Properdin Deficiency Impairs Phagocytosis and Enhances Injury at Kidney Repair Phase Post Ischemia–Reperfusion

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Properdin, a positive regulator of complement alternative pathway, participates in renal ischemia–reperfusion (IR) injury and also acts as a pattern-recognition molecule affecting apoptotic T-cell clearance. However, the role of properdin in tubular epithelial cells (TECs) at the repair phase post IR injury is not well defined. This study revealed that properdin knockout (P/KO) mice exhibited greater injury in renal function and histology than wild-type (WT) mice post 72-h IR, with more apoptotic cells and macrophages in tubular lumina, increased active caspase-3 and HMGB1, but better histological structure at 24 h. Raised erythropoietin receptor by IR was furthered by P/KO and positively correlated with injury and repair markers. Properdin in WT kidneys was also upregulated by IR, while H2O2-increased properdin in TECs was reduced by its small-interfering RNA (siRNA), with raised HMGB1 and apoptosis. Moreover, the phagocytic ability of WT TECs, analyzed by pHrodo Escherichia coli bioparticles, was promoted by H2O2 but inhibited by PKO. These results were confirmed by counting phagocytosed H2O2-induced apoptotic TECs by *in situ* end labeling fragmented DNAs but not affected by additional serum with/without properdin. Taken together, PKO results in impaired phagocytosis at the repair phase post renal IR injury. Properdin locally produced by TECs plays crucial roles in optimizing damaged cells and regulating phagocytic ability of TECs to effectively clear apoptotic cells and reduce inflammation.

Keywords: apoptosis, inflammation, ischemia–reperfusion injury, phagocytosis, properdin, repair

INTRODUCTION

Acute kidney injury (AKI) has high morbidity and mortality and also has a potential of developing chronic kidney disease (CKD) (1–3). Renal ischemia–reperfusion (IR) injury, a major cause of AKI, is characterized by circulatory disturbance, complement activation, tubular injury, interstitial inflammation, and eventual fibrosis (4, 5). Tubular epithelial cells (TECs) are most vulnerable to IR injury but also actively participate in repair through dedifferentiation, proliferation, and clearing...
injured cells by phagocytosis (6–8). However, these functions may be maladaptive in severe and repeated mild injury, resulting in persistent complement activation, inflammation, and fibrosis (9, 10).

Complement activation via the alternative pathway (AP) is key to induce renal IR injury in rodent models. The suppression of the AP promotor, complement factor B, using monoclonal antibodies (mAb) or genetic modification significantly preserved renal function and morphology in mice subject to IR injury (11, 12). Conversely, the depletion of the AP inhibitors, decay-accelerating factor (DAF) and CD59, also exacerbates renal IR injury (13).

Properdin, the only known positive regulator of the AP, mainly produced by inflammatory cells (14–17), stabilizes complement 3 (C3) convertase (C3bBb), thereby providing a C3b amplification loop, and then stabilizes C5 convertase (18, 19). Inhibiting properdin by either antimouse mAb or gene deletion in DAF and CD59 double-knockout (DAF−/−CD59−/−) mice ameliorated early renal IR injury at 24 h (20).

Properdin also functions as a pattern-recognition molecule (PRM) by binding to targets such as damaged cells or bacteria, independent of its main ligand C3b (21–24). Properdin may also bind to the luminal membrane of proximal TECs in the proteinuric kidneys to mediate complement activation (25). However, the precise role of properdin in renal IR injury and repair and its potential function as a PRM in kidneys and TECs are incompletely defined. We hypothesized that properdin may have different roles at different stages of IR injury, either detrimental in the early injury stage by activating AP or beneficial at the late repair stage by facilitating phagocytic clearance of damaged cells.

