BALB/c-\(\text{Fcgr2b}^{-/-}\)-\(\text{Pdcd1}^{-/-}\) mouse expressing anti-urothelial antibody is a novel model of autoimmune cystitis

Yoshio Sugino1, Nobuyuki Nishikawa1, Koji Yoshimura1, Sadako Kuno2, Yukio Hayashi3, Naoki Yoshimura3, Taku Okazaki4, Akihiro Kanematsu1 & Osamu Ogawa1

1Department of Urology, Kyoto University, Kyoto, Japan, 2Department of Urology, National Center of Neurology and Psychiatry, Musashi Hospital, Tokyo, Japan, 3Department of Urology, University of Pittsburgh School of Medicine, Pittsburgh, USA, 4Division of Immune Regulation, Institute for Genome Research, The University of Tokushima, Tokushima, Japan.

We report the impact of anti-urothelial autoantibody (AUAb) on urinary bladder phenotype in BALB/c mice deficient of the FcγRIIb and PD-1. AUAb was present in serum samples from approximately half of the double-knockout (DKO) mice, as detected by immunofluorescence and immunoblots for urothelial proteins including uroplakin IIIa. The AUAb-positive DKO mice showed degeneration of urothelial plaque and umbrella cells, along with infiltration of inflammatory cells in the suburothelial layer. TNFa and IL-1β were upregulated in the bladder and the urine of AUAb-positive DKO mice. Voiding behavior of mice was analyzed by the Voided Stain on Paper method. 10-week-old and older AUAb-positive DKO mice voided significantly less urine per void than did wild type (WT) mice. Furthermore, administration of the AUAb-containing serum to WT mice significantly reduced their urine volume per void. In summary, this report presents a novel comprehensive mouse model of autoimmune cystitis.

An autoimmune mechanism has been postulated as a candidate cause for non-bacterial inflammatory diseases of the bladder such as interstitial cystitis, eosinophilic cystitis and lupus cystitis in human1–3. However, direct mechanistic association between autoimmune processes and functional changes in the bladder has not been established clinically. In experimental model, administration of exogenous molecule has been shown to elicit inflammatory change in the bladder4. Further, recent studies described the mechanism of autoimmune responses in the bladder by analyzing transgenic mice that constitutively express a self-antigen in the bladder epithelium, but did not elucidate whether the autoantibody could be pathogenic for bladder function in behavioral level or not5,6. In short, role of autoimmune processes in bladder pathophysiology is yet incompletely elucidated.

During the storage of urine in the bladder, umbrella cells, which are located on the surface of the bladder epithelium (urothelium), play an important role in the bladder barrier function7. Uroplakin (UPK) Ia, Ib, II, and IIIa are components of urothelial cell plaques, whose function is to reduce the transcellular passage of water through the urothelium8. Mice defective of UPKs show elevated water permeability in the urothelium9, vesicoureteral reflux, hydronephrosis10 and decreased bladder capacity11, demonstrating the importance of these barrier molecules in normal bladder function. Recent study showed that immunization with recombinant mouse UPK II might cause bladder specific inflammation and urinary frequency, though this study failed to describe any histological changes in urothelium, which includes UPK II proteins12.

The BALB/c mouse deficient in low affinity type IIb Fc receptor for IgG (FcγRIIb or Fcgr2b) and programmed cell death 1 (PD-1 or Pdcd1-1) (BALB/c-\(\text{Fcgr2b}^{-/-}\)-\(\text{Pdcd1}^{-/-}\) mouse / the DKO mouse) exhibits autoimmune phenotypes including autoimmune dilated cardiomyopathy13 and gastritis14. Interestingly, serum samples derived from the DKO mice also react to the urothelium and to UPKIIAa, and hydronephrosis is reported as a phenotypic consequence of anti-urothelial autoantibody (AUAb) due to periureteral inflammation14. As an extension of this study, we hypothesized that ablation of urothelium by the autoantibody should also impair bladder function. Therefore we analyzed the bladder of DKO mice to examine impact of the anti-urothelial antibody (AUAb), using biological and physiological techniques including a method reported as “Voided Stain on Paper (VSOP) method”15, and demonstrate here a novel comprehensive mouse model of autoimmune cystitis.
Results

