Research

Toxicity of polymyxins: a systematic review of the evidence from old and recent studies
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Abstract

Background The increasing problem of multidrug-resistant Gram-negative bacteria causing severe infections and the shortage of new antibiotics to combat them has led to the re-evaluation of polymyxins. These antibiotics were discovered from different species of Bacillus polymyxa in 1947; only two of them, polymyxin B and E (colistin), have been used in clinical practice. Their effectiveness in the treatment of infections due to susceptible Gram-negative bacteria, including Pseudomonas aeruginosa and Acinetobacter baumannii, has not been generally questioned. However, their use was abandoned, except in patients with cystic fibrosis, because of concerns related to toxicity.

Methods We reviewed old and recent evidence regarding polymyxin-induced toxicity by searching Pubmed (from 1950 until May 2005).

Results It was reported in the old literature that the use of polymyxins was associated with considerable toxicity, mainly nephrotoxicity and neurotoxicity, including neuromuscular blockade. However, recent studies showed that the incidence of nephrotoxicity is less common and severe compared to the old studies. In addition, neurotoxic effects of polymyxins are usually mild and resolve after prompt discontinuation of the antibiotics. Furthermore, cases of neuromuscular blockade and apnea have not been reported in the recent literature.

Conclusion New evidence shows that polymyxins have less toxicity than previously reported. The avoidance of concurrent administration of nephrotoxic and/or neurotoxic drugs, careful dosing, as well as more meticulous management of fluid and electrolyte abnormalities and use of critical care services may be some of the reasons for the discrepancy between data reported in the old and recent literature.

Introduction

Polymyxins were discovered in 1947 from different species of Bacillus polymyxa [1,2]. Although the effectiveness of polymyxins against most Gram-negative bacteria, including Pseudomonas aeruginosa and Acinetobacter baumannii, has not been questioned, early administration of polymyxins was associated with reports of adverse renal and neurological effects in a considerably large number of patients [3,4]. Thus, compounds of this class of antibiotics were gradually withdrawn from clinical practice as newer antibiotics with the same or broader antibacterial spectra and reportedly lower toxicity were introduced, except for patients with cystic fibrosis who suffer from recurrent pulmonary infections due to multidrug-resistant bacteria [5-7]. However, the emergence of Gram-negative bacteria that are resistant to almost all classes of available antibiotics except polymyxins, especially Pseudomonas aeruginosa and Acinetobacter baumannii strains, and the shortage of new antibiotics with activity against them has led to the re-use of polymyxins [8-12]. The objective of this critical review of the old and recent literature is to elucidate the incidence, mechanisms, prevention, and treatment of adverse events of polymyxins, focusing on patients without cystic fibrosis.

This class of antibiotics consists of five chemically different compounds, polymyxin A, B, C, D, and E (colistin). Only polymyxins B and E have been used in clinical practice. Colistin consists of a cyclic heptapeptide and a tripeptide side-chain acylated at the amino terminus by a fatty acid. The amino acid components in the molecule of colistin are D-leucine, L-threonine, and L-α-γ-diaminobutyric acid. Polymyxin B has the same structure as colistin but contains D-phenylalanine instead of D-leucine [13].

Commercially, colistin appears as colistin sulfate, which is used orally for bowel decontamination and topically as a pow-
der for skin infections, and as colistimethate sodium, which is used parenterally and by inhalation. Colistimethate sodium has been found to be less toxic and to have fewer undesirable side effects than colistin, but is also less potent. Polymyxin B is available for clinical use as polymyxin B sulfate and is used parenterally, topically (ophthalmic and otic instillation), intrathecally, by inhalation, and as an irrigation solution [14,15].

Several attempts to generate less toxic derivatives were made [16]. Most of these derivatives lacked the fatty acid and/or the diaminobutyric acid components of their original molecules. Experimental studies demonstrated that these compounds were much less toxic compared to the parent ones, but at the same time they had considerably reduced antibacterial effect [17,18].

Methods
Data for this review were obtained through literature searches of publications included in PubMed from 1950 until May 2005, references cited in relevant articles, and the world-wide web. The main search terms used in searches of literature databases were 'colistin', 'polymyxin E', 'polymyxin B', 'adverse effects', 'nephrotoxicity', 'colomycin', 'colimycin', 'neurotoxicity' and 'toxicity'. Only English language papers were reviewed.

Results and discussion
In Tables 1 and 2 we summarize the available publications reporting data regarding the incidence of toxicity, including nephrotoxicity, neurotoxicity, and other adverse effects of polymyxins. Specifically, Tables 1 and 2 refer to old (from 1962 to 1977) and recent (from 1995 to 2005) articles, respectively, reporting adverse effects of polymyxins in patients without cystic fibrosis.

Nephrotoxicity
Incidence
Although most of the studies or case reports published until 1983 did not include the definitions of nephrotoxicity, early reported experience with the use of polymyxins, mainly of colistin, revealed a high incidence of nephrotoxicity. The majority of the studies in the older literature referred to intramuscular administration of colistimethate sodium [4,19-25]. Notably, the incidence of nephrotoxicity was 36% in a study of patients with pre-existing acute or chronic renal disease and 20.2% in another large study of 288 patients [4,25]. Additionally, in three studies [26-28], intravenous colistimethate sodium was given for the treatment of patients with Gram-negative bacterial infections, including urinary tract infections, pneumonia, and septicaemia. These studies included 48, 23, and 8 patients, respectively; 10.5% of patients had prolonged increase of blood urea nitrogen levels (average increase of 50 mg/dl) [26], 26.1% of patients experienced renal impairment during therapy [27], and 50% had a fall in creatinine clearance (with a range of 16.5 to 38 ml/min) and an increase in serum creatinine levels (with a range of 0.2 to 2 mg/dl) [28]. Another interesting finding was the relatively high number of case reports that were published in the old literature reporting patients who experienced acute renal failure during treatment with colistimethate sodium. A point that deserves to be stressed, however, is that in most of these cases the total daily dose of colistimethate sodium was considerably higher compared to the currently recommended dose [3,29-34].

