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# 28 Abstract

29 We examined the effects of exposure to chronic intermittent hypoxia (CIH) on 30 baroreflex control of renal sympathetic nerve activity (RSNA) and renal excretory responses 31 to volume expansion (VE) before and after intra-renal TRPV1 blockade by capsaizepine 32 (CPZ). Male Wistar rats were exposed to 96 cycles of hypoxia per day for 14 days (CIH), or normoxia. Urine flow and absolute Na<sup>+</sup> excretion during VE were less in CIH-exposed rats, 33 but the progressive decrease in RSNA during VE was preserved. Assessment of the high-34 35 pressure baroreflex revealed an increase in the operating and response range of RSNA and 36 decreased slope in CIH-exposed rats with substantial hypertension (+19mmHg basal mean arterial pressure, MAP), but not in a second cohort with modest hypertension (+12mmHg). 37 38 Intra-renal CPZ caused diuresis, natriuresis and a reduction in MAP in sham and CIHexposed rats. Following intra-renal CPZ, diuretic and natriuretic responses to VE in CIH-39 40 exposed rats were equivalent to sham. TPRV1 expression in the renal pelvic wall was similar in both experimental groups. Exposure to CIH did not elicit glomerular hypertrophy, renal 41 42 inflammation or oxidative stress. We conclude that exposure to CIH: 1) does not impair the low-pressure baroreflex control of RSNA; 2) has modest effects on the high-pressure 43 44 baroreflex control of RSNA, most likely indirectly due to hypertension; 3) can elicit 45 hypertension in the absence of kidney injury; and 4) impairs diuretic and natriuretic responses 46 to fluid overload. Our results suggest that exposure to CIH causes renal dysfunction, which 47 may be relevant to obstructive sleep apnea.

48

<sup>49</sup> Key words: Intermittent hypoxia, baroreflex, volume expansion, diuresis, TPRV1.

ADH	Anti-diuretic hormone
ANP	Atrial natriuretic peptide
AOPP	Advanced Oxidation Protein Products
AUC	Area under curve
CGRP	Calcitonin gene- related peptide
CIH	Chronic intermittent hypoxia
CPZ	Capsaizepine
DBP	Diastolic blood pressure
GFR	Glomerular filtration rate
HR	Heart rate
MAP	Mean arterial blood pressure
NOX	NADPH oxidase
OSA	Obstructive sleep apnea
ROS	Reactive oxygen species
RSNA	Renal sympathetic nerve activity
SBP	Systolic blood pressure
SOD	Superoxide dismutase
SP	Substance P
TRPV1	Transient receptor potential cation
	channel subfamily V member 1
UFR	Urine flow rate
VE	Volume expansion
IFN-Y	Interferon gamma
IL-1β	Interleukin 1 beta
IL-4	Interleukin 4
TNF-α	Tumor necrosis factor alpha

# 55 Introduction

56 Obstructive sleep apnea (OSA) syndrome is characterized by frequent interruption of 57 ventilation due to repetitive upper airway obstruction during sleep (35). People with OSA 58 have a three-fold higher risk of developing hypertension (1). Chronic intermittent hypoxia 59 (CIH) models the hypoxia and reoxygenation cycles experienced in OSA, due to recurrent apnea. There is evidence from studies in animal models and humans (25) that exposure to IH 60 61 elicits sympatho-excitation and hypertension, revealing that this characteristic feature of the 62 disorder is adequate to cause autonomic dysfunction, notwithstanding that other features of OSA, not modelled in CIH, such as hypercapnia and large intra-thoracic sub-atmospheric 63 pressure swings associated with airway obstruction, could also contribute to cardiovascular 64 dysfunction in OSA. Animal models of CIH develop sympathetic hyperactivity and diurnal 65 hypertension (28, 40, 41, 51). Moreover, exposure to CIH evokes detrimental processes in the 66 67 kidney such as inflammation, fibrosis, glomerulosclerosis and proteinuria (4, 5, 29, 45), which are markers for chronic kidney disease. 68

69 Fluid homeostasis is mainly regulated by the diuretic and natriuretic function of the 70 kidney. Fluid overload activates low-pressure cardiopulmonary baroreceptors causing a reflex 71 suppression of renal sympathetic nerve activity (RSNA) promoting natriuresis and diuresis 72 (33, 37). Previous studies have revealed blunted low-pressure baroreflex control of RSNA in response to fluid overload in disease models that involve kidney injury (3, 19, 20). Indeed, 73 failure of the baroreflex mechanism is associated with long-term elevation of sympathetic 74 75 nerve activity, which initiates and/or exacerbates hypertension (15). The influence of 76 exposure to CIH on the low-pressure baroreflex control of RSNA and kidney function has not 77 yet been examined. In addition, there is little known about the effects of exposure to CIH on 78 the high-pressure baroreflex control of RSNA.

The importance of renal innervation in long-term control of blood pressure has received attention following clinical trials showing that bilateral renal denervation was associated with a significant reduction of blood pressure in people who are hypertensive with OSA (16, 43, 49). The involvement of renal nerves in mediating the derangement of baroreflex control of blood pressure is revealed by renal denervation, which restored the sensitivity of blunted high- and low-pressure baroreflex control of RSNA in models of hypertension that involve renal injury (19). In addition, pharmacological inhibition of renal inflammation, a feature of animal models of CIH (29, 45, 46, 48), was associated with
normalization of baroreflex control of RSNA (3, 21).

88 Previous studies have shown that selective renal deafferentation was associated with attenuation of the hypertensive phenotype (9, 12). Afferent renal nerves are densely located 89 in the renal pelvic wall and to a lesser extent in the cortex, and their activity is mediated by 90 91 the neuropeptides, calcitonin gene related peptide (CGRP) and substance P (SP) (13, 30). SP 92 release is modulated by the activity of other receptors, such as transient receptor potential 93 vanilloid 1 (TRPV1) channels, localized in the renal pelvic wall. TRPV1 receptors are nonselective cation channels that allow the influx of Na<sup>+</sup>, Mg<sup>2+</sup> and Ca<sup>2+</sup> (17). Renal TRPV1 94 activation is associated with an increase in RSNA and blood pressure in rats (10, 36). In 95 96 addition, acute hypoxia increases basal TRPV1 outward current amplitudes, which is 97 suggested to be due to increased intracellular reactive oxygen species (ROS), that can increase intracellular  $Ca^{2+}$ , with excitatory effects on the afferent reflex (10, 22). Thus, 98 exposure to CIH might provoke renal oxidative stress and inflammation, sufficient to activate 99 100 renal afferent discharge via TRPV1 receptors, eliciting excitatory reno-renal reflexes 101 contributing to hypertension.

The current study was designed to examine the effects of exposure to CIH on highand low-pressure baroreflex control of RSNA and renal excretory responses to fluid overload. We hypothesized that exposure to CIH increases TRPV1 signalling via renal afferent nerves modulating baroreflex control of RSNA. We examined high- and low-pressure baroreflexes and renal diuretic and natriuretic responses to fluid overload in control and CIH-exposed rats, before and after intra-renal TRPV1 blockade with capsaizepine (CPZ).

108

# **Materials and Methods**

#### 111 *Ethical approval*

Eight-week old male Wistar rats were purchased from Envigo (Bicester, UK) and were housed in our institution's animal facility under 12-hour light: 12-hour dark cycle with water and standard chow available *ad libitum*. All experimental procedures were conducted under authorization from the Health Products Regulatory Authority [AE19130/P073] in accordance with European Union directive (2010/63/EU) with prior ethical approval from University College Cork (AEEC 2018/002).

