Novel soluble guanylyl cyclase activators increase glomerular cGMP, induce vasodilation and improve blood flow in the murine kidney

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Background and Purpose: Generation of cGMP via NO-sensitive soluble guanylyl cyclase (sGC) has been implicated in the regulation of renal functions. Chronic kidney disease (CKD) is associated with decreased NO bioavailability, increased oxidative stress and oxidation of sGC to its haem-free form, apo-sGC. Apo-sGC cannot be activated by NO, resulting in impaired cGMP signalling that is associated with chronic kidney disease progression. We hypothesised that sGC activators, which activate apo-sGC independently of NO, increase renal cGMP production under conditions of oxidative stress, thereby improving renal blood flow (RBF) and kidney function.

Experimental Approach: Two novel sGC activators, runcaciguat and BAY-543, were tested on murine kidney. We measured cGMP levels in real time in kidney slices of cGMP sensor mice, vasodilation of pre-constricted glomerular arterioles and RBF in isolated perfused kidneys. Experiments were performed at baseline conditions, under L-NAME-induced NO deficiency, and in the presence of oxidative stress induced by ODQ.

Key Results: Mouse glomeruli showed NO-induced cGMP increases. Under baseline conditions, sGC activator did not alter glomerular cGMP concentration or NO-induced cGMP generation. In the presence of ODQ, NO-induced glomerular cGMP signals were markedly reduced, whereas sGC activator induced strong cGMP increases. L-NAME and ODQ pretreated isolated glomerular arterioles were strongly dilated by sGC activator. sGC activator also increased cGMP and RBF in ODQ-perfused kidneys.

Conclusion and Implication: sGC activators increase glomerular cGMP, dilate glomerular arterioles and improve RBF under disease-relevant oxidative stress conditions.
INTRODUCTION

Chronic hypertension leads to end-organ damage, which especially affects blood vessels and kidneys. End-organ damage goes along with gradual increase in proteinuria, impaired kidney function including reduced glomerular filtration rate (GFR) and the development of chronic kidney disease (CKD) (Griffin, 2017). Persisting chronic kidney disease often results in end-stage renal disease and ultimately the need for a kidney transplant (Brenner et al., 2001; Coresh, 2017). Currently, the first-line management of chronic kidney disease consists of inhibiting the renin-angiotensin-aldosterone system, which is known to play a pivotal role in chronic kidney disease development (Siragy & Carey, 2010). Although therapies inducing reduction of systemic blood pressure (BP) have shown beneficial effects on kidney function in patients with renal disease, these effects are not fully maintained in chronic therapy (Breyer & Susztak, 2016) or in patients with type-2-diabetes and advanced end-stage renal disease (Schievink et al., 2016). Recently, sodium-glucose-cotransporter 2 (SGLT2) inhibitors which are approved for the treatment of type-2-diabetes, have been demonstrated to prevent GFR decline in chronic kidney disease patients (Lin et al., 2019; Muskiet et al., 2017). However, despite reduction of cardiovascular risk by SGLT2 inhibitors in chronic kidney disease patients (Briasoulis et al., 2018; Vallon & Thomson, 2017), an initial GFR decrease under SGLT2 inhibitors was reported (Sugiyama et al., 2020). Thus, there is still a need for new chronic kidney disease therapies with a different molecular mode of action, which can overcome limitations of currently used drugs.

The cGMP signalling system is a central regulator of cardiovascular homeostasis and has great potential as a target for new effective pharmacological therapies. Recent studies illustrated that genetic variants in components of this pathway significantly influence BP and the risk of cardiovascular and renal disease (Emdin et al., 2018; Erdmann et al., 2013; International Consortium for Blood Pressure, Ehret et al., 2011; Maass et al., 2015). The soluble guanylyl cyclase (sGC) generates cGMP upon activation by NO. An impairment of the NO/sGC/cGMP pathway and a decline in cGMP contributes to the development vascular dysfunction. Enhanced NO signalling also was associated with a higher GFR as determined via cystatin C and creatinine (Krishnan et al., 2018; Ott et al., 2012). Genetic variants of sGC and cGMP-dependent protein kinase type I, a downstream effector of sGC, are causally associated with altered vascular structure and remodelling, and sGC gain of function is associated with a higher GFR and lower risk for chronic kidney disease (Emdin et al., 2018). Therefore, sGC-enhancing drugs might be beneficial for treating chronic kidney disease patients.

Recent studies indicated that oxidative stress, which is triggered by comorbidities in chronic kidney disease, like hypertension, diabetes or obesity, could be one of the drivers of kidney function decline in chronic kidney disease (Coppolino et al., 2018; Samarghandian et al., 2017; Sinha & Dabla, 2015; Su et al., 2019). Oxidative stress is leading to sGC oxidation and ultimately the loss of its haem group, resulting in the formation of haem-free apo-sGC. Apo-sGC can no longer bind and be stimulated by the endogenous ligand NO (Stasch et al., 2015). Thus, oxidative stress disrupts NO/sGC/cGMP signalling and, thereby, counteracts physiological regulation of kidney function by NO. sGC activators are small molecules that activate sGC in a haem- and NO-independent manner and, thus, have the potential to restore sGC/cGMP signalling under conditions of oxidative stress. Runcaciguat is a novel potent and selective sGC activator, which activates apo-sGC in vitro, ex vivo and in vivo (Hahn, 2018) and has great potential to prevent the decline of kidney function (Benardeau et al., 2020; Hahn et al., 2021). However, the potential kidney-specific mechanisms behind the renoprotective effects of runcaciguat are not well characterised. Therefore, we aimed to investigate where and under what conditions runcaciguat and its close analogue BAY-543 (Rühle et al., 2020) induce cGMP production in the kidney and how these sGC activators influence the diameter of isolated glomerular arterioles as well as renal blood flow (RBF) in isolated perfused whole kidney preparations.

