IS26 Veers Genomic Plasticity and Genetic Rearrangement toward Carbapenem Hyperresistance under Sublethal Antibiotics

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ABSTRACT  Multidrug-resistant Gram-negative carriers of Klebsiella pneumoniae carbapenemases (KPCs) often subvert antibiotic therapy due to inadequate sensitivity in laboratory detection. Although unstable gene amplification has been recognized to crucially contribute to underestimation or misestimation of antimicrobial resistance in clinical isolates, the precise mechanisms underlying carbapenem resistance driven by amplification of blaKPC-2 remain obscure. Here, we reported that IS26-mediated amplification of blaKPC-2 rapidly and robustly gave rise to carbapenem hyperresistant phenotypes in an Escherichia coli clinical strain following sublethal meropenem or tobramycin preexposure. Intriguingly, IS26 also underpinned amplification of a 47 kb multiple drug resistance (MDR) region encompassing nine antibiotic resistance genes and six IS26 insertion sequences. Tandem-repeat analysis and experimental validation demonstrated that blaKPC-2 amplification was indeed mediated by IS26, which was further experimentally shown to involve intricate genetic rearrangement. Such gene amplification arose dynamically under antibiotic stress and subsided upon antibiotic withdrawal. Instead of reducing the amplification of the IS26-flanked MDR region, drug combinations in vitro exacerbated it. Our study, thus, provides valuable insights into how dynamic gene amplification processes can precipitously transform resistance status and complicate diagnosis.

IMPORTANCE  Klebsiella pneumoniae carbapenemases (KPCs) are powerful β-lactamases that enable Gram-negative pathogens to destroy clinically important carbapenems in antibiotic therapies. In particular, KPC-2 is difficult to detect due to a lack of instrument sensitivity in regular laboratory screens, which leads to misdiagnosis and poor treatment outcomes. It remains unclear how blaKPC-2 rapidly induces exceedingly high-level resistance against carbapenems following the challenges of sublethal antibiotics. Here, we demonstrated that, under sublethal doses of antibiotics, insertion sequence IS26 mediated rapid amplification of multiple resistance determinants, including blaKPC-2 and a multiple drug resistance (MDR) region, which was accompanied by intricate genetic rearrangement.

KEYWORDS  antimicrobial resistance, Klebsiella pneumoniae carbapenemase (KPC), insertion sequence IS26, gene amplification, genomic plasticity, genetic rearrangement

The global spread of carbapenem-resistant Enterobacteriaceae poses a relentless menace to public health systems, necessitating urgent concerted actions to innovate strategies for antimicrobial control and treatment (1). Members of Enterobacteriaceae develop carbapenem resistance primarily through the acquisition of carbapenemases with Klebsiella pneumoniae carbapenemases (KPCs) being the most prevalent (2). Typically, multidrug-resistant and
recalcitrant to treatment Enterobacteriaceae KPC-producers have disseminated worldwide and become a major cause of increased morbidities and mortalities in healthcare-associated infections (HAIs) in hospital niches (3). Additionally, KPC-producing opportunistic pathogens have gained notoriety for subverting antibiotic therapy (4, 5) because they often exhibit deceptively low-level resistance to carbapenems in routine laboratory screens, leading to misdiagnosis or underdiagnosis (3, 6). For example, automatic detection systems reportedly identified up to 87% of KPC-producing K. pneumoniae (KPC-KP) as being susceptible to imipenem or meropenem (2). Intrinsically, the difficulty of reliably detecting antimicrobial susceptibility of KPC-KP isolates to carbapenems is due in part to heteroresistance, wherein a subpopulation with greater resistance to antibiotic stress than a dominant population is present, while the proportion of the former gradually rises in the presence of antibiotics (7). Gene amplification has been proposed as one of the drivers of heteroresistance. Notably, amplified blaKPC genes have been detected in K. pneumoniae isolates with profound implications for the reduction of pathogens’ susceptibility to carbapenems (8, 9). To this date, however, the precise mechanisms underlying carbapenem resistance mediated by blaKPC-2 amplification, particularly from the perspective of carbapenem heteroresistance contributed by unstably amplified blaKPC, remain elusive.

Conventionally, clinical use of antibiotics is guided by both empirical experience and laboratory findings whose rationale is to attain the best cure rates with the highest possible nontoxic drug concentrations without provoking the development of de novo resistance within patients (10). In practice, due to a constant gap in diagnostic precision, many human and animal treatment regimens have proved insufficient for achieving such goals because pharmacologically effective antibiotic concentrations often fall below laboratory-determined MICs in body compartments and tissues in vivo, resulting in suboptimal inhibition of target bacteria. Indeed, growing evidence demonstrates that sublethal concentrations of antibiotics are a significant driver for the development of antibiotic resistance (11, 12). Sublethal, subinhibitory concentrations of antibiotics exert their effects basically on three different levels as selectors of resistance, generators of genetic and phenotypic variability, and signaling molecules (10). However, the molecular underpinnings of how sublethal antibiotics drive the processes of heteroresistance formation remain to be elucidated.

