Comparative Evaluation of Different Phenotypic Methods in Detection of Klebsiella pneumoniae Carbapenemase (KPC)

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**ABSTRACT**

In recent years, carbapenem resistance has emerged in Klebsiella pneumoniae (K. pneumoniae) isolates due to acquisition of different carbapenemases enzyme. Carbapenemase producing bacteria are often misidentified by routine microbiological susceptibility testing and incorrectly reported as sensitive to carbapenems; however, resistance to the antibiotic ertapenem is common and a better and sensitive indicator of Carbapenem resistance. Routine laboratory detection of carbapenemase producing K. pneumoniae isolates is crucial, both for a therapeutic management and an efficient infection control. The present study was undertaken to assess the evaluation of different phenotypic methods in detection of K. pneumoniae Carbapenemase (KPC). All clinical isolates of K. pneumoniae was collected from December 2014 to November 2016 was included in the study. Urine, Pus, ET secretions, BAL, Blood and other Body fluids received in the laboratory were subjected to routine processing as per standard operating procedures. Antimicrobial susceptibility testing was performed as per the CLSI guidelines (2013) by Vitek2 automated system to detect carbapenem resistant K. pneumoniae (CRKP) as primary test. Further, confirmatory test was carried by Modified Hodge test, HiChrome agar KPC test and Carba NP test as different phenotypic tests. Total of 1,539 Klebsiella pneumoniae were isolated from various clinical samples. Only, 254 (16.50%) isolates were carbapenem resistant K. pneumoniae (CRKP). In which, 184 (72.44%) were positive with clover-leaf indentation for Modified Hodge Test (MHT) followed by 235 (92.51%) were positive for HiChrome agar KPC test and 239 (94.09%) were positive (Color change within 2hrs of incubation from red to orange/yellow color) for Carba NP test. Among the different phenotypic tests conducted, Carba NP test showed 239(94.09%) were positive, which was rapid (<2 hours), inexpensive technique compared to HiChrome agar KPC test and Modified hodge test. This technique can be implemented in any laboratory for early detection of Carbapenem resistance (CR).

**Keywords**

Carbapenem Resistant-K. pneumoniae, Modified Hodge test, Carba NP test, HiChrome agar KPC test

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**Introduction**

Klebsiella pneumoniae, a gram negative bacilli belonging to the family Enterobacteriaceae are ubiquitously present and reported worldwide. In recent years, K. pneumoniae have become important pathogens in nosocomial infections. An estimated 8% of all nosocomial bacterial infections such as septicemia, pneumonia and urinary tract infections are caused by Klebsiella, mainly by Klebsiella pneumoniae and Klebsiella oxytoca, primarily affecting immuno compromised patients. K. pneumoniae strains that produce extended spectrum β-lactamase enzymes (ESBLs) with
transferable resistance to all β-lactams (except cephamycins and carbapenems) were first detected in the mid-1980s in Western Europe. Currently, there is a worldwide and non-uniform spread of the ESBL-producing *K. pneumoniae* with a prevalence as high as 45.4% of all *K. pneumoniae* isolates. Carbapenem antibiotics are considered as the drugs of choice for the treatment of extended spectrum beta-lactamase (ESBL)-producing Enterobacteriaceae and other multidrug resistant bacteria. The emergence of bacterial strains that produce carbapenemases further limits the therapeutic options available to clinicians. However, resistance to carbapenems is being increasingly detected and is mainly related to the action of carbapenemase-type enzymes. Carbapenem resistant *Klebsiella pneumoniae* is becoming threat globally and also as nosocomial infection because of its resistance to all carbapenem group of antibiotics. This study will help in betterment of patients in ICUs and also help to control nosocomial infections and to know the rapid technique to detect carbapenem resistance.

**Materials and Methods**

This was a hospital based prospective study, undertaken in the department of Microbiology. All clinical isolates of *Klebsiella pneumoniae* were collected from December 2014 to November 2016 was included in the study. Fecal *K. pneumoniae* was excluded from this study and inclusion criteria were only *Klebsiella pneumoniae subspecies pneumoniae*. The samples collected were processed as per standard methods. The study protocol was approved by the ethics committee of the institute.