118

## 119 Animal model of chronic intermittent hypoxia

120 Rats were randomly divided into two experimental groups: CIH and sham (control). 121 Rats assigned to the CIH group were placed in commercial environmental chambers attached 122 to a computer-controlled gas delivery system (Oxycycler<sup>™</sup>, Biospherix, NY, USA). Rats in 123 the CIH group were exposed to recurrent hypoxia cycles, resulting in changes in ambient 124 oxygen concentration every 5 min within the chamber from  $21\pm2\%$  for 210 sec to a nadir value of  $6\pm0.5\%$  over 90 sec), for 12 cycles per hour, for 8h during the daytime (lights on), 125 for 14 consecutive days. Rats of the sham group were housed in the same room and were 126 127 exposed to ambient air (~21% O<sub>2</sub>) for 14 days. On day 15, rats were prepared surgically to 128 assess high- and low-pressure baroreflexes under anesthesia.

129

#### 130 Surgical protocol

131 CIH-exposed (n=20) and sham rats (n=23) were anesthetized by intra-peritoneal 132 injection of a mixture of urethane (416 mg/kg, Sigma-Aldrich, St. Louis, MO, USA),  $\alpha$ -133 chloralose (27 mg/kg, Sigma-Aldrich) and sodium pentobarbital (Euthatal (200 mg/ml), 134 Merial animal health Ltd, UK, 33 mg/kg). Anesthesia was maintained during surgery using a maintenance dose of a mixture of urethane (62 mg/kg) and  $\alpha$ -chloralose (4 mg/kg), given as 135 intravenous bolus injection. Animals were placed on a temperature-controlled heating pad to 136 137 maintain body temperature at 37°C (Harvard Apparatus, Cambridge, UK). Airway patency 138 was facilitated by tracheotomy (PE240, Portex, Kent, UK). A cannula (PE25 attached to 139 PE50) containing heparinized saline (4 U/ml), connected to a fluid-filled pressure transducer

(ADInstruments, Hastings, UK) was inserted into the right femoral artery to measure arterial
blood pressure and heart rate (HR). The pressure transducer was connected to a quad-bridge
amplifier that recorded pulsatile blood pressure using LabChart 7 software (ADInstruments,
Oxford, UK). The right femoral vein was cannulated (PE50) to infuse FITC-inulin (10
mg/kg/hr, TdB consultancy, Uppsala, Sweden) continuously over the duration of the
experiment (32). The urinary bladder was cannulated (PE240) to collect urine samples.

146 Baseline blood pressure was recorded for 2 minutes ~4-5 minutes after femoral artery 147 cannulation (before surgical access of the kidney) and compared between the groups, in 148 essence to confirm the hypertensive phenotype in CIH-exposed rats. In the first of the two 149 cohort studies, CIH-exposed (n=9) and sham animals (n=10) were then placed on their right 150 side and the left kidney was exposed by a retroperitoneal flank incision. A cannula (PE10) 151 was inserted into the rostral part of the left kidney at 3.5-4.5 mm depth, to reach the cortico-152 medullary region of the kidney (3) and was fixed in place using acrylate glue. This cannula 153 was connected to a 25 µl micro-syringe (Hamilton, USA) placed on a micro-infusion pump 154 (KD Scientific, Linton Instruments, Norfolk, UK) to infuse saline or CPZ (5 µg/ml, Sigma-155 Aldrich, St. Louis, MO, USA) at 17  $\mu$ l/min. In the second cohort study, CIH-exposed (n=10) 156 and sham rats (n=13) were placed on their left side to expose the right kidney, which was 157 cannulated using a similar approach to the first cohort.

In both cohorts, the renal nerve bundle running between the renal artery and vein of the left kidney was dissected and placed on a bipolar multi-stranded stainless-steel electrode (Medwire, NY, USA). The nerve bundle was sealed over the recording electrode using silicone glue (Klasse 4 dental, Augsburg, Germany). RSNA was recorded using a high impedance head stage attached to an amplifier (NeuroAMP EX®, ADInstruments, UK) connected to a Power Lab data acquisition system.

164

#### 165 *Experimental protocol*

The experimental protocol of the first cohort study is described in Figure 1. Baseline MAP, HR and RSNA were recorded. After intra-renal infusion of saline to the left kidney high-pressure baroreflex responses were determined. Successive volume expansion (VE) challenges were performed, first during intra-renal infusion of saline and then during intrarenal infusion of the TRPV1 blocker, CPZ. The animals were euthanized at the end of the experiment by an overdose of anaesthetic (mixture of 80 g/kg urethane and 55 mg/kg  $\alpha$ -chloralose), given intravenously. RSNA reaches its maximum level during euthanasia due to the sudden precipitous drop of blood pressure, before the complete disappearance of RSNA. The maximum RSNA was determined to allow normalization of baseline RSNA to the maximum. The minimum electrical activity was measured to determine background noise, which was subtracted from all recordings.

The second cohort study involved exclusive assessment of the high-pressure baroreflex response CIH-exposed (n=10) and sham rats (n=10) during two successive phases. The first phase was after 30 minutes of intra-renal infusion of saline (17  $\mu$ l/min) and the second phase was after 30 minutes of intra-renal infusion of CPZ (5  $\mu$ g/ml at 17  $\mu$ l/min). In this cohort of rats, neither VE challenges nor renal function measurements were performed.

183

#### 184 Measurement of Excretory Parameters

Blood samples (P1-P3 samples, Figure 1) of  $\sim$ 400 µl were collected and underwent 185 centrifugation at 10,956 g for 1 minute. Plasma was stored at  $-20^{\circ}$  for further analysis. To 186 187 minimise any change in hematocrit, the blood cells that remained after centrifugation were 188 suspended in an equivalent volume of saline and returned to the animal via the arterial line. 189 Blood samples (at P1 and P3, Figure 1) were taken from the femoral artery to measure sodium concentration using an i-STAT system (Abbott Laboratories, penAbbott Park, IL, 190 191 USA). Two urine samples were collected (U1 and U2, Figure 1), during intra-renal infusion 192 of saline or CPZ. In addition, urine samples were collected during VE (Figure 1). An 193 additional urine sample was collected during the recovery period i.e. 30 minutes after the end 194 of the VE trial (Figure 1).

Urine flow (UF), glomerular filtration rate (GFR), and absolute and fractional Na<sup>+</sup> 195 196 excretion were measured at baseline and during the VE trial. Urine volume was measured gravimetrically and UF was calculated as: UF = Uv/t; Uv is urine volume, t is time of urine 197 198 collection. GFR was measured as a function of FITC-inulin clearance and was calculated using the formula: GFR= ([Uin]Uv)/[Pin]; [Uin] is urine concentration of FITC-inulin, [Pin] 199 is plasma concentration of FITC-inulin. FITC-fluorescence was measured in urine and 200 plasma samples using a fluorometric microplate reader (Wallac victor<sup>2</sup> 1420 multilabel 201 counter, Perkin Elmer, MA, USA). Urinary Na<sup>+</sup> concentration was measured using flame 202

203 photometry (M410, Sherwood Scientific, Cambridge, UK). Plasma Na<sup>+</sup> concentration at P2 204 was determined using flame photometry. However, plasma Na<sup>+</sup> concentration (P1 and P3) 205 was measured immediately upon sample collection using the i-STAT system. Absolute Na<sup>+</sup> 206 excretion was calculated using the formula:  $[UNa^+]$ . UF;  $[UNa^+]$  is urinary Na<sup>+</sup> 207 concentration. Fractional Na<sup>+</sup> excretion was calculated using the equation: Na<sup>+</sup> 208 clearance/GFR.100% where Na<sup>+</sup> clearance =  $([UNa^+]Uv)/[PNa^+]$ ;  $[PNa^+]$  is plasma Na<sup>+</sup> 209 concentration.

