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CD133 and CD24 expression in renal tissue of patients affected by autosomal dominant polycystic kidney disease

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ABSTRACT

Background: Autosomal dominant polycystic kidney disease is a condition mainly characterized by the progressive development and enlargement of cysts in each kidneys. In this process a high rate of proliferation and apoptosis of tubular cells has been documented and interpreted as a futile attempt of tissue repair. In consideration of the role of stem cells in reparative processes we investigated the presence and localization of CD133 + CD24+ renal progenitors in renal ADPKD tissue and cells. Methods: Two normal kidneys and two ADPKD kidneys were examined. CD133 and CD24 expression was investigated by confocal microscopy and immunoblotting. Furthermore cystic isolated cells and cultured immortalized cells were characterized. Results: CD133 and CD24 have the same localization in ADPKD tissues and in normal kidneys: expression is restricted to a subset of epithelial cells (PEC) of Bowman’s capsule and to tubular cells in a focal and segmental pattern. Furthermore, in ADPKD tissue, cysts diffusely express CD133 and CD24. According to a quantitative analysis in ADPKD tissue CD133 + CD24+ cells are statistically more expressed in tubules (p < 0.001) and less expressed in the Bowman’s capsule (p = 0.0016) compared to the same localizations in control tissue. Conclusions: CD133 and CD24 antigens, typically expressed by renal epithelial progenitors, are more expressed in ADPKD tubules and highly expressed in ADPKD cysts. Whether CD133 and CD24 expression would signify renal progenitor recruitment or alternatively an expression pattern of the dedifferentiation of ADPKD cells remains unclear.

Keywords: Renal Stem Cells; CD133; CD24; ADPKD

1. INTRODUCTION

Autosomal Dominant Polycystic Kidney Disease (ADPKD) can arise from mutations in either PKD1 (85% of affected patients) or PKD2 (15% of affected patients) genes with clinically indistinguishable manifestations [1]. ADPKD affects over 1:1000 of the world wide population and it causes 1:3000 hospitalization. ADPKD is asymptomatic in about 50% of subjects with pkd genes mutated [2]. It is a systemic disease associated with renal failure, intracranial arterial aneurysms, cardiac valvular defects, colonic diverticulosis and cyst formation in other organs such as the liver, spleen and pancreas. Nevertheless ADPKD gives mainly renal manifestations: the patients develop renal cysts which progressively lead to disruption of renal parenchyma, interstitial fibrosis, cellular infiltration and loss of functional nephrons. Renal degeneration finally progresses to end-stage renal disease [1].

PKD1 (located on chromosome 16) codes for a large transmembrane protein of 460 KDa involved in the control of calcium permeation and cell-cell/matrix interaction. PKD2 (located on chromosome 4) codes for a calcium channel of about 110 KDa. The polycystin 1 and 2 (the proteins were coded by PKD1 and PKD2 genes) collaborate together in a functional complex [3,4].

Previous studies have demonstrated a high rate of proliferation and apoptosis in the cysts of ADPKD patients4 and this phenomenon has been interpreted as an unproductive attempt of tissue reparation. In this scenario a
likely involvement of renal progenitors can be hypothesized.

Sagrinati et al. identified, in human adult kidneys, a subset of Parietal Epithelial Cells (PECs) in the Bowman’s capsule that revealed self-renewal potential and high cloning efficiency. This subset of PECs exhibits co-expression of stem cell markers CD24 and CD133 and absence of lineage-specific markers. Under appropriate culture conditions, individual clones of CD24 + CD133 + PEC could be induced to generate mature, functional, tubular cells with phenotypic features of proximal and/or distal tubules [5].

ADPKD is characterized by expansion of cysts and tubular hyperplasia in the context of a sustained proliferative and apoptotic process. In this brief report we explored the pathologic features of presumptive renal progenitor cells in ADPKD renal tissue trying to explain the nature of their involvement in the disease: tissue repair or increase of cystic lesions.

