Neutrophil Gelatinase-associated Lipocalin (NGAL)  
A Promising Biomarker for Early Detection of Nephrotoxic Injury

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Renal toxic injury can neither be predicted nor quantified exactly by the serum creatinine level. NGAL is a promising biomarker for early detection of AKI due to different causes. Our objective was to evaluate its value in the setting of nephrotoxic injury. We performed an experiment in which we administered gentamicin, atropine and fipronil at different doses to the laboratory mice to quantify the evolution of creatinine and NGAL. Compared to creatinine, NGAL increased faster and was detected at elevated levels from the first 4 h after administration of both low and high dose gentamicin. Fipronil, atropine and the combination caused a significant increase in NGAL serum values at 4 hours, an increase that lasted up to 24 and 72 h, respectively, compared to the control group. Its highest levels were recorded at 4 h. Compared to creatinine, NGAL increased faster and was detected at elevated levels from the first 4 h after administration of fipronil, atropine and the combination of them.

Keywords: NGAL, neutrophil gelatinase-associated lipocalin, acute kidney injury, nephrotoxic

Nephrotoxic injury is not always clinically apparent, so laboratory tests are essential for diagnosis. Toxic renal damage may be suspected in the case of urinary volume changes, but oliguria is not always present. Acute kidney injury (AKI) represents the sudden loss of renal function reflected by a decrease in glomerular filtration rate (GFR), increased creatinine and blood nitrogen levels, and failure of the hydroelectrolytic homeostasis regulation of the kidney (1-3).

Functionally lipocalins are transport proteins, they contain a 20 aminoacid signal peptide and the lipocalin domain. The domain helps binding lipocalins to their ligands. This structurally includes an eight stranded β barrel with its loops running in an antiparallel direction. The lipocalin transports low molecular chemicals (4,5).

Crystallographic studies have demonstrated the high affinity of NGAL for binding bacterial siderophores and for the first time NGAL was discovered as a neutrophil granule component. It is demonstrated that NGAL can be found in lots of human tissues like liver, lung, kidney, small intestine, bone marrow, prostate, adipose tissue and macrophages. NGAL main function is acting as a bacteriostatic agent, by binding to bacterial siderophores it restricts the iron needed by bacteria for growth. (6-8) NGAL is also a chemoattractant for neutrophils and an inhibitor of oxidative stress. (9) Toxins and ischemia are very frequent involved in developing AKI. (10) Early during AKI high levels of NGAL are detected. Animal experiments suggest that NGAL could reduce apoptotic tubule cell death (11). NGAL can lessen the renal injury and inhibit epithelial cell apoptosis via Bcl2/Bax signaling pathways (12-16).

A child in the early stages of severe AKI with a significantly reduced GFR could have relatively normal or slightly elevated creatinine values because there was not enough time for it to accumulate. In addition, creatinine is removed by dialysis, it can not properly evaluate the renal function once dialysis has started. Despite these limitations, serum creatinine continues to be the most commonly used laboratory test for the diagnosis of AKI in children (17-18).

Objective
Renal toxic injury can neither be predicted nor quantified exactly by the serum creatinine level. The occurrence of acute renal injury during children poisoning increases mortality, increases the duration of hospitalization and aggravates the long-term prognosis of patients so early and correct recognition of AKI is essential for the correct assessment of intoxication cases. We need novel markers for early detection of AKI in children poisoning. NGAL is a promising biomarker for early detection of AKI due to different causes. Our objective was to evaluate its value in the setting of nephrotoxic injury (19-23).

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**Experimental part**

**Material and methods**

**Experiment 1**

45 Wistar SPF (specified pathogens free) rats were used purchased from the Cantacuzino Institute and divided into 3 groups as follows:
- Group A - control group - 15 rats - which were injected intraperitoneal (ip) with 0.1mL / kg ip solution control - physiological serum (group M)
- Group B - test group - 15 rats - tested with gentamicin 25 mg / kg. ip (group G1) for three days
- Group C - test group - 15 rats - tested with gentamicin 75 mg / kg.ip (group G2) for three days

**Experiment 2**

60 Wistar SPF (specified pathogens free) rats were used, purchased from the Cantacuzino Institute which were divided into 4 groups as follows:
- Group M - control group - 15 rats - which were injected ip with 0.1 mL normal saline / 100g body weight (group M)
- Group B - test group - 15 rats - 7.5 mg / kg fipronil administered by oral gavage (group B)
- Group C - test group - 15 rats - atropine tested 28 mg / kg administered ip. (group C)
- Group D - test group - 15 rats - tested with the combination of oral fipronil 7.5 mg / kg. body + atropine i.p. 28 mg / kg body weight (group COMB)

The rats were housed in groups of three in plexiglass cages whose floor was covered with absorbent specimens and were brought into the laboratory 5 days before the experiment for acclimatization. Throughout the experiment the rats had ad libitum access to food and water. Rat cages were housed in the same room throughout the experiment with maintaining a relatively constant temperature (~ 22 °C).

