Tubular Regeneration: When Can the Kidney Regenerate from Injury and What Turns Failure into Success?

Martin E. Johansson

Center for Molecular Pathology, Department of Laboratory Medicine, Lund University, Skåne University Hospital, Malmö, Sweden

Key Words
Kidney regeneration · Acute kidney injury · Tubular damage

Abstract
Background: The most common intrarenal cause for acute kidney injury/renal failure is tubular damage. The kidney tubules are arranged as compartments of cellular mosaics to perform their functions, and at rest almost a fifth of the human ATP consumption is allotted to the reabsorption of substances from the filtrate, rendering especially the proximal tubules highly sensitive to oxygen and/or nutrient deprivation. Normally mitotically quiescent, the tubular epithelium shows a brisk regenerative response following injury if supportive care is offered, allowing functional restoration. Despite this, the cellular machinery behind the regenerative capacity is still not unequivocally defined. This is at odds with other epithelia such as those of the skin and intestine, where stem cells maintain a continuous flow of new cells from designated niches. Summary: This review discusses the classical concept of renal regeneration, i.e. stochastically surviving cells undergoing dedifferentiation (or epithelial-mesenchymal transition) followed by replenishment of the tubular epithelium. Furthermore however, this view has recently been challenged by the concept of organ-confined stem/progenitor cells, bone marrow-derived stem cells, or mesenchymal stem cells taking part in the regenerative events. Whereas results from animal models support the classical view, morphologically distinct cells have been demonstrated in human kidneys, requiring interpretation. This review presents some of the previous work and techniques and highlights issues that need to be reconciled. Key Messages: In adult humans, the kidney tubules contain scattered cells with a distinct set of markers and properties, such as increased robustness during tubular damage. These cells may be induced by injury or represent a resident progenitor cell pool. To date, animal studies using lineage-tracing methods argue for an inductive scenario. In humans, the situation is less clear and one might speculate that the cellular heterogeneity might reflect elements of cellular reprogramming to a progenitor-like state, perhaps by induction. Due to intense investigational efforts, however, a scientific consensus may soon be reached, which will benefit further research.

Introduction
The term acute kidney injury (AKI) is slowly replacing the less distinct former designation, acute renal failure. The condition denotes a clinical syndrome where the glomerular filtration rate is in acute decline, causing a rise in...
creatinine and, if progressive, the gradual development of uremia. Despite some therapeutic achievements, AKI is still associated with considerable mortality [1]. Of the intrarenal causes for AKI, the overwhelming majority (80–90%) of cases are due to either ischemia or nephrotoxicity or both [2, 3]. The main tissue targets for AKI are the renal tubules, and the corresponding pathological changes are called acute tubular necrosis (ATN). This term is overly narrow, however, since the histological changes do encompass cell necrosis but also apoptosis and sloughing of cellular parts or indeed whole viable cells into the tubular lumen. In humans, actual cell necrosis is relatively rarely seen, as opposed to animal model systems for ATN where necrosis dominates the histological changes. Clinical, after the initiation phase, ATN enters the maintenance phase and, if supportive care is offered, a recovery phase follows where the renal tissue (most importantly the tubules) regenerates and kidney function is reestablished.

**Tubular Injury**

In health, the renal tubular epithelium is mitotically almost quiescent, in stark contrast to rapidly dividing epithelia such as in the small intestine, skin, or colon. The constant regeneration in these epithelia is driven by stem cells residing in clearly defined niches, but the epithelia of parenchymatous organs do not share these cellular kinetics (fig. 1a). Upon injury, however, the kidneys display a forceful regenerative capacity. AKI is seldom an indication for renal biopsy; therefore, human histological data regarding the distribution of injury along the nephron is limited. Instead, animal experimentation has shown that in general the proximal tubules (PT) are most sensitive to injury, followed by the medullary thick ascending limb part of the distal tubules (DT) [4]. Anatomically, the outer stripe of the outer medulla is most severely affected, and this is most probably due to the relatively low oxygen tension of the medullary region, which is constitutively hypoxic [5]. This environment of low oxygenation continues into the cortex by the medullary ray structures, supplied by venous blood from the medullary region. The medullary rays lack glomeruli, instead harboring the pars recta segments (S3) of the PT and the medullary thick ascending limb, descending into the outer medulla.

