CD44-Positive Cells and Hyaluronan are a Hallmark of a Rat Model of Aristolochic Acid Nephropathy

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Abstract

Background: Aristolochic acid (AA) Nephropathy (AAN) is characterized by tubulointerstitial injury leading to fibrosis and tubular atrophy. Defective regeneration of tubular cells was hypothesized as a mechanism leading to Proximal Tubular Epithelial Cell (PTEC) atrophy. Here, we examined the distribution of CD44, an adhesion molecule involved in differentiation, and its main ligand, Hyaluronan (HA), in an experimental AAN model.

Methods: Wistar rats were injected daily with AA or vehicle during 35 days. Tubular enzymuria and tissue expression of CD44 and HA were evaluated at days 3, 7, 10, 21 and 35. Co-expressions of CD44 and PCNA (proliferation), vimentin (mesenchymal phenotype), α-SMA (myofibroblasts), and CD133 (progenitor cells) were investigated.

Results: AA induced acute tubular damage followed by PTEC atrophy. In controls, CD44 was limited to the basolateral membrane of collecting ducts while HA expression was confined to the medullary interstitium. CD44 was overexpressed as soon as 3 days after AA exposure, spreading to the apical and basolateral membranes of dedifferentiated PTECs and appearing on numerous interstitial cells along with marked HA accumulation around necrotic tubules. Both CD44 and HA expressions increased steadily throughout the 1-month follow-up. A large amount of tubular and interstitial CD44+ cells were proliferative (PCNA+) and mesenchymal-like (vimentin+). Few CD44+ cells expressed α-SMA or CD133.

Conclusions: Our results suggest CD44 is a strong marker of dedifferentiation, and thus of regeneration, in a rat model of AAN. Concomitant HA accumulation around necrotic tubules may create a sustained crosstalk between CD44+ cells and their interstitial ligand. PTEC atrophy is accompanied by overexpression, not deficiency, of the CD44-HA axis.

Keywords: CD44; Hyaluronan; Tubular Dedifferentiation; Progressive chronic kidney disease; Aristolochic acid nephropathy

Introduction

Initially called Chinese herb nephropathy, Aristolochic Acid Nephropathy (AAN) was reported in 1992 in young Belgian women after intake of slimming pills containing root extracts of Aristolochia sp. [1,2]. AAN is a progressive chronic Tubulointerstitial (TI) disease characterized by tubular atrophy and dense TI fibrosis with relatively well-preserved glomeruli [1,3]. It is also found in various regions of the world where exposure to similar toxins is prevalent [4]. Clinically, AAN results in a rapid deterioration of renal function leading to end-stage renal disease [2]. Our group has developed an experimental model of AAN in rats [5]. This model recapitulates the structural and functional impairment of proximal tubular epithelial cells (PTECs) shown in humans [5,6]. Increased oxidative stress, prominent collagen deposits and TGF-β expression, impaired tubular regeneration [7], as well as massive inflammatory cells infiltration [3] are hallmarks of experimental AAN. The latter has a biphasic evolution with an acute early phase (3-10 days) of tubular necrosis followed by a chronic phase (after 14 days) of interstitial fibrosis and tubular atrophy [3,5]. Interestingly, histological data suggest a defect of tubular repair leading to the hypothesis that incomplete regeneration of tubular cells could enhance the development of severe tubular atrophy in AAN [5].

Numerous studies have described the renal accumulation of CD44, an adhesion molecule involved in cell differentiation, and its main ligand, Hyaluronan (HA), a matrix glycosaminoglycan, in various renal diseases such as ischemia-reperfusion, lithiasis, or renal allograft rejection [8-13]. CD44 is a glycoprotein involved in cell-cell and cell-matrix interactions, in particular cell migration and tissue repair. CD44 is strongly upregulated in the kidney during proximal tubule regeneration [10,12]. Delayed resolution of inflammation and reduced HA clearance are observed in CD44-knockout mice recovering from renal, liver, or lung injuries [14-16]. Moreover, CD44 is a marker of undifferentiated mesenchymal cells [17]. HA, on the other hand, is a ubiquitous glycosaminoglycan found in the extracellular matrix of many tissues including the renal medulla, but not the cortex, except during inflammatory and ischemic injuries.