Our data show that sGC activators increase the glomerular cGMP concentration under oxidative stress conditions. Moreover, sGC activators induce dilation of pre-constricted glomerular afferent and efferent arterioles in the absence of NO and presence of oxidative stress and they significantly improve RBF under oxidative stress conditions. These findings show that sGC activators have direct effects on the renal vasculature and counteract the oxidative stress-induced decline in RBF. Thus, sGC activators could represent a new therapeutic approach for the treatment of chronic kidney disease patients.
METHODS

2.1 | Materials

The sGC activator runcaciguat (BAY 1101042) (Hahn, 2018; Hahn et al., 2021) and its analogue BAY-543 (Rühle et al., 2020) were synthesised by Bayer AG (Wuppertal, Germany) and dissolved in DMSO (Sigma-Aldrich, Darmstadt, Germany). Structurally, runcaciguat and BAY-543 are closely related, differing only in one side group (Figure S1). In the previous studies cited above, their EC50 to activate purified sGC (11 nM for both compounds) and their ability to dilate pre-contracted rabbit arteries ex vivo (IC50 of 199 and 75 nM for runcaciguat and BAY-543, respectively) were found to be very similar.

The sGC oxidant 1H-[1,2,4]oxadiazolo[4,3-a]quinoxaline-1-one (ODQ) was from Axxora. N-nitroarginine methyl ester (L-NAME) was from Tocris (Minneapolis, MN, USA), angiotensin II (Ang II) and N-nitroarginine methyl ester (L-NAME) from Sigma-Aldrich, and the L- and N-type calcium channel blocker cilnidipine from Camyan Chemical (Ann Arbor, MI, USA). The NO donor diethylamine NONOate (DEA/NO) was from Axxora. (DEA/NO) was stored in 100 mM stock solutions in NaOH (10 mM) and diluted in perfusion buffer immediately before application. Due to its short half-life time (≈16 min at room temperature and pH 7.4), DEA/NO is well-suited to induce fast, transient cGMP increases during real-time cGMP imaging experiments. For analysis of isolated glomerular arterioles and perfused kidneys, the NO-releasing compound 5-nitroso-N-acetylpenicillamine (SNAP; Cayman Chemical) was preferred as it has slower kinetics of NO release (several hours, varies between tissues) as compared with DEA/NO which ensures sustained continuous NO release at 37°C in these experiments.

2.2 | Animals

In the present study, we used mice, because they are amenable to genetic modifications and murine kidneys represent a good model to analyse renal function in mammals. All animal procedures were performed in accordance with the Directive 2010/63/EU, the German Tierschutz-Versuchstierverordnung and the local authorities in Tübingen, Berlin and Regensburg. Animal studies are reported in compliance with the ARRIVE guidelines (Percie du Sert et al., 2020) and with the recommendations made by the British Journal of Pharmacology (Lilley et al., 2020). Mice were housed in groups of up to eight animals in open type III cages at 22°C and 50%–60% humidity in a 12 h light/12 h dark cycle with access to standard rodent chow (ssniff; Soest, Germany) and tap water ad libitum.

For real-time cGMP imaging, 8- to 18-week-old R26-CAG-cGi500(L1) mice were dissected from C57BL/6N mice (R26-CAG-cGi500(L1) mice, and the renal capsule was carefully removed in ice-cold carbogen-gassed Ringer buffer (127.0 mM NaCl, 2.5 mM KCl, 0.5 mM MgCl2, 1.1 mM CaCl2, 1.1 mM NaH2PO4, 26.0 mM NaHCO3, 20.0 mM d-glucose). The pH was adjusted by continuous gassing with carbogen. Kidneys were sectioned with a vibromate (VT1200, Leica, Buffalo Grove, IL, USA) at a thickness of 700 μm. Whole kidneys and slices were incubated in ice-cold carbogen-gassed Ringer buffer for up to 8 h until real-time cGMP imaging was performed as described below.

2.3 | Preparation of kidney slices

For the preparation of acute tissue slices, kidneys were dissected from R26-CAG-cGi500(L1) mice, and the renal capsule was carefully removed in ice-cold carbogen-gassed Ringer buffer (127.0 mM NaCl, 2.5 mM KCl, 0.5 mM MgCl2, 1.1 mM CaCl2, 1.1 mM NaH2PO4, 26.0 mM NaHCO3, 20.0 mM d-glucose). The pH was adjusted by continuous gassing with carbogen. Kidneys were sectioned with a vibromate (VT1200, Leica, Buffalo Grove, IL, USA) at a thickness of 700 μm. Whole kidneys and slices were incubated in ice-cold carbogen-gassed Ringer buffer for up to 8 h until real-time cGMP imaging was performed as described below.

