Challenge to healthcare: Multidrug resistance in Klebsiella pneumoniae

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Abstract. Klebsiella pneumoniae has been associated with different types of infections and one of the important aspects of Klebsiella associated infection is the emergence of multi-drug resistant strains particularly those involved in nosocomial diseases. For our studies we have collected fifty – nine clinical isolates, from different places of India as this study was conducted in India. Morphologically resembled Klebsiella were recovered from respiratory, UTI and pus cases. Out of fifty-nine clinical isolates, twenty were found positive for K.pneumoniae at biochemical characterization. These clinical isolates of K.pneumoniae were further tested for antimicrobial sensitivity and most of them were found to be multidrug resistant. All confirmed K.pneumoniae isolates were resistant to carbenicillin and one among them recovered from sputum sample of a pneumonic patient was resistant to all the antimicrobial agents tested except exhibiting a partial susceptibility to amikacin. Klebsiella pneumoniae strains from clinical cases in our study were found highly susceptible to quinolones, aminoglycoside, amikacin and gentamycin. At the same time over 60% strains were resistant to chloramphenicol and tetracycline. We also found that 28 to 76% of them were resistant to cephalosporins (ceftizoxime and cefotaxime).

Keywords: Klebsiella pneumoniae, Antimicrobial sensitivity, multidrug resistance

1. Introduction

Klebsiella are ubiquitously present and reported worldwide. In recent years, Klebsiellae have become important pathogens in nosocomial infections [1]. The importance of Klebsiella species in the ever increasing number of gram negative aerobic bacillary nosocomial infections in the United States [2] and India [3] has been well documented. Epidemic and endemic nosocomial infections caused by Klebsiella species are leading causes of morbidity and mortality [4]. In addition to being the primary cause of respiratory tract infections like pneumonia, rhinoscleroma, ozaena, sinusitis and otitis, it also causes infections of the alimentary tract like enteritis, appendicitis and cholecystitis. They are frequently associated with the infections of urinary tract, genital tract, and the eyes. Statistical data and evidences from researches prove that multidrug resistance bacteria are emerging worldwide which is a big challenge to healthcare. Multidrug resistant bacteria cause serious nosocomial and community acquired infections that are hard to eradicate using available antibiotics. Moreover, extensive use of broad-spectrum antibiotics in hospitalized patients has led to both increased carriage of Klebsiella and the development of multidrug-resistant strains that produce extended–spectrum beta-lactamase (ESBL). Epidemic strains of cephalosporin resistant K.pneumoniae have been associated with increased morbidity and mortality in hospitalized patients [5]. Since 1983, nosocomial outbreaks of ESBL producing K.pneumoniae infections in Europe [6], the United States and South America have been described [7,8]. Between 1990 and 1992, 5% of K.pneumoniae clinical isolates produced ESBLs [9]. In France, 10 to 30 % of K.pneumoniae strains are reported to produce plasmid mediated ESBLs of the TEM or SHV families [7]. We have started our studies with following aims and objectives:

- To analyze the severity of multidrug resistance of Klebsiella pneumoniae in India
• Challenges faced by healthcare community in India during treatment of multi drug resistant *K. pneumonia* patients to overcome this problem.

• To come up with some suggestions to overcome multidrug resistant problem.

2. Materials and Methods

We have collected standard and clinical isolates of *K. pneumoniae* from different places of India and utilized them for their antibiotic resistant ability.

2.1. Bacterial strains:

Standard culture of *K. pneumoniae* strain 3296 was collected from Yamaguchi University, Japan whereas total fifty nine clinical isolates were collected from different places of India. Out of all, fifty-four clinical isolates of *Klebsiella* species were recovered from pus, urine and sputum samples from Armed Force Medical College, Pune, India, four samples from pneumonic patients and one UTI sample from Patel Chest Hospital, New Delhi, India.

2.2. Biochemical characterization:

All the clinical isolates were examined morphologically for colony characteristics on agar media. Those exhibiting mucoid colonies were processed for biochemical testing. Biochemical test employed were urease production, citrate utilization and fermentation of sugars. Sugar fermentation tests performed were sucrose, glucose, mannitol, lactose, adonitol, dulcitol, melibiose and esculin. Indole test and H$_2$S production on TSI agar, oxidase, catalase and nitrate were also carried out. Besides these tests, motility and growth of organism in potassium cyanide were also checked. For biochemical tests standard procedures were used [10].

2.3. Antibiotic sensitivity testing:

Standard and positive clinical samples of *K. pneumoniae* were grown on nutrient agar. After 24 hours incubation at 37°C, fresh colonies of *K. pneumoniae* were grown in a tube containing 5mL of a tryptic-soy broth followed by overnight incubation at 37°C until it achieves the turbidity. Next day, dipped a sterile non-toxic swab into the broth and rotated the swab several times, pressing firmly on the inside wall of the tube above the fluid level to remove excess inoculum from the swab. Organism was inoculated over the dried surface of a Muller-Hinton agar plate by streaking the swab over the entire sterile agar surface three times, rotating the plate 60° each time. We have placed the antibiotic discs evenly on the surface of the agar plate by using a sterile forceps then plates were placed in incubator at 37°C. After 24 hours of incubation, we examined each plate and reported the organism to be susceptible, moderately susceptible (intermediate), or resistant for antibiotics.