Blood, Urine, CSF, Sputum, Ear and Nose swab, Abscess, Pus, Wound swab aspirates, Tissue were received in the laboratory were subjected to routine processing as per standard operating procedures. Phenotypic Identification of *K. pneumoniae* was carried out on Mac-Conkey Agar with Pink, round, large, dome shaped, mucoid, Lactose fermenting colonies. Followed by Blood Agar showing Grey, round, dome shaped large colonies. Gram Stain was carried indicating Gram negative rod, thick, short, rod shaped, capsulated, non- motile, 1-2 × 0.5-0.8µm arranged singly or in pairs. Colonies were later subjected to biochemical tests such as, catalase, oxidase, Indole production, Urease hydrolysis, utilization of citrate, Triple sugar Iron agar test, Methyl red test, Voges-Proskauer test as per standard CLSI guidelines. Antimicrobial susceptibility testing was performed as per the CLSI guidelines (2013) by Vitek 2 automated system to detect carbapenem resistant *K. pneumoniae* (CRKP) as primary test. Further, confirmatory test was carried by Modified Hodge test, HiCrome Agar KPC test and Carba NP test as different phenotypic tests.

**Modified Hodge test**

Prepare a 0.5 McFarland dilution of the *E. coli* ATCC 25922 in 5 ml of nutrient broth. Dilute 1:10 by adding 0.5 ml of the 0.5 McFarland to 4.5 ml of saline. Streak a lawn of the 1:10 dilution of *E. coli* ATCC 25922 on Mueller Hinton agar plate and allow to dry 3–5 minutes. A 10 µg Ertapenem susceptibility disk (CT1761B-ETP 10mcg, [B. No.-178667] Oxoid, UK.) was placed in the center of the test area. In a straight line, test organism was streaked from the edge of the disk to the edge of the plate. Up to four organisms can be tested on the same plate with one drug. Incubate overnight at 35°C ± 2°C in ambient air for 16–24 hours. After 16–24 hrs of incubation, examine plate for a clover leaf-type indentation at intersection of test organism and *E. coli* 25922, within zone of inhibition of carbapenem susceptibility disk. MHT Positive test: Clover leaf-like
indentation of *E. coli* 25922 growing along test organism growth streak within disk diffusion zone. MHT Negative test: No growth of *E. coli* 25922 along test organism growth streak within disc diffusion zone.

**HiCrome Agar KPC Test:** (HIMEDIA M1831-100G)

Suspend 16.50g in 500ml distilled water. Heated to boiling point to dissolve the medium completely. Sterilized by autoclaving at 15lbs pressure (121°C) for 15 minutes. Cooled to 45-50°C and aseptically the rehydrated contents of one vial of HiChrome KPC selective supplement (FD279) was added. Mixed well and poured into sterile Perti-plates.

Once the media is solidified, media plates are kept for quality control check for 24 hours, if plates do not show any growth, then quality check is approved. Direct plating of the sample is carried out using inoculation loop and incubated for 18-24 hours. After incubation plates are observed for Metallic Blue Colonies of *Klebsiella pneumoniae* which is considered to be positive. Bacterial colonies grown other than *Klebsiella pneumoniae* will be indicated by color differentiation as: *E. coli*-Pink color, *Pseudomonas*- Cream color, Bacteria which are not resistant to carbapenems will be inhibited/ not grown on culture plates.

**Carba NP Test**

One calibrated loop (10 μL) of the tested strain directly recovered from the antibiogram was resuspended in a Tris-HCl 20 mmol/L lysis buffer (S D Fine Chemical Limited-SDFCL [37119 K01]-C16A/1915/0409/13, Mumbai, India). Vortexed for 1 minute and further incubated at room temperature for 30 minutes. This bacterial suspension was centrifuged at 10,000 × g at room temperature for 5 minutes. 30μL of the supernatant, corresponding to the enzymatic bacterial suspension, was mixed with 100μL of a 1-mL solution made of 3 mg of Imipenem and Cilastin injection IP (Lupinem, Himachal Pradesh, India.[B.No.- HZP6001]), pH 7.8, phenol red solution(Nice Chemicals(P) LTD[P40671]-B.No.-409119, Kochi, India), and 0.1 mmol/L ZnSO₄ (S D Fine Chemical Limited-SDFCL [40621 K05]-C16A/2516/0801/08, Mumbai, India). The phenol red solution was prepared by mixing 2mL of a phenol red (Lupinem, Himachal Pradesh, India [B.No.- HZP6001]) solution 0.5% (wt/vol) with 16.6 mL of distilled water. The pH value was then adjusted to 7.8 by adding drops of 1 N NaOH. A mixture of the phenol red solution and the enzymatic suspension being tested was incubated at 37°C for a maximum of 2 hours and the results were interpreted.