2.4 | Real-time cGMP imaging

Förster resonance energy transfer (FRET)/cGMP imaging was performed as described previously (Thunemann et al., 2013). The set-up consisted of an upright Examiner.Z1 microscope (Zeiss, Oberkochen, Germany), a Yokogawa CSU-X1 spinning disc confocal scanner (Yokogawa Denki, Musashino, Japan), three diode lasers (445, 488 and 561 nm), three water immersion objectives (W N-ACHROMAT 10/0.3, W Plan-APOCHROMAT 20/1.0 DIC UV VIS-IR, W Plan-APOCHROMAT 40/1.0 DIC VIS-IR; all from Zeiss) and one air objective (EC Plan-NEOFLUAR 2.5/0.085; Zeiss). Yellow fluorescent protein (YFP) fluorescence of the cGMP sensor cGi500 was detected with a CCD camera (Spot Pursuit, Diagnostic Instruments, Sterling Heights, MI, USA) through a 525/50 nm emission filter after excitation with the 488 nm laser. For FRET-based imaging, the donor fluorophore, cyan fluorescent protein (CFP), was excited with the 445 nm laser, and a Dual-View laser (445, 488 and 561 nm), three water immersion objectives (W N-ACHROMAT 10/0.3, W Plan-APOCHROMAT 20/1.0 DIC UV VIS-IR, W Plan-APOCHROMAT 40/1.0 DIC VIS-IR; all from Zeiss) and one air objective (EC Plan-NEOFLUAR 2.5/0.085; Zeiss). Yellow fluorescent protein (YFP) fluorescence of the cGMP sensor cGi500 was detected with a CCD camera (Spot Pursuit, Diagnostic Instruments, Sterling Heights, MI, USA) through a 525/50 nm emission filter after excitation with the 488 nm laser. For FRET-based imaging, the donor fluorophore, cyan fluorescent protein (CFP), was excited with the 445 nm laser, and a Dual-View beam splitter (Photometrics, Tucson, AZ, USA) with 505 nm dichroic mirror, 470/24 nm and 535/30 nm emission filters was used for simultaneous acquisition of CFP and YFP emission. Signals were recorded with an electron-multiplying charged-coupled device (EM-CCD) camera (QuantEM 512SC, Photometrics) at a frame rate of 0.2 Hz and an exposure time of 0.2 s. The system was controlled by VisiView 4.0.0.12 (VisiRon Systems, Puchheim, Germany). Kidney slices were continuously superfused with carbogen-gassed Ringer buffer with or without drugs at a flow rate of 1 ml/min at room temperature. The custom-built superfusion system consisted of an fast protein liquid chromatography pump (Pharmacia P-500, GE
Healthcare, Chicago, IL, USA), fast protein liquid chromatography injection valves (Pharmacia V-7, GE Healthcare), a magnetic platform (Warner Instruments, Hamden, CT, USA), a superfusion chamber (RC-26, Warner Instruments), a mesh-assisted Slice Hold-Down (SHD-26H/10, Warner Instruments) and sample loops of different sizes (7 ml for ODQ; 2 ml for other drugs). To ensure that drug exposure was comparable between different tissue slices, the same drug volumes were applied for the same time span. A vacuum pump with adjustable vacuum (Laboport N86, KNF Neuberger, Hamburg, Germany) was connected to the system to constantly remove excess buffer.

2.5 | Image acquisition and post-processing

Online image acquisition was performed with VisiView 4.0.0.12 (Visitron Systems, Puchheim, Germany), and offline post-processing and analysis were performed with Fiji software (RRID:SCR_002285) (Rueden et al., 2017; Schindelin et al., 2012). Images were aligned in x/y dimension with the Fiji plugin MultiStackReg v1.45 (RRID: SCR_016098). For further evaluation, Excel (RRID:SCR_016137; Office 16; Microsoft, Redmond, WA, USA) and Origin 2019 (RRID: SCR_014212; OriginLab, Northampton, MA, USA) were used. CFP and YFP emission were used to calculate the CFP/YFP ratio. The relative CFP/YFP ratio change (black traces in the respective graphs; referred to as R (cGMP)), which correlates with the cGMP concentration change, was obtained by normalisation to the baseline recorded for ~3 min at the beginning of each experiment. This normalisation is necessary to account for variations in the basal fluorescence intensity between the preparations. For peak evaluation, R (cGMP) traces were smoothed according to the Savitzky–Golay filtering method (smoothing window = 30 points) and the Peak Analyzer tool of Origin was used to calculate the AUC for each signal.

2.6 | Dissection and perfusion of glomerular arterioles

Kidneys were removed and sliced along the corticomedullary axis. Afferent and efferent arterioles were prepared according to procedures detailed by Liu and colleagues (Liu et al., 2012). In short, afferent and efferent arterioles with attached glomeruli were isolated and transferred into a chamber assembled on the stage of an inverted microscope. Arterioles were perfused using a system of pipettes, which allowed to hold and perfuse the vessels. Both, afferent and efferent arterioles were perfused from the free end, so orthograde in case of afferent arterioles and retrograde in case of efferent arterioles. Perfusion pressure was 100 mmHg for afferent arterioles and 40 mmHg for efferent arterioles, and the perfusion rates were in physiological ranges. DMEM (DMEM/F-12, Gibco, Darmstadt, Germany) containing 0.1% BSA (Carl Roth, Karlsruhe, Germany) was used during vessel dissection. The same solution was present in the experimental chamber, whereas the DMEM perfusion solution which was applied to the arteriolar lumen contained 1.0% BSA.