In this study, properdin knockout (PKO) mice and counterpart wild-type (WT) mice were subject to 30 min of bilateral renal ischemia followed by 72 h reperfusion. TCMK-1 cells (mouse kidney cell line) and TECs isolated from PKO and WT mice, respectively, were exposed to hydrogen peroxidase (H2O2) to mimic the oxidative stress of IR injury. The role of properdin as a PRM and beyond was explored at the repair stage of kidneys post IR and in TECs subjected to IR-related injury, focusing on phagocytosis, apoptosis, and inflammation.

MATERIALS AND METHODS

Animal Model

Properdin-deficient mice were generated by site-specific targeting and maintained at University of Leicester (17). Male C57BL/6 WT and PKO mice aged 8–12 weeks were used in this study. All procedures were performed in accordance with the institutional guidelines reviewed by the Animal Welfare and Ethical Review Body and under the license approved by the UK Home Office (Project License, 70/8169 and Personal License, IA536CDE7). Mice were randomly divided into four groups: (I) WT sham (n = 4), (II) PKO sham (n = 5), (III) WT IR (n = 9), and (IV) PKO IR (n = 7). The animals were anesthetized by inhalation of 2.5% isoflurane in oxygen, and experimental procedures were optimized and refined based upon published protocols (26). The body temperature of animals during surgery was maintained at 36.5°C–37°C. Bilateral renal pedicles were exposed in the lateral position, dissected and clamped with a non-traumatic vascular clip for 30 min. After application of clips, the kidney was observed until patchy blanching developed and then replaced into the abdominal cavity. After removal of clips, gradual appearance of a normal pink color indicated kidney reperfusion.

After 72 h, the animals were bled by cardiac puncture and then sacrificed. The kidneys were harvested and snap frozen in liquid nitrogen or fixed in 4% (w/v) formaldehyde in 10% normal saline for further analysis. The animal experimental design is shown in Figure 1A.

Renal Function

Serum creatinine (SCr) and blood urine nitrogen (BUN) were determined using the QuantChrom™ Creatinine or Urea Assay Kit (BioAssay System, Hayward, USA). All procedures were performed according to the manufacturer’s instructions.

Histology

Paraffin-embedded kidney tissues were sectioned at 4 μm and stained by hematoxylin and eosin (H&E). The scoring of tubulointerstitial damage (TID) and counting of mitotic figures in the cortex was performed blindly to experimental groups by two independent researchers. The criteria of TID scoring included the loss of tubular epithelia, tubular cell vacuolation, tubular dilatation, luminal cast formation, and interstitial expansion (27). A histological score was assigned based on the percentage of affected areas in observed regions: 0, no damage <1%; 1, 1%–25%; 2, 26%–50%; 3, 51%–75%; and 4, >75%. In addition, mitotic figures were counted as condensed chromosomes aligned in metaphase in dividing cells (28). TID and mitotic figures were evaluated in 12 or 20 non-overlapping cortical fields at 200× or 400× magnification, respectively.

In Situ End Labeling of Apoptotic Cells

Apoptotic cells in kidneys were identified by in situ end labeling (ISEL) of fragmented DNA using terminal deoxynucleotidyl transferase-mediated uridine triphosphate provided in ApopTag® Peroxidase Kit (Merck Millipore, Darmstadt, Germany) (27). The ISEL+ cells were counted respectively in the tubular areas, tubular lumens, and interstitial areas of the renal cortex of 20 non-overlapping fields at 400× magnification.

Immunoblotting

Twenty-five micrograms of protein was separated by the electrophoresis of polyacrylamide denaturing gels and transferred onto polyvinylidene fluoride membrane. The primary antibodies, rabbit–antimouse caspase-3 (1:400, 9662, CST, Danvers, USA), rabbit–antimouse high-mobility group box-1 protein (HMGB1, 1:1,000, 3935, CST), mouse–antimouse erythropoietin receptor (EPOR, 1:1,000, PAB18350, Abnova, Taiwan), rabbit–antimouse properdin (1:1,000, AB186834, Abcam, Cambridge, USA) and...
mouse–antimouse β-actin (1:5,000, A5441, Sigma, Dorset, UK), were applied to the membranes overnight at 4°C, followed by the incubation of horseradish peroxidase-labeled secondary antibody (goat–antirabbit/mouse, K4063, DAKO) and developed by enhanced chemiluminescence (Thermo Fisher Scientific, Rockford, USA). The ratio of target protein to β-actin in volume density, as an endogenous loading control, was calculated for each detection, and then, the fold change of detected protein in the experimental group against the WT sham control was obtained as final results (27).

