RESEARCH ARTICLE

MULTIDRUG RESISTANCE PROFILE AND METALLO-β-LACTAMASE PRODUCTION IN PSEUDOMONAS FLUORESCENS ISOLATED FROM HOSPITAL.

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Abstract

Background: Pseudomonas fluorescens is an ubiquitous, Gram-negative, bar molded, mono whipped bacterium, equipped for causing genuine diseases in safe traded off host. It is a standout amongst the most troublesome multidrug safe microorganisms equipped for delivering metallo-β-lactamase (MBL). This examination was gone for assessing the expanding commonness of multidrug opposition profile and MBL generation in P. fluorescens detached from healing center of CIMS, Bhopal India.

Result: In this examination, a sum of 600 examples were haphazardly gathered from the doctor’s facility. They were tried for the nearness of P. fluorescens on cetrimide agar, watching the agar plates for development and pigmentation. Encourage biochemical test and sub-atomic investigation were done to additionally affirm their way of life as P. fluorescens.

Conclusion: From this investigation, it was presumed that multidrug safe P. fluorescens is available in the hospital. The level of opposition was high in both shade and non-color creating disconnects, however the obstruction of non-color delivering P. fluorescens was higher. Medications of decision to be viewed as first for the therapy of infections produced by multidrug safe P. fluorescens are gentamicin, ofloxacin, ciprofloxacin and meropenem.

Introduction:

Pseudomonas fluorescens belongs to the family Pseudomonadaceae and consists of gram-negative, rod shaped, nonfermentive, oxidase positive bacilli [1][2]. It is known to reveal ingrained resistance to several antimicrobial agents. It also produces enzymes, namely β-lactamases, which is responsible for the wide spread β-lactam resistance. These β-lactamases hydrolyze the amide bond of the four-membered characteristic β-lactam ring, thus rendering the antimicrobial ineffective [3][4].

Pseudomonas fluorescens in maximum cases is not perceived as bacterial pathogen in humans; however, miscellaneous culture-based and culture liberated studies has been discovered at low proportion in the primordial microbiota of variegated body sites. While most extensively designed for its contribution in the soil and the rhizosphere. P. fluorescens can induce bacteremia in humans, with nearly all proclaimed cases being allocated either to exchange of contaminated blood products or to use of spoiled equipment integrated with intravenous infusions. There is also tantalizing amalgamation between P. fluorescens and human disease, in that approximately 50% of...
Crohn’s disease patients develop serum antibodies to *P. fluorescens*. Altogether, these reports are genesis to highlight a far more recurrent, tempt, and potentially complex alliance between humans and *P. fluorescens* during health and disease.[5]

Metallo-beta-lactamases (MBLs) are B class enzymes which hydrolyze carbapenems [6]. They have been described as the enzymes which require divalent cations, normally, zinc, as a metal cofactor for chemical movement and are restrained by metal chelators, for example, ethylenediamine tetra acidic corrosive (EDTA).[7]

For severe Pseudomonas infections Carbapenems are the antibiotics of choice [8]. Carbapenems are the B-lactam gathering of medications that are generally utilized as antimicrobials. The steady reaction of *P. fluorescens* to broadened range B-lactamases (MBL) creating strains[9]. Multi-sedate safe (MDR) *P. fluorescens* are equipped for creating chemicals that can inactivate beta-lactams, for example, metallo-B-lactamase (MBL) or, in other words a noteworthy extent of carbapenem opposition in these microbes.

Hence, this examination was intended to research multidrug and carbapenem safe *P. fluorescens* in healing center to prevent its transfer from the creatures to vulnerable human host to cause zoonotic infection. This will be finished by researching the effects of a few anti-toxins on the development of *P. fluorescens* secluded from the hospital of grown-up male and the hospital of CIMS, Bhopal, India.