Here, our investigation focused on an E. coli clinical strain carrying the blaKPC-2 gene, which was isolated from a fatal case of a patient who failed to respond to antibiotic treatment. After exposure to a sublethal dose of carbapenem (2 μg/mL) for 24 h, the strain rapidly and robustly adapted to a range of high-level carbapenem doses during regrowth, accompanied by amplification of blaKPC-2. This amplification was positively associated with an extraordinary increase in antimicrobial resistance (AMR) as exemplified by amplified resistance determinants, including a 47 kb multiple drug resistance (MDR) region flanked by IS26. We monitored the dynamics of genomic plasticity in terms of copy number of amplified gene elements as well as any rearrangement of the MDR region. Tandem-repeat analysis and experimental validation demonstrated that blaKPC-2 amplification was mediated by IS26. Remarkably, combination therapy of meropenem (a carbapenem) and tobramycin (an aminoglycoside) failed to reduce amplification of the IS26-flanked MDR region harboring resistance genes but instead aggravated its progression. Our present work thus fills a critical gap in our understanding of IS26-driven amplification of blaKPC-2 as a driving force of AMR through which Enterobacteriaceae bacteria manage to evade carbapenem assaults, and thus compromise the efficacy of antibiotics in clinical use.

RESULTS

Preexposure to sublethal antibiotics rapidly unfetters carbapenem hyperresistance. In our previous study, three outbreak-associated strains of E. coli (sequence type ST131) were separately isolated from sputum, urine, and blood samples collected within 1 month from a patient (13). These strains exhibited low-level carbapenem...
resistance (MIC for meropenem: 4 μg/mL; and MIC for biapenem: 4 μg/mL) and shared nearly identical genomic sequences, indicating that they likely arose from a single parent clone. Subsequent whole-genome analysis showed that a plasmid (pE0171; GenBank accession no. MK370988) carrying \(\text{bla}_{KPC-2}\), which conferred carbapenem resistance, was present in the isolates. Notably, \(\text{bla}_{KPC-2}\) was in an IS26-based composite transposon (Unit 1), whereas two copies of Unit 1 were found in the MDR region (Fig. S1). During antifective management, the patient had been treated with biapenem at 0.6 g per day for 15 days. Nonetheless, biapenem therapy proved ineffective and the patient later died of complications from \(\text{E. coli}\) infection despite a reported peak plasma concentration of biapenem (9.49 to 25.81 μg/mL) greater than 4 μg/mL (14). Based on observations of this clinical course of the disease, we hypothesized that a sustained low concentration of carbapenem in vivo could lead to detrimental development of resistance.

For experimental proof of concept, one of the above-mentioned \(\text{E. coli}\) clinical isolates, E01-7-1, was selected for further experimentation. Broth containing 0, 0.5, or 2 μg/mL of meropenem was inoculated with the strain followed by culture for 24 h (Fig. 1A). The inoculums were then taken and subcultured individually on agar plates containing a gradient of meropenem concentrations (0 to 512 μg/mL) for regrowth (Fig. 1B). The percentage of bacteria exhibiting meropenem (MEM) resistance was calculated as the number of colonies grown on LB agar plates containing MEM divided by the number of bacterial colonies grown on plain LB agar plates. Our results showed that the highest concentration of meropenem permissive to fithe growth of the control bacteria (i.e., without preexposure to sublethal antibiotics) was 4 μg/mL, while the frequency of meropenem (MEM)-resistant bacteria was about 7.35 \(\times 10^{-2}\) (Fig. 1C). In comparison, the highest growth-permissive meropenem concentration was found to be 128 μg/mL and 512 μg/mL for groups preexposed to sublethal meropenem at 0.5 and 2 μg/mL, respectively, and the corresponding frequencies of meropenem-resistant subpopulations on plates were determined to be 8.45 \(\times 10^{-2}\) and 4.63 \(\times 10^{-2}\), respectively. The frequency of resistant bacteria declined evidently with increasing antibiotic concentration. Strikingly, however, the frequency of resistant bacteria in groups preexposed to 2 μg/mL meropenem and then regrown on meropenem containing agar plates (8, 16, and 32 μg/mL) approached 1. This suggested that the presence of sublethal meropenem concentration (2 μg/mL) was sufficient and potent to cause the emergence of meropenem-resistant subpopulations with exceedingly high-level resistance within a relatively short time (e.g., red line in Fig. 1C).
When bacterial samples preexposed to sublethal meropenem (2 μg/mL, 24 h) were added as an inoculum to broth medium containing meropenem (4 to 512 μg/mL) for regrowth, results similar to those of regrowth in agar plates were obtained (Table S1). Broths inoculated with sublethal antibiotic preexposed bacteria were observed to allow growth in the presence of 512 μg/mL meropenem. In contrast, broths inoculated with unexposed bacteria showed bacterial regrowth in a medium containing hardly more than 4 μg/mL meropenem during regrowth. Interestingly, the efflux pump inhibitor phenylalanine-arginine-β-naphthylamide dihydrochloride (PAβN; 50 μM), had no effects on meropenem MIC (Table S1), suggesting that efflux pumps had little or no contribution to phenotypic outcomes in MICs. These results indicated that sublethal antibiotic concentrations favored the enrichment of bacterial subpopulations that were selectively at an advantage in acquiring resistance.