Tubular cellular injury may be divided into reversible and irreversible processes. Reversible renal injury results in depolarization of the tubular cell, where normally compartmentalized proteins are redistributed in injured cells [6]. Important examples are the redistribution of integrins and Na⁺/K⁺ ATPase to the apical membrane. Irreversible tubular injury results in either apoptosis or necrosis. Animal models most often employ arrest of the renal circulation for a considerable time (20–30 min), resulting in dramatic histological changes, with extensive necrosis and apoptosis. It may be questioned to what degree this correctly emulates the human counterpart. A model more close to human ATN is the animal model of septic AKI, which also produces relatively mild histological changes [7].

---

**Fig. 1.** a Normal human kidney stained for Ki67. No nuclei were positive, and thus no signs of mitotic activity could be seen. b, c Two markers for the scattered cells in the PT. Both were identified by gene expression analyses of ALDEFLUOR-sorted cells. Whereas MYOF is a novel marker for these cells (b), vimentin (VIM) is classically associated with proliferating tubular epithelia (c). Red arrows show the tubular cells and green arrows show the parietal epithelial cells of Bowman’s capsule. Interestingly, most markers for the scattered cell population in the PT are indeed also identified in the parietal epithelial cells. Counterstain: hematoxylin.
Tubular Regeneration

The Classical View

The undifferentiated metanephric mesenchyme has exhausted its capacity at birth, and any remaining nephrogenic rest at birth represents an aberration [8]. This holds true in most animals with the exception of fish, who retain their nephrogenic capacity during adulthood. Regarding the cellular source for tubular regeneration in mammals, the classical view is that the stochastically least injured cells sustain a reversible measure of injury, followed by dedifferentiation. The cells flatten out to cover the denuded basal membranes, divide, and repopulate the tubules [6]. Finally, tubular integrity is restored as the cells differentiate and repolarize. This view has been well substantiated in rodent model systems by thymidine incorporation assays as, for example, performed by Vogtseder et al. [9], who specifically investigated the S3 segment of the PT. It has been suggested that tubular cells of this segment are arrested in the G1 phase via cyclin D1, allowing for a tubular cell population that may rapidly respond to tubular injury. More recently a series of studies employed fate-mapping techniques. Cells derived from the Six2-expressing cap mesenchyme formed from the metanephric mesenchyme were used to tag the tubular epithelium [10]. Infliction of ligation-induced injury showed that only cells intrinsic to the tubules took part in the repair. This does not rule out a tubular progenitor cell category, but in a subsequent study, using an unbiased DNA analog-based serial label retention technique, it was demonstrated that regeneration occurred in a random fashion not consistent with a designated progenitor cell population [11].

Recently the proximal tubular epithelium was labeled using a CreERT2 cassette knocked into the locus for SLC34A1. It was shown that the bulk epithelium drives regeneration and a distinct subset of progenitor cells could not be identified. Furthermore, the regenerating cells displayed CD24 and CD133 along with KIM1, markers ascribed to renal progenitors, indicating that these epitopes are markers for dedifferentiation/injury rather than progenitor markers [12]. As authoritative as these studies are, it is important to bear in mind that they all are derived from rodent systems and do not formally establish the situation in the human kidney. Recently, however, it was shown in a rat model of unilateral ureteral obstruction that CD24+/CD133+/vimentin+ cells seem to be inducible during renal injury [13]. That study also showed that the majority of proliferating cells in human regenerating kidneys were CD24 positive.

Recent Ideas on Tubular Regeneration: The Stem/Progenitor Concept

During the last decade, stem or progenitor cells have been implicated in renal tubular repair. Reports have suggested 3 possible sources: bone marrow-derived stem cells, mesenchymal stem cells, and kidney-confined tubular stem cells [14]. Extrarenal stem cells initially received much interest and these were thought to directly support tubular regeneration by integration into the tubules. Fate-mapping experiments have shown conclusively however, that these cells only contribute indirectly to tubular regeneration. The consensus is that they provide soluble factors that support regeneration. This is underscored by the fact that nonrenal cells derived from granulation tissue ameliorate ischemic renal injury (IRI)-induced AKI by endocrine action [15].

Intrarenal stem or progenitor cells have been identified using two major approaches. Either unique properties ascribed to stem cells have been used to isolate this cell population or previously described markers from other stem cell systems have been used to search for kidney stem cells.

Functional Studies

Renal progenitor cells have been identified in rats using label retention after BrdU staining, where proliferating cells are labeled during the S-phase followed by a washout phase [16]. The method aims at labeling slowly dividing stem cells. The method is based as much on a vigorous washout phase as it is on a labeling phase and, though it is a classical approach in rapidly dividing epithelia, it is unclear to what extent this method may be used in mitotically quiescent parenchymatous organs such as the kidney or liver. Regardless, these cells have been localized to the renal papilla as well as to the renal tubules and it has been shown that the label-retaining cells of the PT account for the majority of cell divisions after renal injury [17]. Label-retaining cells in the renal papillary region have more recently been investigated during searches for mTERT-positive progenitor cells. Genetically labeled mTERT-expressing papillary cells could not be shown to be renal progenitor cells [18].