**Methods:-**
Under sterile condition the cotton sticks were vaccinated into clean supplement stocks and hatched at 35°C for 24h. The stocks were then be streaked on officially arranged cetrimide agar plates which is a strong specific medium utilized for the disengagement and recognizable proof of *P. fluorescens*. [10]. It restrains bacterial development apart from *P. fluorescens* & furthermore improves Fluorescein & Pyocyanin color creation. The cetrimide agar planning was set up as indicated by producer’s directions and disinfected in the autoclave at 121°C for 15 min. Results were pursued after 24h of hatching at 35°C.

**Identification of the Isolates**
Confines were recognized dependent on settlement morphology, generation of pigmentation and non-color creation on cetrimide agar. Gram recoloring catalase test and atomic examination were similarly done on the seclude to additionally affirm the detaches as *P. fluorescens*.

Air dried slides were overwhelmed with Precious stone violet and Gram’s iodine for 1 min and were washed with water. Before being decolorized utilizing 95% liquor or (CH3)2CO for around 10-20sec & washed with water. Before being washed safranin included for 1 min. Air dried slides saw under the magnifying instrument utilizing the oil drenching focal length.

**Catalase test:**
This test was utilized to demonstrate the generation of catalase, a compound that catalyzes the arrival of O2 from H2O2. A sterile circle stick was use to transfer a little measure of state development on the surface of a spotless, dry glass slide, a drop of 3% H2O2 is dropped on the province on the glass slide, the fast advancement of O2 bubbles watched demonstrated a positive outcome for *P. fluorescens*.

**Phenotypic Screening for Metallo-β-lactamase (MBL):**
Screening of MBL-delivering confines was performed utilizing the joined circle dissemination strategy on the detaches that were multidrug safe from both pigmented and non-pigmented secludes. The detaches were phenotypically screened for the nearness of a metallo-β-lactamase (MBL) utilizing the metal chelating specialist EDTA. Distinguishing proof of MBL action was dictated via carbapenem-EDTA consolidated circle technique. Two imipenem and two meropenem IMP (10µg) and MEM (10µg) circles were put on a Muller Hinton agar plate immunized with *P. fluorescens* suspension of 0.5 Mcfarland turbidity principles and 10µ of a clean 0.5 M EDTA (pH 8.0) arrangement was connected on one circle of the 2 imipenem and the meropenem. The plates were brooded at 35°C for 24h. The zones of hindrance around the IMP & IMP-EDTA circles and MEM and MEM-EDTA plates were watched and looked at.
Sub-atomic Analysis
Separates were additionally recognized utilizing the polymerase affix response intensification to affirm that they are P. fluorescens. A few strains of Pseudomonas were chosen arbitrarily and exposed to quality sequencing which were confirmed by grouping of 16S r RNA quality as per the accompanying strategy.

DNA extraction method:-
The DNA extraction was completed on the examples utilizing the Jena Bioscience Bacteria DNA preparation Kit which is a turn segment based bacteria DNA preparation kit. It was intended for fast and high immaculateness seclusion of genomic DNA from Gram-positive and Gram-negative microscopic organisms. The turn section based strategy totally expels PCR inhibitors, for example, divalent cations and proteins bringing about a high virtue arrangement of genomic DNA. Materials included Resuspension Bufr, Lysis Bufr, Binding Bufr, RNase Alysozyme (Before utilize, ‘twofold’ refined water was included as demonstrated the bottle) and put away at -20°C, Proteinase K (before utilize, ‘twofold’ refined water was included as shown the tube) and put away at -20°C, Activation Bufr, Washing Bufr(before utilize, 96-99% Ethanol was included as demonstrated the tube), Elution Bufr, Spin segments and 2 ml accumulation tubes, 96-99% Ethanol, double refined water and microtubes 1.5 or 2.0 ml. (we use any advance technique).