2.7 | Protocols for perfusion of glomerular arterioles

Arterioles were allowed to acclimatise for 10 min after establishing the perfusion. Viability was tested by short-term application of KCl (100 mM). Only vessels which showed full and sustained constriction were used for subsequent experiments. Arterioles were treated for 15 min with the non-specific NOS inhibitor L-NAME to induce NO deficiency, or for 10 min with the sGC-oxidising substance ODQ and vehicle (DMSO), respectively. Ang II, which belongs to the strongest vasoconstrictors particularly in the renal vasculature (Patzk et al., 2001), was applied to pre-constrict the vessels. Here, Ang II was applied in ascending concentrations (2 min for each concentration). After application of the highest Ang II concentration, cumulative concentrations of runcaciguat or of the known vasodilators NO (in the form of SNAP; sGC-dependent) and cilnidipine (sGC-independent) were administered to the Ang II-constricted arteries for 2 min each. In the ‘L-NAME + vehicle’ group, DMSO was applied instead of runcaciguat in corresponding concentrations. To analyse the time course of runcaciguat-induced vasodilation, arterioles pretreated with L-NAME and pre-constricted with Ang II were subjected to a prolonged 10 min treatment with runcaciguat or vehicle.

2.8 | Measurement of arteriole diameters

Perfused arterioles were continuously displayed on the computer screen using a video camera and the respective software (Motacam 2.0, Motic Asia, Kowloon, Hong Kong). Luminal diameters served for the estimation of arteriolar tone and reactivity. For the assessment of concentration–response relationships, pictures were recorded at a frame rate of 1 Hz. Then, the vessel diameter in five pictures was averaged in steady state conditions for each concentration effect to limit the influence of movement artefacts. For long-term investigations of runcaciguat action (10 min), pictures were recorded at a frame rate of 0.1 Hz. Either way, the arteriolar diameters were measured using ImageJ2 (RRID:SCR_003070) (Rueden et al., 2017). For analysis of the Ang II concentration–response relationship, arteriolar diameters were normalised to the initial diameter after L-NAME/ODQ/vehicle and before Ang II application. For evaluation of the dilation of pre-constricted arterioles induced by runcaciguat, SNAP and cilnidipine, diameter changes were normalised to the total constriction induced by L-NAME/ODQ/vehicle + Ang II. Both normalisation procedures are typically applied in vessel physiology to limit the influence of intra- and inter-individual variations of arteriole diameters and thereby focus the evaluation on pharmacological effects. The part of the vessel with the strongest response was chosen for analysis.
2.9 | Preparation and analysis of isolated perfused whole kidneys

Mice were anaesthetised with an intraperitoneal injection of xylazine hydrochloride (10 mg·kg\(^{-1}\)) and ketamine HCl (100 mg·kg\(^{-1}\); betapharm Arzneimittel, Augsburg, Germany) and placed on a warmed table. Isolation and perfusion of mouse kidneys followed a published method (Schweda et al., 2003). In short, the abdominal cavity was opened by a midline incision, and the aorta was clamped distal to the right renal artery. The mesenteric artery was ligated and a metal perfusion cannula (0.8 mm outer diameter) was inserted into the abdominal aorta. Then, the aorta was ligated proximal to the right renal artery and perfusion was started in situ with an initial flow rate of 1 ml·min\(^{-1}\). The right kidney was excised, placed in a tempered humid chamber and perfused at constant pressure (100 mmHg). Finally, the renal vein was cannulated (1.5 mm outer diameter polypropylene catheter). The venous effluent was drained outside the humid chamber and collected for determination of venous blood flow. The basic perfusion medium supplied from a 37°C-tempered 200 ml-reservoir consisted of a modified Krebs–Henseleit solution containing amino acids (10 ml·L\(^{-1}\) Aminoplasmal B. Braun 10%), 8.7 mM d-glucose, 0.3 mM pyruvate, 2.0 mM l-lactate, 1.0 mM ketoglutarate, 1.0 mM l-malate and 6.0 mM urea. The perfusate was supplemented with 60 g·L\(^{-1}\) BSA, 10 μM L-1 vasopressin 8-lysine and freshly washed human red blood cells (10% haematocrit). Ampicillin (30 mg·L\(^{-1}\)) and flucloxacinil (30 mg·L\(^{-1}\)) were added to inhibit possible bacterial growth in the medium. To improve the functional preservation of the preparation, the perfusate was continuously dialysed against a 10-fold volume of the same composition, but lacking erythrocytes and BSA. For oxygenation of the perfusion medium, the dialysate was gassed with 94% O\(_2\)/6% CO\(_2\).

2.10 | Determination of renal blood flow and cGMP secretion

Perfusate flow from kidneys isolated as described above was calculated by collection and gravimetric determination of the venous effluent. After establishing a constant perfusion pressure (100 mmHg), perfusate flow rates stabilised within 12–15 min. Stock solutions of the indicated drugs were added to the perfusate. For evaluation, the amount of venous effluent was normalised to the perfusate flow under control conditions or after application of 30 μM ODQ as indicated in the respective graphs to exclude differences in baseline perfusion between the preparations. For determination of renal cGMP production, venous effluent was collected over a period of 1 min during four intervals along the study and its cGMP concentration was determined after acetylation of samples using a cGMP enzyme immunoassay Kit (Cat# 581021, Cayman Chemical, Hamburg, Germany). The cGMP secretion was calculated by multiplying cGMP concentration and perfusate flow.