2. RESULTS

2.1. Expression of CD133 and CD24 in Normal and ADPKD Kidney

In normal kidney at the glomerular level the confocal microscopic analysis shows that CD133 and CD24 are co-expressed by a subset of parietal epithelial cells (PEC) in the Bowman’s capsule (Figure 1: on line publication): CD24 has a generalized and diffuse presentation, CD133 is generalized and segmental. At tubular level the distribution of the expression of both CD133 and CD24 is focal and global. In the normal kidney specimen the overall intensity of CD24 + cells have the similar median value compared to CD133 + cells but the distribution of the intensity for the CD24 + cells is larger (Figure 2(a): on line publication). In the ADPKD kidney specimen the overall intensity of CD133 is higher, but not statistically significant, than the intensity of CD24 + cells (Figure 2(b): on line publication). It is interesting to note that at cellular level CD133 is polarized to the luminal side of the tubular cell surface as well to the urinary space side in positive PECs, whereas CD24 is not clearly polarized but shows a diffuse distribution at the membrane and cytoplasm level (Figure 1: on line publication).

Analysis of confocal microscopy of ADPKD kidneys revealed that CD133 and CD24 were expressed in a subset of glomerular (Figures 3(a)-(c)) and tubular (Figures 3(d)-(f)) cells and they showed the same tissue of normal kidney [5]: CD133 presents a generalized and segmental pattern while CD24 is generalized and diffuse in the Bowman’s capsule. The expression of both CD133 and CD24 is focal and global in the tubules.

The cysts of ADPKD samples showed a segmental co-expression pattern of CD133 and CD24. CD24 has a generalized distribution in the samples while CD133 is segmental. The cells of large cysts show a flattened morphology and a co-localization of CD133 and CD24 can be appreciated in some segments (Figures 3(g)-(i)).

The quantitative comparison between normal kidney and ADPKD kidney revealed an increased CD133 expression in tubules while glomerular expression appears
Figure 3. CD133 and CD24 expression in ADPKD samples. The first lane shows glomerular expression of CD133 (a) and CD24 (b). CD133 and CD24 are co-localized and co-expressed by a subset of PEC in the Bowman’s capsule (c). The second lane shows tubular expression of CD133 (d) and CD24 (e). CD133 and CD24 are co-expressed in these tubules (f). The third lane shows cystic expression of CD133 (g) and CD24 (h). CD133 and CD24 are co-localized and co-expressed by the cystic cells (i).

reduced in ADPKD samples (Figure 4). After image analysis Mann-Whitney test resulted statistically significant for these comparisons (ADPKD tubules vs Normal tubules: z = −7.681 p < 0.001; ADPKD Glomerula vs Normal Glomerula: z = 3.162 p = 0.0016). CD 24 expression in glomerular and tubules does not show quantitative difference between normal and ADPKD samples (Figure 4).

2.2. Expression of CD133 and CD24 in Cultured Cells

Cells deriving from the cystic wall of ADPKD tissue were scraped and expanded in culture. The epithelial nature of these cells was confirmed by cytocheratin immunohistochemical characterization (Figure 5 and Table 1: on line publication). In the subsequent analyses were used the clones of isolated cells that showed the complete expression of the cytokeratins (CAM 5.2, CK 7, MNF 116) usually expressed by cystic cells in tissue specimens.

Isolated cystic cells show a characteristic expression pattern for CD133 and CD24 (Figures 6(A)-(C): on line publication). CD133 is localized on the cellular membrane (Figure 6(A): on line publication) while CD24 is globally diffused in the cytoplasm (Figure 6(B): on line publication).
Table 1. Immunohistochemical analysis of cytokeratin expression in ADPKD tissue. (1) Focal and global distribution; (2) Focal and segmental distribution.