Sublethal meropenem preexposure elicits hyperresistance via dynamic processes of gene amplification. To scrutinize this phenomenon, we analyzed the sequences of carbapenem resistance genes (bla<sub>KPC-2</sub>, ompC, and ompF) and their promoter regions in strains grown with or without stress from sublethal meropenem and in strains regrown with high-dose meropenem. No mutations were detected among the strains. However, an increase of 1.6-fold in copy number of bla<sub>KPC-2</sub> under sublethal meropenem (2 μg/mL) relative to untreated conditions was observed. To better understand the mechanism permitting regrowth in high-dose meropenem, we determined the production of β-lactamases, expression of bla<sub>KPC-2</sub> and copy number of bla<sub>KPC-2</sub> in bacterial cells exposed to high concentrations of meropenem (ranging from 16 to 512 μg/mL) following sublethal meropenem preexposure (2 μg/mL). Remarkably, β-lactamase levels rose with increasing concentrations of antibiotics (Fig. 2A). Similarly, the qRT-PCR results showed statistically significant increases in expression of the bla<sub>KPC-2</sub> gene, although without an increase in repA of plasmid pE0171, which was a carrier of bla<sub>KPC-2</sub> (Fig. 2B; original data in Table S2). Finally, we confirmed that the copy number of bla<sub>KPC-2</sub> was positively correlated with antibiotic concentration (ranging from 16 to 512 μg/mL) with respect to the normalized levels of the housekeeping gene purA encoding adenylosuccinate synthase (Fig. 2C).

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The relative copy number of bla<sub>KPC-2</sub> was 1.2 in the absence of meropenem stress, which dramatically rose to 28.9 at 512 μg/mL meropenem (Fig. 2C; data in Table S2). These observations strongly suggested that an augmentation in bla<sub>KPC-2</sub> copy number could account for the rapid increase in resistance to meropenem. Predictably, the copy number of repA was unaltered, implying that amplification of bla<sub>KPC-2</sub> was not tied to an increase in the copy number of the plasmid carrying bla<sub>KPC-2</sub>. This novel finding lends support to the notion that rapid emergence of resistance could sequentially arise from preexposure to a
single sublethal-dose antibiotic (such as meropenem) and subsequent reexposure to antibiotics at higher concentrations.

**Genomic plasticity through amplification of an IS26-flanked MDR region with bla<sub>KPC-2</sub> and complex genomic arrangements.** To dissect the mechanisms underlying amplification of bla<sub>KPC-2</sub> genes, we applied Illumina sequencing to the analysis of DNA extracted from strains subjected to various levels of meropenem stress at the end of the logarithmic phase during regrowth (clean data size and mean coverage as shown in Table S3). We then mapped short reads onto the complete genome of the strain E01-7-1. Analysis on read-mapping depths relative to the housekeeping gene purA revealed a 47 kb MDR region containing bla<sub>KPC-2</sub> with significantly higher coverage of reads in strains exposed to meropenem relative to untreated control strains (Fig. 3 and Table S3). The read-mapping depth ratio of this region was positively associated with antibiotic concentration with changes in bla<sub>KPC-2</sub>-mapping depth being 1.63-fold higher in the meropenem preexposed (2 μg/mL) strains and 1.96 to 29.94-fold higher in the meropenem regrowth (64 to 512 μg/mL) stains than in the case where bacteria were unexposed. The aforementioned 47 kb region turned out to be flanked by IS26 and IS15DI, which share 99% nucleotide identity and 100% coverage with IS26. Changes in IS26-mapping depths were found to increase correspondingly by 1.28 to 21.12-fold. Furthermore, the amplified region contained 16 antibiotic resistance genes (ARGs) underpinning resistance to macrolides, aminoglycosides, carbapenems, sulfonamides, and diaminopyrimidines. Considering the existence of two Unit 1 sequences, 7 types of ARGs were identified. The results implicate IS26 as a prominent mediator of amplification of bla<sub>KPC-2</sub> genes during the rapid emergence of hyperresistance following sublethal antibiotic challenges. Furthermore, unexpected amplification of the MDR region under single-drug stress underscored the complexity of such gene amplification.