PCR Amplification of the PGSS Gene
Polymerase chain response was done to enhance the quality of the microbes utilizing the preliminary match PGSS F (GACGGGTGAGTAATGCCTA) and PGSS R (CACTGGTGTTCCTTCTATA). The PCR response was done utilizing the Solis Biodyne 5x hot firepol mix ace blend. PCR was performed in 25 µl of a response blend. The response focus was brought down from 5x fixation to 1x focus containing 1x mix ace blend bufr, 1.5 mm MgCl2, 200 µm of each deoxynucleoside triphosphates (dNTP), 25pMol of every groundwork, 2 unit of hot firepol DNA polymerase, Proofreading Enzyme. Five microlitre (5 µl) of the extracted DNA and sterile refined water was utilized to make up the response blend. Warm cycling was led in an Eppendorf Vapo ensure warm cycler for an underlying denaturation at 95°C for 5 min pursued by 30 enhancement cycles of 30 s at 95°C; 1 min at 58°C and 1 min or 30 s at 72°C. This was trailed by a last expansion venture of 10 min at 72°C. The intensification item was isolated on a 1.5% agarose gel and electrophoresis was completed at 80V for 1 h 30 min. After electrophoresis, DNA groups were pictured utilizing ethidium bromide recoloring. Hundred base sets DNA stepping stool was utilized as DNA sub-atomic weight marker. Observation of band in the 600bp locale indicated the enhancement of 16S rRNA quality. PCR items were sent for Sanger sequencing. All sequenced information were exposed to BLAST examination on (http://www.ncbi.nlm.nih.gov/Blast.cgi). [11]

Atomic Detection of Resistance Genes
Add up to DNAs of various bacterial segregates were removed by the DNA extraction unit (Bioneer Company, Korea, Cat. number K-3032-2). The presence of blaIMP, blaVIM, blaDIM, blaAIM, blaGIM and blaNDM qualities were controlled by PCR utilizing the accompanying appropriate groundworks: IMP-F(GGAATAGAGTGGCTTAAYTCTC), IMP-R (GGTTTAAYAAAACAACCACC),(232bp) for blaIMP, VIM-F(GATGGTGTTTGGTGCATA), VIM-R(CGAATGCCGCACCCAG) (390 bp) for blaVIM, DIM-F(GCTTGTCTTCGTTCTACAC), DIM-R(CGTTCCGCGATTGATTTT) (699 bp) for blaDIM, AIM-F(CTGAAGGTGTACGGAAACAC), AIM-R(GTTCGGCCACCTCGAATTTG),(322bp) for blaAIM, GIM-F(TGACACACCGCTGCTGAA), GIM-R(AACTTCCAACCTTTGCCATGC) (477 bp) for blaGIM, NDM-F(GGTTTGGCGATCTGTGTTTTC) and NDM-R(CGAAATGCCGTACATCACGATC) (621 bp) for blaNDM (15). Responses were performed on a warm cycler (Eppendorf, Master Cycler Gradient) and PCR programs utilized in this examination were as beforehand depicted (19). PCR item groups were examined after electrophoresis on a 1.5% agarose gel at 95 V for 45 minutes in 0.5x Tris/Borate/EDTA (TBE) containing ethidium bromide, and the outcome was checked under UV illumination. P. fluorescens PA53 (ACCESSION: KM359726) for IMP-1 and P. fluorescens Psa1 (ACCESSION: KT313641) for VIM-1 qualities were utilized as the control strains. DNA sequencing was performed on the cleansed PCR items by the Bioneer Company (Korea).

Statistical Analysis:-
To break down the outcomes, MINITAB16 programming was utilized. P esteem and certainty interims (CI) were < 0.05 and 95%, individually.
**Result:**

**Source Distribution of Isolates**
In this examination, a sum of 600 examples were gathered arbitrarily with sterile swab sticks from the doctor's facility. The disengages were additionally distinguished and affirmed by development and color generation on cetrimide agar. Both color creating separates indicating darker, dull green and yellowish green pigmentation and the non-shade delivering disengages were additionally exposed to Gram recoloring, catalase test, and polymerase tie response examinations to additionally affirm the confines as *P. fluorescens*. 439 examples out of the 600 examples gathered were observed to be sure for Pseudomonas.