<table>
<thead>
<tr>
<th></th>
<th>CAM 5.2</th>
<th>CK 7</th>
<th>CK 20</th>
<th>MNF 116</th>
</tr>
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<tbody>
<tr>
<td>Bowman’s Capsule</td>
<td>-</td>
<td>+ (2)</td>
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<tr>
<td>Tubules</td>
<td>+ (1)</td>
<td>+ (1)</td>
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<td>Cysts</td>
<td>+ (2)</td>
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Figure 6. CD133 and CD24 expression in cystic cells in culture. CD133 is closely expressed by the cellular membrane (A). CD24 is expressed in the cytoplasm (B). CD133 and CD24 are co-expressed by the cystic cells (C).

The expression of CD24 is more diffuse while few cells are CD133+; only a small subset of isolated cystic cells are both CD133 and CD24 positive.

The same expression pattern is appreciable in HK-2 cell line while in HEK-293 cell line this peculiar pattern is lacking, both antigens presenting a cytoplasmatic distribution (Figure 7: on line publication).

By western blot analysis CD133 is highly expressed by cystic cells, HK-2 presents an intermediate level of expression while HEK-293 cell line shows an almost absent expression (Figure 8). Analogous amounts of CD24 can be detected in the three cellular populations (Figure 8).

3. DISCUSSION

The high rate of proliferation and apoptosis of cystic cells is a well known phenomenon in ADPKD [4]. It has been interpreted as an unsuccessful proliferative repair response of abnormal tubular cells degenerating into cyst formation. The abnormal behavior of the tubule in ADPKD patients originates from genetic ablation of a critical function of the polycystin proteins and primary cilia [6,7]. This system is supposed to act as a sensor for renal injury monitoring the urinary intratubular flux. According to this model an abnormal constitutive activation of this pathway in ADPKD leads to the futile (proliferative) attempt to repair an essentially non existing injury [4].

In this scenario where proliferation and renal tissue repair are active we could speculate a possible role for renal progenitors. The target of our report has been an explorative analysis of surface antigens related to stemness in ADPKD tissue. A multipotent progenitor cell population constituted by CD24+ and CD133+ parietal epithelial cells (PECs), in adult human glomerula has been described. In this report a murine model of acute renal failure caused by glyceral induced rhabdomyolysis has been generated and injection of CD24+ and CD133+ PECs differentiated into tubular cells significantly ameliorating kidney damage 5. CD133 is a pentaspan transmembrane glycoprotein suspected to have a role in maintaining stem cell properties by suppressing differentiation and promoting proliferation. CD133 expression is also associated to several types of cancer [8-10]. CD133 function is not clear but its presence on early and undifferentiated cells is suggestive of a growth factor receptor, and the presence of five tyrosine residues on the 50-aa cytoplasmic tail may indicate that the protein is phosphorylated in response to ligand binding and triggers a signal transduction [11]. CD24 is a sialoglycoprotein expressed on mature granulocytes [12], B and T lymphocyte subsets [13,14], normal and cancer stem cells...
Knowing this element will be important to understand their origin and their hypothetical migration. In our specimens we could appreciate the presence of a rich CD24+ and CD 133+ cell population. In ADPKD, the distribution of these cells is qualitatively similar to normal tissue with the notable exception of a significant expression in cystic cells. Quantitatively, CD 133+ cell population appears globally increased in ADPKD tissue. In particular, we found a statistically significant increase of CD 133 expression in tubes while glomerular expression appears reduced in ADPKD compared to control tissue. In this scenario, the speculative hypothesis of an active migration of progenitors from the glomerular “niche” [17] to the tube interstitial compartment is intriguing.

The abundant presence of CD133+ and CD24+ cells in ADPKD renal tissue sustains the hypothesis of an important role of this cell population in the disease. Inflammation and tubular damage are common features of ADPKD and CD133+ and CD24+ cells could likely play a role in the reparative process. Furthermore, the diffuse presence of these elements in the cystic wall suggests that they could contribute to cystic expansion and disease progression. Any way as far as now, no evidence is available to sustain the hypothesis that CD133+ and CD24+ cystic cells are progenitors migrated into the cystic wall rather than the alternative hypothesis that they represent dedifferentiated [18] tubular cells that express “de novo” CD133 and CD24.