2.11 | Data and statistical analysis

The data and statistical analysis comply with the recommendations of the British Journal of Pharmacology on experimental design and analysis in pharmacology (Curtis et al., 2018). Variances in group sizes within individual comparisons are due to exclusion of defective preparations and yields, which were lower than expected. Data are presented as mean ± SD or mean ± SEM as specified in the figure legends. P values <0.05 were considered significant. Randomisation and blinding were not applicable because the experimental setups demanded application of substances in distinct succession. Instead, analysis and evaluation were performed uniformly and were based exclusively on objective parameters.

Statistical analysis was performed with Origin 2019 (OriginLab, Northampton, MA, USA; FRET/cGMP imaging) or GraphPad PRISM 8 (RRID:SCR_002798; GraphPad Software, San Diego, CA, USA; analysis of isolated perfused kidneys). For not normally distributed data sets, statistical differences were analysed non-parametrically by Mann–Whitney U-test. In case of normally distributed data, statistical differences were analysed parametrically by Student’s t-test (equal variances) or Welch’s t-test (unequal variances). The DEANO concentration–response curve was calculated with the sigmoidal ‘DoseResp’ fitting function of origin.

Time- and concentration-dependent differences of diameter changes of arterioles were statistically assessed with the free software R (RRID:SCR_001905; version 3.6.3) (R Core Team, 2020) using the Brunner test (Brunner & Langer, 1999). Sample sizes subjected to statistical analysis were at least five animals per group (n = 5), with n = number of independent values. This test is a non-parametric counterpart of the two-factorial ANOVA and tests the main hypothesis of a global difference between two groups. It is appropriate for the comparison of serial repeated measurements without normal distribution.

2.12 | Nomenclature of targets and ligands

Key protein targets and ligands in this article are hyperlinked to corresponding entries in the IUPHAR/BPS Guide to PHARMACOLOGY http://www.guidetopharmacology.org and are permanently archived in the Concise Guide to PHARMACOLOGY 2019/20 (Alexander et al., 2019).

3 | RESULTS

3.1 | Renal glomeruli express a functional NO/cGMP signalling pathway

To measure the spatiotemporal cGMP dynamics in the kidney, we used transgenic cGMP sensor mice expressing the FRET-based cGMP indicator cGi500 (Russwurm et al., 2007; Thunemann et al., 2013).
Specifically, we used the R26-CAG-cGi500(L1) mouse line that expresses the cGMP sensor globally in all tissues. By FRET imaging of live kidney slices ex vivo, cGMP signals were recorded in real time in response to the endogenous sGC ligand NO and in response to the sGC activator BAY-543 alone and in combination with NO. The sGC-oxidising agent ODQ was used to mimic chronic kidney disease-related oxidative stress conditions. Slices were kept in carbogen-gassed Ringer buffer during preparation and measurements to keep the tissue vital.

Visual inspection of the sensor-derived fluorescence confirmed that the cGMP sensor was broadly expressed in the kidney of cGMP sensor mice (Figure 1a). Note that the yellow fluorescence seen in the photomicrographs indicates expression of the sensor protein but not the cGMP concentration. The latter is determined by ratiometric analysis of the sensor’s CFP and YFP fluorescence as described in Section 2.5. Individual glomeruli were selected as regions of interest and several glomeruli were averaged to quantify the cGMP concentration in the absence and presence of various drugs.

First, we tested the capacity of renal glomeruli to generate cGMP in response to the NO-releasing compound DEA/NO. DEA/NO concentration-dependently increased the cGMP concentration in the glomeruli of cGMP sensor mice with an EC_{50} of 4.4 ± 0.1 μM (Figure 1b,c). Application of ODQ (50 μM) slightly reduced the basal cGMP production and, as expected, abolished NO-induced cGMP generation (Figure 1b). As a control, atrial natriuretic peptide (ANP), which increases cGMP via simulation of the particulate guanylyl cyclase A, was applied to the kidney slices at the end of each experiment. ANP (100 nM) potently increased glomerular cGMP levels independent of ODQ application (Figure 1b,d), indicating that sGC-independent cGMP generation was not affected by ODQ and that the tissue was still vital at the end of each experiment. Together, these data demonstrated the presence of a functional and oxidation-
sensitive NO/sGC/cGMP signalling pathway in the glomeruli of murine kidneys. In subsequent FRET imaging experiments, 10μM DEA/NO was used as a concentration that induces non-saturated cGMP responses.

3.2 The sGC activator BAY-543 increases glomerular cGMP under oxidative stress

Next, we analysed modulation of glomerular sGC/cGMP signalling by the sGC activator BAY-543, a close analogue of runcaciguat. As expected from our previous results (Figure 1b,c), under baseline conditions (i.e. in the absence of ODQ), application of DEA/NO alone reproducibly induced an increase of cGMP in glomeruli (Figure 2a,c).

In the absence of ODQ, application of BAY-543 did not alter the cGMP concentration and did not affect NO-induced cGMP signals. Interestingly, in the presence of ODQ, BAY-543 (10 μM) increased the cGMP concentration to a similar extent as 10 μM DEA/NO (Figure 2b,c).

Under the same oxidative stress conditions, that is, in the presence of ODQ, NO no longer increase cGMP (Figure 2b,c), consistent with the findings reported in Figure 1b. Altogether, our data shown in Figures 1 and 2 indicated that the sGC activator BAY-543 stimulates cGMP production in mouse glomeruli under oxidative stress conditions, which have impaired NO-induced cGMP signalling.