To advance a more detailed mechanistic picture of the bla<sub>KPC-2</sub> amplification involved, we applied Oxford Nanopore Technologies (ONT) sequencing to the acquisition of long reads of the E. coli strain exposed to 512 μg/mL meropenem. In total, 5,295 reads containing bla<sub>KPC-2</sub> were obtained, among which 1,287 reads indicated carriage of more than two copies of bla<sub>KPC-2</sub> (Table 1). An unexpected level of diversity in bla<sub>KPC-2</sub> copy numbers was observed among the reads, which underlined the complexity of the gene amplification machinery. To illustrate, the amplification of bla<sub>KPC-2</sub> possibly occurred through two processes (Table 1) with the first being tandem IS26-based bla<sub>KPC-2</sub> transposon amplification. For example, seven identical bla<sub>KPC-2</sub> transposons in a tandem array were found within a...
59 kb read (Fig. 4A). In addition to direct repeats in tandem, two inverted blαKPC-2 transposons linked together were present in read 7 (Fig. 4B). The second mechanism presumably occurred via amplification of a composite transposon with blαKPC-2 flanked by IS26. For instance, read 155 was composed of three 31 kb tandem units bordered by two IS26s with nine ARGs and six IS26s in each unit (Fig. 4C). Read 25 likely included tandem units containing two blαKPC-2 copies (Fig. 4D). These read data further confirmed that amplification of blαKPC-2 was mediated by IS26. Accordingly, we analyzed the copy number of IS26 transposase by qPCR. In searching for a possible correlation between copy number and transcription level of blαKPC-2 and IS26 genes, it was found that the copy number of IS26 and blαKPC-2 (r = 0.92) and the transcription level of IS26 and blαKPC-2 (r = 0.97) were strongly correlated (Fig. S2; original data in Table S2). Of note, these results showed a trend similar to that of blαKPC-2 under antibiotic stress but with higher copy numbers of IS26 transposase, which is consistent with the E. coli genome possessing more basal copies of the genes (12 copies).

By leveraging a hybrid assembly of ONT long-sequencing reads and Illumina short-sequencing reads, we obtained a complete genome of strain E01-7-1 cultured in the presence of 512 μg/mL meropenem. The assembled genome was 5.1 Gb in length with a GC content of 50.89% and contained a 137 kb plasmid. In contrast, there was again no blαKPC-2 amplification mediated by IS26 found in the MDR region of the plasmid. This intriguing finding suggested that strain E01-7-1 could indeed generate subpopulations with multiple copies of blαKPC-2 upon carbapenem, but these subpopulations may carry diverse structural variants of the MDR region or may not be dominant in the population. Phenotypically, strains with such properties were deemed to exhibit heteroresistance. Furthermore, the circular genomic DNA molecule did not appear to contain an IS26-mediated translocatable unit (TU).

**Experimental validation of blαKPC-2 amplification mediated by IS26.** To experimentally interrogate the roles of IS26 in amplification of blαKPC-2, we constructed a plasmid consisting of the vector pUC57 and IS26-blαKPC-2-IS26 (Fig. 5A). The IS26-blαKPC-2-IS26 sequence was designated Tn7094 (~2.9 kb) (LSTM, [https://transposon.lstmed.ac.uk/tm-registry](https://transposon.lstmed.ac.uk/tm-registry)). Next, E. coli Top10 (recA deficiency) was transformed with the pUC57: Tn7094 plasmid by using kanamycin and meropenem as selective markers. In observation, the recombinant plasmid intensified the resistance of its E. coli host against meropenem, from < 0.25 to 2 μg/mL. Subsequently, an inoculum (1 x 10^7 colony forming unit [CFU]) of E. coli with the recombinant plasmid was incubated in a liquid medium containing 1 μg/mL meropenem (or 1/2 MIC) for 24 h. These sublethal antibiotic preexposed bacterial cells were then challenged with various concentrations of meropenem (ranging from 0 to 64 μg/mL) during the regrowth phase. To monitor the dynamics of Tn7094 under various levels of meropenem stress, plasmids were extracted from culture. Analysis on plasmid patterns corroborated that more bands of higher molecular weights were detectable in extracts from bacterial cells incubated at higher levels of

| blαKPC-2 copy no. | Read no. | Amplification process |
|-------------------|----------|-----------------------|
| 7                 | 2        | IS26-based blαKPC-2 transposon |
| 6                 | 2        | Mode 1               |
| 5                 | 5        |                      |
| 4                 | 24       | Composite transposon with blαKPC-2 flanked by IS26 |
| 3                 | 159      | Mode 2               |
| 2                 | 1095     |                      |

*Amplification of blαKPC-2 occurs in two ways: tandem amplification of IS26-based blαKPC-2 transposon (Mode 1), and amplification of a composite transposon with blαKPC-2 flanked by IS26 (Mode 2), as detailed in Fig. 4.
meropenem compared with that at 0 μg/mL, which could be a direct consequence of gene amplification mediated by IS26 (Fig. 5B).