The current study was designed to examine the distribution and time course of CD44 and HA expression during experimental AAN.

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Materials and Methods

Experimental protocols

The experimental protocol approved by the local Ethic Committee for Animal Care was performed in 4-week-old male Wistar rats (Elevage Janvier, Le Genest Saint-Ilaire, France). Experimental procedures were conducted according to the National Guidelines for the Care and Use of Laboratory Animals. Weight-matched rats were randomly assigned to two groups. One group (n=30) received s.c. injections of AA (a mixture containing 40% AA1 and 60% AAII, Acros Organics Co.) at 10 mg/kg bw daily for 35 days. AA was dissolved in polyethylene glycol (PEG) 400 (Fluka Chimie) to a final concentration of 10 mg/ml and diluted in distilled water before injection. The SHAM group (n=30) was treated with a v/v solution of PEG 400 and distilled water only. Control rats (n=4) did not receive any treatment and were euthanized at day 0. Body weights were measured weekly for adjustment of drug dosages. Six rats in each group were euthanized on days 3, 7, 10, 21 and 35, following a 24-h period in a metabolic cage to collect urine. Blood sample was collected, and the kidneys were excised and immediately processed for further analysis.

Biochemical evaluation of urinary tubular markers

Urinary creatinine values were determined by a pseudokinetic Jaffe’s method using a Creatinine Diagnostic Kit (Sigma-Aldrich) [5]. Total proteinuria was quantified by the Bradford binding assay as described in [18]. Urinary excretion of lysosomal enzyme N-Acetyl-β-D-Glucosaminidase (NAG) was measured by a colorimetric assay following the manufacturer’s protocol (Roche Diagnostic). Urinary excretion of Leucine Aminopeptidase (LAP) was detected by a spectrofluorimetric assay as detailed previously [5].

Histology

Paraffin-embedded kidney sections were stained with PAS, hemalun, and Luxol fast blue to assess morphological alterations and with Sirius Red to detect collagen I and III deposits.

Immunohistochemical procedures

CD44 (mouse anti-rat antibody OX49, Pharmingen), vimentin (chicken anti-vimentin antibody, Chemicon), CD133 (rabbit anti-CD133 antibody, Biorbyt), and α-smooth muscle actin (α-SMA) (mouse anti-α-SMA, DAKO) were detected on paraffin sections using a streptavidin-biotin immunoperoxidase method (ABC method) as detailed in previous publications [3,7,8]. Detection of HA was performed using a biotinylated HA-Binding Protein (HABP, Calbiochem). Kidney sections were labelled sequentially at room temperature between the secondary antibodies, antisera raised in different species. Thereafter, sections were incubated for 1 h at room temperature with the second primary antibody, followed by FITC-conjugated secondary antibody for 30 min. After final rinses in distilled water, sections were mounted in ProLong® Gold antifade reagent (Invitrogen).

Morphometric analysis of marker immunoreactivity in renal tissue

The relative area positively stained for HA and CD44 was assessed on a semi-quantitative basis by adapting a single-blind analysis of paraffin sections (1 section per animal). Slides were coded and analysed by two of us who was kept blinded of the treatment. To standardize the evaluation procedure, an additional lens engraved with a 1-cm² square grid was inserted in one of microscope eye pieces. Each paraffin section was scanned at x400 magnifications, and 15 square fields were evaluated. The degree of stained area was graded along an arbitrary scale.