![Figure 4.2](image_url)

**Figure 4.2:** Varying pigment production by *P. fluorescens* isolated from the hospital

In the recent study, the most fruitful antibiotic against the studied isolates was colistin. The results of disc diffusion test with different antibiotics for *P. fluorescens* isolates.

| Antibiotics                | Resistance in (%) |
|----------------------------|-------------------|
| Gentamicin                 | (95)              |
| Amikacin                   | (95)              |
| Imipenem                   | (96)              |
| Carbenicillin              | -                 |
| Cefepime                   | (96)              |
| Meropenem                  | (96)              |
| Ciprofloxacin              | (97)              |
| Piperacillin/tazobactam    | (95)              |
| Ceftazidime                | (85)              |
| Aztreonam                  | (97)              |
| Piperacillin               | (95)              |
| Tobramycin                 | -                 |
| Colistin                   | (1)               |
| Ticarcillin                | (99)              |
| Doripenem                  | (95)              |

In the study, among the 439 *P. fluorescens* clinical isolates, 178 (64.02%) were determined as MBL producers by the CDDT test.
Molecular Identifications of the Isolates

A sum of 15 disengages were chosen haphazardly from the segregates (5 pigmented confines and 10 non-pigmented separates) for atomic recognizable proof utilizing the Polymerase Chain Reaction (PCR) to enhance the quality of the microbes. Figure 4.3 demonstrates the enhanced DNA groups at 600 bp, visually seen by ethidium bromide recoloring on 100bp DNA stepping stool utilized as the DNA sub-atomic weight marker. PCR results of 16S rDNA which was particular for Pseudomonas species were distinguished in all examples in 1.5% agarose gel electrophoresis for identification of qualities in Pseudomonas from the 15 haphazardly chose disengages, 13 were discovered positive of Pseudomonas (4 positive pigmented out of 5 and 9 positive non-pigmented out of 10), path 3 and 10 did not demonstrate any intensification which demonstrates that they are negative for Pseudomonas.

Identification of MBL Genes

PCR strategy demonstrated the presence of blaIMP-1 and blaVIM-1 qualities in 30 (16.8%) and 52 (29.2%) separated strains of P. fluorescens, separately; while the other quality was not distinguished. The nucleotide arrangement information accessible to the GenBank succession database and relegated promotion no. KM359726, KM359725, KT313640, KP780165, KP765726, KP765725, JX648311, KR703251, for blaIMP and KT313641 for blaVIM in P. fluorescens strains.

Discussion:

The disclosure of antibiotic resistant bacteria is forbidding the potency of many antimicrobial agents. It has increased the hospital stay of the patients, thus leading to an increased economic burden on them. This examination was gone for assessing the expanding commonness of multidrug safe P. fluorescence including metallo-beta-lactamase creating P. fluorescens separated from the career college Bhopal, Madhya Pradesh, India.

P. fluorescens is an omnipresent, multi-tranquilize safe creature fit for causing genuine infections which are hard to treat because of the nearness of the harmfulness and opposition quality, low penetrability of its external layer [12], constitutive articulation of different efflux pumps with wide substrate specificity [13] and generation of compound fit for causing safe e.g metallo-beta-lactamase [9]. Shade creation and non-color generation by the segregates on Cetrimide agar was one of the elements considered for the recognizable proof of the isolates. Color generation from Pseudomonas fluorescens ranges from blue-green pigmentation called pyocyanin, yellow-green fluorescent shade called pyoverdine, darker color called the pyomelanin, and red color called pyorubin, disconnects from this
Metallo-beta-lactamases (MBLs) are catalysts that makes microscopic organisms impervious to a wide scope of beta-lactam anti-toxins one of which incorporates the carbapenem family [15]. A blend of imipenem-EDTA and meropenem-EDTA was utilized to test for the creation of MBLs in this investigation following the techniques. The shade creating P. fluorescence disengaged from the hospital show the most noteworthy creation of MBLs (83.12%) while the non-shade makers got from the hospital had the minimum generation (72.31%) utilizing the meropenem-EDTA blend. Utilizing the imipenem-EDTA mix, Non-color creating P. fluorescence disengaged from the hospital show the most astounding generation of MBLs (60.77%) while the color delivering P. fluorescence disengaged from the hospital show the slightest generation (32.65%). This outcome demonstrates that the creation of MBLs is higher in non-color delivering P. fluorescens and this could be a motivation behind why they are impervious to anti-infection agents.