**Immunohistochemistry**

Antigen retrieval was performed using 20–40 µg/ml proteinase K (Sigma) digestion at 37°C for 15 min. The sections were then blocked with 10% goat serum and 0.5% bovine serum albumin (BSA) in 3% milk for 1 h at room temperature, followed by primary antibody labeling, mouse–antimouse F4/80 (1:100, ab100790, Abcam) or rabbit–antimouse erythropoietin receptor (EPOR, 1:800, PAB18350, Abnova) at 4°C overnight. The next day, DAKO secondary (K4063) was applied to the sections for 30 min. The antibody binding was then revealed by

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**FIGURE 1 |** P^K^ alleviated kidney structural damage at 24 h but aggravated renal functional and structural injury at 72 h post IR. (A) Schematic illustration of mouse experimental design. Bilateral kidney ischemia for 30 min and reperfusion for 24 h or 72 h were performed in male WT and P^K^ C57BL/6 mice. (B, C) SCr, BUN, and semiquantitative score of TID in both WT and P^K^ mice at 24 h were increased by IR; but only TID was improved by P^K^ (sham: n = 3–4; IR: n = 6–8). (D, E) At 72 h, SCr, BUN, and TID score in both WT and P^K^ mice were also elevated by IR, but all parameters were further increased by P^K^ (sham: n = 4–5; IR: n = 7–9). Data were shown as means ± SEMs, and analyzed by one-way ANOVA and LSD test. *p < 0.05; **p < 0.01. (F) Representative H&E stained images showing the cortex structure of the kidney at 72 h post IR. Scale bar: 100 µm.
transcription (Thermo Fisher Scientific). Real-Time Quantitative PCR

Total RNA was extracted using Trizol reagent (Thermo Fisher Scientific). Three micrograms of total RNA was used for reverse transcription (Thermo Fisher Scientific). One microliter of complementary DNA (cDNA) product was amplified with SYBR Green reaction system (Bioline, London, UK) containing 250 nM forward and reverse primers (Sigma, Table 1) at 95°C for 10 min followed by 40 cycles of 95°C for 20 s, 60°C for 30 s, and 72°C for 20 s. The level of glyceraldehyde 3-phosphate dehydrogenase (GAPDH) messenger RNA (mRNA) was detected as an endogenous control.

Double Labeling of Apoptotic TCMK-1 Cells With Properdin

TCMK-1 cells (American Type Culture Collection, Manassas, USA), mouse kidney epithelial cell line, were grown in Dulbecco’s modified Eagle’s medium (DMEM)/F-12 medium (Gibco, Carlsbad, USA) with 10% (v/v) fetal calf serum (Gibco), 2 mM L-glutamine (Gibco), 100 U/ml penicillin G, and 100 mg/ml streptomycin (Sigma, Dorset, UK) at 37°C in a 5% CO2 humidified atmosphere.

TCMK-1 cells were cultured in serum-free DMEM/F-12 medium (Gibco, Carlsbad, USA) and stimulated with 200 µM H2O2 for 24 h. The cells were then fixed and subjected to ISEL and developed with 3,3′-diaminobenzidine (DAB) chromogen (Vector). Antiprotein C8 antibody (ab186834, Abcam) diluted 1:100 was applied followed by biotinylated goat–anti-rabbit IgG (1:300, BA-1000, Vector) and alkaline phosphatase streptavidin (1:200, SA-5100, Vector). Binding was detected by Fast Red (Sigma) for 30 min and developed with 3,3′-diaminobenzidine (DAB, Vector, Burlingame, USA) and hematoxylin counterstaining.