Cell counts

The frequency of CD44 or CD133-positive cells in the interstitial spaces was evaluated by a semi-quantitative analysis as described previously [8]. Briefly, the distribution of positive cells in the interstitium was performed on one section per animal. For each section, 10 square fields (0.084 mm²/field) were observed at x400 magnification. The incidence of CD133-positive collecting ducts was also evaluated at x400 magnification. As a whole, 100 collecting ducts were evaluated in each specific renal area (OSOM and ISOM). Of note, the presence of one single positive cell makes the tubule positive. Results were expressed as the percentage of CD133-positive collecting duct according to their distribution (OSOM, ISOM).

mRNA quantification via real-time reverse transcriptase PCR

Total RNA was extracted with the High Pure RNA Tissue Kit (Roche Diagnostics) according to the manufacturer’s protocol. The mRNA quantification was performed using a 2-step real-time reverse-transcriptase PCR (LightCycler, Roche Diagnostics). β-actin was used as the housekeeping gene and was purchased as ready to use (Life technologies). TGF-β1 primers and probes were designed and purchased from Eurogentec. Relative gene expressions were calculated using the 2-ΔΔCT method.

Calculations and statistic

Results are presented as mean values ± SEM. The level for statistical significance was defined as P<0.05. Mann-Whitney unpaired test was applied to compare AA, SHAM and control groups at each corresponding time point.

Results

Time course of renal damage and TI lesions

A significant decrease in body weight was observed from day 10 to day 35 in AA-treated rats compared to SHAM rats at corresponding time-points (day 10: 283 ± 6 vs 307 ± 10; day 21: 322 ± 10 vs 364 ± 8; day 35: 346 ± 11 vs 422 ± 9 g respectively, P<0.05). However, the kidney-to-
body weight ratio significantly rose at days 21 and 35 in AA-treated rats, revealing renal hypertrophy (Figure 1A).

Proteinuria increased significantly as early as day 3 and persisted to day 35 in AA-treated rats (Figure 1B). In parallel, NAG enzymuria significantly increased from day 3 to day 35 (Figure 1C), reflecting structural impairment of PTEC [5]. A transient increase in LAP, a membrane-bound brush border enzyme, was detected at days 7 and 10, and then significantly decreased during the last days of observation (Figure 1D), confirming progressive PTEC atrophy. Conventional microscopy using PAS staining revealed morphological alterations as early as day 3: swelling of tubular cells, progressive loss of tubular brush border and a significant increase in infiltrating interstitial cells, mostly in the outer stripe of the outer medulla (OSOM) and in the medullary rays. As illustrated in (Figures 1F-H), patchy zones of necrotic PTECs associated with significant cell infiltration were observed during the acute phase. Necrotic cells and cellular fragments were found in the lumen of the proximal tubules. Later, at day 21 (Figure 1I), numerous atrophic tubules were detected with a maximal degree of severity at day 35 (Figure 1J).

**Time course of fibrosis**

As observed in Figure 2, the renal mRNA level of TGF-ß, a marker of fibrosis, significantly and progressively increased from day 10 to 35 in AA-treated rats (Figure 2A). On the other hand, there was no significant difference in the density nor distribution of the collagen components of the extracellular matrix, as stained by Sirius Red, from day 3 to 21. Collagen accumulated at day 35, mostly in OSOM and medullary rays, in the form of thickened basal membrane of altered PTECs and extensive fibrous deposits around necrotic tubules (Figures 2C and D). Overall, the time course of PTEC lesions, atrophy, and fibrosis is similar to what has been described in this AAN model [3,5,7].

**HA density and distribution**

In control or SHAM rats, HA was distributed in the three areas of the kidney (cortex, OSOM, and inner stripe of the outer medulla ISOM) with a strong cortico-papillary gradient (Figure 3A). In the normal cortex, HA concentration was very low and mainly concentrated in the connective tissue surrounding large blood vessels. A thin frame of HA staining was also present in the interstitial matrix around a few glomeruli or occasionally in the peritubular spaces (Figure 3A). In OSOM, as in the cortex, HA was