Expression of AUAb in DKO mice. The presence of AUAb in serum samples was detected using mouse serum as primary antibodies for immunofluorescence (IF) of WT mouse bladder (Supplementary Fig. 1A) and immunoblotting for UPKIIIa, a component of urothelial plaque (Supplementary Fig. 1B)14. AUAb in serum of the DKO mice could be detected as early as 7 weeks of age by the IF. At 10-week-old, approximately half of DKO mice (7/13; 53.8%) expressed AUAb (Supplementary Fig. 1A right), but no WT mice did (0/10; 0%, Supplementary Fig. 1A left). In addition, the DKO mouse serum with positive immunoreactivity for the urothelium by IF, also recognized uroplakin 3A (UPKIIIa) by immunoblotting, indicating that UPKIIIa is one of the target proteins of AUAb (Supplementary Fig. 1B).

These AUAb-positive DKO mice remained positive until 16-week-old, while no new AUAb-positive mice appeared after 10-week. The positive rate of AUAb in the serum samples from DKO mice varied by the colony as shown in the former and current studies (35.3% and 53.8%, respectively)14.

DKO mice expressing AUAb have degenerated urothelium. We then evaluated the impact of AUAb on bladder morphology, by comparing wild-type mice, DKO mice with AUAb (AUAb-positive DKO mice), and those without AUAb (AUAb-negative DKO mice). There was marked disarrangement of urothelial plaque on the surface of the bladder of AUAb-positive DKO mice, but not in WT or AUAb-negative DKO mice (Fig. 1A–I). Hematoxylin and eosin (H&E) staining, and more prominently scanning electron microscopy (SEM), showed a disarranged cellular structure of the bladder epithelium in AUAb-positive DKO mice at 16-week-old compared to WT mice or AUAb-negative DKO mice (Fig. 1A, B, and D, E). The surface of bladder epithelium from AUAb-positive DKO mice exhibited poorer plaque formation, exposing bare surface of smaller epithelial cells beneath it (Fig. 1E). This morphological difference correlated with weaker staining of UPKIIIa in the bladders of AUAb-positive DKO mice compared to bladders from WT or AUAb-negative DKO mouse (Fig. 1G, H, and I).

These morphological features showed a close similarity to that of UPKIII KO mice reported by Hu et al.15, suggesting that AUAb may cause immunological ablation of UPKIII and apical cell layer of the urothelium.

Infiltration of inflammatory cells at the suburothelial layer of AUAb-positive DKO mice. The other remarkable morphological feature in the AUAb-positive DKO mice was massive infiltration of inflammatory cells in the suburothelial layer, which was never observed in WT or AUAb-negative DKO mice (Fig. 2A–C). There was marked infiltration of c-kit-positive inflammatory cells into the suburothelial layer of AUAb-positive DKO mice (Fig. 2D–F). The c-kit-positive cell count was significantly elevated in AUAb-positive DKO mice compared with AUAb-negative DKO or WT mice (Fig. 2G). Although c-kit has been known as a marker for mast cells16, these c-kit-positive cells did not have typical morphological feature of mast cells. There were also inflammatory cells positively stained for CD4 (T cell marker), CD11c (dendritic cell marker) and B220 (activated B cell marker) (Fig. 2I, K, M), while immunoreactivity for F4/80, which is the marker of macrophage was not elevated (Fig. 2O).