For double labeling of properdin and F4/80 in the kidneys of WT mice, proteinase K at 40 µg/ml was used for antigen retrieval at 37°C for 30 min. After the staining of F4/80 was obtained, the sections were then incubated with primary antibody rabbit–antimouse properdin (1:200, ABF185, Merck Millipore) at 4°C overnight. Afterwards, biotinylated secondary goat–antirabbit immunoglobulin G (IgG) was applied to the slides for 30 min at 37°C (1:300, BA-1000, Vector), followed by alkaline phosphatase streptavidin (1:200, SA-5100, Vector) for 30 min at 37°C, and then developed by Fast Red (Sigma) for 8 min.

Real-Time Quantitative PCR

Total RNA was extracted using Trizol reagent (Thermo Fisher Scientific). Three micrograms of total RNA was used for reverse transcription (Thermo Fisher Scientific). One microliter of complementary DNA (cDNA) product was amplified with SYBR Green reaction system (Bioline, London, UK) containing 250 nM forward and reverse primers (Sigma, Table 1) at 95°C for 10 min followed by 40 cycles of 95°C for 20 s, 60°C for 30 s, and 72°C for 20 s. The level of glyceraldehyde 3-phosphate dehydrogenase (GAPDH) messenger RNA (mRNA) was detected as an endogenous control.

Table 1

| Genes     | Primers (5′-3′)                          |
|-----------|-----------------------------------------|
| Properdin | Forward: TTCAOCCAGATGTGAGGAGT            |
|           | Reverse: GCTGACATTCTGAGAAGCTC           |
| GAPDH     | Forward: CCTGAAGAAACGGTGTTGGAAGTGT      |
|           | Reverse: AGAATGGAGAGTTCGTTTGAAAGTC      |
with 500 µl of E. coli bioparticles (0.5 mg/ml, suspended in DMEM/F12 medium) for 2 h. Non-phagocytosed E. coli bioparticles were removed, and the cells were trypsinized and resuspended. On a fluorescence-activated cell sorting (FACS) system (BD), a total of 10,000 gated cells were analyzed for the fluorescent intensity of FITC. Due to the different level of auto fluorescence in WT TECs and PKO TECs, a different threshold was set up for each phenotype. To make two phenotypes comparable, the fold change of positive cells or FITC fluorescent intensity of total cells was calculated against that of the control—E. coli group for each phenotype.

The primary cultured WT TECs and PKO TECs at passage 2 or 3 were also seeded onto glass coverslips precoated with poly-D-lysine (0.1 mg/ml; Sigma) and cultured in the above-described culture medium, either without serum or with 2% serum from WT or PKO mice without heat inactivation for 24 h. The cells were stimulated with 200 µM H2O2 for another 24 h. The TECs were fixed by 1% paraformaldehyde for 10 min and then by an ice-cold mixture of ethanol and acetic acid (2:1) at −20°C for 5 min, followed by ISEL staining. The number of ISEL+ cells, phagocytosed ISEL+ cells (normal TEC overlapping with ISEL+ cells), and total cells were counted in 20 random fields at 400× magnification.

**Statistical Analysis**

Data are expressed as mean ± standard error of the mean (SEM). One-way ANOVA was used to check the homogeneity of variance, followed by post-hoc least significant difference (LSD) test for multiple comparisons or unpaired Student’s t-tests for comparison between two groups, using SPSS Statistics Standard V26.0 software (IBM, New York, USA). p < 0.05 was considered statistically significant.