![Figure 1: Time course of TI lesions. Ratio of kidney weight to body weight demonstrates kidney hypertrophy at days 21 and 35 (A). Urinary protein level indicates a rapid elevation in AA group as soon as day 3 (B). Urinary N-acetyl-ß-D-glucosaminidase (NAG) level in AA group shows a significant increase by day 3 and remains elevated during the 35 days period (C). Decreased urinary leucine aminopeptidase (LAP) level (D) attests tubular damage. Representative photographs of tissue injury (x200) at different time points: control condition (E); day 3 - increase of interstitial cells (F-arrow); day 7 - patchy zones of necrotic proximal tubular epithelial cells (PTECs) (#) associated with significant cell infiltration (arrow) (G); day 10 - increase in necrotic PTECs (H); day 21 - increased tubular necrosis with appearance of atrophic tubules and significant cell infiltration (arrow) - (I); day 35 - extended zone of atrophic tubules (star) associated with infiltration of interstitial cells (arrow) - (J). Values are means ± SEM. N=6 in each group. *p ≤ 0.05 versus SHAM rat at corresponding time-point.](image-url)
found mostly in specific clusters around blood vessels and as a weak discontinuous network between tubular structures (Figure 3B). In ISOM, a continuous HA network extended between all tubular and vascular structures (Figure 3A).

AA intoxication induced significant changes in the distribution and density of HA in the renal tissue. In AA-treated rats, HA increased progressively and markedly in the extracellular matrix of outer medulla and medullary rays along tubular damages. During the acute phase, a highly HA-enriched matrix developed in the patchy zones of tubular necrosis (Figure 3C-F). Later, during the chronic phase, HA progressively invaded the whole interstitium and was associated with atrophic and undifferentiated tubules (Figure 3G-J). HA density was also associated with interstitial cell infiltration during the acute phase and with fibrotic areas in the chronic phase. As illustrated in Figure 3K, the time-course of HA density in the outer medulla showed a gradual and significant increase throughout the one-month observation period.

Time-course of CD44 expression

In control or SHAM rats (Figure 4A-C), CD44 immunoreactivity was mostly present on the basal membrane of collecting ducts. In ISOM, the vast majority (98%) of collecting ducts expressed CD44 at their basal pole (Figure 4C). In addition, the thin descending limbs of Henle’s loop were also uniformly positive for CD44. In the cortex and OSOM, a few collecting ducts expressed CD44 (Figure 4A&B). CD44 was also present, to a lesser extent, on isolated cells in proximal tubules and on some glomerular cells (Figure 4A). In all areas, a very small amount of CD44+ cells was observed within the interstitium (Figure 4A & B).

In AA-treated rats, at day 3, the density of CD44+ cells rose within the interstitium (Figure 4D-F). These cells were either isolated or concentrated in focal areas, between apparently intact tubules. At this time point, tubular CD44 expression was restricted to collecting ducts. At days 7 and 10 (Figure 4G-I), patchy zones of PTEC necrosis were observed together with a significant increase in CD44 expression. CD44 staining was now abundant on both interstitial cells around the damaged tubules and along apical and basolateral plasma membranes of necrotic PTECs. During the chronic phase, at days 21 (Figure 4J-L) and 35 (Figure 4M-O), TI injury extended dramatically and CD44 density remained high in both interstitium and atrophic or undifferentiated tubules.

Figure 4P illustrates the overall time course of CD44 immunostaining while Figure 4Q illustrates the number of CD44-positive cells in the interstitium in OSOM. Higher magnified photographs of CD44 expression on interstitial cells are shown in Figure 4R-W. In addition to a fibroblast-like phenotype, a subpopulation of strongly positive flat cells was found, forming a cuff surrounding the tubules (Figure 4U-W). These CD44+ cells were further characterized.

Characterization of CD44+ cells

**PCNA:** As it has been already demonstrated in our previous study [7], PCNA+ tubular and interstitial cells were seen in the renal tissue with a maximal expression during the acute AAN phase (1 to 10 days). As illustrated in Figure 5A-D, PCNA-positive cells emerged in significant numbers three days after AA intoxication (Figure 5A), to reach a maximum value at day 10 Figure 5B and progressively decreased at days 21 and 35 Figure 5C. Here, we showed that a significant proportion of PCNA+ cells was also CD44+, and thus proliferating, at days 7 to 21 with a peak at day 10 (Figure 5E-H). These proliferative cells were observed in both tubular and interstitial compartments. As