Expression of inflammatory cytokines in the suburothelial layer. In addition to morphological analyses, we analyzed the inflammatory process by identifying inflammatory factors using PCR analyses. TaqMan low-density array (TLDA) showed that mRNA expression levels of six molecules (Il2ra, Tnf, Icos, Il1b, Socs1, Vcam1) were significantly upregulated in the bladder of AUAb-positive DKO mice, as compared to WT mice (Supplementary Fig. 2A: p<0.01). Meanwhile, ELISA assay for urinary cytokines revealed that the protein level of seven molecules (GM-CSF, TNFα, IL-1α, G-CSF, IL-1β, RANTES, MIP-1β) was significantly upregulated in the urine of AUAb-positive DKO mice, as compared to WT mice (Supplementary Fig. 2B: p<0.05). Consequently, two molecules, TNFα and IL-1β were elevated in both bladder mRNA and urine protein levels (Fig. 3A). Of these, TNFα is known as a major mediator of inflammatory response.

Figure 1 | DKO mice expressing AUAb have degenerated urothelium. Histological analyses of bladder tissue of WT (A, D, G), AUAb-positive DKO (B, E, H), and AUAb-negative DKO (C, F, I) mice are displayed. (A–C) Hematoxylin and eosin (H&E) stain showed smaller epithelial cells including umbrella cells in the bladder of AUAb-positive DKO mice (B) than WT (A) and AUAb-negative DKO mice (C). (D–F) Scanning electron microscopy (SEM) revealed epithelial plaque in the bladder of WT (D) and AUAb-negative DKO mice (F); however, the size of plaques was decreased in AUAb-positive DKO mice (E). (G–I) The intensity of staining for UPKIIIa was reduced in the urothelial layer of AUAb-positive DKO mice (H) compared to WT (G) and AUAb-negative DKO mice (I). Bars indicate 30 μm in A–C, G–I, 10 μm in D–F.
of inflammation and an important therapeutic target of autoimmune diseases such as rheumatoid arthritis or inflammatory bowel disease\(^1\)\(^7\),\(^1\)\(^8\),\(^1\)\(^9\),\(^1\)\(^0\), and is also reported to play a crucial role in the pathology of inflammatory diseases of the bladder in animal models of neurogenic cystitis\(^1\)\(^1\)\(^9\),\(^1\)\(^2\). In the AUAb-positive DKO mice, the \(\text{tnf} \) mRNA expression level was upregulated in the stomach and the bladder, suggesting that the inflammatory process is not systemic, but involved limited number of organs, against which specific antibody is expressed, the stomach\(^1\)\(^4\) and the bladder (Fig. 3B).

Another separate ELISA study replicated the finding that urine of AUAb-positive DKO mice contained a significantly higher level of TNF\(\alpha\) compared to WT mice (Fig. 3C: \(p<0.05\)). This elevated urine TNF\(\alpha\) (Fig. 3C) may not derive from the kidney, but should account for overexpression of this molecule in the diseased bladder, since it was neither elevated in the kidney nor in the serum, (Fig. 3B).

In summary, the elevation of the inflammatory mediators including TNF\(\alpha\) indicated the significance of inflammation process in the bladder of AUAb-positive DKO mice, which made us to focus on the phenotype in context of the bladder function.

**Presence of AUAb in the serum is associated with impaired bladder storage function.** To evaluate the behavioral consequence of the inflammation in AUAb-positive mice, we analyzed urination of the mice by a method reported as ‘VSOP method’ by us\(^1\)\(^5\). From 10 to 16-week-old, the AUAb-positive DKO mice (\(n=7\)) voided significantly less urine per void than did WT mice (\(n=10, p<0.01\)), while there were no significant differences between AUAb-negative DKO (\(n=6\)) and the WT mice (Fig. 4A upper). Because AUAb-positive DKO mice had lower body weight than WT mice, we also examined the data with correction by body weight, and confirmed that the urine volume per void of AUAb-positive DKO mice was still significantly smaller than that of WT mice (Fig. 4A middle). The significant difference was also noted when the urine volume per void was adjusted as the ratio against the urine volume per void in 4-week-old (urine volume per void index, Fig 4A lower). Urine volume per void of AUAb-negative DKO mice showed slight and transient decrease compared to WT, but did not reach statistical significance except on the weight-adjusted data at 10-week-old (Fig. 4A middle), suggesting that presence of AUAb had greater impact on the bladder function than the genetic modification itself.