**RESULTS**

**PKO Increased Inflammation and Its Mediators in Kidneys Post 72-h IR**

PKO-associated more severe IR kidney damage at 72 h was further examined closely by assessing infiltrated inflammatory cells in kidneys by the immunostaining of F4/80, a marker of macrophages, and examining injury markers including 17 kDa active caspase-3 and HMGB1 in kidney homogenates using Western blot. F4/80+ macrophages in tubular lumina and tubulointerstitial areas were remarkably increased by IR compared to sham groups, and further elevated by PKO (Figures 3A, B). The expression of 17 kDa active caspase-3, an executor of apoptosis (32) and inflammation (33, 34), was increased by IR compared to sham controls of WT and PKO mice, and further raised in IR kidneys by PKO compared to WT (Figure 3C). The same change pattern was also revealed in HMGB1 protein (Figure 3D), a recruiter of inflammatory cells and activator of complement (35–38).

**IR Raised Mitosis and PCNA With PKO Increased EPOR Post 72-h IR**

As repair was initiated at 72 h post IR, the compensatory response under the absence of properdin was assessed by mitotic cells and the proliferative marker PCNA. Mitotic cells in H&E-stained sections, reflected kidney repair, were increased by IR in WT and PKO kidneys compared to the phenotype sham (Figure 4A), while a similar change was seen in PCNA protein detected by Western blot, without significant differences between PKO and WT (Figure 4B).

To further understand the role of PKO in the repair phase post IR, the expression and localization of EPOR, a part of innate repair receptor (39) EPOR/fkR and essential for the phagocytic function of macrophages (40), were evaluated by Western blot and immunostaining. EPOR was increased by IR in WT and PKO kidneys compared to respective sham controls, further elevated by PKO compared to WT (Figure 4C). The weak staining of EPOR was evenly distributed in the cytoplasm of TECs in the cortical areas of sham kidneys, greatly enhanced by IR, mainly localized around severely damaged tubules, in WT and PKO mice (Figure 4D). The level of EPOR was positively correlated with mitotic figures and PCNA (Figure 4E).

**Elevated Properdin in IR Kidneys Was Localized in TECs and Macrophages**

To observe the involvement of properdin in IR injury and whether properdin tags apoptotic inflammatory cells, the
expression of properdin mRNA and protein in IR kidneys was first examined and then followed by its double immunostaining with F4/80 proteins. Exons 3–7 of properdin DNA was absent in PKO mice (17), so properdin mRNA was detected by quantitative PCR (qPCR) using primers recognizing properdin inside the exons 3–7. The level of properdin mRNA was raised by IR compared to sham controls in WT kidneys (Figure 5A), in which properdin protein was also elevated, but absent in PKO kidneys (Figure 5B).

The immunostaining of properdin was mainly localized on the apical surface of TECs and raised by IR in WT kidneys (Figure 5C). Properdin and F4/80 double-stained cells with typical apoptotic morphological features were observed in tubular areas, interstitial areas, and tubular lumina in IR WT kidneys (Figure 5D).

**Properdin Tagged Apoptotic Cells Phagocyted by Healthy Cells**

To further examine the effect of properdin as a PRM on apoptotic renal parenchymal cells, the double staining of properdin and ISEL was performed in H2O2-treated TCMK-1
cells in vitro. Patchy properdin staining was observed mainly in the cytoplasm of cells, without any staining in negative control cells incubated with normal rabbit IgG (Figure 6A). Some properdin+ cells having typical apoptotic features, shrunken nuclei with halos and condensed cytoplasm, or properdin and ISEL double+ cells, and ISEL+ cells were adjacent to or phagocytosed by healthy cells.

The effect of properdin on TECs was further verified by siP in H$_2$O$_2$-treated TCMK-1 cells, in which siP downregulated properdin expression but further increased HMGB1 (Figure 6B) and the number of early and late apoptotic cells in contrast to the negative control (siNC, Figure 6C).

