Figure Legends

Figure 1: Prevalence of NDM increased over time in certain vSNFs. (A) Time window of when facilities were tested for each survey. Observance of NDM+ CRE in vSNFs and LTACHs in survey 1 led to targeted follow-up surveys (2, 3, and 4) in these facilities. (B) Proportion of NDM and KPC positive samples across surveys and facilities. vSNF O was not included as there was uneven sampling across surveys. vSNF = ventilator-capable skilled nursing facility. LTACH = long-term acute care hospital, ICU = intensive care unit. ICUs are not shown in panel B because of very low prevalence (see Table 1).

Figure 2: The majority of NDM+ isolates are Klebsiella pneumoniae ST147 and carry NDM on an IncF plasmid. Number of sequenced isolates of various species and sequence types, what carbapenemase(s) they contain, and whether they have the IncF plasmid.

Figure 3: Study isolates are clonally separated from all publicly available isolates outside the Chicago region. Maximum likelihood phylogeny of study and public isolates annotated by geographic region and genomic element.

Figure 4: NDM+ ST147 Klebsiella pneumoniae was introduced into the region around 2015. Dated phylogeny generated by bactdating. Grey bar on the root is the lower and upper bounds of the confidence interval (2015.09 to 2016.17). vSNF = ventilator-capable skilled nursing facility. LTACH = long-term acute care hospital, ICU = intensive care unit.
Figure 5: Intra-facility transmission is driving prevalence at high-prevalence vSNFs. (A) Number of isolates in the largest subclade of the maximum likelihood phylogeny containing ≥90% of isolates from the given facility (see methods for more details). Note that the y axis is log10-scaled. (B) Pairwise SNV distances of isolates from the same and different facilities across surveys. The grey diamond at a pairwise SNV distance of 12 indicates the threshold for closely related isolates (see methods for details). vSNF = ventilator-capable skilled nursing facility. LTACH = long-term acute care hospital, ICU = intensive care unit.