**Substances used:**
- normal saline solution
- gentamicin solution for injection 80 mg / 2 mL (Biochemi)
- fipronil solution 0.25g / 100mL (Merial)
- sulfuric atropine (Sigma-Aldright)
- the combination of fipronil-atropine (COMB)

**Parameters monitored**

Rats were weighed and monitored daily from the point of view of general appearance (fur, mucous membranes, snout, ears) in the event of appearance or behavior changes. According to the protocol of work in the 3 moments of the experiment blood samples were collected and serum creatinine (crea) and NGAL (neutrophil gelatinase-associated lipocalin) were measured. Blood samples were collected after 12 h of fasting. Biochemical determinations were performed by MedCenter SRL (possessing RENAR accreditation).

The rats were anesthetized with diethyl ether to ensure rapid installation and lasting anesthesia as well as easy recovery from anesthesia. The blood samples were collected by puncture of the retroorbital sinus with the help of Pasteur glass pipettes with 145 mm length. With the tip of the pipette, the puncture of the retroorbital venous plexus was performed collecting 1.5 mL of blood. The blood was harvested in serum separator tubes. Serum determinations of NGAL were done by ELISA method.

Statistical results were calculated by computing standard averages and deviations. Statistical significance testing was performed using the t-Student test. For statistical analysis of the data we used Microsoft Excel 2010.

All the experiments described in this article were carried out in 2012 in the experimental laboratories of UMF Bucharest in accordance with The European Directive 86/609/EEC/24.11.1986 and The Romanian Government Ordinance 37/30.01.2002 regarding the protection of animals used for experimental and other scientific purposes.

**Results and discussions**

**Experiment 1**

Gentamicine (C21H41N6O7) is an antibiotic that belongs to the aminoglycoside group. The renal toxic effect of aminoglycosides is not fully understood. Aminoglycosides form reactive oxygen species that cause lipid peroxidation. They destroy phospholipids contained in the brush border membrane of proximal tubules, the lumen is obstructed with cellular debris and renal impairment occurs. Early after administration we can observe a raise in the dimensions and number of the lysosomes, they contain lamellar structures that represent undegraded phospholipids. This process is called phospholipidosis and appears to be the basis of nephrotoxic injury (24-27).
In the control group, the average weight of the rats at baseline was 181g, then at 24 h, they have presented an average of 173g, 72 h an average of 169 g, and at 7 days the average was of 172g.

The fact that all the animals in this group showed a weight loss during the 7th day can be explained by the blood collection maneuver that is quite stressful for the laboratory animals.

Compared to the control group, the small dose gentamicin treated group had an average body weight of 187 g at baseline, as at 24 h this was 182 g at 72 h 178 g, and at 7 days averaged 182 g. We can conclude that low dose gentamicin determined a decrease in body weight parallel to the control group.

The group receiving gentamicin 75 mg / kg.c (G2) had an initial mean weight of 182g reaching 176 g at 24 hours, 172 g at 72 h and 173 g at 7 h days. As with small doses, high dose gentamicin caused a decrease in weight compared to the control group. The evolution of the three lots was parallel to control group.

**Determination of serum creatinine level and NGAL in the three studied groups**

**Serum creatinine**

Compared to the control group, the G1 group had an average serum creatinine of 0.72 mg / dL at 4 h , at 24 h was 0.96 mg / mL, at 72 h was 0.94 mg / mL. We can conclude that gentamicin administered at low dose resulted in a significant increase in serum creatinine from the second day of administration compared to the control group. The lot receiving gentamicin at high dose (G2) had a mean serum creatinine at the start of the experiment of 0.98 mg / mL to reach 0.92 mg / mL at 24 h and 1.05 mg / mL at 72 h. As in the case of the previous group G1, in the case of group G2 the large-dose gentamicin, produced a significant increase in creatinine levels compared with the control group from 24 h and up to 72 h.
Serum determination of NGAL for the three studied lots

Thus, it was observed that the group receiving gentamicin at low dose had a significant increase of mean serum value of NGAL of 899 U / mL at 4 h (p<0.05) after the administration, at 24 h this was 653U / mL (p<0.05), and at 72 h 482 U / mL (p<0.05). We can conclude that the low dose of gentamicin determined a significant increase in NGAL starting at 4 hours after comparative administration with control group.

The lot receiving gentamicin at high dose had a significant increase of the mean value of NGAL at the start experiment of 1773 U / mL (p<0.05) to reach 1333 U / mL at 24 h (p<0.05) and 631 U / mL at 72 h (p<0.05). As in the previous case, in the G2 group it can be seen that gentamicin has led to an important increase in serum levels of NGAL starting 4 h after administration and maintaining high levels 72 hours compared to the control group.

Compared to creatinine, NGAL increased faster and was detected at elevated levels from the first 4 h after administration of both low and high dose gentamicin.