The expression of CD133 and CD24 is maintained in cystic cells isolated and cultured in vitro. Furthermore, we evaluated the expression of these antigens in two tubular immortalized cellular lines, HEK293 and HK2.

4. CONCLUSION

In conclusion, we demonstrated an abundant population of CD133+ and CD24+ cells in renal tissue obtained from ADPKD patients. This cell population is present in the cystic wall and abundant in tubules, which conversely seems to be reduced in ADPKD glomerular in comparison to normal tissue. The role of this cell was not clear yet, but it could be interesting to evaluate more cases of ADPKD, in different stages (from asymptomatic to late stage), and clarify if CD133+ and CD24+ cells were prognostic factors for a poorer or better evolution of the disease. Additionally, we cannot univocally conclude that the CD133 + CD24 + cystic cells represent migrated renal progenitors rather than tubular dedifferentiated cells 18, but with a deeper biomarker analysis we could understand their origin and their hypothetical migration. Knowing this element will be important to understand more in detail pathophysiology of ADPKD and find new target to delay the progression of the disease.

5. MATERIAL AND METHODS

5.1. Antibodies

CD133 monoclonal (293C3) antibody for immunofluorescence was provided by Miltenyi Biotec (Miltenyi Biotec, Bergisch Gladbach, Germany). CD133 monoclonal (W6B9C1) antibody for Western blot was provided by Miltenyi Biotec (Miltenyi Biotec, Bergisch Gladbach, Germany). CD24 monoclonal (SN3) antibody was provided by Santa Cruz (Santa Cruz Biotechnology Inc., Santa Cruz CA, USA). PKD1 rabbit polyclonal (H-260) antibody was provided by Santa Cruz (Santa Cruz Biotechnology Inc., Santa Cruz CA, USA). Goat anti mouse IgG2b-Alexa 568 conjugate, goat anti mouse IgG2b-Alexa 633 conjugate, goat anti mouse IgG1-Alexa 568 conjugate, goat anti rabbit IgG-Rhod conjugate, goat anti mouse IgG-HRP conjugate were used as secondary antibodies. Cytokeratins monoclonal (CAM 5.2, CK7 [RCK105], CK20 [SPM140] and MNF 116) antibody was provided by Becton Dickinson (only CAM 5.2) (Becton Dickinson, CA, USA) and Abcam (the others) (Abcam, Cambridge, UK).

5.2. Tissues

Normal kidney fragments were obtained from a cadaveric kidney donor. The kidney was not suitable for transplantation due to the incidental discovery of a neoplastic nodule of 5 cm at the lower pole upon macroscopic examination. The pathologic examination diagnosed a clear cell carcinoma (grade 2 according to Fuhrman classification; T1bN0M0 according to the International Union Against Cancer classification); the lower pole was excised and the remaining parenchyma was used in the following procedures. The remaining parenchyma did not reveal any evidence of clear cell carcinoma.

Polycystic kidney tissue was obtained from 2 ADPKD patients underwent to nephrectomy in preparation for receiving renal transplantation. All procedures were preceded by the approval of our Local Ethical Committee on human experimentation and informed consent was obtained from the patients.

5.3. Culture

Human Kidney 2 cells (HK-2) were maintained in DMEM F-12 (Invitrogen/GIBCO, Cralsbad, California) supplemented with 10% FBS (PAA laboratories GmbH, Pasching, Austria), 2000 U/ml penicillin (PAA laboratories GmbH, Pasching, Austria), 2000 ug/ml streptomycin (PAA laboratories GmbH, Pasching, Austria), and 20 mM L-glutamine (Invitrogen/GIBCO, Cralsbad, California).

Human Embryonic Kidney 293 cells (HEK-293) were maintained in DMEM (Invitrogen/GIBCO, Cralsbad,
Cystic cells were obtained by fragments of renal cysts deprived of external membrane. Each cyst was picked up, was divided in some little fragments and were digested by collagenase (Invitrogen/GIBCO, Cralsbad, California) for one hours. Every pieces of a single cysts were seeded on a plate, previously treated by collagen type I (Invitrogen/GIBCO, Cralsbad, California). Cells harvested were, in part, seeded on chambers slides and the others were used to extract proteins.