During the past seven years, colistimethate sodium has been re-introduced to clinical practice for the treatment of multidrug-resistant bacterial infections, mainly in the intensive care unit setting [9,10,12]. Data from recent studies do not corroborate the previously reported high incidence of polymyxin induced nephrotoxicity [11,35]. Although, the definition of nephrotoxicity was not standardized between the studies, two of them, which were conducted exclusively in intensive care units and used colistimethate sodium, reported that the observed nephrotoxicity was 14% [11] and 18.6% [12]. Notably, in one study that compared two therapeutic approaches – intravenous colistimethate sodium versus intravenous imipenem/cilastatin for the management of patients with ventilator-associated pneumonia due to *Acinetobacter baumannii*, nephrotoxicity occurred in 24% and 42% of patients, respectively [9]. Of note, polymyxin B was reported in the old literature to be associated with a relatively increased incidence of toxicity compared to colistimethate sodium. However, these data were not verified in two recent studies that showed that the incidence of nephrotoxicity was 14% [36] and 10% [37] among patients receiving polymyxin B therapy. Our experience is similar to that of the investigators of the previous studies [35,38].

Mechanisms
It has been suggested that the toxicity of polymyxins may be partly due to their D-amino acid content and fatty acid component. The proposed mechanism by which polymyxin B induces nephrotoxic events is by increasing membrane permeability, resulting in an increased influx of cations, anions, and water, leading to cell swelling and lysis [39,40]. An experimental study showed that colistin increased the transepithelial conductance of the urinary bladder epithelium [41]. The magnitude of the conductance’s increase was dependent on concentration and length of exposure to polymyxins as well as the divalent cation concentration. The basic molecular mechanisms by which polymyxin B increases the transepithelial conductance in the urinary tract has been proposed to be the same as that of colistin [41]. Renal toxicity associated with the use of polymyxins is considered to be dose-dependent.

Clinical manifestations
Renal insufficiency, manifested by an increase in serum creatinine levels and decrease in creatinine clearance, represents a major adverse effect of the use of polymyxins. Occurrence of haematuria, proteinuria, cylindruria, or oliguria may also be associated with the administration of polymyxins. In addition,
### Table 1

**Old studies (from 1962 to 1977) reporting data on polymyxin-induced toxicity in patients without cystic fibrosis**

| Year [ref] | Setting | Medication used | Number of patients | Demographics | Dosage of colistin/duration | Nephrotoxicity | Neurotoxicity | Other toxicities |
|------------|---------|----------------|-------------------|--------------|-----------------------------|----------------|---------------|----------------|
| 1 1962 [26] | Medical wards | Colistimethate sodium (IV) | 48 | Adults: 150 mg q12 h; Children: 5 to 10 mg/kg/day. Duration: at least 10 days | 12 pts had transient mild elevation of BUN (average increase 14 mg/dl) and returned to normal. 5 pts had prolonged elevation of BUN (average increase 50 mg/dl) and returned to normal | 13/48 pts paresthesias; 3/48 pts ataxia | 3/48 pts pruritus. No drug fever, hepatic or bone marrow toxicity |
| 2 1963 [19] | Medical wards | Colistimethate sodium (IM) | 1 | 64 year old male | 6.5 mg/kg/day (150 mg q8 h) for 12 days (he received concurrently kanamycin IM for 2 days and after colistin therapy chloramphenicol) | BUN increased from normal baseline values to 44 mg/dl (drug was stopped). The BUN continued to rise and then began to return to normal. Postmortem examination of the kidney revealed findings compatible with drug induced nephrotoxicity | Possible hepatotoxicity |
| 3 1963 [66] | Medical and surgical wards | Colistimethate sodium (IM and topically) | 62 | Topically: 1% or 2% solution q4h or q12h. Duration (range): 2 to 7 d Intramuscularly (range): 150 to 300 mg/day. Duration (range): 1.5 to 19 d | Topically: no side effects | Topically: no side effects Intramuscularly: 16/55 pts reported one or more of the following: lethargy, dizziness, nausea, confusion, slurred speech, numbness, paresthesias, pruritus, pain at the injection |
| 4 1963 [20] | Medical wards | Colistimethate sodium (IM) | 11 | Dosage: 1.5 MIU q12h for a week and continued for a further week if the pt was improving. 2 pts received 2 MIU q8h for 5 days and then 3 MIU q8h | No renal toxicity | 2 pts trigeminal paresthesia | 1 pt developed follicular rash of the face |
| 5 1964 [28] | Medical wards | Colistimethate sodium (IV) | 8 | Age (range): 25 to 69 years | Dosage: 2 to 2.5 mg/kg q12 h. Duration (range): 8 to 14 days. | 4/8 pts fall in creatinine clearance (range: 16.5 to 38 ml/ min) and increase in serum creatinine (range: 0.2 to 2 mg/dl) | No neurotoxicity | No pruritus |
| 6 1964 [21] | Children’s hospital | Colistimethate sodium (IM) | 36 newborns | Age (range): 6 hours to 12 days | Dosage: 2.5 to 5 mg/kg/day in 2 to 4 doses. Total dose (range): 10 to 240 mg (1 newborn (3.3 kg) received 160 mg of colistin (overdosage) in 7 days) | 16 pts had renal epithelial tubular cells on urinalyses; 14 pts had urinary protein excretion | No neurotoxicity |
| 7 1964 [22] | Medical wards | Colistimethate sodium (IM) | 1 | 50 year old male | Dosage: 900 mg/day for 5 days, then 200 mg/day for 4 days | Urinary retention, rise in blood urea nitrogen | Difficulty in breathing, dysphagia, generalized weakness, hallucinations, apnea requiring intubation |
| 8 1965 [50] | Medical wards | Colistimethate sodium (IM) | 1 | 66 year old female with azotemia | Dosage: 150 mg q 12 h for 8 days. Cumulative dose: 2,550 mg | 7th day of colistin: circumoral paresthesias; 8th day: vomiting, difficulty in breathing, moving, speaking, and became apneic; 10th day: grand mal seizures followed by transient right facial and arm weakness |
| 9 1965 [24] | Medical wards | Colistimethate sodium (IM) | 17 (19 courses) | Age (range): 33 to 90 years | Total cumulative dose (range): 0.56 gr to 2.4 gr | 8 pts dizziness – vertigo (1 pt discontinued), 5 pts oral paresthesias | 3 pts pain at site of injection, 3 pts nausea/vomiting, 2 pts pruritus/rash |
Table 1 (Continued)