As a next step, we conducted Nanopore sequencing to characterize the amplification of Tn7094. Plasmids from strains grown at 64 μg/mL meropenem were extracted and subjected to Nanopore sequencing analysis. The numbers of reads containing pUC57:Tn7094 and pUC57:IS26 were 121,166 and 113,722, respectively. Amplification

![Gene organization of long reads containing multiple copies of blaoPC-2 as revealed by Nanopore sequencing.](image)

FIG 4 Gene organization of long reads containing multiple copies of blaoPC-2 as revealed by Nanopore sequencing. (A) Seven identical blaoPC-2 transposons in a tandem array in read 1. (B) Two direct and two inverted blaoPC-2 transposons linked together in read 7. (C) Three tandem units, each harboring nine ARGs and six IS26s, bordered by two IS26s in read 155. (D) Amplification of a composite transposon with blaoPC-2 flanked by IS26. Red triangles denote IS26, blue triangles denote blaoPC-2, and gray triangles denote other genes.

![Analysis on dynamics of Tn7094 evolution in E. coli strains preexposed and reexposed to various concentrations of meropenem.](image)

FIG 5 Analysis on dynamics of Tn7094 evolution in E. coli strains preexposed and reexposed to various concentrations of meropenem. (A) Structural design of plasmid pUC57:Tn7094. (B) Plasmids profiles analysis on the dynamics of Tn7094 evolution in strains exposed to 0 μg/mL, 1 μg/mL, 16 μg/mL, or 64 μg/mL meropenem, respectively. (C) Gene organization of long reads containing four tandem arrays of Tn7094 on the plasmid as revealed by Nanopore sequencing. (D) Tandem array of the plasmid pUC57:Tn7094 in a direct-repeat configuration. (E) Tandem array of the plasmid pUC57:Tn7094 in an inverted-repeat configuration. (F) Composite structure from combined amplification of Tn7094 and the plasmid.
of Tn7094 was found in 3,820 reads, which corresponded to the carriage of at least two blaKPC-2 sequences (Table 2). We reasoned that such amplification may take place via three mechanisms, namely, tandem Tn7094 amplification (Fig. 5C and Table 2), plasmid amplification, in which the plasmid pUC57:Tn7094 was aligned in a direct or inverted repeat configuration (Fig. 5D and E), and amplification of Tn7094 and a plasmid of the composite structure (Fig. 5F). Overall, in addition to Tn7094 amplification being mediated by IS26, we also observed intricate IS26-mediated multimerization of plasmids. Nanopore sequencing confirmed our results for plasmid analyses and provided detailed sequence information for elucidating the dynamics of IS26-mediated amplification (Table 2).

**Amplification of IS26-flanked MDR region with blaKPC-2 is unstable.** Because gene amplification typically results in protein hyperproduction, we intuitively conjectured that this process imposed a fitness cost, at least initially, on the bacterium in response to a forced increase in metabolic activities (15). Accordingly, we assessed the stability of amplified genes in the absence of any antibiotic stress. After transferring a strain E01-7-1 previously exposed to high concentrations of meropenem (128, 256, or 512 μg/mL) back to fresh medium without antibiotics and further culturing it for 24 h, the blaKPC-2 and IS26 copy numbers fell sharply relative to that of repA (Fig. 6A; original data in Table S2). Specifically, upon antibiotic withdrawal, the copy numbers of blaKPC-2 decreased by 2.2, 2.5, or 5.8-fold in the cultural medium compared to groups exposed to 128, 256, or 512 μg/mL meropenem, respectively. Similarly, the copy numbers of IS26 also dropped by 1.4, 1.6, or 3.5-fold, respectively. Further Illumina sequencing analysis demonstrated that the 47 kb MDR region, which was rapidly amplifiable during antibiotic treatment, declined in copy number after 24 h of growth in a medium free of meropenem (Fig. S3A and Table S3). Collectively, these results demonstrated that amplification of these genetic elements was unstable, resulting in regression to phenotypes of lower copy numbers in the absence of antibiotic stress. This agrees with the notion that bacteria tend to maximize their survival by responding to environmental cues and ingeniously switching their gene expression programs in a rapid, resource-limiting manner.

**Amplification of the MDR region also occurred in response to tobramycin stress.** According to genome sequencing analysis, six types of ARGs along with blaKPC-2 were located within the MDR region. Two of these, aac(3)-Ila and aph(3’)-Ia, are known to be resistance determinants for aminoglycosides and were found to be flanked by IS26. A logical question that followed was whether blaKPC-2 amplification can occur under stress from preexposure to antibiotic classes other than β-lactams or carbapenems. To this end, we incubated the E. coli strain E01-7-1 in liquid medium containing 8 μg/mL tobramycin (or 1/2 MIC) for 24 h. We then inoculated the medium with antibiotic pretreated bacterial cells in the presence of various concentrations of tobramycin (from 16 to 128 μg/mL). Strikingly, the cells could grow well at 128 μg/mL tobramycin during regrowth. The copy numbers of aac(3)-Ila, aph(3’)-Ia, blaKPC-2, and IS26 increased relative to that of repA in response to a gradient of antibiotic concentrations. However,

| Table 2: Analysis on blaKPC-2 amplification in pUC57: Tn7094* |
|---|---|---|
| **bla**<sub>KPC-2</sub> copy no. | Read no. | Amplification process |
| 4 | 7 | Tn7094 → pUC57 (Mode 1) |
| 3 | 80 | Tn7094 → pUC57 (Mode 2) |
| 2 | 3733 | Tn7094 → pUC57 (Mode 3) |
| 1 | 273838 | Tn7094 → pUC57 (Mode 3) |