While *vimentin* was only expressed in few interstitial cells in control or SHAM rats (Figure 6A), its expression rose gradually and significantly throughout the observation period (Figure 6C). At day 3, no obvious sign of tubular expression was observed, vimentin was expressed in interstitial cells mostly located in the outer medulla and medullary rays (Figure 6B). At days 7 and 10, the proportion of vimentin+ cells significantly rose in PTEC area (Figure 6D). Moreover, damaged PTEC began to express vimentin, demonstrating their dedifferentiated phenotype (Figure 6D). During the chronic phase, both the dedifferentiated PTECs and an impressive proportion of interstitial cells expressed vimentin (Figure 6E&F). Vimentin/CD44 double staining revealed that after 10 days of AA intoxication, 75.6 ± 2.0 % of CD44+-cells were also vimentin+ (Figure 6I). This observation was confirmed at subsequent time-points. At day 35, for instance, atrophic tubules were colonized by both markers (Figure 6J-L).

α-SMA. In control or SHAM rats, α-SMA localization was limited to the media of vessels (Figure 7A). However, a small amount of α-SMA-positive cells was also located in the interstitial space. At day 3, a few clusters of α-SMA-positive interstitial cells were detected around tubules in the cortex and OSOM Figure 7B. Thereafter, the density of positive cells rose progressively and significantly in the zones of necrotic PTEC (Figure 7D&E). At day 35, α-SMA-positive cells gathered around atrophic tubules (Figure 7F). Altogether, α-SMA was never expressed at a tubular level and acquisition of the myofibroblast phenotype predominated during the fibrotic phase. (Figure 7C) illustrates the relative area occupied by α-SMA in the outer medulla over the course of the experiment.

Double fluorescent immunostaining for CD44 and α-SMA showed that only a small amount (10.0 ± 4.0 %) of interstitial α-SMA-positive cells were CD44-positive during the early phase (Figures 7G-H); that percentage increased at day 35 (23.3 ± 5.0 %), when both markers coincided in the patchy zones of PTEC necrosis (Figures 7I-J).

**Discussion**

Human AAN is a "pure" example of progressive T1 nephritis leading to fibrosis and end-stage renal disease. Our group has unraveled several features of this disease using a specific rat model [3,6,7], including the illustrated in (Figure 5 E-G), CD44 (red) was expressed on the apical and basolateral membrane of the cells while PCNA (green) was located in the nuclei. The onset of this proliferation is concomitant with the acute tubular necrosis (Figure 1F-H), suggesting an attempt of PTECs to regenerate Vimentin.
Figure 4A: Time course of CD44 expression. Representative photomicrographs of CD44 in renal tissue of control rat (A=Cortex x200; B=OSOM x400; C=ISOM x400) or AA-treated rats in OSOM at days 3 (D x200; E-F x400), 10 (G x200; H-I x400), 21 (J x200; K-L x400), and 35 (M x200; N-O x400). CD=collecting ducts; *=Atrophic tubules; #=necrotic proximal tubular epithelial cells.
presence of deficient tubular epithelium regeneration. Dedifferentiation markers such as vimentin were abundant in PTECs during the acute phase of AAN [7]. An imbalance between differentiation and proliferation was suggested as one causal mechanism of severe atrophy development.

In the current study, we demonstrate for the first time the temporal expression of CD44 and its major ligand, HA, in the rat model of AAN. CD44 and HA expressions increased dramatically and concomitantly throughout the course of the disease, in parallel with PETC necrosis and interstitial fibrosis. Both markers were tightly linked to the areas of TI lesions, suggesting a crosstalk between CD44 and HA, and between these markers and the AAN lesions. CD44 has been associated with regenerating cells in the liver and kidney [8,10,12,20,21] but from the results of our study there is no sign that a lack of CD44 might be involved in the poor tubular regeneration.