3.3 Runcaciguat induces dilation of renal arterioles

To further explore the physiological effects of cGMP production induced by sGC activators on kidney function, we assessed their effect on the diameters of glomerular vessels in ex vivo experiments. Afferent and efferent glomerular arterioles were isolated from mouse kidneys, mounted on pipettes and perfused. The diameter of afferent and efferent arterioles were measured under resting conditions and in the presence of vasoactive agents. For both afferent and efferent arterioles, three groups of vessels were randomly assigned as ‘L-NAME + runcaciguat’, ‘ODQ + runcaciguat’ and ‘L-NAME + vehicle’, which correspond to the groups used for pretreatment with L-NAME or ODQ and subsequent application of runcaciguat or...
The perfusion conditions were strictly identical in these three groups. In one set of experiments, vessels were pretreated with the NOS inhibitor L-NAME (10^{-4} M, 15 min) to induce NO deficiency. Then, increasing concentrations of Ang II were added on top of L-NAME to pre-constrict the vessels, followed by vehicle or runcaciguat ('L-NAME + vehicle' or 'L-NAME + runcaciguat' group). Ang II concentration-dependently decreased the vessel diameter by approximately 60% of the initial diameter at concentrations between 10^{-8} M and 10^{-6} M in both afferent (Figure 3c) and efferent arterioles (Figure 3d). The absolute afferent and efferent arteriole diameters under resting conditions as well as before and after Ang II application were similar throughout the three different groups (Table 1). In addition, superimposition of the Ang II concentration–response curves produced from afferent (Figure 3c) and efferent arterioles (Figure 3d) before application of vehicle or runcaciguat showed similar sensitivity of the vessel preparations to Ang II.

Together, these results indicated a good reproducibility of the measurements and similar initial conditions for the subsequent application of vehicle or runcaciguat.

Addition of increasing concentrations of runcaciguat (10^{-12} M to 10^{-6} M) to the pre-constricted afferent (Figure 3e) or efferent arterioles (Figure 3f) for 2 min induced strong vasodilation. ODQ pretreated
afferent arterioles were significantly more sensitive to runcaciguat than L-NAME pretreated afferent arterioles (Figure 3e,). ODQ pretreated efferent arterioles also showed a trend towards increased sensitivity to runcaciguat as compared with L-NAME pretreated efferent arterioles, but the difference did not reach statistical significance (Figure 3f). The dilating effect of runcaciguat on L-NAME pretreated vessels started at significantly lower concentrations in efferent arterioles than in afferent arterioles but was not significantly different between ODQ pretreated efferent and afferent arterioles (Figure 3e, f). These observations suggested a greater sensitivity for sGC activator of efferent over afferent arterioles under NO deficiency. The highest applied dose of runcaciguat (10^{-6} M) induced significantly stronger dilation of L-NAME pretreated efferent than afferent arterioles (afferent: 110%, afferent: 49%, Figure 3e,f). Maximal dilatations did not differ significantly in vessels pretreated with ODQ (afferent: 91%, afferent: 83%, Figure 3e,f). The dilation induced by runcaciguat (10^{-6} M) in the presence of ODQ was comparable with the maximal dilation that could be achieved by application of the NO donor SNAP or the L- and N-type calcium channel blocker cilnidipine (Figure S2). Further experiments showed that SNAP-induced but not cilnidipine-induced dilation was efficiently blocked by ODQ (Figure S2). Together, these results indicated that runcaciguat is a potent vasodilator of glomerular arterioles under conditions of NO deficiency and oxidative stress.

To further evaluate the time course of sGC activator action, we evaluated how afferent and efferent arteriole diameters changed during prolonged 10 min bolus applications of runcaciguat (10^{-7} M). In these experiments, arterioles were pretreated with L-NAME (10^{-4} M) and pre-constricted with Ang II (10^{-12}–10^{-6} M, not shown). Again, the diameters before application of runcaciguat or vehicle were

| TABLE 1 | Diameters of glomerular arterioles at different time points during the experiment |
|----------------|---------------------------------|----------------|----------------|----------------|
|                | Afferent arteriole diameter (μm) | Efferent arteriole diameter (μm) |
|                | I (n = 7) | II (n = 7) | III (n = 7) | I (n = 7) | II (n = 7) | III (n = 7) |
| L-NAME + vehicle | 7.7 ± 2.8 | 7.3 ± 3.0 | 3.3 ± 1.7 | 7.7 ± 1.4 | 6.7 ± 1.6 | 3.4 ± 1.7 |
| L-NAME + runcaciguat | 9.1 ± 1.6 | 8.6 ± 2.1 | 3.2 ± 1.0 | 7.5 ± 1.2 | 7.0 ± 1.5 | 2.6 ± 2.0 |
| ODQ + runcaciguat | 7.9 ± 1.4 | 6.7 ± 0.7 | 2.8 ± 0.8 | 6.6 ± 1.1 | 6.2 ± 2.5 | 2.9 ± 1.1 |

Note: Afferent and efferent arterioles were pretreated with L-NAME (10^{-4} M; ‘L-NAME + vehicle’ and ‘L-NAME + runcaciguat’ groups) or ODQ (10^{-5} M; ‘ODQ + runcaciguat’ group) and pre-constricted with incremental concentrations of Ang II (10^{-12}–10^{-6} M). Shown are absolute diameters of afferent and efferent arterioles under resting conditions (I), after application of L-NAME or ODQ (II) and after addition of the highest Ang II concentration (III). An example for the procedure is shown in Figure 3a. Data represent mean ± SD. n values are indicated in parentheses.

