Side effects of aminoglycosides on the kidney, ear and balance in cystic fibrosis

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ABSTRACT
Aminoglycoside antibiotics are a central component of the treatment of pulmonary exacerbations of cystic fibrosis (CF) and slow the decline in lung function which ultimately causes the death of most patients. The prognosis of CF has improved, and thus side effects of treatments have become increasingly important. Observational studies suggest that the morbidity from side effects of aminoglycosides is disturbingly common, and that aggressive treatment may lead to more side effects. This review of the current literature on side effects of aminoglycosides considers the pathophysiological mechanisms, epidemiology and risk factors, investigation of side effects and preventative strategies. Treatments which have shown early promise are identified and areas of future research are discussed.

INTRODUCTION
Cystic fibrosis (CF) is the most common life-limiting autosomal recessive disorder in the Caucasian population with approximately 7000 affected individuals in the UK.1 A mutation in the CF gene leads to a defective or absent CF transmembrane conductance regulator (CFTR) which, in most patients, ultimately leads to bronchiectasis and chronic infection with Pseudomonas aeruginosa. Intravenous aminoglycosides are used for exacerbations owing to their efficacy against P. aeruginosa. Patients often require several courses of intravenous antibiotics per year. The survival of patients with CF has improved, and side effects of treatment have become increasingly important.

We review the current literature on the side effects of aminoglycosides commonly used to treat pulmonary exacerbations of CF. The major side effects of aminoglycosides are kidney injury, hearing impairment and vestibular toxicity.

RENAL TOXICITY IN CYSTIC FIBROSIS
Acute kidney injury
Studies vary in their definition of toxicity, but approximately 5–10% of (non-CF) adult patients receiving an aminoglycoside have a significant increase in serum creatinine.2 Although toxicity is an aminoglycoside class effect, experimental data suggest that gentamicin is more toxic than tobramycin and amikacin.3 A quantitative overview of randomised controlled trials (RCTs) reached broadly the same conclusions.4 The tenacity of infection and altered pharmacokinetics in CF necessitate long courses of high-dose antibiotics.5 Since the introduction of regular 3-monthly intravenous antibiotic therapy for patients with chronic P. aeruginosa in Danish CF clinics in the mid-1970s,6 the management of pulmonary infection in CF has become more aggressive. However, case reports of acute kidney injury (AKI; previously termed acute renal failure) associated with aminoglycoside antibiotics in CF were not published until the late 1990s.

The first report of AKI secondary to intravenous aminoglycosides in CF7 was followed by several other cases and mini-series. Inhaled tobramycin has also been implicated.8 Bertenshaw et al9 conducted a national survey of AKI in patients with CF (AKI was pragmatically defined as an increased serum creatinine for age) and found that the incidence was 4.6–10.5/10 000 CF patients/year. A subsequent case–control study10 established that an aminoglycoside in the previous week increased the risk of AKI >80-fold. Risk factors for renal impairment (including CF-related diabetes, non-steroidal anti-inflammatory drugs and acute dehydration) were strongly associated with AKI (OR 24.0). Exposure to gentamicin in the previous year was associated with AKI while exposure to tobramycin was not.

Chronic kidney disease
Regular courses of aminoglycoside antibiotics may cause subclinical kidney damage leading to chronic kidney disease (CKD). This can manifest at the level of the glomerulus (causing decreased glomerular filtration rate, GFR) and the tubules (causing altered excretion of electrolytes). The stages of CKD are shown in table 1.

Pederson et al11 described a series of 46 patients aged 10–35 years who had received a mean of 20 courses of tobramycin and used 24 h urinary creatinine clearance to assess renal function. The cumulative dose of tobramycin was not related to the creatinine clearance; 39% of patients had reduced GFR (defined as <96 ml/min/1.73 m2). However, 24 h urinary creatinine collections are known to be unreliable, especially in children.

The largest study investigating CKD in CF was reported by Al-Aloul et al.12 Creatinine clearance was measured in 24 h urine collections in 80 adult patients with CF chronically infected with P. aeruginosa. Multiple urinary collections per patient were taken to ensure accuracy. The median number of courses of antibiotics per patient was 40 (range 1–150); 42% of patients had a creatinine clearance <80 ml/min/1.73 m2. Creatinine clearance declined with increasing number of antibiotic courses (r=0.59). Colistin was not associated with decreased kidney function when administered with a β-lactam but, when administered with an aminoglycoside, the two agents together were associated with more kidney damage than an aminoglycoside alone. Creatinine clearance can be predicted using equations such as the modification...
of diet in renal disease (MDRD) and Cockroft–Gault formulae. In this study, the Cockroft–Gault formula to predict creatinine clearance performed poorly and missed cases of CKD.