The color of the vials turned from red to orange or yellow for tested strains that were producing carbapenemases, whereas vials corresponding to bacterial extracts of isolates that did not produce carbapenemase remained red. The color changed from red to yellow as early as 5–10 minutes after incubation for KPC producers began, 30 min-VIM, 15-60 min- MBL (NDM-1), 30-60 min – OXA-48 may be present. In most cases, incubation for 30 minutes is sufficient for obtaining a frank color change for carbapenemase producers.

**Results and Discussion**

A total of 1,539 *Klebsiella pneumoniae* was isolated from various clinical samples over a period of two years from December 2014 to November 2016. Out of 1,539 *Klebsiella pneumoniae* isolates collected, 254 (16.50%) isolates were carbapenem resistant *Klebsiella pneumoniae* (CRKP) (resistant to Imipenem/ Ertapenem/Meropenem/Doripenem antibiotics) by Vitek 2.
Modified Hodge Test (MHT) result

Out of 254 carbapenem resistant *Klebsiella pneumoniae* (CRKP) isolates, 184(72.44%) were positive with clover-leaf indentation for Modified Hodge Test (MHT) and 70(27.55%) were negative without clover-leaf indentation respectively (Fig. 1).

HiCrome KPC agar test result

Among 254 carbapenem resistant *Klebsiella pneumoniae* (CRKP) isolates, 235(92.51%) were positive (Metallic blue) for HiCrome KPC agar test and 19(7.48%) were negative by HiChrome KPC agar test (Fig. 2).

Carba NP test result

Out of 254 carbapenem resistant *Klebsiella pneumoniae* (CRKP) isolates, 239(94.09%) were positive (Color change within 2hrs of incubation from red to orange/yellow color) for Carba NP test and 15(5.90%) were negative without color change (showing red color) (Fig. 3).

Among 254 CRKP isolates, 181(71.25%) were True Positive in all the test like HiCrome KPC agar test, Modified Hodge Test and Carba NP Test and 15(5.90%) were True Negatives.

The potential of *Klebsiella pneumoniae Carbapenamase* (KPC) production among gram negative bacteria along with the development of resistance to carbapenem antibiotics have made difficult and important problem in the treatment of infection.

Detection of KPC producing strains by phenotypic method allows fast, cost effective and accurate results that can prevent spread of this mechanism of resistance. Many different phenotypic methods for detection of KPC have been suggested. Reduced susceptibility to carbapenems are indirect indicators, on which rapid identification of KPC can be followed. All *Klebsiella pneumoniae* isolates were inoculated on Mac Conkey agar medium. Antibiotic susceptibility testing was performed and CRKP was detected by the Vitek 2. In our study, a total of 254(16.50%) *Klebsiella pneumoniae* isolates were Carbapenem resistant from Vitek 2 result. The first commercially available chromogenic medium HiCrome agar KPC was designed for isolation of carbapenem-resistant (CR) gram negative bacteria. Carbapenem resistant *Klebsiella pneumoniae* (CRKP) were excellently detected by HiCrome agar KPC medium by color and morphological characters, allowing rapid differentiation of the bacterial colonies. HiCrome agar KPC showed sensitivity and specificity when related to PCR (gold standard) were 100% and 98.4% respectively.

In our study, from 254 CRKP isolates, 235(92.51%) were positive (Metallic blue) for Crome agar KPC test and 19(7.48%) were negative by KPC test. 35% positive were seen in the test performed by Zmirasamra et al., with metallic blue colony, which is showing less positive rate when compared to our result and Crome agar medium can also be used for fast and direct identification of CRKP gram negative pathogen from clinical isolates.

To rapidly identify carbapenemase producers in *Enterobacteriaceae*, Patrice Nordmanett al., developed the Carba NP test based on the colorimetric detection of hydrolysis of the β-lactam ring of a carbapenem molecule. The test is based on in vitro hydrolysis of a carbapenem, ertapenem /imipenem and uses isolated bacterial colonies for identification. It was 100% sensitive and specific compared with molecular-based techniques. This rapid (<2 hours), inexpensive technique may be implemented in any laboratory.
**Fig.1** Modified Hodge test

Test Negative without Clover-leaf indentation for Modified Hodge Test.