210

#### 211 Atrial natriuretic peptide assay

Atrial natriuretic peptide (ANP) was quantified in plasma samples at baseline (P2) and during VE (P3) of the saline phase in CIH-exposed (n=7) and sham rats (n=7) using a quantitative sandwich ELISA kit (My BioSource, San Diego, CA, USA) as per the manufacturer's instructions.

216

### 217 *Tissue preparation*

218 In separate studies, male Wistar rats were exposed for 14 days to normoxia or CIH as described previously. On day 15, rats were euthanized by an overdose of sodium 219 220 pentobarbital (Dolethal (200 mg/ml), Vetoquinol, France, 60 mg/kg), given intra-peritoneally. 221 The left kidney was collected and halved along its sagittal plane. Each half was cut 222 transversely into three sections: top, middle (containing the pelvic wall) and bottom. Sections 223 were preserved at -80°C for later use. Immediately after harvesting the left kidney, the left renal artery and vein were closed by a bulldog clamp. Intra-cardiac puncture via the left 224 225 ventricle was performed and warmed heparinized saline (~ 63 U/ml) was infused. The rate of 226 infusion was 35 ml/min, which was raised gradually to 60 ml/min until 250-300 ml of 227 heparinized saline was infused into each animal. This was followed by infusion of 4% 228 paraformaldehyde at an infusion rate of 60 ml/min (Formalin solution, neutral buffered, 10%, 229 Sigma-Aldrich, St. Louis, MO, USA). The upper and lower poles of the right kidney were 230 removed, and the remnant kidney was either post-fixed for 24 hours for studies employing 231 immunofluorescence techniques, or wax-embedded in paraffin. Some kidneys that were 232 harvested from sham and CIH-exposed rats that underwent surgical procedures were also 233 perfusion-fixed and wax-embedded in paraffin. In separate CIH-exposed (n=6) and sham rats

(n=6) that were not perfusion-fixed, the right kidneys were excised and after removal of the
upper and lower poles, were directly post-fixed in formalin solution for 24 hours for
subsequent use in studies employing immunofluorescence techniques.

237

#### 238 Biochemical assays

239 The middle part of the left kidney was homogenized (32) for use in different assays: 240 Advanced Oxidation Protein Products (AOPP) assay (Cell Biolabs, San Diego, CA), 241 superoxide dismutase (SOD) and catalase activity assays (Cayman Chemical, Ann Arbor, MI, 242 USA) and a TRPV1 ELISA assay (Mybiosource, Tokyo, Japan) (sham, n=12; CIH, n=8). The 243 same homogenization protocol (32) was used to prepare samples for the assessment of 244 NADPH oxidase (NOX) activity (39) and to determine the concentrations of inflammatory 245 cytokines in renal tissue (sham, n=11; CIH, n=10), except for the use of a lysis buffer made 246 of Tris-HCL (20 mM, pH 7.5, Sigma-Aldrich), sodium chloride (150 mM) and 1% Triton-X-247 100 (Molekula, Dorset, UK) instead of RIPA during homogenization. Absorbances were 248 measured using a microplate reader (SpectraMax® M3, molecular devices, California, USA).

249

## 250 Renal inflammatory cytokines

Pro-inflammatory and anti-inflammatory cytokines (IFN- $\gamma$ , IL-1 $\beta$ , IL-4, IL-5, IL-6, keratinocyte chemoattractant/growth related oncogene, IL-10, IL-13, and TNF- $\alpha$ ) were analysed by sandwich immunoassay using a V-plex proinflammatory panel 2 rat kit (Meso Scale Discovery, Rockville, MD, USA), as per the manufacturer's instructions. Plates were analysed using a QuickPlex SQ 120 plate reader (Meso Scale Discovery). IL-1 $\beta$ , TNF- $\alpha$  and keratinocyte chemoattractant/growth related oncogene were successfully measured. The concentration of all other cytokines in kidney were below the detection limits of the kit.

258

#### 259 *Immunofluorescence*

Immunofluorescence for TRPV1 protein expression in the renal pelvic wall was carried out on 3 CIH-exposed and 3 sham kidneys that were perfusion-fixed and post-fixed, in addition to 6 CIH-exposed and 5 sham kidneys that were only post-fixed. Frozen, formalin-fixed kidneys were sectioned to 20 µm sagittal sections using a cryostat (Leica 264 Biosystems, Wetzlar, Germany). Three sections that contained the renal pelvic wall were 265 randomly selected from each kidney by systematic random sampling. Non-specific binding 266 was blocked by incubating sections in 5% goat serum and 1% bovine serum albumin (with 267 0.5% triton-X) for one hour. Then, sections were incubated with rabbit polyclonal anti-268 TRPV1 antibody (1:200, ACC-030, Alomone labs) diluted with 5% goat serum, 1% bovine 269 serum albumin and 0.5% triton-X for two hours at 37°C. Sections were incubated with goat 270 anti-rabbit FITC secondary antibody (1:100, Sigma-Aldrich, St. Louis, MO, USA) for two 271 hours at room temperature. Sections were covered with anti-fade mounting medium 272 (Vectashield, Vector laboratories, Burlingame, CA, USA).

273 Images were captured using a laser scanning confocal microscope (Olympus FV1000-274 IX71, Tokyo, Japan) provided with filters for FITC and DAPI. Sections were initially 275 visualized using 20X magnification under the DAPI filter and three regions of the pelvic wall were randomly selected from each section. After selection, the FITC filter was switched on to 276 277 capture images. Positive staining was detected in the uroepithelial layer and submucosa/ 278 muscularis propria layers of the pelvic wall. Images were transferred to black and white 279 images and the threshold was adjusted to highlight the area occupied by fluorescent labeling 280 of TRPV1 using Image J software. Threshold area was calculated and normalized to the total 281 area of the uroepithelium within the image to quantify the expression of TPRV1 in that region. Similarly, threshold area in the submucosa and muscularis propria layers were 282 283 calculated and normalized to the total area identified in the image.

284

#### 285 *Renal histopathological staining*

286 Kidneys were taken from CIH-exposed (n=8) and sham rats (n=8) that underwent 287 surgical procedures as well as from rats euthanized following exposure to gasses without 288 undergoing surgery. Paraffin blocks were cut to 10 µm tissue sections using a rotary 289 microtome (Leica RM2135, Germany). Two sections were randomly selected by random 290 number generator each from the upper, middle and bottom parts of the paraffin block (a total 291 of 6 sections from each kidney block). Sections were de-waxed using Neo-Clear® xylene 292 substitute (Merck, Darmstadt, Germany) and washed with down-graded levels of ethanol. 293 Tissue sections were then stained in hematoxylin (Hematoxylin Solution, Harris Modified, 294 Sigma-Aldrich, St. Louis, MO, USA) and eosin (Surgipath Europe LTD, Cambridgeshire, 295 UK or with Sirius red stain (0.1% Direct red 80, Sigma-Aldrich, India). Sections were cleared

in Neo-Clear® xylene substitute, and mounted with DPX mounting medium (Sigma-Aldrich,St. Louis, MO, USA).

298 Hematoxylin and Eosin staining was used to examine total glomerular tuft area. A 299 virtual grid comprised of 91 squares was created to enable random sampling of cortical areas 300 of each kidney section. Selected squares were magnified using a 40X magnification lens to 301 image selected glomeruli (only when the juxtaglomerular apparatus was evident). As some 302 squares did not include any glomeruli and/or the juxtaglomerular apparatus was not visible in 303 some glomeruli, other squares were randomly chosen so that a total of 3 glomeruli per kidney 304 section were analyzed. As 6 sections were randomly selected from each animal, a total of 18 305 glomeruli per animal were chosen to measure glomerular tuft area. Images were taken using a 306 40X magnification lens (Olympus inverted BX53F microscope, Tokyo, Japan) and tuft area 307 was measured using ImageJ software.