Investigators have used biomarkers to obtain indirect evidence of subclinical kidney injury in many research settings. N-acetyl-β-D-glucosaminidase (NAG), one of the best characterised biomarkers, is a proximal tubular enzyme excreted in the urine in response to kidney damage. Paediatric patients with CF administered intravenous tobramycin have increased urinary NAG in response to kidney damage. Paediatric patients with CF may have a cumulative tobramycin dose. In comparison, Al-Aloul et al found that tobramycin was associated with a larger cumulative colistin dose rather than gentamicin (perhaps indicating chronic toxicity) prior to antibiotics was administered in a subsequent meta-analysis.28 In the paediatric group receiving colistin, NAG increased.28

Table 1. Grades of chronic kidney disease categorised by glomerular filtration rate (after a table by Rees et al26)

| Chronic kidney disease grade | Glomerular filtration rate (ml/min/1.73 m²) |
|-----------------------------|---------------------------------------------|
| Grade I                     | >90                                         |
| Grade II                    | 60–90                                       |
| Grade III                   | 30–60                                       |
| Grade IV                    | 15–30                                       |
| Grade V                     | <15                                         |

In general, serum biochemical anomalies may occur in grade II, symptoms appear in grade III, become increasingly severe in grade IV and renal replacement therapy is required in grade V.

The literature on direct comparisons of alternative intravenous agents in CF is complex. Comparing the relative toxicities of tobramycin and colistin in an observational study, Etherington et al22 found that tobramycin was associated with a larger acute rise in NAG. However, increased NAG levels (perhaps indicating chronic toxicity) prior to antibiotics was associated with a higher cumulative colistin dose rather than tobramycin level. In comparison, Al-Aloul et al12 found that, when administered with a non-nephrotoxic agent, colistin was cumulatively less nephotoxic than tobramycin.

Symptomatic hypomagnesaemia associated with aminoglycosides in CF is well documented. Holben et al described a series of 192 patients with CF with a prevalence of hypomagnesaemia of 5%.17

**Mechanism of renal toxicity**

Despite extensive study, the exact mechanisms of aminoglycoside nephrotoxicity remain elusive. Care must be taken when interpreting animal models of aminoglycoside kidney injury as these studies often use doses higher than those used in clinical practice. Recent technologies allow new methods of exploring mechanisms including metabolic and transcriptome analysis,18 proteomic and microarray techniques.19

Results from gentamicin (the most extensively studied aminoglycoside) are often extrapolated to other aminoglycosides. After glomerular filtration, approximately 15% of a filtered load of an aminoglycoside is reabsorbed into the kidney.20 The uptake of aminoglycosides into the proximal tubule (a key step in the pathogenesis of kidney injury) is saturable, although the level of this effect depends upon the aminoglycoside.21 Excretion of aminoglycosides from this compartment is a slow process, taking many days.22

Testing the hypothesis that uptake of drugs into the renal cortex is saturable in humans is difficult but has been studied in a series of studies in patients scheduled for nephrectomy.23

Different dosing schedules of aminoglycosides were used and kidney cortical drug concentrations were directly measured after nephrectomy. The renal cortical concentration of amikacin and gentamicin was lower after a single dose of amikacin compared with the same dose infused over 24 h, despite the area under the curve (AUC) of serum concentration versus time being higher. Thus, uptake of these drugs is saturable in vivo in humans. Although there was a trend to higher cortical levels when tobramycin was continuously infused, statistical significance was not achieved but the numbers of patients were limited.23

Aminoglycosides can affect the nephron at the level of the glomerulus and the tubule, although the latter has been more extensively studied. Insights into the potential mechanisms of decreased glomerular filtration have been gained from ultrastructural studies.24 Exposure of the glomerulus to gentamicin decreases the surface area available for ultrafiltration, the basement membrane increases in thickness, the glomerulus changes in shape and the number and diameter of fenestrae decrease.

The effect of gentamicin on the proximal tubular cell is summarised in figure 1. After uptake, a number of cellular processes are activated, culminating in apoptosis. This contributes to loss of the renal tubular epithelium and thus kidney dysfunction but, conversely, shedding and urinary excretion of apoptotic bodies may excrete aminoglycosides.