Test positive with Clover-leaf indentation for Modified Hodge Test.

**Fig.2** HiChrome agar KPC test

Test sample showing Metallic blue Carbapenem resistant Klebsiella pneumoniae colonies. Positive test for HiChrome KPC agar.

**Fig.3** Carba NP test

Control tube showing negative result with Red colour change for Carba NP Test

Test tube showing Positive result with Yellow colour change for Carba NP Test
Table 1: Comparison of HiCrome KPC agar test, Modified Hodge test and Carba NP test result

<table>
<thead>
<tr>
<th></th>
<th>KPC-P</th>
<th>KPC-N</th>
<th>MHT-P</th>
<th>MHT-N</th>
<th>CNP-P</th>
<th>CNP-N</th>
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<tbody>
<tr>
<td>KPC-P</td>
<td>235(92.51%)</td>
<td>-</td>
<td>177(69.68%)</td>
<td>58(22.83%)</td>
<td>234(92.12%)</td>
<td>01(0.39%)</td>
</tr>
<tr>
<td>KPC-N</td>
<td>-</td>
<td>19(7.48%)</td>
<td>04(1.57%)</td>
<td>15(5.90%)</td>
<td>05(1.96%)</td>
<td>14(5.51%)</td>
</tr>
<tr>
<td>MHT-P</td>
<td>180(70.86%)</td>
<td>04(1.57%)</td>
<td>184(72.44%)</td>
<td>-</td>
<td>184(72.44%)</td>
<td>00(0.0%)</td>
</tr>
<tr>
<td>MHT-N</td>
<td>55(21.65%)</td>
<td>15(5.90%)</td>
<td>-</td>
<td>70(27.55%)</td>
<td>55(21.65%)</td>
<td>15(5.90%)</td>
</tr>
<tr>
<td>CNP-P</td>
<td>234(92.12%)</td>
<td>05(1.96%)</td>
<td>183(72.04%)</td>
<td>56(22.04%)</td>
<td>239(94.09%)</td>
<td>-</td>
</tr>
<tr>
<td>CNP-N</td>
<td>01(0.39%)</td>
<td>14(5.51%)</td>
<td>00(0.0%)</td>
<td>15(5.90%)</td>
<td>-</td>
<td>15(5.90%)</td>
</tr>
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</table>

In the current study, 254 CRKP isolates were tested for Carba NP test. Test result showed, 239(94.09%) were positive (Color change within <2 hours of incubation from red to orange/yellow color) for Carba NP test and 15(5.90%) were negative without color change (showing red color). Our results were similar to the result of Patrice Nordmann et al.,\(^9\). The strains give non interpretable results when bacterial colonies recovered from Drigalski lactose agar or MacConkey agar, regardless of carbapenemase production. Such results might be a consequence of the accumulation of lactic acid in the bacterial isolates that were able to ferment lactose. Therefore, the Carba NP test is not suitable to identify carbapenemase producers grown on Drigalski lactose agar and MacConkey agar plates.\(^10\)

In the present study, 254 CRKP were further tested for KPC and MBL production by Modified Hodge Test (MHT). Recently CLSI accepted MHT as specific and sensitive phenotypic method for carbapenemase detection\(^4, 5\). Out of 254 carbapenem resistant *Klebsiella pneumoniae* (CRKP) isolates, 190(74.80%) were positive with clover-leaf indentation for Modified Hodge Test (MHT) and 64(25.19%) were negative without clover-leaf indentation respectively. Our results are similar with many other studies like, 75% positive result showed by McGettigan et al.,\(^6\) followed 69% by Amjad et al.,\(^11\), Cury et al.,(MHT 71% positive)\(^4\) and little higher rate was seen in Galani et al., with 98% \(^12\), respectively.