**FIGURE 3** | $P^{KO}$ increased the infiltration of inflammatory cells and the expression of injury markers post IR at 72 h. (A, B) Representative images of F4/80 immunostaining in the indicated groups, and the semiquantitative analysis, showed more F4/80+ cells in the interstitium and tubulointerstitium of $P^{KO}$ kidneys than WT controls (sham: $n = 4$; IR: $n = 6–7$). The images were taken from the cortical area of the kidney. Scale bar: 50 μm. (C, D) Seventeen kilodaltons active caspase-3 and HMGB1 protein in the kidney determined by Western blot was increased by IR and furthered by $P^{KO}$ (sham: $n = 4–5$; IR: $n = 7–9$). Data were shown as means ± SEMs and analyzed by one-way ANOVA and LSD test. *$p < 0.05$; **$p < 0.01$.

### Phagocytic Ability of TECs Reduced by $P^{KO}$

Lastly, whether properdin affects the phagocytic ability of TECs was assessed by flow cytometry using pHrodo E. coli bioparticles. The threshold of fluorescent intensity for WT TECs was set lower than that for $P^{KO}$ TECs, as $P^{KO}$ had higher auto fluorescent intensity (Control - E. coli, Figure 7A), which was not changed by H$_2$O$_2$ treatment (H$_2$O$_2$ - E. coli). The fold change of FITC fluorescent intensity in all gated cells was increased by H$_2$O$_2$ only in WT TECs compared to the controls but decreased by $P^{KO}$ (Figure 7A, i), which were also seen in the fold change of positive cells (Figure 7A, ii). Moreover, compared to WT TECs, $P^{KO}$ decreased the fold change
of FITC fluorescent intensity treated by H2O2 and also decreased the fold change of positive cells without H2O2.

Apoptotic TECs were approached, engulfed, and phagocytosed by adjacent healthy-appearing TECs (Figures 7B, C). The percentage of apoptotic cells against the number of total cells were increased by PKO compared with WT TECs, regardless of serum conditions (Figure 7D). However, the percentage of phagocytosed apoptotic cells against the number of apoptotic cells in the pKO TECs was lower than that in WT TECs under all serum conditions.

DISCUSSION
Properdin has diverse and disease-dependent effects that may be modified by the genetic background of animal models (20, 41).
In 24-h post renal IR mice, depleting negative regulators of the AP increased properdin and AP activation and TEC injury, whereas inhibiting properdin ameliorated renal IR injury (20, 42). Conversely, pKO aggravates C3 glomerulopathy in mice (43, 44). In addition, emerging evidence indicates that properdin tagging apoptotic/necrotic T cells leads to their uptake by phagocytes via C3b opsonization or independent of C3b (22). The present study using sole pKO mice demonstrates that properdin has beneficial effects at 72-h kidney repair stage post IR injury, which was attributed to properdin regulating the phagocytic activity of TECs apart from as a PRM optimizing damaged cells (Figure 8).

More severe damage in the pKO IR kidneys at 72 h was associated with increased apoptotic cells in tubular luminal areas, and elevated active caspase-3 and HMGB1. Caspase-3 executes apoptosis and activates inflammatory mediators, while HMGB1 activates the Toll-like receptor 4 (TLR4)/nuclear factor kappa B (NF-κB) signaling pathway (45) and cytokine/chemokine production (35–38). Pko resulted in persistent apoptosis and inflammation in the IR kidneys due to absence of properdin-assisted phagocytic clearance of apoptotic cells (22, 46, 47). The accumulation of apoptotic cells in the tubular lumen of pKO IR kidneys may represent an adaptive mechanism of direct