*Amplification of bla<sub>KPC-2</sub> occurs in three ways: amplification of tandem Tn7094 (Mode 1), tandem amplification of the plasmid pUC57: Tn7094 (Mode 2), and a composite structure that combines amplification of Tn7094 and the plasmid (Mode 3). For further details, see Fig. 5.
the copy numbers of \( \text{aph}(3')\text{-Ia} \) and \( \text{bla}_{\text{KPC-2}} \) decreased at 128 \( \mu g/mL \) tobramycin (Fig. 6B; original data in Table S2). These results suggested that unequal amplification of a unit flanked by IS26 and another unit carrying \( \text{aph}(3')\text{-Ia} \) most likely contributed to survival advantages against tobramycin stress. Illumina sequencing confirmed our results for the qPCR assays in which the read-mapping depths of the MDR region relative to that of repA increased with antibiotic concentration except for the \( \text{bla}_{\text{KPC-2}} \) transposon, which exhibited a marked decrease at 128 \( \mu g/mL \) tobramycin (Fig. S3B; original data in Table S3). Collectively, we furnished the first evidence that preexposure to tobramycin instead of meropenem was also sufficient to trigger amplification of the MDR region, which harbored multiple ARGs other than \( \text{bla}_{\text{KPC-2}} \).

**Drug combination in vitro fails to check amplification of MDR region with \( \text{bla}_{\text{KPC-2}} \).** Conventionally, combinations of multiple antibiotics have been used in clinical practice to slow the emergence of AMR with some success (16). Therefore, we asked whether a combination of antibiotics could help retard or halt the amplification of the MDR region. To this end, \( E. \text{ coli} \) strain E01-7-1 was cultured in a liquid medium containing 8 \( \mu g/mL \) tobramycin and 2 \( \mu g/mL \) meropenem for 24 h. Subsequently, the pre-treated cells were subcultured correspondingly in the presence of various antibiotic concentrations. The relative copy numbers of \( \text{aac}(3)\text{-IIa} \), \( \text{aph}(3')\text{-Ia} \), \( \text{bla}_{\text{KPC-2}} \), and IS26 were determined to be 18.22, 29.55, 13.08, and 76.76, respectively, at 32 \( \mu g/mL \) tobramycin and 32 \( \mu g/mL \) meropenem, which were higher than those for exposure to single drugs at the same concentrations (Fig. 6C; original data in Table S2). The copy numbers of \( \text{aph}(3')\text{-Ia} \), \( \text{bla}_{\text{KPC-2}} \), and IS26 decreased at higher levels of antibiotic stress while that of \( \text{aac}(3)\text{-IIa} \) continued to rise. Evidence from Illumina sequencing confirmed our results for qPCR (Fig. S3C and Table S3). These observations indicated that combination therapy not only failed to retard amplification of resistance genes flanked by IS26 but also escalated IS26-driven amplification to a greater extent than any of the monotherapy drugs tested during antibiotic preexposure.

**DISCUSSION**

Sublethal or subinhibitory concentrations of antibiotics do commonly occur in vivo in clinical treatment contexts, which provided a basis for the evolution of adaptively
resilient clones exhibiting heteroresistance (17). \textit{bla}_{KPC-2} has been recognized as an alarming contributor to carbapenem resistance among Gram-negative pathogens worldwide (3, 18–20). Heteroresistant KPC-producers are highly challenging to detect in the laboratory due to their apparent yet deceptively low levels of carbapenem resistance (3, 5, 19). In this study, we provide the first detailed evidence for IS26-mediated amplification of \textit{bla}_{KPC-2} as a driving force for the rapid emergence of carbapenem hyperresistance in \textit{E. coli}, whose intensity is strongly coupled to antibiotic stress following sublethal dose preexposure. However, this amplification is unstable and transient, adding to the difficulty in clinical identification.

In terms of mechanisms, our results demonstrated that IS26-dependent amplification of \textit{bla}_{KPC-2} rapidly drives the emergence of carbapenem hyperresistance following preexposure to sublethal antibiotics, implying that bacterial subpopulations with amplified \textit{bla}_{KPC-2} must have been enriched or induced under a low or sublethal concentration of meropenem (or even other antibiotics) to effectively counter higher-level antimicrobial assaults. In addition, copy numbers of \textit{bla}_{KPC-2} and the IS26 transposon were linked to the strength of antibiotic stress. In the presence of antibiotic-induced selective pressure, resistant bacterial cells endowed with amplification capacities became progressively enriched, whereas, in the absence of antibiotics, susceptible bacterial cells with lower copy numbers acquired a growth advantage. This growth pattern is characteristic of heteroresistance (15). As sublethal antibiotic concentrations can naturally occur \textit{in vivo} in some pharmacodynamic contexts and across tissues of different depths among patients and livestock (12), the phenomenon reported herein could partially account for the failure of antimicrobial intervention against KPC-producers which exhibit deceptively low-level resistance to carbapenems in routine laboratory screens.