In the present study, we have got 181(71.25%) true positives and 15 (5.90%) true negative result for Crome agar KPC test, Modified Hodge Test and Carba NP test. This was similar to the study of Vanessa BleyRibeiro et al., with 70% true positives and showing slightly high, true negative results of 29%. KPC carbapenemase alone were seen, which may not be applied for other types of carbapenemase, hence MHT is not considered to be specific for KPC detection and sensitivity and specificity may differ in the similar way with other carbapenemases. With minimum carbapenemase activity, MHT initially aims to recognize and differentiate the presence of carbapenemase production with that of other enzymes.\(^13\) The Carba NP test has many advantages. It is not expensive, fast, can make multiple copies, and quick to detect and specific. It completely removes the need for using other proficiency to identify carbapenemase producers that are long-delayed and less receptive or specific. Using this distinct test would enhance the detection of patients infected or colonized with carbapenemase producers.\(^9\)
In our study; out of 235 Crome agar KPC Test positive isolates; 234 (92.12%) were positive for Carba NP test, which was similar to the result of HiCrome agar KPC test positive isolates and 01(0.39%) were negative for Carba NP test. Were as, 177 (69.68%) isolates were found to produce carbapenemase enzyme by MHT and 58(22.83%) were found negative for MHT. The reason for negative result of bacterial strains which were carbapenem resistant but negative by MHT could be explained as the overproduction of ESBL or Amp C enzyme with porin loss. The processing of MHT showed that it is easy, simple with minimum infrastructure and of less cost, reliable test and this can be implemented in routine microbiology laboratories for detection of carbapenemase producers. One isolate (Sample No. -02) was negative by Carba NP Test and Modified Hodge Test but was positive by HiChrome KPC agar test. This isolate will be tested further for the presence of KPC gene by Molecular methods (PCR).

MHT was not specific for production of KPC as per Anderson et al., \(^{15}\); where as in other study by Girlich et al., \(^{16}\) showed 77.4% sensitivity and 38.9% specificity for MHT test in detecting KPC in overall test. This low sensitivity and specificity may be because, MHT detects other carbapenemase enzymes in addition to KPC as it is the only indicative enzymatic activity of carbapenemase and cannot differentiate class A carbapenemases from class B MBLs. And MHT cannot be used as a confirmatory test for recognition of the KPCs because of the difficult elucidation and false positive results. False-positive results are more common in isolates producing AmpC and CTX-M b-lactamase. \(^{17,18,19}\)

Isolates with reduced susceptibility to one or more carbapenems undergo further testing with the Modified Hodge Test (MHT), in order to detect the presence of a carbapenemase enzyme as recommended by CDC. \(^{20,9}\). And the testing of MHT is not necessary, when the isolate is found to be intermediate or resistant to all carbapenems tested by CLSI guidelines, since use of the recently reduced breakpoints should preclude the possibility of misclassification of CR-KP as carbapenem susceptible. They do, however, suggest its use for epidemiological investigation. \(^{21,9}\)

In our study, out of 184 (72.44%) MHT positive isolates; 4 (1.57%) isolates were found Crome agar KPC test negative (with no metallic blue colony). This may be because, the remaining MHT positive isolates which were negative for KPCs and MBLs presumably had the other types of carbapenemase enzymes as the oxacillinase enzyme, as the OXA encoding gene has been found to be highly disseminated in the last few years. Not only do KPC-producing organisms hydrolyze carbapenems, but also, they are often resistant to multiple other antibiotics. \(^{22}\). But, all the 184(72.44%) MHT positive isolates were seen positive for Crome agar KPC test (Table 1).

In the current study, among 254 CRKP isolates, 239(94.09%) Carba NP test positive isolates, 234(92.12%) were positive for Crome agar KPC test and 183(72.04%) were positive for Modified Hodge Test. And 15(5.90%) isolates showed negative result for Carba NP test out of 254 CRKP isolates. This was very much similar to the result of Crome agar KPC with 14(5.51%) negative test isolates and Modified Hodge Test showing 15(5.90%) negative result.

Resistance to carbapenems is being increasingly detected and is mainly related to the action of carbapenemase-type enzymes and becoming threat globally and also as nosocomial infection because of its resistance.
to all carbapenem group of antibiotics. In our study, 254 (16.50%) isolates were carbapenem resistant *Klebsiella pneumoniae* (CRKP). Among the different phenotypic tests conducted, Carba NP test showed 239 (94.09%) were positive (Color change within 2hrs of incubation from red to orange/yellow color), which was rapid (<2 hours), inexpensive technique compared to HiChrome agar KPC test and Modified hodge test. This technique can be implemented in the Microbiology laboratory for early detection of Carbapenem resistance (CR). Further, the test results need to be compared with Molecular results, as PCR is gold standard for detection of Carbapenem resistance and genes (KPC’s and MBL types) coding for carbapenem resistance can be detected.

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