308 Sirius red stain was used to semi-quantify areas occupied by collagen in the cortex 309 and outer medulla. A virtual grid with 25 squares was used to enable random selection of 310 cortical and medullary areas. Selected squares were magnified using a 20X magnification 311 lens to capture images. Three cortical areas and two outer medullary areas per kidney section 312 were randomly chosen. As 6 sections were stained from each animal, a total of 18 cortical 313 areas and 12 areas from the outer medulla were selected for analysis. Glomeruli and 314 perivascular fibres were omitted from images before analysis. The fibrotic area of each image 315 was normalized to the total area of the captured image to generate the % fibrotic area (Figure 316 2).

#### 317 Statistical analysis

Data in tables and within text are presented as mean  $\pm$  SD. Line figures are shown as mean  $\pm$  SE. Baseline MAP, HR and RSNA were compared between groups using independent sample *t*-tests. Baseline RSNA was expressed as absolute values ( $\mu$ V.s) and also was normalized to the maximum RSNA value recorded during euthanasia, expressed as RSNA (% of max). Values were compared between groups using repeated-measures two-way ANOVA.

Baseline values for UF, GFR, and absolute and fractional Na<sup>+</sup> excretion were compared using repeated-measures two-way ANOVA. RSNA sympatho-inhibitory response to VE was expressed as % of baseline (baseline=100%) and as RSNA (% of max). The change in functional parameters and the decrease in RSNA in response to VE were analysed using repeated-measures two-way ANOVA followed by a Bonferroni *post hoc* test. The area
under the curve (AUC) of RSNA (% of baseline), RSNA (% of max), and each functional
parameter was compared between groups and between the saline VE phase and CPZ VE
phase using independent sample *t*-test and paired *t*-test, with significance taken at P<0.0125,</li>
accounting for multiple comparisons.

333 High pressure baroreflex function curves were fitted according to a logistic sigmoidal 334 function equation that shows the relationship between MAP and RSNA or HR 335 y=A1/(1+exp(A2(x-A3)))+A4 where y is the RSNA or HR; A1, response range over which baroreceptors operate; A2, gain coefficient; A3, mid-point blood pressure; A4, minimum 336 337 response of RSNA or HR. Parameters were compared between the sham and CIH-exposed 338 groups using independent sample *t*-test and significance was taken at P<0.05.Baroreflex gain 339 curve parameters during the first (saline) and second (CPZ) phase were compared using 340 paired *t*-tests. Significance was taken at P<0.0125, accounting for multiple comparisons.

Data from biochemical assays, histology and immunofluorescence from CIH-exposed and sham tissues were analysed using independent sample *t*-tests or Mann-Whitney tests, and significance was taken at P<0.05. All statistical analyses were performed using SPSS software (SPSS Statistics for Windows, v25.0. IBM corp., NY, USA). All graphs were plotted using GraphPad® Prism (v6, GraphPad software, San Diego, CA, USA).

346

347

# 349 **Results**

#### 350 Baseline MAP, HR, RSNA and renal excretory parameters

351 A representative recording of baseline parameters is shown in Figure 3. In the first 352 cohort study, MAP was greater in CIH-exposed rats compared with sham rats (Table 1). 353 Baseline RSNA was greater in CIH-exposed rats compared with sham rats (RSNA (% of max),  $36\pm15\%$  vs.  $17\pm7\%$ , p=0.006). MAP was greater in CIH-exposed rats of the second 354 355 cohort compared with sham rats, but to a lesser extent compared with rats of the first cohort 356 study (Table 1 and Supplementary Table 1 https://figshare.com/s/7bc496e361e5e47043d7). 357 HR was similar in CIH-exposed and sham rats of both cohorts. Baseline renal function data are shown in Supplementary Table 2 (https://figshare.com/s/7796b802afe6c02d54f0). 358 359 Exposure to CIH had no significant effect on any of the basal renal excretory parameters.

360

#### 361 *Cardiovascular responses to VE*

Baseline ANP concentration in plasma prior to VE challenge was not significantly different between CIH-exposed and sham rats (CIH vs. sham; 157±17 vs. 167±36 pg/ml). MAP and HR data during each VE trial are shown in Figure 4a and 4b. During the last 2 minutes of each VE trial, MAP was not significantly different from respective baseline values recorded prior to VE (Table 2). HR was significantly elevated during the last 2 minutes of each of the VE trials, both in CIH-exposed and sham rats (Table 2).

368

#### 369 *Renal excretory responses to VE*

VE in CIH-exposed and sham rats resulted in significant increases in GFR, UF and 370 371  $Na^+$  excretion (Figure 4d-g). The response pattern and magnitude of GFR and fractional  $Na^+$ 372 excretion during VE were similar in sham and CIH-exposed rats. Diuresis in response to VE was of significantly lesser magnitude in CIH-exposed rats compared with sham rats after 25 373 and 30 minutes (Figure 4d). Similarly, absolute Na<sup>+</sup> excretion was significantly less in CIH-374 exposed rats after 30 minutes of VE (Figure 4f). Analysis of AUC, reflecting the temporal 375 376 cumulative response to VE, revealed that UF was significantly less in CIH-exposed rats 377 compared with sham rats (Figure 4d, f).

379

#### 380 *RSNA response to VE*

VE caused a progressive significant decrease in RSNA, which reached its maximum decline after 28-30 minutes (Figure 4c). The decrease in RSNA during VE with concomitant intra-renal infusion of saline was not significantly different in CIH-exposed and sham rats (Figure 4c).

385

#### 386 *High-pressure baroreflex*

387 Representative recordings from high-pressure baroreflex trials are shown in Figure 5a. 388 The slope of the RSNA baroreflex function curve of CIH-exposed rats was less than that of 389 sham rats (A2 parameter: Figure 6a). This was related to a significant elevation in response 390 range (A1 parameter) and operating range (Table 3) in CIH-exposed rats compared with 391 sham rats. A rightward shift in the pressure at which the baroreceptors operate (A3 parameter), by ~20 mmHg (p=0.005), in CIH-exposed rats was observed (Figure 6 and Table 392 393 3).. Conversely, data from the second cohort study (with a more modest increase in MAP in 394 CIH-exposed rats) revealed no difference in any of the parameters between CIH-exposed and 395 sham rats (Figure 6).

These apparently conflicting results obtained from the two cohorts are likely explained by the moderate correlation (Figure 7) that we found between baseline blood pressure and the slope (A2,  $R^2=0.386$ , p=0.017) and mid-point pressure (A3,  $R^2=0.639$ , p<0.0001) of RSNA baroreflex function curves, combining all rats in both cohorts. Parameters of the baroreflex control of HR were not significantly different between CIHexposed and sham rats in either of the two cohort studies (Table 3).

402

403 Baseline MAP, HR, RSNA and renal excretory parameters during intra-renal TRPV1
404 blockade

405 Intra-renal infusion of saline had no significant effect on blood pressure, HR and **RSNA** in either 2 406 cohort (Table and Supplementary Table 1 https://figshare.com/s/7bc496e361e5e47043d7). In contrast, MAP was lower during intra-407 renal infusion of CPZ both in CIH-exposed and sham rats (of both cohorts), compared with 408

409 respective baseline values measured prior to CPZ infusion, whereas RSNA and HR were 410 unchanged (Table 2 and Supplementary Table 1 411 https://figshare.com/s/7bc496e361e5e47043d7). During intra-renal infusion of CPZ there 412 was a significant diuresis and natriuresis, both in CIH-exposed and sham rats compared with 413 respective values measured prior to CPZ infusion (Supplementary Table 2 https://figshare.com/s/7796b802afe6c02d54f0). 414

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### 416 *Renal excretory responses to VE during intra-renal TRPV1 blockade*

417 During intra-renal infusion of CPZ, significant increases in UF and absolute Na<sup>+</sup>
418 excretion in response to VE were observed in CIH-exposed and sham rats, with no significant
419 difference between the experimental groups (Figure 4).