Old studies (from 1962 to 1977) reporting data on polymyxin-induced toxicity in patients without cystic fibrosis

| Year | Setting | Agent | Dose | Observation | Toxicity |
|------|---------|-------|------|-------------|----------|
| 10  | 1965    | Medical wards | Colistimethate sodium (IM) | 1 | 75 mg q12 h | Episodes of ptosis, muscular weakness of the face and of the extremities |
| 11  | 1965    | Medical wards (renal department) | Colistimethate sodium (IM) | 25 | 12 males, 13 females. Age (range): 14 to 66 years. All with impaired renal function | Dosage (range): 2 MIU to 4.4 MIU/day. Duration (average): 8.5 days | 9/25 pts had an increase in plasma creatinine levels |
| 12  | 1966    | Medical wards | Colistimethate sodium (IM) | 1 | 47 year old female | 100 mg q8h | Perioral paresthesia, numbness in the hands, weakness, ataxia, light-headedness, shortness of breath, apnea |
| 13  | 1966    | Medical wards | Colistimethate sodium (IM) | 21 | All had urinary tract abnormalities or had undergone prostatectomy | Dosage: 120 mg (1.5 MIU) q8h for 7 days | No constant effect on creatinine clearance was observed |
| 14  | 1966    | Medical wards | Colistimethate sodium (IM) | 4 who developed acute renal failure | Age (range): 41 to 75 years. All with pre-existing renal disease | Dosage: 5 to 6.3 mg/kg/day. Duration (range): 3 to 12 days | Acute tubular failure (3 pts acute tubular necrosis, 1 pt recovered) |
| 15  | 1966    | Medical wards | Colistimethate sodium (IM) | 1 | 48 year old female | 75 mg q12h (she also received chloramphenicol 500 mg q6h po) | Diplopia and bilateral eye ptosis, weakness of neck flexion, difficulty in raising her arms |
| 16  | 1966    | Department of anaesthesiology | Colistimethate sodium (IM) | 1 | 49 year old female with nephrolithiasis | 75 mg q12 h (she also received chloramphenicol 500 mg q4h po and sulfisoxazole 1 g q4h po) | Post-operative apnea |
| 17  | 1967    | Medical and surgical wards | Colistimethate sodium (IV) | 23 | Males, moderately to severely ill | Dosage (range): 1.1 to 5 mg/kg/day q2h for 6 to 7 days (in 2 cases the treatment was discontinued after 2 and 3 days) | 6/23 pts renal impairment; 7/23 pts albuminuria | 1 pt circumoral paresthesia | 5/23 pts mild itching |
| 18  | 1968    | Medical wards | Colistimethate sodium (IV) | 7 | Age (range): 28 to 48 years. 4 females, 3 males; all had terminal and irreversible renal failure | 2 to 3 mg/kg (1 dose) | 2 pts mild dizziness and instability |
| 19  | 1968    | Medical wards | Colistin sulfate (PO) | 93 (48 cases E. coli and 45 cases Shigella spp.) | E. coli: 100,000 IU/kg/day in adults and 150,000 IU/kg/day in children for 7 days Shigella: 200,000 IU/kg/day in adults and 300,000 IU/kg/day in children for 8 to 10 days | No toxic symptoms | 1 pt generalized rash, 1 pt vomiting |
| 20  | 1968    | Medical wards (respiratory care unit) | Colistimethate sodium (IM) and Polymyxin B (IM or IV) | 11 | Age (range): 36 to 74 years. 4 females, 7 males; all had acute or chronic renal disease | Dosage of colistimethate sodium (range): 100 to 400 mg/day. Dosage of polymyxin B: 50 mg (1 dose) IM (1 pt) and 100 mg (1 dose) IV (1 pt) | No toxic symptoms | All pts at their admission had apnea that recovered in all cases. Paralysis 2 pts, diplopia 3 pts, difficulty in swallowing 3 pts, paresthesia 2 pts, generalized weakness 3 pts, blurring of vision 1 pt, slurred speech 1 pt, lethargy 1 pt, coma 1 pt |
| 21  | 1969    | Medical wards | Colistimethate sodium (IV) | 1 | 14 year old male with acute leukemia | Dosage: 5 mg/kg/day for 6 days, then increased to 7 mg/kg/day on day 6, 10 mg/kg/day on day 7, and 17 mg/kg/day on day 9. Duration: 14 days | Acute tubular necrosis |
| 22  | 1969    | Medical wards (pediatrics) | Colistimethate sodium (IM) | 1 | 4 year old female with appendicitis | Dosage: 30 mg/kg q6h (total dose received 1,050 mg during 42 h) | Acute renal failure | Neuronal hyperactivity, seizure-like episodes, uncoordination, disorientation, flaccid quadriplegia, respiratory arrest, apnea |
| Year | Department | Route | Age | Duration | Dose | Toxic Symptoms | Neurotoxicity | Renal Failure | Other Adverse Effects |
|------|------------|-------|-----|----------|------|----------------|---------------|--------------|----------------------|
| 1970 | Medical and surgical wards | Colistimethate sodium (IM) | 288 (317 courses) | 205 courses received a total of <1 gr, 69 courses 1 to 2 gr, 43 courses > 2 gr. All courses were administered IM q12 h | Total: 64/317 courses (renal insufficiency 63 pts, acute tubular necrosis 6 pts, hematuria 1 pt) | Total: 23/317 courses (paresthesias 15 pts, respiratory insufficiency and apnea 6 pts, nausea and vomiting 4 pts, dizziness 3 pts, muscular weakness 2 pts, peripheral neuropathy, confusion, psychosis, convulsive seizure 1 pt each) | Total allergic reactions: 7/317 (drug fever 3 pts, eosinophilia 2 pts, macular eruption 1 pt) |
| 1970 | Medical wards and ICU | Colistimethate sodium (aerosol) | 20 | Age (range): 23 to 81 years | Group 1: 50 mg q8h for 7 days. Group 2: 100 mg q8h for 7 days | No toxic symptoms | 1 pt experienced palpitations and a sensation of chest tightness (treatment was discontinued) |
| 1970 | Department of pediatrics | Colistimethate sodium (IM) | 1 | Age: 10 months (male) | 15 mg q6h (2 doses) and then 250 mg (38.5 mg/kg) (3 doses) | Acute renal failure | No neurotoxicity |
| 1970 | ICU, neurosurgical department | Colistimethate sodium (IV, IM, and aerosol) | 14 | Age (range): 31 to 71 years | Mean duration: 9.7 days. Dosage: 26 MIU/day: 10 MIU IM, 10 MIU IV, and 6 MIU aerosol | In all pts a considerable fall in creatinine clearance and rises in blood urea and serum creatinine levels were observed. 5 pts developed acute tubular necrosis (histological confirmed). In 6 pts renal function returned to normal |
| 1970 | Department of renal disease | Colistimethate sodium (route of administration not reported) | 1 | Age: 41 year old | Duration: 7 days. Dose: 6.3 mg/kg/day | Severe oliguric renal failure |
| 1970 | Department of pediatrics | Colistimethate sodium (IM) | 1 | Age: 3 year old | 150 mg q8h (she received 3 injections) | No renal toxicity | No neurotoxicity |
| 1970 | Medical wards (urology department) | Colistimethate sodium (IM) | 1 | Age: 33 year old male with a solitary kidney | 25 mg q8h for 5 days and 250 mg q8h for 1 day | Increase in serum creatinine levels compared to baseline levels (1.1 mg/dl to 3 mg/dl). Returned to approximately normal values after 6 months |
| 1971 | Department of neurology | Colistimethate sodium (IM) | 1 | Age: 70 year old male with myasthenia gravis | 150 mg (one injection) | Muscular weakness, generalized paresthesias, speech disturbances, ptosis, hypotonia, areflexia, ataxia, difficulty in breathing |
| 1971 | Department of respiratory diseases | Polymyxin B (aerosol) | 2 | Case 1: 51 year old female. Case 2: 57 year old male | Case 1: 15 mg Case 2: 10 mg | 2 hours after the injection: muscular weakness; 30 minutes later he developed respiratory arrest |
| 1973 | Medical wards (Hemodialysis Centre) | Colistimethate sodium (IM) | 2 | Case 1: 16 year old female. Case 2: 23 year old female | Case 1: 150 mg q6h 1st day, 150 mg q4h 2nd day (20 mg/kg/day) Case 2: 180, 240, 180, 120 mg in divided doses on 1st, 2nd, 3rd, 4th day, respectively | Both pts developed acute renal failure | Case 1: neuromuscular blockade that resulted in quadriplegia, apnea, cardiac arrest Case 2: circumoral – acral paresthesias |
| 1974 | Medical wards | Colistimethate sodium (IM) | 1 | Age: 66 year old male | 6 MIU/day for 60 days | No renal toxicity | Total ophthalmoplegia, flaccid paralysis of both upper limbs, reduced speech fluency, difficulty in finding words, apathy |
| 1977 | Department of pediatrics | Colistimethate sodium (IM) | 1 | Age: 5 year old male | 200,000 IU/kg/day for 8 days | Acute oliguric renal failure | Muscular weakness, speech disturbances |