CD44 is an early marker of AAN. While its expression in control kidneys was low and confined to the basal membrane of the collecting ducts cells and a few interstitial cells, strong CD44 signals appeared as soon as 3 days and continued to increase remarkably thereafter, especially in interstitial cells gathering around necrotic PTECs but also on damaged tubules. A large number of these CD44+ cells was PCNA+, suggesting a role of CD44 in the regeneration process of AAN. Indeed, Pozdzik et al. [7] demonstrated a significant increase of PCNA+ tubular and interstitial cells in a similar AAN model. This rise of proliferating cells was maximal during the acute AAN phase, suggesting an attempt of PTECs to regenerate. During the chronic phase, Pozdzik et al. [7] observed a decline of proliferating cells in PTECs concomitant with an increase of tubular atrophy. This latter demonstrated a defect in the
regenerative process of PTEC and a development of TI fibrosis, which were also observed in the present study. Indeed, PCNA/CD44+ cells were maximal at day 10 after AA intoxication and progressively decrease thereafter. However, in our study, CD44 density remained very high during the chronic fibrotic phase, possibly in relation with sustained TI injury. Previously, we reported the involvement of HA and CD44 during the recovery phase of a renal ischemic insult [8]. We showed that CD44 was upregulated in regenerative PTECs concomitantly with an increase in HA density [8]. Therefore, the CD44-HA pair seems to be an intrinsic component of many renal diseases, with actions ranging from the repair process to late fibrosis development. During nephrogenesis, it has been shown that the HA-CD44 couple is critically involved in the formation and branching of the ureteric tree [22]. Similarly, the absence of CD44 worsened tubular injury consecutive to ureteral obstruction [14] as well as lung injury after bleomycin administration [16]. However, a study using CD44 knockout mice subjected to mild bilateral renal ischemia-reperfusion injury revealed that tubular regeneration was not altered and that kidneys recovered normal physiological functions [23]. Therefore, CD44 may not be crucial for all renal disease recoveries, depending on the severity of the lesions. A murine model may be useful to test the hypothesis that CD44 deficiency could lead to more severe tubular atrophy in AAN.

On the other hand, the implication of HA-CD44 in the progression of renal fibrosis has been highlighted in e.g. experimental glomerulonephritis [11] and cyclosporin nephropathy [24]. In the current model of AAN both CD44 and HA are strongly linked to fibrotic islets. What is the most likely link between CD44-HA and renal fibrosis?

To answer that question, our results first showed that only a small population of CD44+ cells co-expressed α-SMA, a marker of myofibroblasts, which are central to fibrogenesis. The best demonstration that HA-producing myofibroblasts and CD44 are principal actors in fibrosis has recently been provided by Li et al. [25]. These authors
have shown that CD44 is upregulated after bleomycin-induced lung injury in transgenic mice that overexpress the HA synthase HAS2 in myofibroblasts. In these mice, the administration of anti-CD44 blocking antibodies prevents the accumulation of collagen. On the contrary, in the rat AAN model, α-SMA expression is a late event limited to the chronic phase and the co-localization of α-SMA with CD44 is weak. Although some CD44+ cells may possess a fibroblastic phenotype, an important role of the HA-CD44 pair in fibrogenesis appears unlikely.

A second hypothesis is that CD44+ cells are mostly regenerative, stem cell-like cells in AAN. Indeed, the contribution of resident, or even infiltrating, stem/progenitor cells in renal regeneration and repair processes has been reported [26-28]. CD133+ cells derived from normal kidney tissue have been shown to differentiate into epithelial and endothelial cells in vitro [29]. Ye et al. demonstrated that CD133+ stem/progenitor cells residing in the human renal cortex and medulla exhibit a proliferative capacity after acute tubular necrosis [30]. In normal human kidney, CD133 seems to be mostly located in the parietal layer of Bowman capsule, brush border of PTEC, collecting ducts, and interstitium [27,30,31]. In many forms of experimental kidney injury, an upregulation of CD133 was reported [28,30,32]. However, this was not reproduced in our rat model of AAN since CD133 was mostly expressed in the collecting duct cells. Moreover, the co-expression of CD44 and CD133 revealed that only 10% of CD44+ cells are double-labeled in both control and AA-treated rats. In addition, we did not observe any evidence of CD133 expression in damaged or dedifferentiated PTEC. Overall, we do not believe that CD44 is a strong marker of stem/progenitor cells involved in renal tissue repair in AAN.