Side population studies using extrusion of the fluorescent marker Hoechst 33342 exploit the fact that stem cells extrude potentially noxious substances via the ABC transporter system. The dye-excluding population was shown to express CD133, CD44, and PAX2 to a higher extent than the bulk cells. Also they demonstrated a higher proliferative capacity and an enhanced capacity to form spheres. Flow-based cell sorting according to ALDH1 ac-
tivity using the ALDEFLUOR system has allowed enrichment of progenitor-like cells [19]. Gene expression analysis followed by histology identified markers such as myoferlin (fig. 1b) and vimentin (fig. 1c) along with CD133, CD24, and CD44, to mention a few. Another approach used the prolonged culture of renal epithelium by serial passage to select for a cell population that was shown to differentiate into renal tubules when injected subcortically and expressed the markers CD90, PAX2, and Oct 4. Finally, a method employing serial ischemic insults to cultivated kidney epithelium utilized the resistance to apoptosis associated with these cells [20, 21].

Marker Studies
CD133 is a protein of unknown function related to stem cells in other organs. In an early study, CD133+ cells were localized to the renal interstitium [22]. Later studies demonstrated a CD133+ progenitor cell population localized to the parietal epithelial cells of Bowman’s capsule [23]. Subsequently, CD133+ cells were also found to be scattered throughout the PT [19]. This cell population seems to share the expression of CD133, CD24, and CD44 [24, 25]. It was also reported that CD106 acts as a differential marker for progenitor cells localized in Bowman’s capsule as opposed to tubular origin, since CD106 is only being expressed by cells at the urinary pole of Bowman’s capsule. Less is known regarding the regulation of these cells in relation to both activation and stemness maintenance, but they have been shown to possess multilineage potential in vitro. Also, they have an activated antiapoptotic circuitry and integrate into injured kidney tubules upon IRI. With the probable exception of Bowman’s capsule, at present no clear-cut niche has been defined for this cell category. Very recently, however, a study pointed to TLR2 as a receptor involved in the activation of these cells. Also miRNA seems to be involved in cellular maintenance, where miRNA 1915 controls the expression of CD133 and PAX2 in these cells [24, 26].

Tubular Regeneration at Organ Level
Tubular regeneration occurs in anatomical proximity to other structures. As pointed out above, the PT and DT are localized in direct proximity to each other as well as to the vasa recta, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medulla operates under relative hypoxia, a circumstance adding anatomical complexity and the potential for cross talk in regeneration. This theme was recently reviewed by Gobe and Johnson [27]. Briefly, the medi...
haps the term state cell would be an alternative to stem or progenitor cell since the cells obviously seem to have a different metabolic state encompassing antiapoptosis, sphere-forming capacity, and a low mitochondrial count compared to the bulk epithelium. Dozens of markers are available today for these cells [13, 19, 23]. If the cells are indeed induced by injury, then we need to explain why there seems to be a perfect marker fit with the parietal cells of Bowman’s capsule. These cells cannot be regarded as chronically injured cells, and the matching markers point to a common cellular function. Regardless of their origin, another issue pertains to the choice of models (most renal injury models employ relatively acute and strong injury, such as in IRI models or unilateral ureteral obstruction models). We know little about how the tubular regeneration capacity changes during long-standing chronic injury. There are other interesting phenomena with a possible connection to the issue, such as ischemic preconditioning. It is known that the exposure of cells to hypoxic conditions in various organs, including the kidney, protects against subsequent ischemic insults, an effect mediated in part via the hypoxia-inducible factors HIF-1α and HIF-2α [30]. It is tempting to suggest that this mechanism may be connected to the scattered cells with progenitor-like properties. The concepts of induced plasticity and induced stem or progenitor cells are areas of intense investigation and maybe further research will prove the renal cellular heterogeneity to be a long sought-after in vivo counterpart of these phenomena mostly studied in an in vitro setting. This might bring a consensual end to the game of terms on the issue of tubular regeneration.

Acknowledgements

Dr. Karl Swärd is thankfully acknowledged for reading the manuscript and providing insightful comments and suggestions. This research was supported by the Marianne and Marcus Wallenberg Foundation, the National Association against Kidney Diseases, Regional ALF Funds, Malmö University Hospital Research and Cancer Funds, and the strategic Cancer Research Program BioCare.

References