*1 mg of colistimethate sodium is approximately equal to 12,500 IU. BUN, blood urea nitrogen; ICU, intensive care unit; IM, intramuscularly; IV, intravenously; MIU, million international units; po, per os; Pt(s), patient(s); ref, reference.
acute tubular necrosis can also develop [14]. Histological findings of colistin-induced renal damage usually involve focal irregular dilatation of tubules, epithelial and polymorphonuclear cell cast formation, and degeneration and regeneration of epithelial cells. In addition, separation of tubules by loose collagenous tissue, suggestive of edema, has also been reported. The basement membrane is usually intact, as well as the glomeruli [19,42].

Risk factors
Nephrotoxicity resulting from the use of colistimethate sodium appears to be less compared with that associated with polymyxin B. It is unclear whether there are independent factors that predispose patients to the development of nephrotoxic events. Children seem to experience less polymyxin-induced toxicity, probably in part because prescription of polymyxins, and generally all medications, is based on individual body weight in this patient population [4]. Concomitant administration of potential nephrotoxic agents, such as diuretics and some antimicrobial agents, increases the likelihood of development of renal adverse effects [4,43].

Treatment
When primary signs of renal dysfunction are present, early discontinuation of polymyxins is necessary. Quick diuresis by intravenously administered mannitol has also been proposed to enhance renal clearance of the drug and thus to reduce serum drug levels [32]. Meticulous supportive care, including close monitoring of fluid intake and output, frequent determinations of electrolytes, and appropriate management to maintain balance of fluids and electrolytes, is required when renal adverse effects of polymyxin use are detected. The influence of hemodialysis and peritoneal dialysis in decreasing serum levels of polymyxins has not been clarified. Older reports suggested that the amount of drug that is removed from blood by these two methods is relatively small [44,45]. Patients that underwent peritoneal dialysis lost approximately 1 mg of colistimethate sodium per hour [45]. Thus, in cases of polymyxin-induced renal failure, both therapeutic approaches have been used, not to decrease serum drug levels but in order to manage renal complications. Exchange transfusions have been proposed as an effective method for the removal of polymyxins [3].