Experiment 2

Fipronil (C12H4Cl2F6N4OS) is a new class phenylpyrazole insecticide used in many countries against insects that gained resistance to conventional pesticides. The toxicity of fipronil appears by acting on the GABA chloride channel in nerve membranes resulting in increased excitability of the neurons by inhibition of chloride passage. The nephrotoxic effect of fipronil is caused mainly by inducing oxidative stress and generation of free radicals, the effects being dose dependent. The nephrotoxic injury causes cellular necrosis in the proximal tubule and interstitial fibrosis (28-31).

Acute atropine overdose can induce an anticholinergic syndrome which manifests as both peripheral and CNS symptoms. Toxic effects can appear at therapeutic concentrations, atropine blocks the action of acethycholine at the muscarinic receptor. The smooth muscles of the bladder are considerably affected producing urinary retention. Severe atropine poisoning can produce metabolic acidosis, rhabdomyolysis and acute kidney injury. (32)

As can be seen from the above tables, for the control group, the average weight at baseline was 180g. Then at 24 hours, they have showed an average of 179g, at 72 hours an average of 178g and at 7 days the average was 181g.

Compared to the control group, the fipronil treated group had an average body weight of 180 g at the initial time, at 24 h this was 174 g, at 72 h 170 g, and at 7 days have an average weight of 173 g. We can conclude that fipronil produced a more pronounced decrease in body weight compared to the control group.

The group that received atropine had an initial average weight of 188 g that it would reach 184 g at 24 h, respectively 182 g at 72 h and 187 g at 7 days. In this case the group retained a parallel evolution with the control group, the weight loss being less pronounced than with fipronil.

Regarding the combination of fipronil - atropine it was observed that initially this group had an average weight of 185g, at 24 h to 178 g, at 72 h an average weight of 172 g, at the end of the experiment reached an average weight of 172 g.

In conclusion, the combination resulted in a significant decrease in weight of the treated animals compared to the other groups.
Determination of serum creatinine level in the studied groups

Compared to the control group, the fipronil-treated group had an initial average serum creatinine of 0.76 mg/dL at 24 h 0.8 mg/dL and at 72 h its value was 0.85 mg/dL. We can conclude that fipronil has led to an increase in creatinine starting from day 2 compared to the control group.

The group that received atropine had a mean serum creatinine value at the beginning of the experiment of 0.76 mg/dL, to remain at the same value at 24 h, and to increase at 0.86 mg/dL at 72 h. As with fipronil, in the atropine treated group we observed a slight increase in creatinine compared to the control group.

Regarding the combination of fipronil - atropine it is noticeable that initially this group had a mean serum creatinine of 0.88 mg/dL, at 24 h raised to 0.92 mg/dL, and at 72 h recorded an average value of 0.9 mg/dL.

Although this combination determined the highest creatinine levels, however these values are not far exceeding the normal values.

Serum determination of NGAL for the studied groups

The group treated with fipronil had a significant increase of serum NGAL value of 486 U/mL at 4 h (p<0.05), as at 24 h this was 287 U/mL (p<0.05) at 72 h of 198 U/mL (p<0.05). We can conclude that fipronil caused an increase in NGAL serum values at 4 h, an increase that lasted up to 24 hours and 72 h, respectively, compared to the control group. Its highest levels were recorded at 4 h.

The group that received atropine had a mean value of NGAL at the beginning experiment at 653 U/mL (p<0.05) to reach 554 U/mL at 24 h (p<0.05) and 262 U/mL (p<0.05) respectively at 72 h. As with fipronil, in the case of the atropine treated group, it can be observed that atropine determines increases in serum NGAL starting at 4 h after administration and are maintained for up to 72 h after administration compared to the control group.

As far as the combination of fipronil - atropine is concerned, it was observed that this group had initial a significant increase reaching levels of 742 U/mL NGAL (p<0.05), at 24 hours the level was 598 U/mL (p<0.05), and at 72 h we recorded an average of 219 U/mL (p<0.05). In conclusion, the combination caused an increase of NGAL values throughout the experiment. Compared to creatinine, NGAL increased faster and was detected at elevated levels from the first 4 h after administration of fipronil, atropine and the combination of them.
Conclusions

NGAL remains a very promising biomarker for early detection of AKI. In children poisoning it is very important to prevent the development of AKI, because it is demonstrated that renal failure worsens the patient prognosis. Our study suggests that in poisoned animal models serum NGAL raises rapidly and remains persistent elevated in severe nephrotoxic injury. Compared to creatinine it reaches higher levels early during acute poisoning. The results must be validated in clinical practice. In pediatric practice AKI is frequently underestimated and sometimes underdiagnosed. It is very important to have an early marker for AKI in pediatric practice because of better hydration and better control of nephrotoxic drugs we could prevent the onset of toxic renal injury during hospitalization (33-37).

References

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