The time of medication advancement and its utilization in human and creature wellbeing and farming was started by the revelation of anti-toxins over 70 years back. These disclosures were practical against microorganisms thus were viewed as effective against pathogenic organisms however this achievement was brief as they were tempered with in all cases by the rise of safe organisms [16]. Multidrug-safe (MDR) P. fluorescence are life forms resistant to one antimicrobial specialist in three or more antipseudomonal antimicrobial classes (carbapenems, fluoroquinolones, penicillins/cephalosporins and aminoglycosides) [17]. MDR P. fluorescence are exceptionally dangerous on account of its inalienable protection from many medication classes and ability to secure protection from all effective antimicrobial operators [18].

A total of 10 anti-infection agents {Ceftazidime (CAZ), Cefroxime (CRX), Gentamicin (GEN), Ciprofloxacin (CPR), Ofloxacin (OFL), Augmentin (AUG), Nitrofurantouin (NIT), Ampicillin (AMP) Imipenem (IMP) and Meropenem (MER)} were employed to procure the antibiogram of the separates. All the non-color creating detaches showed the largest amount of obstruction, trailed by the shade delivering secludes from the hospital while the color creating disengages acquired from the hospital showed minimal level of opposition. Secludes demonstrated most noteworthy protection from Ampicillin, Ceftazidime, Cefroxime and Augmentin separately while the segregates were most helpless to Gentamicin (Aminoglycoside), trailed by ciprofloxacin and ofloxacin (Floroquinolones). From this investigation, these antimicrobials are suggested as medications of decision for the treatment of infections caused by P. fluorescens. Carbapenem gathering of anti-toxins, for example, Imipenem and Meropenem are right now considered if all else fails for the treatment of infections caused by multidrug safe P. fluorescens [9]. This investigation appear, that P. fluorescens is quick getting to be safe and these medications and steadily ending up less effective. This might be a result of the capacity of the bacterium to deliver metallo-beta-lactamase which is a catalyst equipped for hydrolyzing β-lactam anti-microbials. From this examination, the effect of meropenem was seen to be higher than imipenem on P. fluorescens in view of the adjustment on the structure of meropenem and its solid restricting liking to penicillin restricting protein which is significantly found in P. fluorescens.

At long last, however this investigation bolsters the confirmation that gentamicin is a medication of decision for treatment of infection caused by multidrug safe P. fluorescens, carbapenems gathering of anti-microbials, for example, imipenem and meropenem which are considered if all else fails for treatment of infections caused by multidrug safe P. fluorescens are quick lessening in its effectiveness as the outcomes got in this examination are not viewed as great.

**Conclusion:-**

Non-shade creating P. fluorescens should not be ignored as they could be more impervious to antimicrobials than shade delivering P. fluorescens. Multidrug safe P. fluorescens can be zoonotic and its protection from antimicrobials is on the expansion. While carbapenem gathering of anti-microbials are as of now the final resort for infections caused by multidrug safe P. fluorescens, its opposition is on the expansion. It is anyway vital that options, for example, the aminoglycosides and the fluoroquinolones could be looked for as it might lessen the reliance on carbapenem gathering of anti-toxins and help control MDR.
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