Neurotoxicity
Incidence
The incidence of neurotoxicity related to the use of polymyxins reported in the old literature was considerably less compared to nephrotoxicity. Specifically, the most frequently experienced neurological adverse effects were paresthesias that occurred in approximately 27% and 7.3% of patients receiving intravenous and intramuscular colistimethate sodium, respectively [4,26]. Furthermore, at least eight cases were published between 1964 and 1973 correlating the intramuscular administration of polymyxins with the development of episodes of respiratory apnea [22,33,46-51]. However, recently performed studies in patients without cystic fibrosis are not in accordance with the previously reported data regarding the incidence of polymyxin-induced neurotoxicity [11,12,38]. No episodes of neuromuscular blockade or apnea induced by polymyxins have been reported in the literature over the past 15 years or more.

Mechanisms
The interaction of polymyxins with neurons, which have a high lipid content, has been associated with the occurrence of several neurotoxic events. In addition, the probability of development of neurotoxicity has been directly associated with the concentration of the active form of polymyxins in the blood [14]. Neuromuscular blockade induced by polymyxins has been attributed to a presynaptic action of polymyxins that interferes with the receptor site and blocks the release of acetylcholine to the synaptic gap [33,52]. Other investigators have suggested a biphasic mechanism to explain this neurotoxic event; a short phase of competitive blockade between acetylcholine and polymyxins is followed by a prolonged phase of depolarization associated with calcium depletion [51,53,54]. Neurotoxicity resulting from the use of polymyxins is also considered to be dose-dependent.

Clinical manifestations
The reported neurological toxicity is associated with dizziness, generalized or not muscle weakness, facial and peripheral paresthesia, partial deafness, visual disturbances, vertigo, confusion, hallucinations, seizures, ataxia, and neuromuscular blockade. The last of these usually produces a myasthenia-like clinical syndrome, as well as respiratory failure or apnea due to respiratory muscle paralysis [33]. Paresthesias appear to be usually benign, and their mechanism seems to be unrelated to the interference with nerve transmission. An old study that assessed the safety of intramuscularly administered colistimethate sodium during 317 courses revealed that neurological adverse effects were manifested during the first four days of therapy in 83% of the patients who experienced neurotoxic events [4].

Risk factors
Risk factors that may potentially trigger the development of neurotoxicity include hypoxia and the co-administration of polymyxins with muscle-relaxants, narcotics, sedatives, anesthetic drugs, or corticosteroids [22,55]. A patient's gender may influence the likelihood of development of adverse effects. Specifically, neurotoxicity seems to be more common in women, although nephrotoxicity seems to be gender-independent [4]. Patients with impaired renal function or myasthenia gravis are at higher risk of developing neuromuscular blockade and respiratory paralysis [47].
Treatment
Mild neurological manifestations of polymyxins usually subside after prompt cessation of the drugs. In the presence of neuromuscular blockade, immediate discontinuation of polymyxins and other neurotoxic agents is also the first-line approach. Further management consists of mechanical respiratory support if apnea has been developed. The intravenous administration of calcium and cholinesterase inhibitors, such as neostigmine and edrophonium, has led to conflicting results [33,48]. Hemodialysis is indicated only in patients with co-existing acute renal failure.

Other adverse events
Incidence
In studies published in the old literature, the reported incidence of allergic reactions related to colistimethate sodium use was approximately 2% [4]. Mild itching that did not require discontinuation of the drug was reported by approximately 22% of the patients receiving colistimethate sodium intravenously [27]. In addition, a few patients with episodes of rash were also reported [20,56]. In the recent literature, a few patients with episodes of contact dermatitis (eczema and erythematous eruption) have been reported in connection with topical use of colistin sulfate and ophthalmic administration of colistimethate sodium [57,58].

Mechanisms
Several milder adverse reactions, including pruritus, dermatitis, and drug fever, probably represent the result of the irritative effects of the active forms of polymyxins [14] and their histamine-releasing action, especially polymyxin B.

Clinical manifestations
Pruritus, contact dermatitis, macular rash or urticaria, ototoxicity, drug fever, and gastrointestinal disturbances may develop, although rarely, during treatment with polymyxins [26,57,59]. After intramuscular administration, pain may occur at the injection site [24]. Moreover, the development of pseudomembranous colitis represents a rare side effect of polymyxins. Intraventricular or intrathecal administration of polymyxins, especially in high doses, may lead to convulsions and signs of meningismus. During repeated ophthalmic application of polymyxin, low-grade conjunctivitis may develop [14].

An old case report suggested that the administration of colistimethate sodium intramuscularly in a patient with Gram-negative rod bacteremia was possibly associated with hepatotoxicity because an observed rise in serum glutamic oxaloacetic transaminase levels returned to normal after the drug was discontinued; in addition, post-mortem histological examination of the liver revealed non-specific changes (focal vacuolization of hepatic cells in the centrilobular fields with areas of focal necrosis), which were interpreted as drug-induced toxicity [19]. However, no other cases of liver toxicity have been reported in experimental or clinical studies on the use of polymyxins [38,60].

Risk factors
Patients with known allergy to bacitracin are also at higher risk of developing hypersensitivity reactions with the use of polymyxins, as cross-reaction between bacitracin and polymyxins exists [58].

Treatment
In most instances, withdrawal of polymyxins in combination with appropriate supportive treatment is adequate for the treatment of such adverse effects.