A third hypothesis is that a majority of CD44+ cells are mesenchymal-like cells. This is exactly what we found using vimentin as a recognized marker of mesenchymal cells and cell differentiation. Previous studies, focusing on ischemic kidney injury, have demonstrated a transient upregulation of vimentin in dedifferentiated S3 segment of proximal tubules [33,34]. We confirm there is a massive accumulation of CD44+/vimentin+ cells throughout the period of AA injections. These cells are not confined to the interstitial compartment but also belong to damaged tubules, suggesting a role of CD44 in unmasking the dedifferentiation

Figure 6: Representative photomicrographs of vimentin (x400) in renal tissue of control rat (A, arrow) or AA-treated rats at days 3 (B), 10 (D), 21 (E), and day 35 (F). Semiquantitative analysis of vimentin-positive staining in control, SHAM- or AA-treated rats at different time-points after treatment (C). Double immunostaining (x400) for CD44 (red (G-J)) and vimentin (green (H-K)) in AA-treated rats at day 10 (overlay I) and day 35 (overlay L). *p ≤ 0.05 versus SHAM; #p ≤ 0.05 versus control
process. In normal conditions of regeneration, the restoration of brush border is accompanied by a loss of CD44 and vimentin markers [8,33,34]. The persistence of these markers at the late stage of the AAN model may signal factors interfering with the recovery phase such as sustained interstitial inflammation or an irremediable profibrotic environment underscored by high TGF-β and α-SMA expressions. In conclusion, strong activation of the HA-CD44 axis in AAN may signal a continuous attempt by kidney cells to dedifferentiate and regenerate the normal tubular structure through the mesenchymal-like phenotype. These attempts may fail due to the uninterrupted exposure to toxic AA.

Conclusions

Aristolochic acid nephropathy is a progressive tubulo-interstitial nephritis. AA-induced nephropathy consists in a biphasic evolution with an acute phase (3-10 days) characterized by the impairment of the proximal tubules associated with proteinuria followed by a chronic phase (after 14 days) characterized by a progressive development of the interstitial fibrosis. Nevertheless, signs of tubular regeneration during the acute phase were previously shown [7]. Similarly to our data, a transient decrease in proximal tubular enzymuria (LAP) was observed along with a transient increase in proliferating (PCNA+) cells reflecting an attempt of PTEC regeneration. In our study, we demonstrated the temporal expression of CD44 and hyaluronan in the rat model of aristolochic acid nephropathy. We highlighted an early and sustained activation of the HA-CD44 axis along with impaired PTEC regeneration as confirmed by the localization of CD44 on proliferative cells as well as the high prevalence of mesenchymal-like phenotype in tubular structures. However, the regenerative process failed, probably due to the sustained toxic exposure to AA, resulting in an irreversible proximal tubular atrophy and TI fibrosis.

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Figure 7: Representative photomicrographs of αSMA (x200) in renal tissue of control rat (A) or AA-treated rats at days 3 (B), 10 (D), 21 (E), and day 35 (F). Semi-quantitative analysis of α-SMA-positive staining in control, SHAM- or AA-treated rats at different time-points after treatment (C). Double immunostaining for αSMA (green) and CD44 (red) (arrowhead) in AA-treated rats at day 10 (G, H x400 + insert to better illustrate the co-localization) and at day 35 (I, x400 – J, x1000). *p ≤ 0.05 versus SHAM; #p ≤ 0.05 versus control.

References

Figure 8: Representative photomicrographs of CD133 (x400) in renal tissue of control rat (OSOM, A – ISOM, B) or AA-treated rats at day 21 (OSOM, C – ISOM, D). The arrow in the insert (C) shows the CD133-positive cells in the interstitium. Double immunostaining for CD133 (green) and CD44 (red) in AA-treated rats at day 21 (E, x400; F-G, x1000). Semi-quantitative analysis of CD133-positive staining in the collecting ducts (OSOM, H – ISOM, I) and in the interstitium (OSOM, J – ISOM, K) for control, SHAM- or AA-treated rats at different time points after treatment.