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5.4. Immunofluorescence

Three-micrometer frozen kidney samples mounted on Super Frost Plus slides (Menzel-Gläser, Thermo Scientific, Waltham, MA) were cut by a cryostat (Leica 1720, Leica Mycrosystems, Heerbrog, Germany). The sections, or cellular spots, were fixed with PFA for 10 min, washed with PBS. Slides were coated (3% Bovine Serum Albumin in PBS buffer) for 1 h at 20°C. Slides were incubated by primary antibody for 1 h at 37°C, washed with PBS and successively incubated by secondary antibody for 1 h at 20°C. Primary and secondary antibodies were diluted in coating solution. Slides were closed with Vectashield® Mounting Medium with DAPI (Vector Laboratories GmbH, Pasching, Austria) and were trypsinized (Invitrogen/GIBCO, Cralsbad, California). Cells harvested were, in part, seeded on chambers slides and the others were used to extract proteins.

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5.5. Immunohistochemical

The same type of sample mounted on slides used for immunofluorescence analysis was handled for immunohistochemical evaluation. Slides were coated (3% Bovine Serum Albumin in PBS buffer) for 1 h at 20°C. Slides were incubated by primary antibody over night at 4°C, washed with PBS and successively incubated by secondary antibody (conjugated horse radish peroxidase) by for 30 min at 20°C. Hydrogen peroxide and diaminobenzidine (DAB) (SIGMA-ALDRICH, St. Louis, MO, USA) were used to obtain chromogen reaction. Olympus BX41 microscope was used to evaluate the samples.

5.6. Image Analysis

Images of immunofluorescence-marked kidney were sampled by an acquisition system composed by Olympus BX41 microscope, Olympus XC30 and CELL B 3.0 analysis image processing (Olympus Europa GmbH, Hamburg Germany). Images were saved in tagged image file format (.tiff). The files were analyzed with Inspector Matrox (Matrox Electronic System Ltd., Quebec Canada). ROI (region of interest) defines a portion of image that contains areas to analyze. ROI of each renal component (tubules, glomeruli and cysts) were sampled and converted in a list of single grey tone amount, from white (highest value) to black (zero). Background fluorescence was obtained sampling spot of negative structure, such mesangium or negative tubules, in every image.

5.7. Statistical Analysis

Total brightness/positive area ratio was calculated for each ROI. The following analysis of expression were performed: CD24+ in tubules of normal tissue, CD24+ in glomeruli of normal tissue, CD24+ in tubules of ADPKD samples, CD24+ in glomeruli of ADPKD samples, CD24+ in cysts of ADPKD samples, CD133+ in tubules of normal samples, CD133+ in glomeruli of normal tissue, CD133+ in tubules of ADPKD samples, CD133+ in glomeruli of ADPKD samples, CD133+ in cysts of ADPKD samples. Normal and ADPKD tissue components were compare by Mann-Whitney test. Results were considered statistically significant for p < 0.01. All statistical analysis were performed with the software SPSS Statistics 19.0.1 (IBM corporation, New York, USA).

5.8. Western Blot Analysis

Total proteins were extracted from cellular pellets with Triton X-100 based lysis buffer. Proteins were loaded on SDS-PAGE (3% stacking, 7.5% resolving), previously denatured with 4X Reducing Loading Buffer (200mM TrisHCl, 20% β-Mercapto Ethanol, 8% SDS, 0.4% Bromoefenol Blue, 40% glycerol). After SDS-PAGE run and transblot, nitrocellulose membranes were previously incubated by antibodies anti-CD133 and anti-CD24 and subsequently by horse radish peroxidase conjugated secondary antibodies. The detection was obtained by exposition of radiological plates to peroxidase substrate activated by the enzyme.

6. DISCLOSURES

The authors certify that there is no conflict of interest with any financial organization regarding the material discussed in the manuscript.
7. ACKNOWLEDGEMENTS

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