Adverse events related to aerosolised colistin
Treatment with aerosolized colistin may be complicated by sore throat, cough, bronchoconstriction, and chest tightness. The nature of bronchoconstriction that develops during nebulization of polymyxins has been proposed to be associated with several mechanisms. Among them are direct chemical stimulation, the liberation of histamine, allergy in the airway, irritation from chemicals or from the foam that is produced during nebulization, and hyperosmolarity in the airway [61]. Nebulized polymyxins can cause bronchoconstriction even in patients with no history of asthma or atopy, although if these conditions exist the risk is greater [61]. Bronchoconstriction usually requires discontinuation of the medication, the administration of bronchodilators and supplemental oxygen.

Prevention of adverse events
Early and correct adjustment of the dose of polymyxins in the presence of impaired renal function, frequent urinalyses and serum urea or creatinine measurements, close daily monitoring of urinary output and of neurological status, and the avoidance of concurrent administration of other agents with known nephrotoxicity or neurotoxicity may help prevent the development of adverse effects. Bronchoconstriction usually responds to treatment with bronchodilators; thus, pre-treatment of patients receiving inhaled colistimethate sodium with these medications could prevent the occurrence of this adverse event [61].

Recommendations regarding the dosage of polymyxins differ between various manufacturers. Colistin manufactured in the United States contains colistimethate sodium equivalent to 150 mg colistin base activity in each vial. The recommended dosage is 2.5 to 5 mg/kg per day, divided into 2 to 4 equal doses in adult patients with normal kidney function [62]. Manufacturers in the United Kingdom recommend a dosage of 4 to 6 mg/kg (50,000 to 75,000 IU/kg) intravenous colistimethate sodium per day, in 3 divided doses for adults and children with body-weight ≤ 60 kg, and 80 to 160 mg (1 to 2 million IU) every 8 hours for body-weight >60 kg [63]. The recommended dosage for intravenous polymyxin B sulphate is 1.5 mg to 2.5 mg/kg/day (15,000 IU to 25,000 IU/kg/
### Table 2

**Recent studies (from 1995 to 2005) reporting data on polymyxin-induced toxicity in patients without cystic fibrosis**

| Year | Setting | Medication used | Number of patients | Demographics | Dosage of colistin/duration | Definition of nephrotoxicity | Nephrotoxicity | Neurotoxicity | Other toxicities |
|------|---------|----------------|--------------------|--------------|-----------------------------|-------------------------------|----------------|--------------|----------------|
| 1 1995 | Department of dermatology | Colistin sulfate (ointment/topically) | 1 | 45 year old | 50,000 IU for 10 days | Edematous eczema |
| 2 1998 | Department of dermatology | Colistimethate sodium (ophthalmic solution) | 1 | 4 year old male with bilateral ocular prosthesis | Case 1: 5 mg (62,500 IU) q12h for 19 days Case 2: 5 mg (62,500 IU) q12h for 5 days then 10 mg (125,000 IU) q12h for 12 days | No adverse reactions |
| 3 1999 | Neurosurgical wards | Colistimethate sodium (intraventricular) | 2 | 16 year old male and 34 year old female | | No adverse reactions |
| 4 1999 | ICU (52%), transplant unit (13%), surgical and medical wards (35%) | Colistimethate sodium (IV) | 59 (60 cases) | Mean age: 42.1 years. Mean (± SD) APACHE II: 13.1 (± 7.0) | Mean duration: 12.6 d (2 to 34 d). Mean daily dose: 152.8 mg (approximately 2 MIU) (60–300 mg) | 22 pts (37%); 11/41 with normal baseline renal function had worsening during treatment (mean increase in serum creatinine 0.9 ± 0.6 mg/dl) and 11/19 with abnormal baseline renal function had worsening during treatment (mean increase in serum creatinine 1.5 ± 1.4 mg/dl). Nephrotoxicity did not cause discontinuation |
| 5 2000 | Medical wards | Colistimethate sodium (aerosol) | 3 | 67 year old male, 45 year old male, 59 year old male | 150 mg (2 MIU) q12h for 13 days, 100 mg (approximately 1.5 MIU) q12h for 14 days, 150 mg (2 MIU) q12h for 11 days | No nephrotoxicity |
| 6 2002 | Neurosurgical wards | Colistimethate sodium (IV) | 1 | 14 year old male | 1 MIU q8h for 30 days | No adverse reactions |
| 7 2003 | Abdominal organ transplantation ICU | Colistimethate sodium (IV) | 23 (20 had received organ transplantation, 3 abdominal surgery) | Mean age: 52 years | Mean duration: 17 days (7 to 38 days) | Renal failure was defined by a requirement either for intermittent hemodialysis or for continuous venous hemofiltration | 1/2 pts developed renal failure requiring artificial kidney support (the other 21 pts were already receiving artificial kidney support) |
| 8 2003 | ICU | Colistimethate sodium (IV) | 24 with sepsis, 26 courses of colistin | Mean age: 44.3 years. Mean APACHE II: 20.6 | Mean duration: 13.5 days (4 to 24 days). Dosage: 3 MIU q8h | Renal failure was defined as an increase in serum creatinine >1 mg/dl during treatment | 3 pts (1.4%). Only 1 pt required continuous venovenous hemofiltration |
| 9 2003 | Tertiary care hospital | Polymyxin B (parenterally) | 60 receiving polymyxin B | Mean age: 61 years | Mean duration: 13.5 days (1 to 56 days). Mean daily dose: 1.1 MIU | Renal failure was defined as doubling of serum creatinine value of ≥ 2.0 mg/dl | 7/50 pts (14%) | No clinically apparent neuromuscular transmission blockade |
### Table 2 (Continued)