3. Results

The standard and clinical isolates of *Klebsiella* were examined by battery of biochemical tests. Standard *K. pneumoniae* cultures showed positive reactions for urease production, citrate utilization, catalase reaction and fermentation of sugars like glucose, lactose, sucrose, mannitol, adonitol, melibiose and esculin. Organisms showed negative test for indole production. There was no H$_2$S production on Triple Sugar Iron agar but growth of organism was seen in KCN. All fifty-nine clinical isolates exhibiting colonies similar to *Klebsiella* species were tested biochemically. Thirty-six clinical isolates showed typical common biochemical reaction pattern similar to the one seen with *Klebsiella* species, being positive to glucose, lactose, sucrose, mannitol and negative to oxidase and indole. When tested by second battery of biochemical reactions that included some rare sugars (adonitol, melibiose, esculin and dulcitol), twenty of them exhibited reactions attributable to majority of *K. pneumoniae* sub species pneumonia strain, being positive to adonitol, melibiose, esculin, urease and citrate. They also showed growth in KCN. The rest of the isolates had variable reactions with these tests.
Antibiotic sensitivity testing of twenty confirmed *K. pneumoniae* clinical isolates was done on Muller – Hinton agar plates. On the basis of resistance to antibiotic, strains were categorized in three groups. One group had strains which were susceptible (over 85%) to quinolones and aminoglycosides. The second group had strains which were moderately susceptible (intermediate) to antiribosomal antibiotics (chloramphenicol and tetracycline) with 62% resistant strains. The third group contained strains which were resistant to semi-synthetic penicillins, ampicillin, carbenicillin (76-100%) and to co-trimoxazole (76%). Results are shown in fig 1.

Fig 1: Antibiotic resistance pattern of Klebsiella pneumoniae isolates

### 4. Discussion

In vitro data showed a wide range of beta-lactams, aminoglycosides, quinolones and other antibiotics are useful for treatment of klebsiellae infections [11, 12, and 13]. The clinical isolates of *K. pneumoniae* were tested for antimicrobial sensitivity and most of them were found to be multidrug resistant. All the *Klebsiella pneumoniae* isolates were resistant to carbenicillin and one among them recovered from sputum sample of a pneumonic patient was resistant to all the antimicrobial agents tested except exhibiting a partial susceptibility to amikacin. In such cases the disease is prone to progress to permanent debilitation or death of the patient if, isolation and identification of the causative agent and the subsequent antimicrobial susceptibility testing is not carried out at the early stage of the disease.

In our studies, *K. pneumoniae* strains from clinical cases were found highly susceptible to quinolones and aminoglycoside, amikacin and gentamycin. At the same time over 60% strains were found resistant to chloramphenicol and tetracycline. Twenty-eight to 76% of them were resistant to cephalosporins (ceftizoxime and cefotaxime). Cephalosporins have been widely used as monotherapy and in combination with aminoglycosides for the treatment of Klebsiella infection.

Plasmid encoded resistance to broad spectrum cephalosporins is becoming a widespread phenomenon in clinical medicine. These antibiotics are inactivated by an array of different extended spectrum beta lactamases (ESBLs) which have evolved by stepwise mutation of TEM/SHV type beta lactamases. Plasmid encoding these enzymes has been encountered in several members of the family enterobacteriaceae, but are, for unknown reasons, most often harboured by *K pneumoniae* [14]. Epidemic and endemic nosocomial infections caused by ESBL producing *K pneumoniae* represent a persistent problem in many parts of the world, especially in ICUs [15 and 16]. The emergence of multidrug resistant strains particularly those involved in nosocomial diseases and the alarming rise in resistance to SHV and ESBL producing groups of
antibiotics result in high morbidity and mortality. Early identification of agent, therefore, is important for timely management of patients.

Klebsiella has been associated with different types of infections and one of the important aspects of Klebsiella associated infection is the emergence of multi-drug resistant strains particularly those involved in nosocomial diseases. The alarming rise in resistance to SHV and ESBL producing groups of antibiotics result in high morbidity and mortality. TEM- and SHV type ESBL producing Klebsiella pneumoniae were extensively reported worldwide after it was first identified in enterobacterial isolates from India [17]. The high prevalence of these drug resistant strains has further necessitated the requirement of a rapid and accurate identification system for K.pneumoniae.

In our studies, we have found that out of fifty-nine clinical isolates, twenty clinical isolates were found positive for K.pneumoniae at biochemical characterization. When these clinical isolates of K.pneumoniae were further tested for antimicrobial sensitivity and most of them were found to be multidrug resistant. The isolates were highly susceptible quinolones and the aminoglycosides. Over sixty percent strains were resistant to chloramphenicol and tetracycline but all the isolates were resistant to carbenicillin.

5. Conclusion

Statistical data and evidences from researches prove that multi drug resistant bacteria are emerging worldwide which causes many public health problems and challenges to healthcare. Moreover, uses of broad spectrum antibiotics, insufficient aseptic condition and technique with inadequate control of infections spread had aggravated this problem. Recently, WHO warned society during press release and stated that antibiotics may lose their power to cure disease if action is not taken now against antimicrobial resistance problem [18]. WHO also recommended six ways to overcome multi drug resistant problem, those are:

- Committing to a comprehensive, financed national plan with lines of accountability and community engagement;
- Strengthening surveillance and laboratory capacity;
- Ensuring a regular supply of good-quality medicines;
- Regulating and promoting rational use of medicines and proper patient care;
- Enhancing infection prevention and control in health settings; and
- Fostering innovation, research and development

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7. References