**FIGURE 5** | Properdin expression and localization in kidneys post IR at 72 h, as well as its co localization with F4/80+ macrophages. (A) The expression of properdin mRNA in kidneys analyzed by qPCR was raised by IR in contrast to the sham control in WT mice. (B) The expression of properdin protein in kidney homogenates detected by Western blot was increased by IR in WT mice but was not seen in pKO mice (sham: n = 3–4; IR: n = 7). Data were shown as means ± SEMs. Significance was assessed by unpaired t-test for qPCR result and one-way ANOVA and LSD test for Western blots. *p < 0.05. (C) Representative images of properdin immunostaining in the cortex of WT sham and IR kidneys, with the boxed area enlarged and properdin staining on the apical surface of tubular cells was pointed by an arrow. All scale bar represent 50 µm. (D) Properdin and F4/80 double-stained cells were visualized in tubular areas, interstitial areas, and tubular lumina (pointed by arrows), with the boxed area enlarged. Most of double staining positive cells had typical apoptotic features including reduced cell volume, and the cellular membrane becoming ruffling and blebbing were also shown. Scale bar: 50 µm (upper row) and 20 µm (lower row).
elimination, while recruited macrophages may phagocytose apoptotic cells and also penetrate through tubular epithelia into the lumen. Properdin-labeled apoptotic macrophages were shown in the different compartment of IR kidneys of WT mice in this study, thus rendering them susceptible to phagocytes in vivo.

EPOR is expressed on TECs, where it forms heterodimers with βcR (EPOR/βcR), an innate repair receptor (39), and ameliorates renal damage and promotes tissue self-repair following IR (27, 48). Distributed mainly around damaged tubules (49), EPOR maintains tissue homeostasis by inhibiting the proinflammatory function of macrophages and enhancing their phagocytic function (40, 50). Increased EPOR, therefore, may also represent the compensatory response of repair in PκO IR kidneys at 72 h.

FIGURE 6 | Properdin tagged apoptotic TCMK-1 cells and inhibiting properdin increased apoptotic cells and HMGB1 expression in H2O2-treated TCMK-1 cells. (A) Properdin protein was visualized in TCMK-1 cells after H2O2 treatment for 24 h (indicated by a circle and two arrows nearby), in which the staining of properdin was negatively controlled by normal rabbit IgG. ISEL+ apoptotic TCMK-1 cells and its colocalization with properdin protein were also seen (pointed by another two arrows, n = 3). Scale bar: 5 µm. (B) The expression of properdin and HMGB1 detected by Western blot was increased by H2O2 treatment for 24 h, whereas siP-reduced properdin expression but increased HMGB1 (n = 3). (C) The early and late apoptotic TCMK-1 cells detected by flow cytometry analysis were both increased by H2O2 treatment for 24 h and furthered by siP (n = 3). FITC-fluorescein was tagged to Annexin V, thus can reveal the binding of Annexin V to the cellular surface of apoptotic cells that exposed phosphatidylserine (PS). Propidium iodide (PI) passed leaky necrotic cells to stain the DNA. Data were shown as means ± SEMs and analyzed by one-way ANOVA and LSD test. siP, siRNA targeting properdin; siNC, negative control siRNA. *p < 0.05; **p < 0.01.

TECs serve as major local semiprofessional phagocytes (8) and express various complement components, with C3 and factors H, B, I, and D being the most predominant (42, 51). Here, properdin was expressed in renal TECs being protective in terms of limiting HMGB1 and apoptosis under oxidative stress. Properdin, released from TECs, binds to the glycosaminoglycan chains of cell surface proteoglycans in apoptotic TECs to facilitate healthy TECs uptake these cells (22). Effectively clearing apoptotic cells by phagocytosis is crucial to limit renal injury and at the same time promote repair/remodeling post IR (8, 52, 53), in which the process of opsonizing damaged cells was absent in PκO kidney resulting in higher level of apoptosis and severer damage.
Additionally, the effect of properdin on the phagocytic ability of TECs was detected by the uptake of fluorescence labeled *E. coli* particles, and oxidative-stress-induced apoptotic cells. Under unstressed and oxidative-stress condition, *pKO* significantly reduced the number of phagocytic cells, which indicates that properdin might affect the transformation of TECs to phagocytes. Moreover, *pKO* significantly decreased the fluorescence intensity of gated *pKO* TECs under oxidative stress, which was increased in WT TECs, indicating that properdin is essential for upregulating the phagocytic ability of TECs under stress.