420

#### 421 Low- and high-pressure baroreflex control during intra-renal TRPV1 blockade

422 During intra-renal infusion of CPZ, the decrease in RSNA (% of max) in response to 423 VE was equivalent in CIH-exposed and sham rats (Figure 4c). However, an apparent 424 potentiation of the RSNA sympatho-inhibitory response to VE was observed in both groups 425 when RSNA (% of baseline) was considered (Supplementary Figure S1, 426 https://figshare.com/s/9659e62049b971e25ab7). Intra-renal CPZ did not significantly affect 427 high-pressure baroreflex of RSNA or HR in CIH-exposed and sham rats of cohort 2 428 (Supplementary Table 3 https://figshare.com/s/870e5f260f49995a4d8d, and Figure 6c and 429 6d).

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#### 431 *TRPV1 immunofluorescence in renal tissue*

TRPV1 channels are mainly expressed in the fibres located between the muscular layer and the uroepithelium of the renal pelvic wall (Figure 8). Positive staining was also expressed in the uroepithelium (Figure 8). TRPV1 fluorescence occupied  $5.1\pm1.0\%$  of the area of the muscularis propria of the pelvic wall of CIH-exposed rats, which was equivalent to sham rats ( $6.4\pm1.9\%$ , Figure 8f). TRPV1 expression in the uroepithelium was equivalent between CIH-exposed and sham rats ( $3.7\pm0.9\%$  vs.  $3.9\pm1.1\%$ ). TRPV1 protein concentration in kidney tissue (Figure 8f, bottom) was not significantly different between CIH-exposed andsham rats.

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#### 441 *Renal histopathological assessment*

The average glomerular tuft area was equivalent in CIH-exposed and sham rats (Figure 9). A slightly greater degree of fibrosis was observed in the cortex of CIH-exposed kidneys compared with sham kidneys (Figure 9b and d). In contrast, there was no significant difference in collagen expression in the outer medulla between CIH-exposed and sham kidneys (Figure 9c and d).

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#### 448 *Renal oxidative stress, inflammation and protein oxidation*

The activity of NOX, SOD and catalase enzymes was measured in whole kidney tissue and there was no significant change in the activity of these enzymes after exposure to CIH. There was no significant difference in NOX, SOD and catalase enzyme activities between CIH-exposed and sham rats (Table 4). Furthermore, there was no evidence of greater protein oxidation in the kidneys of CIH-exposed rats (Table 4). Cytokine concentrations (IL-1 $\beta$ , keratinocyte chemoattractant/growth related oncogene and TNF- $\alpha$ ) were not significantly different between CIH-exposed and sham rats (Table 4).

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# 465 **Discussion**

This study provides several novel findings: 1) In CIH-exposed rats, diuretic and natriuretic responses to VE were impaired, notwithstanding that the sympatho-inhibitory response of RSNA to fluid overload was preserved; 2) the high-pressure baroreflex was modestly affected in CIH-exposed rats, an outcome that was related to the magnitude of CIH-induced hypertension, suggesting an indirect effect of exposure to CIH; 3) CIH-induced hypertension can manifest in the absence of renal injury.

VE produced reductions in RSNA similar to the findings of previous studies (8, 19).
Exposure to CIH, sufficient to cause hypertension, did not adversely affect the sympathoinhibitory response of RSNA to VE. This revealed that altered low-pressure baroreflex
control of RSNA did not contribute to CIH-induced hypertension.

476 Basal renal function was equivalent in sham and CIH-exposed rats. However, the 477 renal excretory response to VE in CIH-exposed rats was blunted with resultant sodium and 478 fluid retention. These findings suggest that compensatory mechanisms in the kidney ensured 479 maintenance of normal basal kidney function in the face of CIH-related stress and 480 sympathetic over-activity. Dysfunction was revealed however, during VE challenge, with 481 evidence of blunted diuretic and natriuretic responses, which appear independent of neural 482 control of the kidney. Similarly, in streptozotocin-induced diabetic rats, basal diuresis and 483 natriuresis were equivalent to non-diabetic rats, but excretory responses were impaired during 484 VE (44). The underlying mechanisms that contribute to impairment of VE-dependent excretory responses are multifactorial and do not depend solely on changes in RSNA. Such 485 486 mechanisms include alterations in renin-angiotensin system (RAAS) activity, renal perfusion 487 pressure, and GFR. Thus, our finding of blunted sodium and fluid excretion during VE, 488 despite preserved reflex reduction in RSNA, highlights that other mechanisms contribute to 489 diuretic and natriuretic responses to VE and reveals that one or more of these mechanisms are 490 perturbed by exposure to CIH.

Impaired cardiac baroreflex function was previously reported in people with OSA and in rats exposed to CIH (24, 38). Accordingly, we hypothesized that exposure to CIH disrupts the high-pressure baroreflex control of RSNA (3, 18, 20). CIH-exposed rats of the first cohort showed a significant shift in mid-point blood pressure by almost 20 mmHg, which was associated with an increase in the operating range of the RSNA baroreflex. Moreover, a 496 marked increase in response range (A1 parameter) of the RSNA baroreflex was evident in 497 CIH-exposed rats. Elevation in the operating and response range of RSNA resulted in a decrease in the slope of the baroreflex (A2 parameter). These changes were concomitant 498 499 with a substantial hypertensive phenotype (Table 1). Conversely, CIH-exposed rats of the 500 second cohort with modest hypertension displayed baroreflex responsiveness that was 501 equivalent to the respective sham group. Interestingly, a significant correlation between basal 502 MAP and the baroreflex slope (A2) and between MAP and mid-point pressure (A3) was 503 evident (Figure 7). This indicates that resetting of RSNA baroreflex and apparent decreased 504 slope of the curve is associated with elevated blood pressure. It is likely that baroreflex 505 adjustments following exposure to CIH in this study and marked attenuation of baroreflex 506 function observed in severe CIH protocols, are secondary to the elaboration of hypertension 507 following exposure to CIH, as the second cohort of rats were hypertensive (modest compared 508 with cohort 1), but displayed no changes in baroreflex parameters revealing that the latter is 509 not obligatory for the manifestation of CIH-induced hypertension.

The baroreflex control of HR was unchanged in CIH-exposed rats in agreement with previous studies utilizing 5, 7 and 10 days of exposure to CIH (24, 47, 50). Lai et al (24) analysed day-to-day arterial blood pressure and baroreflex sensitivity of the heart in conscious rats. MAP was higher by 15 mmHg after 5 days of exposure to CIH with no alteration in baroreflex function, followed by a decline in baroreflex sensitivity starting 10 days after the onset of exposure to CIH (24). Indeed, long-term exposure to CIH was associated with decreased cardiac baroreflex gain (11, 14, 27).