Changes in proximal tubular function are manifested as proteinuria, glycosuria and urinary loss of brush border enzymes. Altered tubular function has been attributed to multiple mechanisms including alteration in production of membrane transporters15 and decreased parathyroid hormone-mediated cAMP formation and subsequent decreased magnesium uptake.25

**Strategies to reduce nephrotoxicity of aminoglycosides**

Many groups are currently investigating strategies to reduce the nephrotoxicity of aminoglycosides. Potential approaches include blocking tubular uptake, reducing the formation of reactive oxygen species (ROS), inhibiting cellular signalling leading to apoptosis, and enhancing the ability of the kidney to regenerate. These were outlined 10 years ago but have not yet been translated into clinical practice.26

Strategies to reduce nephrotoxicity already exist. Attention to the dose prescribed and monitoring serum levels are paramount in this. Patients with CF who develop AKI are more likely to have received gentamicin in the preceding year than tobramycin when compared with controls,19 so tobramycin is the preferred aminoglycoside in CF. Avoidance of risk factors such as concurrent non-steroidal anti-inflammatory drugs and dehydration would seem prudent.

Manipulation of the dosing regimen provides a cost-effective and simple method of reducing kidney injury. Given the saturable uptake of aminoglycosides, it follows that a single daily dose would be expected to be less nephotoxic than the same daily dose in three divided doses (see figure 2). The TOPIC study, a large randomised trial of tobramycin for patients with CF, established that there is equal efficacy with a single daily dosing regimen as with a multiple daily dosing regimen,27 a finding confirmed in a subsequent meta-analysis.28 In the paediatric group receiving a single daily dose, the serum creatinine level decreased during the course of treatment compared with a rise in the group receiving three divided doses. In further support of renal safety, in the once daily group the rise in NAG was 33% less than in the group receiving a three times daily regimen in both adults and children.

The pharmacokinetics of aminoglycosides are complicated by a circadian rhythm in elimination. In the once daily group of the TOPIC study, most of whom received their antibiotics in the evening, there was a lower elimination rate of tobramycin than in the three times daily group.29 There may be a diurnal variation in renal clearance of the drug, with decreased clearance occurring at night. This would lead to increased exposure of the kidneys to aminoglycoside during the course of the illness if the
drug is administered at night compared with the morning. This hypothesis requires testing in clinical trials.

**OTOTOXICITY**

**Otoxicity in patients with CF**

Estimates of the prevalence of otoxicity in patients with CF depend on the methods used and definitions of hearing impairment. The most straightforward measurement of hearing impairment is the pure tone audiogram (PTA). In this, the patient listens to a series of different frequencies and the quietest level, measured in decibels (dB)).

A selection of the numerous studies reporting PTAs in CF which clearly report the population studied, the definition of hearing impairment and the sample prevalence is given in table 2. A frequently cited report of the prevalence of hearing impairment is that of Mulheran et al. They studied 70 patients recruited from a CF clinic and, using the definition of thresholds of hearing impairment of 17%.

Oto-acoustic emissions (OAE; sounds generated by the cochlea during hearing) and distortion product OAEs (DPOAE; in which the ear generates a sound of a third pitch when presented simultaneously with two sounds of differing pitch) have been proposed as an early screening tool for cochlear damage. Patients with CF can have a normal PTA but abnormal oto-acoustic emissions which may be an indicator of early toxicity.31

As in the kidney, interactions between iron and aminoglycosides leading to the formation of ROS play a role in toxicity. Administering iron in animal models increases otoxicity.37 Gentamicin causes upregulation of intermediaries in the apoptosis signalling pathways between the cell surface and the nucleus. Blocking these and other intermediaries in the apoptosis cascade causes reduced apoptosis in response to gentamicin.38

Several mitochondrial ribosomal mutations are associated with otoxicity. The most well known is the A1555G mutation in the MT-RNR1 mitochondrial gene described by Prezant et al.39

**Mechanism of otoxicity**

While the mechanisms of otoxicity and nephrotoxicity are similar, there are several key differences. Otoxicity and nephrotoxicity are separate events, so patients with otoxicity need not have nephrotoxicity and vice versa. Megalin is responsible for the uptake of aminoglycosides into the proximal tubular cells but is not found in the hair cells of the organ of Corti.35 After aminoglycoside administration, the concentration in the hair cells of the organ of Corti is static over several weeks (the kidney gradually excretes it).34

Aminoglycoside otoxicity causes degeneration of hair cells in the organ of Corti, predominantly in the basal turn which is required to sense high-frequency sounds. Extracellular signalling normally causes influx of calcium into hair cells. After gentamicin exposure, calcium influx no longer occurs in basal hair cells but persists in apical hair cells.36 Thus, a clear relationship exists between the clinical features of high-frequency hearing loss, pathophysiological and histological findings.