**FIGURE 7** | Properdin contributes to the phagocytic efficacy of primary isolated TECs. (A) Flow cytometry analysis of phagocytosed FITC-labeled *E. coli* by primary isolated WT and *pKO* TECs with or without H2O2 treatment for 24 h (n = 3). The fluorescent intensity of FITC conjugated *E. coli* reflected the phagocytic function of TECs. (i) The fold change of the average intensity of FITC fluorescence among total cells, calculated against the control (−*E. coli*) group, was increased in WT TECs but decreased in *pKO* TECs by H2O2. (ii) The fold change of *E. coli* positive cell counts (P1 area) in each group against the control (−*E. coli*) group was increased in WT TECs by H2O2 but stayed low in *pKO* TECs of either the control or H2O2 groups compared to corresponding WT groups. (B) Representative images of apoptotic WT and *pKO* TECs detected by ISEL (*ISEL*+ cells indicated by arrows, n = 3). Scale bar: 50 μm. (C) Healthy WT TECs approaching, partially engulfing, and phagocytosing adjacent ISEL+ cells were demonstrated and indicated by arrows. Scale bar: 5 μm. (D) Semiquantitative analysis showed that the percentage of ISEL+ cells against the total number of cells was increased by *pKO* regardless of serum conditions, while the percentage of phagocytosed ISEL+ cells against the total number of ISEL+ cells was decreased by *pKO* after H2O2 treatment for 24 h (n = 3). Data were shown as means ± SEMs and analyzed by one-way ANOVA and LSD test. *p < 0.05; **p < 0.01.
PKO may affect the expression of PRM including properdin per se and pattern-recognition receptors (PRRs) on TECs. Apart from certain "eat me" signals expressed on damaged cells, the PRM further facilitates their recognition by phagocytes. Properdin might also be a constitutive "eat me" marker as it was expressed on the plasma membrane of TECs. For instance, the molecule of kidney injury molecule 1 (KIM-1) is one of the PRRs, expressed on TECs (as semiphagocytes) subjected to injury. KIM-1 as a receptor binds to its ligand, phosphatidylserine, as an "eat me" marker exposed on damaged cells (8). The evidence of reduced phagocytosis in PKO TECs indicates that properdin might not only tag apoptotic cells mediating phagocytosis but also directly affect the phagocytic ability of TECs. In addition, the phagocytic feature of WT and P^KO^ TECs was not affected by additional serum with or without properdin, indicating that only locally produced properdin was essential for phagocytosis. Taken together, this study demonstrates for the first time that properdin has a key direct role in the phagocytic ability of TECs.

There are certain limitations in this study. First, the interaction and mechanism of properdin, KIM-1 and EPOR on the phagocytosis of TECs, and macrophages in renal injury and repair are worthy of further investigation. Second, a time-course model of renal IR in P^KO^ mice will be useful to disclose the dynamic change and the precise role of properdin in injury, recovery, or chronic progression.
In conclusion, this study showed that lack of properdin had harmful effects on kidneys at the repair stage post IR injury, which was attributed to properdin not only osonizing damaged cells but also affecting the phagocytic ability of TECs to effectively clear apoptotic cells and subsequent inflammation. Therefore, a novel mechanism in IR injury and repair was found, in which properdin is crucial for phagocytosis in the repair stage after IR-induced injury.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding author.

ETHICS STATEMENT

All animal procedures were subject to institutional review by the Animal Welfare and Ethical Review Body and approved under UK Home Office Project License PPL 70/8169.

AUTHOR CONTRIBUTIONS

BY and CS designed and supervised the study. YW, ZZ, XZ, and HW carried out experiments. YW, ZZ, CS and BY analyzed the data and constructed the figures. RC supervised cell culture work. YW, BY, CS and NB wrote and revised the paper. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2021.697760/full#supplementary-material
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