Based on these findings, we examined whether AUAb-containing serum is a pathogenic factor by itself in animals without genetic alteration. Intraperitoneal administration of the AUAb-containing serum resulted in a significant reduction in urine volume per void as compared to the WT serum-treated mice (shown as urine volume per void index, Fig. 4B), but no reduction in body weight, demonstrating that AUAb inhibits the growth of the bladder capacity. The bladder from mice 1 week after last of 4 injections of AUAb-positive sera
Fig. 4C upper right) showed disarrangement of urothelial plaque, which was not observed in mice after injection of control serum (Fig. 4C upper left). The intensity of staining for UPKIIIa was reduced in the urothelial layer of the bladder of mice after injection of AUAb-containing serum (Fig. 4C lower left) compared to the bladder of mice after injection of control serum (Fig. 4C lower right).

**Discussion**

The present study demonstrates that BALB/c-Fcgr2b−/− Pkd1−/− mouse (the DKO mouse) expressing AUAb is a novel animal model for autoimmune cystitis. AUAb-positive DKO mouse is a novel disease phenotype exhibiting three distinct features: disarrangement of urothelial layer, inflammation in the suburothelial layer and reduced functional bladder capacity.

The association of AUAb and these three findings could be explained by various possible mechanisms. First intriguing association is found between disarranged urothelial layer and inflammation in the suburothelial layer. Infiltration of the inflammatory cells could be explained either as inflammation directly following antibody-antigen reaction in the urothelial layer, or as consequence after disruption of impermeable barrier function by the urothelium against ions, water and urea21, which entails infiltration of urine into interstitium of the bladder. The target of AUAb, uroplakin is known as a critical factor for urothelial barrier function7–9. Upk3a knockout mice have a degenerated leaky urothelium9, and Upk2 knockout mice show a frequent urination pattern with reduced urine volume per void11. Since AUAb-positive DKO mice show similar phenotypic changes such as degenerated urothelium and reduced urine volume per void in this study (Fig. 1 and Fig. 4A, B and C), alterations in the morphological and functional properties of AUAb-positive DKO mice could be attributable to degenerated, leaky bladder urothelium associated with AUAb.

The other interesting phenotype is reduction in functional bladder capacity of AUAb-positive DKO mice. Although these mice are genetically modified, the reduction of the capacity does not seem to be congenital, but most likely attributable to appearance

| Upregulated in PCR array of cDNA from the bladder tissue †† | Upregulated in cytokine array of the urine † |
|------------------------------------------------------------|------------------------------------------------|
| \( \text{Tnf} \) / \( \text{Tnf}α \)                      | GM-Csf                                         |
| \( \text{Il}1\beta / \text{Il}1\beta \)                    | \( \text{Il}1\alpha \)                         |
| \( \text{Il}2\alpha \)                                     | \( \text{G-Csf} \)                            |
| \( \text{Icos} \)                                         | \( \text{Socs}1 \)                            |
| \( \text{Socs}1 \)                                        | \( \text{Vcam}1 \)                            |
| \( \text{Rantes} \)                                       | \( \text{Mip}-1\beta \)                       |