It is noteworthy that in addition to \textit{bla}_{KPC-2}, many other resistance genes have been found on compound transposons bounded by IS26, while MDR regions often include multiple IS26 transposons containing several ARGs (21–23). In our study, we compellingly demonstrated efficient amplification of an MDR region containing 16 ARGs mediated by IS26 in response to a single antibiotic or a combination of antibiotics. Previously, evidence has shown that combinatorial use of antibiotics can effectively inhibit amplification caused by a single antibiotic, supporting the wisdom of empirical practice (16). However, our present study has demonstrated situations wherein a drug combination \textit{in vitro} not only failed to halt but exacerbated IS26-mediated amplification of the MDR region. Therefore, we reason that combination therapy may not be universally applicable to all cases of bacterial infections and that greater caution is needed for its use against HAIs caused by multidrug-resistant strains armed with multiple IS26 transposons because insufficiently informed combination therapies may carry their risks of inadvertently complicating resistance development.

Through analysis of long reads, our study also shed new light on the intricate nature of \textit{bla}_{KPC-2} amplification. Multiple IS26 sequences are present in the MDR region of the clinical strain E01-7-1, and various amplified units flanked by IS26 were recomposed from this diverse MDR region landscape during amplification. In \textit{E. coli} carrying the constructed plasmid pUC57:Tn7094 and amplification of Tn7094, the plasmid became cointegrated through homologous recombination, resulting in multiple copies. These data suggest a crucial role of IS26 in mediating MDR region activity and plasmid evolution under selective pressure. The amplified units were frequently rearranged in a tandem array. Consistent with our observations, a recent study showed that tandem amplification of \textit{bla}_{TEM-18} is achieved through excision of a TU from an IS26-flanked transposon and reinserion into an adjacent IS26 (24). However, we did not obtain an assembled TU because \textit{bla}_{TEM-18} amplification was mediated by IS26. On the other hand, an IS26 transposon provides a direct-repeat sequence that is ideal for observing unequal crossing over because two direct repeats on sister chromatids may be incorrectly paired during replication and recombine with each other, resulting in duplication of intervening sequences (25–27). However, we demonstrated with the aid of a constructed plasmid (pUC57 and IS26-\textit{bla}_{KPC-2} 1IS26) that amplification of \textit{bla}_{KPC-2} could
occur in a recA-deficient host (E. coli Top10), indicating that amplification is RecA-independent. Gene duplications arise through nonequal homologous recombination between long, direct-repeat sequences, such as insertion sequences previously considered to be at work in recA-dependent mechanisms (10). Therefore, the mechanisms underlying blaKPC-2 amplification by IS26 require further resolution.

Tandem amplification is generally unstable and transient, making it difficult to trace. Fortunately, genome sequencing has greatly empowered strategies for assessing amplification. To illustrate, tandem repeats of ARGs flanked by IS26 were found in the genomes of E. coli, K. pneumoniae, Acinetobacter baumannii, and Proteus mirabilis (Table S4). Furthermore, ARG repeats mediated by IS1, IS5, IS10, ISCR1, ISCR2, IS1236, IS1216E, IS466, and IS50 (Table S4) have been detected in E. coli, K. pneumoniae, Acinetobacter baylyi, Haemophilus influenzae, Enterococcus spp., Salmonella typhimurium, Streptomyces coelicolor, and Raoultella planticola. These data suggested that tandem gene amplification mediated by IS occurred frequently on bacterial genomes, allowing researchers to gain insights into the pathogenic roles of transposons in the resistance development of bacteria under antibiotic stress.

In summary, our work highlights blaKPC-2 amplification as key molecular underpinnings of bacterial adaptation to the sublethally challenge of single or multiple antibiotics (such as carbapenems and aminoglycosides), which is rapidly and robustly mediated by IS26 under selective pressure. Our evidence also strongly implicates genomic plasticity (such as dynamic changes in blaKPC-2 copy numbers during antibiotic exposure) and intricate genetic rearrangement as critically important to the emergence of carbapenem hyperresistance. Overall, our work has advanced a theoretically useful perspective on identifying bacterial targets for combating the insidious threat from multidrug-resistant pathogens armed with gene amplification machinery. To help improve clinical outcomes, further investigation on novel, accessible detection measures against such under-examined pathogens are urgently needed.

MATERIALS AND METHODS

Bacterial strains, culture, and antibiotics. E. coli E01-7-1, a clinical strain, was isolated from a hospital in Beijing, China, as described previously (13). E. coli Top10 was used for plasmid cloning and amplification experiments. All strains were cultured at 37°C in a lysogeny broth (LB) medium. Meropenem (MEM), tobramycin (TOB), and kanamycin (KAN) were used in the experiments (MEM, TOB, and KAN were purchased from Solarbio). The antimicrobial susceptibility of strains was determined by broth microdilution assays in cation-adjusted Mueller-Hinton broth. MIC determination was performed with a serial dilution technique in 96-well microtiter plates and interpreted in accordance with guidelines set out by the Clinical and Laboratory Standards Institute (28th edition) (http://www.clsi.org/).