517 In renal injury, there is activation of renal pelvic chemoreceptors causing activation of afferent renal nerves, sympatho-excitation and hypertension (7, 23, 31, 34). In renal failure 518 519 and high fat-induced obesity models, impairment of high- and low-pressure baroreflex 520 function was linked to increased renal afferent nerve activity (3, 20). Therefore, renal 521 denervation and/or suppression of renal inflammation is associated with restoration of the 522 baroreflex control of RSNA (19, 21). Similarly, in some animal models of CIH, increases in 523 systemic and renal inflammatory biomarkers and ROS have been reported together with impaired baroreflex control of HR (11, 24, 47). Afferent renal nerve fibres are abundant in 524 the renal pelvic wall (23). A subset of these afferent nerves contains TRPV1, which is co-525 localized with neurotransmitters such as SP (17). In rats, cortico-medullary infusion of the 526 527 TRPV1 agonist capsaicin induces an excitatory reno-renal reflex and an increase in blood 528 pressure (36). It is also known that acute hypoxia increases the tonic activity of TRPV1

receptors. Hypoxia-induced activation of TRPV1 was suppressed using CPZ and Tiron, a membrane permeable ROS scavenger (22). Moreover, *in vitro* studies showed that TRPV1 activation decreases cell viability and induces apoptosis (42). This effect is mediated by an increase in intracellular calcium influx followed by ROS production, which causes mitochondrial depolarization and DNA fragmentation (42).

534 CPZ, at a similar dose to that used in the present study, was utilized in previous 535 studies to block TRPV1 resulting in marked changes in RSNA (26). The infusion of CPZ 536 over 30 minutes in the present study diffused into the pelvic wall where renal afferent nerves 537 are located. This was confirmed histologically by observing the location of lissamine green 538 stain infused for 30 minutes (not shown). Intra-renal infusion of CPZ caused a significant 539 diuresis and natriuresis accompanied by significant reduction in MAP, but without an 540 attendant decrease in RSNA. In addition, the renal sympatho-inhibitory response to VE after 541 CPZ administration was unaffected, both in CIH-exposed and sham rats. Nevertheless, 542 following intra-renal infusion of CPZ, diuretic and natriuretic responses to VE were 543 equivalent in CIH-exposed and sham rats, which suggests a potential role of renal TRPV1 544 receptors in CIH-induced impairment of renal excretory function, although we cannot 545 exclude the possibility that the intra-renal CPZ-induced diuresis and natriuresis per se (which 546 was independent of RSNA) may have contributed to the recovery of renal excretory responses to VE in CIH-exposed rats, masking the original mechanism contributing to 547 548 impaired renal excretory function. Therefore, our results should be viewed as preliminary and 549 the molecular mechanisms contributing to CIH-induced impairment in renal excretory 550 function requires further study.

551 Of interest, there was no difference between CIH-exposed and sham rats in TRPV1 552 expression in the renal pelvic wall. Histological analysis in the CIH-exposed rats showed 553 modest cortical fibrosis, which may relate to decreased cortical oxygen tension and reduced 554 renal blood flow in CIH-exposed rats, which was previously reported (32). Interestingly, 555 neither inflammation nor altered pro- or anti-oxidant enzymatic activity was evident in CIHexposed kidneys. Moreover, inflammatory cytokine concentrations were not different 556 between experimental groups. Variable renal outcomes at the molecular level in animal 557 models of CIH are likely related to differences in the pattern, duration and intensity of CIH, 558 as recently reviewed (6). Glomerular hypertrophy and/or inflammation was reported in other 559 560 studies (45), but were not observed in this study, suggesting that these changes, when they

561 562 occur, are more likely to be secondary to hypertension, rather than a direct effect of exposure to CIH, although they may also arise due to synergistic effects of hypertension and hypoxia.

563 This study has several limitations. Data were collected from anesthetized rat 564 preparations. As such, cardiovascular, renal, and autonomic nervous system function were 565 altered compared with the conscious state. It will be important to determine if exposure to 566 CIH adversely affects renal excretory function in conscious animals. Intra-renal CPZ 567 administration was unilateral, which might not reflect the full biological effect of TRPV1 568 antagonism. Unilateral infusion of drugs has been used effectively in other studies of 569 baroreflex function (2, 3). In the present study, unilateral CPZ produced renal and 570 cardiovascular effects, nevertheless the effects of bilateral TPRV1 inhibition is worthy of 571 further investigation. For logistical reasons, high-pressure baroreflex function was not 572 assessed after intra-renal CPZ in rats of cohort 1, where baroreflex alterations were 573 subsequently revealed. The effect of intra-renal CPZ was assessed in rats of cohort 2, but 574 high-pressure baroreflex control of RSNA was preserved in these rats with modest 575 hypertension. Therefore, we cannot discount a possible role for TRPV1 signalling in 576 modulating high-pressure baroreflex control of RSNA in CIH-exposed rats and this warrants 577 further attention. Our study design involved the assessment of RSNA and renal excretory responses to VE in successive challenges, before and after intra-renal TRPV1 blockade. We 578 579 acknowledge that restoration of renal excretory responses to VE in CIH-exposed rats, which 580 was attributed to blockade of TRPV1 receptors, might relate to the performance of a second 581 VE trial per se, independent of antagonism. Moreover, renal excretory responses to the 582 second VE challenge should be viewed in the context of the profound diuresis and natriuresis 583 caused by intra-renal CPZ. Our data on GFR during VE should be viewed cautiously given 584 that measurements were not taken in steady-state conditions and blood sampling was limited 585 as we did not wish to interfere with MAP, and hence reflex control of RSNA, during VE 586 challenges.

In summary, a relatively modest paradigm of exposure to CIH was sufficient to induce hypertension, without evidence of renal inflammation and/or oxidative stress. Exposure to CIH blunted sodium and fluid excretion in response to VE with preserved suppression of RSNA. Exposure to CIH induced modest changes in high-pressure baroreflex control of RSNA, which was dependent on the magnitude of blood pressure elevation in CIHexposed rats suggesting that modulation of high-pressure baroreflex control of RSNA in this model is a consequence of hypertension, but may in turn proceed to a pathological level and 594 contribute to disease progression. Our results suggest that exposure to CIH causes renal 595 dysfunction, which may be relevant to obstructive sleep apnea.



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# 761 **Figure legends:**

762 Figure 1: A schematic representation of the experimental protocol. In sham and 763 CIH-exposed rats of cohort 1, following a 1 hour stabilization period, baseline cardiovascular 764 and RSNA were determined. After intra-renal infusion of saline to the left kidney for 30 765 minutes, phenylephrine (PE, 50 µg/ml, 0.2 ml) and sodium nitroprusside (SNP, 50 µg/ml, 0.2 766 ml) were infused intravenously to elicit high-pressure baroreflex responses. The first volume 767 expansion (VE) trial (saline phase) was then performed, which was followed by a 90-minute 768 recovery period. Then, intra-renal infusion of CPZ was initiated for 30 minutes into the left 769 kidney and a second VE trial was performed. Two urine samples (U1 and U2) were collected 770 during baseline before each VE trial. The "Urine collection" phase represents the collection 771 of a urine sample 10 minutes after the start of the VE trial followed by subsequent collection 772 of urine samples every 5 minutes until the end of the VE challenge. Arterial blood was 773 sampled for plasma (P1 and P2). Each VE trial was accompanied by infusion of inulin (FITC-774 inulin in saline, 10 mg/kg) for 30 minutes via the femoral vein at 0.25 ml/min/100g body 775 weight using a syringe pump (Graseby syringe pump 8100, Dublin, Ireland). Each VE trial 776 period was followed by 30 minutes of recovery during which infusion of saline through the 777 intravenous line was stopped, but with continuous intra-renal infusion of saline or CPZ. P, plasma sample; VE, volume expansion; I.R., intra-renal; CPZ, capsaizepine; PE, 778 phenylephrine; SNP, sodium nitroprusside. 779

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Figure 2: Assessment of renal fibrosis. A virtual grid was used for the random selection of
kidney regions. Sirius red stained images were converted to grey scale and a colour threshold
function was used to highlight collagen labelling in red.