† † p<0.05
†† p<0.01

Figure 3 | Higher level of TNFα is detected in the bladder and the urine of AUAb-positive DKO mice. (A) TLDA indicated that mRNA expression level of six inflammatory molecules was significantly upregulated in the bladder of AUAb-positive mice compared to WT mice (†† p<0.01). Urine protein assay also revealed that seven inflammatory proteins were significantly upregulated in the urine of AUAb-positive DKO mice compared to WT mice († p<0.05). IL-1β and TNFα were expressed higher in both. (B) TNFα mRNA was elevated in the stomach and bladder of AUAb-positive DKO mice (*p<0.05). (C) ELISA assay for mouse TNFα has revealed that AUAb-positive DKO mice urine samples contain significantly higher level of TNFα than WT mice (*p<0.05).
of the AUAb in the serum and conspicuous bladder pathology associated with it for three reasons. First, the capacity of the mice is similar until 4 week, after which AUAb appears. Second, there are distinct difference between AUAb-positive and -negative DKO mice. Last, the AUAb-positive serum alone could induce similar functional impairment in WT mice. The possibility of neurogenic bladder dysfunction caused by inflammation in the spinal cord seemed negative based on results of the histological analysis of spinal cord sections (data not shown), and the inflammatory process producing various cytokines including TNFα in the bladder itself seems to impair bladder function. In this process although its functional role is not determined, TNFα level seems to reflect the localized inflammatory process. It is likely that AUAb causes this process by antigen-antibody reaction, although precise following mechanism is unclear. It is rather unlikely that antibody-dependent cellular cytotoxicity could be the cause for the urothelial destruction, since histologically there is no infiltration of inflammatory cells into urothelium in spite of massive inflammation of the bladder, while the activation of complement pathway may be a subject for future study. However, as a limitation of our study, we cannot conclude that AUAb is the only causative factor for the bladder inflammation, as there may be additional factor in sera of AUAb-positive DKO mice causing cystitis, and we could not purify the immunoglobulin for sera transfer experiments for technical reason, although important cytokine such as TNFα was undetectable in the serum of AUAb-positive mice.

Figure 4 | AUAb-containing serum is pathogenic to bladder storage function. (A) Urine volume per void of female AUAb-positive DKO mice (●: n=7), female AUAb-negative DKO mice (▲: n=6), and female WT mice (■: WT, n=10) was recorded from 4-week-old to 16-week-old by the VSOP method. From 10 to 16-week-old, there were significant differences (**p<0.01) in urine volume per void between AUAb-positive DKO mice and WT mice, although there were no significant differences between AUAb-negative DKO mice and WT mice (upper panel). The data adjusted by body weight are shown in the middle panel and the data shown as urine volume per void index (adjusted at 4-week’s urine volume per void) are in the lower panel. There were also significant differences between AUAb-positive DKO (●) and WT (■) from 10 to 16-week-old (**p<0.05), although there were no significant differences between AUAb-negative DKO (▲) and WT (■) except the urine volume adjusted by body weight at 10-week-old (#p<0.05). (B) Serum samples from AUAb-positive DKO mice (●, n=5) and serum samples from WT mice (■, n=5) were injected into WT female mice intraperitoneally (300 μl/body at 4-, 7-, 10-week-old; arrow). The urine volume per void recorded by VSOP method revealed a significant decrease of urine volume per void by administration of the serum from AUAb-positive DKO mice, compared with the serum from WT mice (shown as urine volume per void index). Data are expressed as means ± SE for each group. (C) H&E staining of the mouse bladder after injection of AUAb-containing serum (upper right) showed the disarrangement of urothelial plaque, which was not observed in the bladder after injection of control serum (upper left). The intensity of staining for UPKIIIα was reduced in the urothelial layer of the bladder of AUAb-containing serum injected mice (lower left) compared to the bladder of control serum injected mice (lower right). Bars indicate 100 μm.
In behavioral level, the inflammation does not inhibit growth of the bladder itself, since the net wet weight of the retrieved bladders was not different between each group (data not shown). The other more likely possibility is that the inflammation makes animals feel urinary fullness or pain more keenly and frequently, as seen in the clinical patients with bladder inflammation, although there was no sign of bladder pain such as lower abdominal licking or freezing on the AUAb-positive DKO mice.