Population analysis profile (PAP). E. coli E01-7-1 was initially cultured on LB agar without antibiotics. A single colony was picked from an agar plate and pregrown in 4 mL LB medium overnight (12 h) with shaking at 200 rpm. During the antibiotic preexposure phase, bacterial cells were transferred to 4 mL LB with 0.5 and 2 μg/mL MEM or 4 mL LB without antibiotics followed by incubation at 37°C for 24 h (i.e., 12 h times two). After adjusting bacterial density to an optical density at 600 nm (OD600) of 1.0, plating was performed by serial dilutions on LB agar with or without various concentrations of MEM (0, 0.125, 0.25, 0.5, 1, 2, 4, 8, 16, 32, 64, 128, 256, and 512 μg/mL), CFU was enumerated after 24 h culture at 37°C. The percentage of bacteria exhibiting MEM resistance was calculated as the number of colonies growing on MEM agar plates divided by the number of bacterial colonies grown on plain LB.

In vitro gene amplification experiments. Cells cultured overnight (12 h) as above in PAP were pre-exposed to 2 μg/mL MEM or 4 mL Antibiotic-free LB was added to a bacterial inoculum of 5 × 10^7 CFU/mL followed by incubation at 37°C for 24 h (with transferring once at 12 h). Cells from the two conditions were then separately transferred to a 4 mL LB medium containing various concentrations of MEM (16 μg/mL, 32 μg/mL, 64 μg/mL, 128 μg/mL, 256 μg/mL, or 512 μg/mL) with an inoculum size of 5 × 10^5 CFU/mL. Subsequently, the strains were incubated at 37°C for 24 h during the regrowth phase. Efflux pump inhibition was performed by using phenylalanine-arginine β-naphthylamide (PAβN) as a supplement at a final concentration of 50 μM in LB. A >4-fold decrease in MIC in the presence of PAβN was considered significant. PCR was performed for targeted resistance genes and their promoter, and the PCR products were sequenced by using an ABI3730 sequencer. E. coli Top10 E. coli strain transformed with pUC57:Tn7094 was cultured on LB agar with 50 μg/mL KAN. A single colony was picked from an agar plate and pregrown in a 4 mL LB medium. After 12 h of incubation, bacterial cells were transferred to 4 mL LB with 1 μg/mL MEM or LB without antibiotics as a control followed by further incubation at 37°C for 24 h. Bacterial cells from the two conditions were then transferred to a 4 mL LB medium containing 1 μg/mL or 16 μg/mL MEM. The same methods were used in experiments for TOB and combination drug treatment. Each experiment was performed in triplicate in independent cultures. Bacteria were later harvested for the nitrocefin assay and RNA/DNA or plasmid extraction.
Determination of gene copy numbers through qPCR. Changes in gene copy numbers of bla<sub>KPC</sub>, aac(6)'-Ile, aph(3')-Ia, I526, and aph4 were calculated through qPCR by using the 2<sup>-ΔΔCT</sup> method to determine copy numbers against the single-copy housekeeping gene purA. Primers used here were the same as those for qRT-PCR (Table S5). All experiments were performed with a minimum of three biological replicates and three technical replicates per run. At least three replicates with Ct value standard deviation (SD) < 0.3 were used to determine to mean Ct values.

Additional methods were detailed in Text S1, including β-lactamase enzyme assays, determination of transcription levels by qRT-PCR, whole-genome sequencing, and bioinformatics analysis, evaluation of amplification stability, and construction of pUC57:Tn7094.

Availability of data. All sequence data are available for direct download at the China National Microbiology Data Center (NMDC) with the following accession number: NMDCX0000112.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

TEXT S1. DOCX file, 0.02 MB.
FIG S1. EPS file, 0.1 MB.
FIG S2. TIF file, 0.3 MB.
FIG S3. TIF file, 0.4 MB.
TABLE S1. XLSX file, 0.01 MB.
TABLE S2. XLSX file, 0.02 MB.
TABLE S3. XLSX file, 0.02 MB.
TABLE S4. XLSX file, 0.02 MB.
TABLE S5. XLSX file, 0.01 MB.

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J.F. and D.-W.W. conceived the study. J.F., D.-W.W., and N.-K.W. designed the experiments. D.-W.W., G.Z., C.W., and J.L. performed molecular biology and bacteria experiments. Y.S. and D.-W.W. performed bioinformatic analysis. J.F., N.-K.W., and D.W. wrote the manuscript with inputs from other contributing authors. J.F. and N.-K.W. edited and finalized the manuscript. J.F. supervised the project. All authors read and approved the manuscript.

We declare no conflict of interest.

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