Figure 3: Original recordings of cardiovascular parameters and RSNA in an anesthetised rat. Representative original recordings of baseline arterial pressure (AP), heart rate (HR) which was derived from the AP recording, and raw and integrated RSNA (Int. RSNA) recordings from a sham rat.

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Figure 4: Cardiovascular, RSNA and renal excretory responses to VE. In sham (n=8) and 790 CIH-exposed rats (n=9), MAP, HR, RSNA, UF, GFR, absolute and fractional Na<sup>+</sup> excretion 791 responses to VE challenges were determined. In the left-hand panels, the time course of 792 793 responses to and recovery from VE during intra-renal infusion of saline or CPZ (TRPV1 794 blocker) are shown. Data (mean±SE) were averaged over each 2-minute period during VE 795 trials. Two-way ANOVA (time x exposure) was used to statistically compare the data. # 796 p < 0.05 for time;  $\Phi p < 0.05$  for time points significantly different from respective baseline 797 (post-hoc analysis);  $\Psi$  p<0.05 for exposure; \* p<0.05 compared with corresponding time 798 point in sham rats. In the right-hand panels, area under the curve (AUC) analysis for the respective responses is shown (expressed in arbitrary units, AU), calculated using the 799 800 trapezoidal rule, by the summation of the VE period. Data are shown as individual data points 801 for each rat, with box and whiskers plots representing the interquartile range and maximum 802 and minimum values. Data were compared with independent *t*-tests. Significance was taken 803 at p<0.0125, accounting for multiple comparisons. MAP, mean arterial pressure; RSNA, 804 renal sympathetic nerve activity; UF, Urine flow; GFR, glomerular filtration rate; MAP, 805 mean arterial blood pressure; HR, heart rate; CPZ, capsaizepine.

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807 Figure 5: Original recordings illustrating high- and low-pressure baroreflex control of RSNA in anesthetised rats. Representative original recordings of arterial blood pressure, 808 809 raw and integrated RSNA responsiveness to blood pressure manipulations evoking the highpressure baroreflex (a) and VE evoking the low-pressure baroreflex (b). Recordings are 810 811 shown for representative sham and CIH-exposed rats showing responses during intra-renal infusion of saline and intra-renal infusion of CPZ, (TRPV1 blocker). AP, arterial blood 812 813 pressure; RSNA, renal sympathetic nerve activity; Int. RSNA, integrated RSNA; VE, volume expansion; CPZ, capsaizepine. 814

816 Figure 6: High-pressure baroreflex control of RSNA. Data (mean±SE) are shown for 817 RSNA (% of baseline) as a function of arterial blood pressure in sham and CIH-exposed rats 818 of the first (a; sham, n=10; CIH, n=8) and second cohort (b; sham, n=10; CIH, n=10). In 819 graphs (c) and (d), data (mean±SE) are shown for RSNA (% of baseline) as a function of 820 arterial blood pressure during intra-renal infusion of saline and CPZ (TRPV1 blocker) of 821 sham (c) and CIH-exposed rats (d) of the second cohort. Data for baroreflex parameters are 822 shown in table 3 and supplementary table 3 (https://figshare.com/s/870e5f260f49995a4d8d). 823 RSNA, renal sympathetic nerve activity; BP, blood pressure.

824

Figure 7: Correlations between baseline MAP and baroreflex parameters. The relationship between mean arterial blood pressure and the baroreflex parameters, A3 and A2, are shown for all sham (n=20) and CIH-exposed rats (n=18). Linear regression analysis was performed. Significance was taken at P<0.05. MAP, mean arterial blood pressure; A3, midpoint pressure of the response; A2, gain coefficient.

830

Figure 8: TRPV1 immunofluorescence in the renal pelvic wall. Immunofluorescence 831 832 labeling of renal pelvic wall showing TRPV1-positive labeling within the muscularis propria 833 (white arrows) and uroepithelium layer (red arrows). (a) is a representative image from a 834 sham kidney and (b) is a representative image from a CIH-exposed kidney. Negative controls 835 demonstrate specificity of the antibodies used in the study (c, primary antibody omitted; e, 836 secondary antibody omitted); Blue labelling of nuclei by Hoescht. (d) Immunofluorescence 837 labeling of TRPV1 channels in neuronal cell bodies of lumbar dorsal root ganglia as a 838 positive control. Scale bar 50 µm. Data for TRPV1-positive staining (% of total area) in sham 839 (n=8) and CIH-exposed rats (n=9) are shown as individual data points for each rat, and box 840 and whiskers plots representing the interquartile range and maximum and minimum values (f, 841 top). Data for TRPV1 protein concentration of kidney tissue homogenate in sham (n=12) and 842 CIH-exposed rats (n=8) are shown as individual data points for each rat, with box and 843 whiskers plots representing the interquartile range and maximum and minimum values (f, 844 bottom). Data were compared with independent *t*-tests. Significance was taken at p < 0.05.

**Figure 9: Renal histology.** Representative images from a sham and CIH-exposed rat are shown. (a) shows hematoxylin and eosin staining; (b) shows Sirius red staining of renal cortex; (c) shows Sirius red staining of renal outer medulla. Scale bar 50  $\mu$ m. Data for glomerular tuft area and percentage fibrotic area are presented in (d) as individual data points for each rat, with box and whiskers plots representing the interquartile range and maximum and minimum values. Data from sham (n=8) and CIH-exposed rats (n=8) were compared using independent *t*-tests. Significance was taken at p<0.05.





# Figure 2



# Figure 3





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# Figure 8



# Figure 9



1 Table 1. Baseline cardiovascular and RSNA parameters in sham and CIH-exposed

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rats

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Parameter	Sham	CIH	p-value	Sham	CIH	p-value		
After arterial cannulation (before renal surgical instrumentation)								
<u>First cohort</u> <u>Second cohort</u>								
MAP (mmHg)	124±	13 143±7*	0.001	115±11	127±9*	0.014		
SBP (mmHg)	143±	17 163±11*	0.008	137±13	144±12	0.154		
DBP (mmHg)	114±	12 133±5*	<0.000	<b>1</b> 104±11	118±10*	0.005		
HR (beats/min)	432±2	28 461±39	0.083	411±29	425±41	0.355		
After stabilization (after renal surgical instrumentation)								
MAP (mmHg)	84±9	97±11*	0.028	103±12	107±18	0.565		
SBP (mmHg)	116±	14 125±15	0.262	134±15	132±22	0.871		
DBP (mmHg)	69±8	82±11*	0.043	88±13	94±19	0.365		
HR (beats/min)	400±3	52 445±47	0.084	436±19	450±34	0.204		
RSNA (% of max)	) 17±7	36±15*	0.006	24±10	32±18	0.387		
RSNA (uV.s)	0.66±0.3	30 1.34±0.7	0.231	1.39±0.84	1.67±0.65	0.878		
Maximum RSNA ( uV .s)	4.70±2.0	03 4.03±2.1	8 0.349	7.04±5.29	6.60±3.54	1.000		

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7 Data are presented as mean±SD and were collected from the first (sham, n=10; CIHexposed, n=9) and second cohorts of animals (sham, n=13; CIH, n=10). Baseline 8 cardiovascular parameters were recorded for 2 minutes ~4-5 minutes after arterial 9 cannulation i.e. before retroperitoneal incisions were performed to expose the kidneys. 10 Cardiovascular parameters and RSNA were also recorded for 2 minutes after the 11 stabilization period, 1 hour following renal surgical instrumentation. Parameters were 12 analysed using independent samples t-test or Mann-Whitney test. \* p<0.05 between 13 exposure groups. CIH, chronic intermittent hypoxia; MAP, mean arterial blood pressure; 14 SBP, systolic blood pressure; DBP, diastolic blood pressure; HR, heart rate; RSNA (% of 15 max), renal sympathetic nerve activity normalized to maximum activity recorded during 16 baroreflex activation during euthanasia. 17