**Recent studies (from 1995 to 2005) reporting data on polymyxin-induced toxicity in patients without cystic fibrosis**

| Year | Study Details | Cohort | Mean Age | APACHE II | Duration | Dosage | Nephrotoxicity | Comments |
|------|---------------|--------|----------|-----------|----------|--------|---------------|----------|
| 10 2003 [74] | ICU Colistimethate sodium (IV) | 35 (21 received colistin (CO group) and 14 IM group) | Mean age: CO group 56.9 years, IM group 64.9 years, Mean APACHE II: CO group 19.6, IM group 20.5 | CO group: mean duration 14.7 days (10 to 21 days), Dosage: 2.5 to 5 mg/kg/day | In patients with normal renal function (creatinine <1.2), renal failure was defined as creatinine value >2 mg/dl, as a reduction of creatinine clearance of 50% relative to antibiotic initiation, or need for renal replacement therapy. In patients with normal renal function, renal failure was defined as increase of 50% of the baseline creatinine level, as a reduction of creatinine clearance of 50% relative to antibiotic initiation, or need for renal replacement therapy | 5/21 pts (24.4%; CO group), 8/14 pts (42%; IM group) |
| 11 2004 [75] | ICU Colistimethate sodium (IV) | 1 | 41 year old male | 2 MIU/day continuous infusion | No adverse reactions |
| 12 2004 [76] | ICU Colistimethate sodium (IV) | 1 | 48 year old male | 9 MIU/day (2.5 mg/kg/day) for 13 days | No adverse reactions |
| 13 2004 [37] | Tertiary care hospital, ICU (92%) Polymyxin B (IV) | 25 (29 courses: 21 IV, 6 aerosol, 2 combination) | Mean age: 55 years, Mean APACHE II: 21 | Loading dose on day 1 with 2.5 to 3 mg/kg IV polymyxin B. Aerosolized: approximately 2.5 mg/kg/day (approximately 1.75 MIU). Mean duration: 19 d (2 to 57 d) | Nephrotoxicity was defined as the doubling of serum creatinine during therapy | 3/29 courses (10%) |
| 14 2005 [12] | ICU Colistimethate sodium (IV) | 43 | Mean age: 56.5 years, APACHE II: 23.8 ± 3.7 | 3 MIU q8h | Acute renal failure was defined as a rise of ≥ 2 mg/dl in serum creatinine level in patients with previously normal renal function. In patients with a history of renal insufficiency, acute on chronic renal failure was defined as at least doubling of the baseline serum creatinine level (defined as the creatinine level at the initiation of colistin treatment) | 8/43 pts (18.6%); 3/35 pts with normal renal function (8.6%) and 5/8 pts with chronic renal failure (62.5%) |
| 15 2005 [77] | ICU (84%), medical (11%), surgical (5%) Colistimethate sodium (aerosol, IV, IM, intrathecal) | 80 (85 courses: 71 aerosol, 12 IV or IM, 2 intrathecal) | Mean age: 57 ± 15 years | Mean duration of aerosol: 12 ± 8 d. Mean duration of IV or IM: 11 ± 6 d. Mean duration of intrathecal: 8 d and 10 d | Nephrotoxicity was defined as a serum creatinine increase of 50% or 1 mg/dl with respect to the baseline level during treatment | 12 courses of IV or IM were recorded. Mean ± SD baseline serum creatinine: 1.25 ± 0.79 mg/dl. Mean ± SD final serum creatinine: 1.20 ± 0.64 mg/dl. Mean ± SD baseline BUN: 8.95 ± 8.96 µmol/l. Mean ± SD final BUN: 8.39 ± 8.06 µmol/l |
| 16 2005 [38] | Mainly ICU pts Colistimethate sodium (IV) | 17 (19 courses) | Median age: 51 years, Median APACHE II: 14 | Mean ± SD duration: 43.4 ± 14.6 days, Mean ± SD daily dose: 4.4 MIU (352 mg) ± 2.1 MIU (168 mg) | Renal failure was defined as an increase more than 50% of the baseline creatinine level to a value higher than 1.3 mg/dl or a decline in renal function requiring renal replacement therapy | Median baseline serum creatinine: 0.6 mg/dl. Slight increase of the median of values of creatinine at the end by 0.1 mg/dl. Median baseline BUN: 42 mg/dl. Median final BUN: 41 mg/dl. 1 pt had an increase of more than 50% of the baseline creatinine level to a value higher than 1.3 mg/dl at the end of colistin treatment |

No paresthesias, vertigo, muscle weakness, or apnea were observed.
| Year | Reference | Setting                                                                 | Study Group | Details                                                                                   | Toxicity                                                                 |
|------|-----------|-------------------------------------------------------------------------|-------------|-------------------------------------------------------------------------------------------|-------------------------------------------------------------------------|
| 2005 | [35]      | ICU (80%), medical and surgical wards (20%)                             | Colistimethate sodium (IV) | 50 (54 episodes) Mean age: 59.2 years. Mean APACHE II: 16.1 Mean duration: 21.5 days. Mean daily dose: 4.5 MIU Renal failure was defined as an increase more than 50% of the baseline creatinine level to a value higher than 1.3 mg/dl or as a decline in renal function requiring renal replacement therapy | 4/50 pts (8%) 1 pt polyneuropathy (not confirmed) resolved without discontinuation |
| 2005 | [78]      | ICU                                                                     | Colistimethate sodium (aerosol) | 8 Mean age: 59.6 years. Mean APACHE II: 16.8 Dosage (range): 1.5 to 6 MIU/day. Duration (mean): 10.5 days Worsening of renal function: 1 pt No neurotoxicity |
| 2005 | [79]      | ICU                                                                     | Colistimethate sodium (IV) | 1 57 year old male 250 mg q6h for 4 days Acute renal failure (on the 4th day of colistin therapy) | No adverse reactions |
| 2005 | [80]      | Neurosurgical wards                                                     | Colistimethate sodium (intraventricular) | 1 23 year old female 125,000 IU q12h for 3 weeks No adverse reactions |
| 2005 | [81]      | ICU                                                                     | Colistimethate sodium (IV) | 55 Mean age: 49 ± 16 years. Mean APACHE II: 21 ± 7 Duration (mean): 13 ± 5 days Renal failure was defined as a decrease in the estimated creatinine clearance rate of 50%, compared with the rate at the start of therapy, or as a decline in renal function that necessitated renal replacement therapy No adverse reactions. Mean creatinine levels before treatment: 2.3 ± 0.5 mg/dl. Mean creatinine levels after treatment: 2.5 ± 0.6 mg/dl |
| 2005 | [82]      | ICU                                                                     | Colistin (IV) | 1 35 year old male 6 MIU/day for 12 days, 3 days, and 1 day Acute renal failure occurred at the 2nd and 3rd introduction of colistin. Renal function returned to normal values within 3 and 5 days after colistin withdrawal |
| 2005 | [83]      | ICU                                                                     | Colistimethate sodium (IV) | 14 Mean age: 49 years Mean dose: 6 MIU/day. Mean duration: 12 days 1 pt experienced deterioration of renal function (serum creatinine up to 2.8 mg/dl) |
| 2005 | [84]      | ICU, medical wards                                                      | Colistimethate sodium (aerosol) | 21 Mean age: 60.6 ± 15 years. Mean APACHE II: 23.1 ± 9.1 19 pts received 2 MIU/day, 1 pt 3 MIU/day, and another pt 4 MIU/day. Median duration: 14 days Renal failure was defined as a decrease in the estimated creatinine clearance rate of 50%, compared with the rate at the start of therapy, or a decline in renal function that necessitated renal replacement therapy No episodes of acute renal failure No symptoms of neurotoxicity 1 pt experienced bronchospasm that resolved on discontinuation of colistin therapy |