By any mechanism, the AUAb was shown to play a key role in inducing the bladder dysfunction seen in DKO mice, but translational potential of this model to human pathology is still elusive. There are features in this model resembling pathological processes of human disease like interstitial cystitis or lupus cystitis. For example, AUAb-positive DKO mice has increased number of c-kit positive cells and activated B cells in the bladder. C-kit has been known as a marker for mast cells16, and degranulation of mast cells results in urinary fullness or pain more keenly and frequently, as seen in the more likely possibility is that the inflammation makes animals feel the bladder itself, since the net wet weight of the retrieved bladders were different from clinical interstitial cystitis in human. Beside this, the most different point from the human pathology is the presence of autoantibodies against urothelium, which have not been reported in human patients. At the same time, it is also unlikely that AUAb exists in human at the detrimental level as seen in the DKO, since development of hydroureteronephrosis itself is rarely seen in non-bacterial inflammatory diseases of the urinary tract. We can predict from our findings that, if AUAb exists in the human bladder diseases, it should appear either in a extremely limited number of severest cases, or just transiently at a lower level than seen in DKO mice.

Experimentally, the pathophysiology of AUAb-positive DKO mice presents a novel animal model of bladder pathology; bladder inflammation associated with autoantibody and behavioral phenotype. Our system is distinct from reported autoimmune cystitis models in several aspects1-6,12. The reported models failed to show either change in behavioral phenotype6, histological changes in the urothelium32, or a characterized pathogenic factor as AUAb4, all of which are seen in our AUAb-positive DKO mice. Moreover, the appearance of AUAb does not require stimulation of exogenous material such as ovalbumin for sensitization, showing that defective tolerance mechanism itself is enough to induce spontaneous sensitization against urothelium, generating potent autoantibody to alter the micturition behavior. In summary, the AUAb-positive DKO mouse demonstrates that autoimmunity can induce significant impairment of the bladder function.

In conclusion, this report characterizes a novel comprehensive mouse model of autoantibody-associated cystitis, which provides various insights about establishment of the bladder autoimmunity.

Immunohistochemistry (IHC) and scanning electron microscopy. Bladder tissues were collected after emptying the bladder of mice by pressing their lower abdomen. The following antibodies were used: anti-UPIIIA (PROGEN, Heidelberg, Germany) for two hours at room temperature and visualized by F(ab’)2 fragment of goat anti-mouse IgG (Jackson Immuno Research, PA, USA) with enhanced chemiluminescent detection system (GE Healthcare, Buckinghamshire, UK).

RNA isolation and quantitative RT-PCR. RNA was isolated from mice organs using the RNeasy mini kit (Qiagen, CA, USA). Whole bladders were used. RNA (1 µg) was reverse-transcribed using the First-Strand cDNA synthesis kit (Amersham Biosciences, NJ, USA) following manufacturer’s protocol. Real-time PCR was performed using SYBR Green PCR Master Mix (Applied Biosystems, CA, USA) and the relative standard curve method, at 95°C for 15 seconds and 60°C for 1 minute. PCR efficiency was examined by serially diluting the template cDNA and the melting curve data were collected to the PCR specific band. None of the PCR product was detected in control samples in which reverse transcriptase was omitted. An internal loading control of glyceraldehyde-3-phosphate dehydrogenase (GAPDH) expression was used for all genes of interest. Results were quantified using the relative standard curve method. The primers used were as follow: GAPDH (452 bp), sense: 5’-ACCAAGTTGCTGATCCTAC-3’ and antisense: 5’-TTTCCACCACTGGTTGCTGA-3’, TNF-α (60 bp) sense: 5’-CCCTCACTCAGATCTCATCC-3’ and antisense: 5’-GTTAGAGGTGGCCTGACA-3’. Each PCR regimen involved 5 minutes of initial denaturation step at 95°C followed by 38 cycles at 94°C for 60 seconds, 60°C (TNFα) or 65°C (GAPDH) for 60 seconds, and 72°C for 1 minute and 30 seconds.

Urinary cytology assay. Inflammatory cytological concentrations in urine samples from AUAb-positive DKO and WT mice were analyzed by Bio-Plex Mouse Cytokine Group I 23-Plex (Bio-Rad, PA, USA) according to the manufacturer’s instruction. TNFα ELISA assay. TNFα concentration in mouse urine samples was analyzed by another ELISA assay, Mouse TNFα ELISA (Bioscience, CA, USA), according to the manufacturer’s instruction.