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**Review**

**Figure 1** Potential cellular mechanism of toxicity. This diagram summarises the major themes of the literature, much of which is in vitro work and so has to be interpreted as a model. Gentamicin enters the tubular cell via the multiligand receptor megalin (shown in blue) and then uptake is via a clathrin-coated pit. Gentamicin passes in a retrograde manner via the lysosomes, Golgi and endoplasmic reticulum into the cytoplasm where it forms complexes with iron which catalyse the formation of reactive oxygen species (ROS). In the lysosomes, gentamicin causes release of the pro-apoptotic cathepsins and, in the endoplasmic reticulum, gentamicin causes release of the pro-apoptotic stress protein von Hippel–Lindau binding protein 1 (VBP1). Cytosolic gentamicin either directly or indirectly (via ROS) causes the release of cytochrome C from the mitochondria, a key step in apoptosis. ROS cause an increase in the gene expression of oxidative stress-inducible genes, stress-inducible chaperones and oxido-reductive enzymes. Gentamicin may have intracellular effects without entry to the cell. It can cause a rapid increase in intracellular Ca2+ and an early increase in the anti-apoptosis and pro-proliferative signals Akt and Erk; this may be mediated by the membrane Ca2+ receptor CaR (shown in purple).
The cost-effectiveness of screening for the A1555G mutation in patients with CF has been modelled. Making the assumptions of a low prevalence of the mutation (0.086%), a high false detection rate, and that clinicians would switch antibiotics while the study by Scheenstra et al had a higher cumulative exposure to tobramycin.

### Strategies to prevent ototoxicity

Measures to reduce the generation or effects of ROS due to aminoglycosides have been extensively investigated and there are several promising avenues of research. A brief report of a randomised trial of aspirin versus no aspirin for 14 days in 195 patients treated with gentamicin showed a lower incidence of hearing impairment with aspirin (5% vs 15%; p = 0.013). Additionally, a randomised trial comparing N-acetylcysteine (NAC) versus no NAC in adult patients on dialysis treated with gentamicin reported fewer cases of hearing impairment in the NAC group (20% vs 55%, p = 0.022). Both of these preliminary results require confirmation but are nevertheless encouraging.

### VESTIBULOTOXICITY

Vestibulotoxicity due to aminoglycosides is secondary to damage of the hair cells in the vestibular system and does not necessarily occur with cochleotoxicity. Early symptoms of vestibulotoxicity, often insidious in onset, include vague dizziness and feeling light-headed. These may easily go unrecognised. Partial recovery can occur but can take several months.

There is little published literature regarding the prevalence of vestibulotoxicity in CF owing to the difficulty of defining and measuring it, particularly in children. A number of possible tests are available to screen for vestibular dysfunction, many of which are technically challenging for the non-specialist. Two studies have systematically screened for vestibulotoxicity in CF. Thomsen et al reported a study of 55 patients with a mean age of 11.8 years who had received a mean of 3.2 courses of tobramycin. A differential caloric test was performed and, in this series, no patients had an abnormal result.

Scheenstra et al reported a detailed investigation of 25 patients with CF in the Netherlands, 30% of whom had peripheral vestibular loss diagnosed by electronystagmography (often considered the gold standard test). Of three patients who reported symptoms which suggested horizontal vestibular function loss such as rotatory dizziness, one patient had severe bilateral vestibular dysfunction. There was no association between cumulative exposure to tobramycin and vestibulotoxicity.

Comparison of the aminoglycoside exposure of the two studies is difficult as the study by Thomsen et al reported mean courses of antibiotics while the study by Scheenstra et al reported cumulative area under the curve. It is likely that the patients in the study by Scheenstra et al had a higher cumulative exposure to tobramycin.
CONCLUSION

Patients with CF have a large lifetime exposure to multiple medications. Aggressive treatment with intravenous antibiotics slows the progression of lung disease, decreases morbidity and delays mortality, with a cost of an increased risk of side effects. Minimising the risk of side effects is of paramount importance within the ‘first, do no harm’ principle.

Recent single-centre studies have suggested that the prevalence of side effects such as chronic kidney disease, ototoxicity and vestibulotoxicity are disturbingly high in the CF population. If these findings are confirmed, clinicians will need strategies to minimise the risk of toxicity. CKD has implications for lung transplantation as it is well recognised that further nephrotoxicity occurs after transplantation.

Currently, there are limited options for monitoring the early signs of toxicity. Serum creatinine-based formulae are poor at predicting renal function in the CF population; a biomarker which reliably predicts GFR (or screens patients for renal impairment) is urgently needed in the clinical setting. Similarly, once the PTA has become abnormal due to aminoglycoside ototoxicity, hearing is not expected to improve. Further research evaluating the role of DPOAEs as an early marker of cochlear damage is required.

Some strategies have already been widely adopted, such as once daily aminoglycosides, the use of tobramycin rather than gentamicin and careful monitoring of drug levels. Other approaches such as the optimum time of day of administration of antibiotics, screening for mutations which predispose to otoxicity and the use of aspirin and NAC in patients with CF require careful consideration and further research.

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