BUN, blood urea nitrogen; ICU, intensive care unit; IM, intramuscularly; IV, intravenously; MIU, million international units; Pt(s), patient(s); ref, reference; SD, standard deviation.
day), divided into 2 equal doses for adults and children older than 2 years with normal renal function; 1 mg of polymyxin B is equal to 10,000 IU. Infants with normal renal function may receive up to 4 mg/kg/day (40,000 IU/kg/day) in cases of lifethreatening infections [64].

Overdoses

Overdoses with polymyxins, mainly with colistimethate sodium, have been reported several times in the old literature. Although, one case of a three year old child who received intramuscularly 450 mg (approximately 5.5 million IU) of colistimethate sodium reported no adverse effects, the majority of cases with polymyxin overdose resulted in acute renal failure and various manifestations of neurotoxicity, including neuromuscular blockade and apnea [3,31,33,34]. It should be emphasized that cases of polymyxin overdose with fatal consequences are scarce [29]. There is no antidote for polymyxin overdose. Management requires early cessation of the medication and appropriate supportive treatment. In the presence of established acute renal failure, haemodialysis and peritoneal dialysis can only manage renal complications, since they have little influence on the elimination of polymyxins, as discussed above. If apnea occurs, mechanical ventilation support is needed.

Drug interactions

The concurrent use of polymyxins with curariform muscle relaxants and other neurotoxic drugs such as ether, tubocurarine, succinylcholine, gallamine, decamethonium, and sodium citrate must be avoided, since these agents may trigger the development of neuromuscular blockade [55]. Co-administration of sodium cephalothin and polymyxins may enhance the development of neurotoxicity, so this combination of antimicrobial medication should also be avoided [4]. In addition, antimicrobial agents with known neurotoxic effect, such as aminoglycosides, should generally be avoided or given with great caution in patients who receive polymyxins. In such instances, close monitoring of the patients receiving these antibiotics is mandatory. Experimental studies showed that application of polymyxins in combination with glutamic acid to a peripheral nerve could cause transgaglionic degenerative atrophy [65].

Conclusion

The data from the recent literature suggest that the incidence of toxicity resulting from the use of polymyxins is less frequent and severe compared to what has been previously reported. Possible explanations for the observed discrepancy include the fact that the available formulation of colistimethate sodium for intramuscular administration was used intravenously in the old studies until a new formulation was prepared. In addition, the intramuscular formulation also contained dibucaine hydrochloride, which could potentiate the neurotoxic effect of colistimethate sodium. It should be highlighted that the dosages of polymyxins used in most of the studies published in the old literature were considerably higher compared to the recommended dosages administered nowadays. In fact, several reported cases of polymyxin-induced toxicity were associated with overdose. Thus, this may account for the observed difference in the incidence of polymyxin-induced toxicity noted between the old and recently published studies. A major limitation in the interpretation of polymyxin-induced nephrotoxicity and neurotoxicity in the intensive care unit setting, however, is the frequent existence of multiple organ failure, septic shock, and mechanical ventilation support. These conditions may considerably influence the assessment of polymyxin-induced toxicity. Dosage adjustment of polymyxins in the presence of impaired renal function and prompt discontinuation of polymyxins after development of early signs of their toxicity were not always performed in a timely fashion. Furthermore, the already reported experience regarding the toxicity of polymyxins in the old literature has led to more correct use of these antibiotics by physicians nowadays. In addition, the avoidance of co-administration of potential nephrotoxic and/or neurotoxic agents with polymyxins, as well as the development of critical care supplies, may also explain the observed differences. In the coming years further research is needed to assess the safety profile of polymyxins, clarify several aspects of their toxicity, and investigate the benefits of different dosing regimens, including the administration of these antibiotics in fewer daily doses.

Key messages

- Polymyxins are valuable antibiotics for use in patients in the intensive care setting.
- Polymyxins have been recently re-introduced in clinical practice for the treatment of patients with multidrug-resistant Gram-negative bacterial infections.
- Nephrotoxicity and neurotoxicity represent the major adverse effects of polymyxins.
- Data from the recent literature suggest that the use of polymyxins is associated with lower and less severe toxicity compared to that reported in the old literature.
- Caution is needed when polymyxins are administered, particularly in patients with renal dysfunction.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

MEF conceived the study idea. Both authors contributed to the reviewing of the articles and writing of the manuscript. Both authors approved the final manuscript. MEF had full access to all the data in the study and takes responsibility for the integrity of the review of the data.

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