(a) Time course of runcaciguat-induced AA dilation

(b) Time course of runcaciguat-induced EA dilation

**FIGURE 4** Time course of runcaciguat-induced dilation of angiotensin II (Ang II) pre-constricted glomerular arterioles. Glomerular arterioles were pretreated with L-NAME (10^{-4} M) and pre-constricted with Ang II (10^{-12} to 10^{-6} M), as shown before. Then, runcaciguat (10^{-7} M; ‘L-NAME + runcaciguat’) or DMSO (‘L-NAME + vehicle’) were applied and diameter changes of (a) afferent arterioles and (b) efferent arteriole were monitored over 10 min. Data represent mean ± SD. n values are indicated in parentheses. The colour code given in panel (b) corresponds to the experimental groups indicated in panel (a). *P < 0.05
similar (data not shown). The maximum vasodilation achieved 10 min after application of runcaciguat was 50% of constriction in afferent arterioles (Figure 4a) and 60% in efferent arterioles (Figure 4b). Maximal vasodilation and kinetics of diameter changes were not significantly different between afferent and efferent arterioles.

Overall, these data indicated a reduction in renal resistance triggered by the sGC activator runcaciguat under conditions of oxidative stress and NO deficiency. At the tested concentration range, runcaciguat induced dilation of afferent and efferent arterioles. The vasodilating effect of runcaciguat was stronger after pretreatment with ODQ than with L-NAME, suggesting that runcaciguat activates the sGC more potently under conditions of oxidative stress.

3.4 Runcaciguat increases cGMP and improves RBF in perfused mouse kidneys

To complement the cGMP imaging data generated in kidney slices (Figures 1 and 2) and vascular reactivity measurements with isolated glomerular arterioles (Figures 3 and 4), we went on to analyze RBF and cGMP production under close-to-native conditions in whole kidneys. To do so, mouse kidneys were isolated and perfused at constant pressure (100 mmHg) as described previously (Schweda et al., 2003). Once perfusion of the kidney had stabilised (i.e. after 20 min perfusion), ODQ was infused to mimic NO deficiency and oxidative stress as observed in chronic kidney disease patients (Elshamaa et al., 2011; Martens & Edwards, 2011; Stasch et al., 2015).

**FIGURE 5** Renal blood flow (RBF) and cGMP secretion in perfused mouse kidneys. Kidneys were isolated and perfused at constant pressure (100 mmHg). (a) RBF was analysed by measuring the amount of venous effluent. After stabilization of perfusion (i.e. after 20 min), ODQ (30 µM) followed by incremental concentrations of runcaciguat (0.01, 0.1, 1 and 10 µM) and S-nitroso-N-acetylpenicillamine (SNAP; 10 µM) were applied sequentially, while the venous effluent was continuously quantified. Filled circles indicate time points at which the cGMP concentration in the venous effluent was determined (n = 6). (b) The cGMP concentration in the venous effluent was determined via enzyme immunoassay and multiplied by the perfusate flow to determine cGMP secretion as a measure of intrarenal cGMP formation. (c, d) Similarly, the effects of SNAP (10 µM) in the absence or presence of ODQ (30 µM) on RBF (c) and on renal cGMP secretion (d) were determined (n = 5). Data represent mean ± SD. *P < 0.05; n.s., not significant; ctrl, control.
ODQ (30 µM) strongly reduced RBF to approximately 50% of control (Figure 5a). In line with this finding, ODQ also reduced the cGMP content in the venous effluent as compared with baseline conditions (Figure 5b). Reduction of RBF by ODQ plateaued roughly 10 min after the start of infusion. Under these oxidative stress conditions, runcaciguat (10 µM) increased the cGMP content from 1.21 ± 0.6 to 13.1 ± 0.7 pmol·min⁻¹·per g kidney (Figure 5b). At the same time, runcaciguat dose-dependently increased RBF to a level above the baseline recorded in the absence of ODQ at the beginning of the experiment (Figure 5a). After this increase in RBF by runcaciguat, addition of the NO donor SNAP did not significantly change cGMP production (13.1 ± 0.7 versus 15.3 ± 1.2 pmol·min⁻¹·per g kidney) (Figure 5b) and did not further change RBF (Figure 5a). In addition, we analysed the effects of SNAP on RBF and cGMP secretion under basal versus oxidative stress conditions (Figure 5c,d). Under basal conditions, SNAP potently increased renal perfusion and cGMP content, while it failed to do so in the presence of ODQ. In summary, these data showed that in the presence of ODQ mimicking chronic kidney disease-related oxidative stress conditions, runcaciguat increases the renal cGMP concentration and RBF in perfused mouse kidneys, whereas NO is no longer able to do so under these conditions.

4 | DISCUSSION

This study showed that sGC activators increase the glomerular cGMP concentration, induce dilation of pre-constricted glomerular arterioles and improve RBF under conditions of NO deficiency and oxidative stress. It is known that oxidation of sGC from its native to the haem-free form, apo-sGC, prevents binding of NO and thereby shuts down the NO/sGC/cGMP signalling pathway (Stasch et al., 2015). This leads to reduced production of cGMP, which is important for the regulation of body homeostasis and kidney function (Krishnan et al., 2018; Wang-Rosenke et al., 2008). As sGC activators act in an NO- and haem-independent manner, these compounds might be able to restore cGMP signalling under pathophysiological conditions involving oxidative stress. Because chronic kidney disease is associated with oxidative stress (Matsuda & Shimomura, 2013; Moon & Won, 2017) and oxidation of sGC might be a critical driver of chronic kidney disease, sGC activators represent a promising new option for chronic kidney disease treatment. However, the mode of action of sGC activators in renal tissue and specifically their impact on renal blood vessels is poorly understood.