Voids Stain on Paper (V SOP) method. Voiding behavior of mice was analyzed as described previously33. Briefly, the mice were fed with 50 µl of distilled water via gastric tube and placed on a screen above the filter paper, and their urination was recorded over two hours. The recorded stain areas were copied to another paper and calculated to the volume by computer software Photoshop (Adobe, USA). Age-dependent changes in bladder volume were also evaluated, either as net volume, volume adjusted to body weight, or using the urine volume per void index calculated as the ratio of urine volume per void of 4,7,10,13 or 16-week-old mice against that of 4-week-old mice.

Pathogenicity test of the AUAb-containing serum. Test serum samples were pooled from 20 DKO mice at 16 to 20-week-old, and confirmed to have AUAb by IF. Control serum samples were pooled from 15 age-matched BALB/c wild-type mice. These serum samples were preserved at 80°C. The pooled serum (300 µl/body) was then injected intraperitoneally to WT female mice at 4,7,10 and 13-week-old. On the other session, the serum was injected on the WT mice at 4,7,10, and

Methods

Animals. Wild-type BALB/c mice were purchased from Clea Japan, Inc. (Tokyo, Japan). BALB/c−/− and BALB/c−/− mice were kindly provided by Drs. Tasuku Honjo (Kyoto University), Jeffrey V. Ravetch (Rockefeller University), and Toshiyuki Takai (Tohoku University)13,23. BALB/c−/− and BALB/c−/− mice were purchased from Clea Japan, Inc. (Tokyo, Japan).

Immunofluorescence (IF). The existence of AUAb in the serum of mice was examined by utilizing it as a primary antibody of IF for mouse bladder section. The bladder tissues were collected from wild-type BALB/c mice, which were snap frozen in the OCT compound. Cryosections were fixed with CytoFix (BD Biosciences, CA, USA) and incubated with x100 diluted serum from mice, followed by FITC labeled anti-mouse IgG Ab (Jackson ImmunoResearch Laboratories, Inc., PA, USA). Sections were examined with an Eclipse E1000M microscope (Nikon, Tokyo, Japan).

Immunoblotting. Mouse UPKIIIa partial cDNA (corresponding to W19 to A125) was cloned into pGE6-43 vector (GE Healthcare, Buckinghamshire, UK) and GST-fusion protein of mouse UPKIIIa (GST-UPKIIIa) was expressed and purified according to the manufacturer’s instruction17. The purified solution (20 µg) was boiled for 5 min in a SDS sample buffer solution and separated by SD PAGE on a 10% Tris-HCl mini gel and transferred onto a polyvinylidene difluoride membrane following standard methods. Membranes were incubated with x100 diluted mouse serum or mouse monoclonal antibody (mAb) against UPKIIIa (PROGEN, Heidelberg, Germany) for two hours at room temperature and visualized by F(ab’)2 fragment of goat anti-mouse IgG (Jackson Immuno Research, PA, USA) with enhanced chemiluminescent detection system (GE Healthcare, Buckinghamshire, UK).
13-week-old and the bladder tissues were collected one week after the last injection and evaluated histologically.

Statistical analysis. Results are expressed as means ± SE. Statistical comparisons between two experimental groups were performed by Student’s paired and unpaired t-test using Statview 5.0 (SAS institute, Inc., NC, USA), and P value < 0.05 was considered as significant.

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Author contributions
Drs. Sugino Y, Nishikawa N and Kameyama A have planned and performed the study, and prepared majority of the manuscript. Drs. Yoshimura K, Kuno S, Yoshimura N and Okazaki T gave us contributions to conception and design of the study. Dr. Ogawa O gave us a general guidance and support of the study as laboratory director. All authors reviewed the manuscript.

Additional information
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