18

# 21 Table 2. MAP, HR and RSNA before and during VE challenges in sham and CIH-

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- 22 exposed rats.
- 23

First VE (saline phase)				Second VE (CPZ phase)				
Parameter	MAP	HR	RSNA	RSNA	MAP	HR	RSNA	RSNA
	(mmHg)	(beats/min)	(% of max)	(uV.s)	(mmHg)	(beats/min)	(% of max)	(uV.s)
		Before intra-r	renal infusion	of saline	Before intra-renal infusion of CPZ			
Sham	90±9	409±35	15±5	0.71±0.34	94±11	430±41	18±6	0.87±0.43
CIH	99±10	444±49	$32\pm18^{\phi}$	1.19±0.82	98±11	$448{\pm}42$	23±6	0.95±0.72
		After intra-ren	al saline (befo	re VE)	After intra-renal CPZ (before VE)			
Sham	90±8	408±49	18±8	0.86±0.50	83±5 <sup>#</sup>	412±45	18±6	0.86±0.41
CIH	90±4	428±46	$30\pm14^{\phi}$	1.14±0.78	90±5 <sup>#</sup>	431±46	22±8	0.94±0.74
Last two minutes of VE				Last two minutes of VE				
Sham	92±15	451±35*	15±6*		76±10	443±33*	12±6*	
CIH	91±9	458±31*	18±5*		90±9	450±25*	11±8*	
		2x2 .	ANOVA (befo	ore vs. after in	tra-renal sa	lline/CPZ)		
Exposure	P=0.075	P=0.131	P=0.035	P=0.180	P=0.225	P=0.165	P=0.158	P=0.567
Time	P=0.102	P=0.084	P=0.894	P=0.330	P=0.024	P=0.100	P=0.722	P=0.396
Exposure	P=0.065	P=0.007	P=0.044	P=0.137	P=0.502	P=0.748	P=0.975	P=0.838
x time								
		2x2 ANOVA	(after intra-re	nal saline/CP	Z vs. last tv	wo minutes of V	Έ)	
Exposure	P=0.516	P=0.404	P=0.077		P=0.045	P=0.205	P=0.581	
Time	P=0.072	P<0.001	P=0.01		P=0.168	P=0.002	P=0.001	
Exposure	P=0.786	P=0.182	P=0.115		P=0.073	P=0.218	P=0.252	
x time								

24

25 Data were collected from sham (n=8) and CIH-exposed rats (n=9) of the first cohort. Data

are expressed as mean±SD. Baseline values over a 2-minute period were averaged before

and after the infusion of saline or CPZ and were analysed using repeated-measures two-

28 way ANOVA. The last two minutes of each VE trial were averaged and compared with

baseline values collected immediately before VE using repeated-measures two-way
 ANOVA. \* p<0.05 vs. corresponding value after intra-renal infusion of saline/drug. #</li>

- p<0.05 vs. corresponding value before intra-renal infusion of drug. <sup>\$\overline\$</sup> p<0.05 vs. sham.</li>
   MAP, mean arterial blood pressure; HR, heart rate; VE, volume expansion; CPZ, capsaizepine; RSNA (% of max), renal sympathetic nerve activity normalized to maximum activity recorded during baroreflex activation during euthanasia.

38 Table 3. High-pressure baroreflex parameters during intra-renal infusion of saline

39 in sham and CIH-exposed rats (cohort 1 and cohort 2).

40

Parameter	Sham	СІН	p-value	Sham	CIH	p-value			
RSNA baroreflex									
	First co	hort		Second cohort					
A1 (%)	73±26	104±32*	0.043	114±30	100±22	0.280			
A2	$0.29 \pm 0.27$	$0.09 \pm 0.05*$	0.016	0.10±0.03	0.15±0.14	0.912			
(mmHg <sup>-1</sup> )									
A3 (mmHg)	109±6	129±14*	0.005	123±8	121±17	0.757			
A4 (%)	35±20	22±13	0.131	14±13	24±10	0.062			
Max. gain	5.0±4.9	2.4±1.3	0.237	2.8±0.9	3.4±2.5	0.739			
(mmHg <sup>-1</sup> )									
Operating	27±23	53±25*	0.016	44±16	40±18	0.646			
range (mmHg)									
HR baroreflex									
	First co	hort		Second coh	ort				
A1 (beats/min)	59±25	65±44	0.732	88±31	92±40	0.843			
A2	$0.16 \pm 0.18$	$0.19{\pm}0.19$	0.673	$0.12 \pm 0.08$	$0.11 \pm 0.04$	0.674			
(mmHg <sup>-1</sup> )									
A3 (mmHg)	116±9	122±15	0.341	125±17	125±15	0.999			
A4 (beats/min)	370±46	410±59	0.139	387±38	397±76	0.707			
Max. gain	1.8±1.5	2.1±1.5	0.423	$2.2 \pm 0.8$	2.2±0.9	0.821			
(mmHg <sup>-1</sup> )									

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Data are expressed as mean±SD and represent baroreflex parameters of cohort 1 (sham, n=10; CIH, n=8) and cohort 2 (sham, n=10; CIH, n=10). Datawere analysed using
independent sample *t*-tests. \* p<0.05 vs. sham group. CIH, chronic intermittent hypoxia;</li>
RSNA, renal sympathetic nerve activity; HR, heart rate; A1, response range of
RSNA/HR; A2, the gain coefficient; A3, midpoint pressure of input; A4, the minimum
response.

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- 50

52	Table 4: Biomarker	concentrations in	kidney tissue	homogenates o	f sham and CIH

- 53 exposed rats.
- 54

Parameter	Sham	СІН	p-value
AOPP (µM)	15.6±2.2	14.8±3.1	0.728
	(n=12)	(n=8)	
NADPH oxidase activity (mU/min)	3.54±0.87	4.37±1.27	0.167
	(n=8)	(n=9)	
SOD activity (U/ml)	803.5±125.0	792.3±120.8	0.846
	(n=11)	(n=8)	
Catalase activity (U/ml)	15939±2888	17448±2539	0.105
	(n=12)	(n=8)	
TNF-α (pg/mg)	0.22±0.22	0.16±0.14	0.790
	(n=8)	(n=10)	
IL-1β (pg/mg)	5.04±1.34	6.15±1.31	0.070
	(n=11)	(n=10)	
Keratinocyte chemoattractant/growth	1.59±0.55	$1.98 \pm 0.78$	0.137
related oncogene (pg/mg)	(n=11)	(n=9)	

55

56 Data are expressed as mean $\pm$ SD. AOPP, advanced oxidation protein products; SOD, 57 superoxide dismutase. SOD activity, keratinocyte chemoattractant/growth related 58 oncogene and IL-1 $\beta$  concentrations were analysed using independent samples *t*-test. 59 Other biomarkers were analysed using Mann-Whitney test.

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Chronic intermittent hypoxia impairs diuretic and natriuretic responses to volume expansion in rats with preserved low-pressure baroreflex control of the kidney

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# OUTCOME

- **RSNA High-pressure** Baroreflex sensitivity
- Mid-point blood pressure



Low-pressure baroreflex control of RSNA

**CONCLUSION** Exposure to CIH blunts the high-pressure but not low-pressure baroreflex control of RSNA, most likely indirectly due to hypertension. CIH also impairs diuretic and natriuretic responses to fluid overload, which may involve Downloaded from journals in triag. renal/aj TRAPat/12 signaling 102.069) on November 19, 2020.