To explore the spatiotemporal dynamics of NO- and sGC activator-induced cGMP signals in the kidney, we performed real-time cGMP imaging in kidney slices of transgenic cGMP sensor mice. To our knowledge, this is the first demonstration of cGMP signals in live kidney tissue. We showed that NO and sGC activator increase the cGMP concentration in glomeruli in the absence and presence of oxidative stress, respectively. Furthermore, runcaciguat induced dilation of glomerular arterioles and improved RBF under disease-relevant conditions of oxidative stress and NO depletion. Although the present study did not experimentally distinguish between specific renal cell types, the combined results strongly suggest that runcaciguat acts on apo-sGC in vascular smooth muscle cells of glomerular arterioles, leading to an increase of cGMP, activation of cGMP-dependent protein kinase type I and vasodilation. This model is consistent with the cellular distribution of sGC in the kidney. Using a highly specific antibody against the β1 subunit of sGC, expression of the protein was detected in renal vascular cells, including glomerular arterioles (Theilig et al., 2001). Expression data, however, do not show the functionality of sGC in a specific compartment. It is well known that enzyme activity is regulated by post-translational mechanisms including, in the case of sGC, protein oxidation. Our cGMP imaging data link sGC expression in glomeruli with respective enzyme activity in the absence and presence of oxidative stress. Because the R26-CAG-cGi500(L1) mice used in this study express the cGMP sensor in all cell types, we cannot exclude the possibility that cGMP signals measured in our FRET imaging experiments were also derived from non-vascular smooth muscle cells that express sGC, such as interstitial fibroblasts (Theilig et al., 2001). However, it is unlikely that these cells were involved in sGC activator-induced dilation of renal arterioles. To analyse which specific renal cell types are capable of generating cGMP in response to NO and sGC activators, additional experiments could be performed using transgenic mice with cell type-specific expression of the cGMP sensor (Thunemann et al., 2013). Furthermore, it would be interesting to analyse the effects of sGC activators on cGMP signals in chronic kidney disease kidneys.

Real-time cGMP imaging illustrated that BAY-543 induces cGMP generation in glomeruli specifically in the presence of oxidative stress. This finding complements previous in vitro experiments showing that apo-sGC has a higher sensitivity for sGC activators than native sGC (Schmidt et al., 2009). To further dissect the mechanism of sGC activator action on kidney function, we investigated the effects of runcaciguat on the diameter of murine glomerular afferent and efferent arteries. It is known that pre-glomerular resistance and glomerular haemodynamics contribute to the control of the murine GFR (Patzak et al., 2004). Under pathological conditions, such as hypertension and type-2-diabetes, an increase in vascular resistance provokes a reduction of vascular diameter and arterial remodelling in the kidney, leading to impaired vascular reactivity and kidney perfusion (Polichnowski et al., 2013; Touyz et al., 2018). In the present study, we showed that runcaciguat improves vascular reactivity of pre-constricted murine glomerular afferent and efferent arterioles under conditions of NO deficiency and oxidative stress. Runcaciguat induced a concentration-dependent relaxation of pre-constricted glomerular arterioles with an apparent higher sensitivity for ODQ over L-NAME pretreated arterioles. This finding is consistent with a predominant action of sGC activator under oxidative stress as measured in our cGMP imaging experiments with kidney slices. Interestingly, the stronger action of runcaciguat in efferent compared with afferent arterioles indicate an influence on the GFR. This must be investigated in further experiments, for example, via cannulation of the ureter during experiments on isolated perfused kidneys.

To validate the cGMP imaging and vascular reactivity data obtained in kidney slices and isolated arterioles in a more physiological
Within the scope of this study, the effect of sGC activators on the renal vasculature was analysed in detail. In addition to their vasodilative effects, sGC activators may also be renoprotective via antifibrotic, antiproliferative and anti-inflammatory effects on vascular and non-vascular compartments of the kidney. In this context, it is interesting to note that the sGC activator BI 703704 (Boustany-Kari et al., 2016) and the sGC stimulator praliciguat (Liu et al., 2020) inhibited the progression of diabetic nephropathy in obese ZSF1 rats in doses that did not alter BP, perhaps via suppression of inflammation and apoptosis in tubular cells (Liu et al., 2020). sGC activators may inhibit glomerular remodelling also via direct effects on vascular smooth muscle cells independent of vasodilation. It is well known that NO/sGC/cGMP signalling in vascular smooth muscle cells regulates vascular plasticity and remodelling during diseases like atherosclerosis (Lehnners et al., 2018).

In summary, our results indicate that sGC activators increase cGMP production in vascular smooth muscle cells in glomerular arterioles and, thereby, improve kidney perfusion under disease-relevant conditions of oxidative stress and NO depletion. The unique selectivity of sGC activators for oxidised apo-sGC has great potential to limit off-target effects on healthy tissue, where sGC is mainly present in its native reduced form (Sharkovska et al., 2010). Most of the comorbidities in chronic kidney disease are associated with increased oxidative stress burden, leading to endothelial dysfunction with low NO production as well as NO-unresponsive apo-sGC. It is an intriguing concept that sGC activators could overcome the pathophysiological blockade of the endogenous NO/sGC/cGMP in renal tissue. Thus, sGC activators like runcaciguat could provide a novel and effective chronic kidney disease treatment.

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