RESEARCH ARTICLE

The CORM ALF-186 Mediates Anti-Apoptotic Signaling via an Activation of the p38 MAPK after Ischemia and Reperfusion Injury in Retinal Ganglion Cells

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Abstract

Purpose

Ischemia and reperfusion injury may induce apoptosis and lead to sustained tissue damage and loss of function, especially in neuronal organs. While carbon monoxide is known to exert protective effects after various harmful events, the mechanism of carbon monoxide releasing molecules in neuronal tissue has not been investigated yet. We hypothesize that the carbon monoxide releasing molecule (CORM) ALF-186, administered after neuronal ischemia-reperfusion injury (IRI), counteracts retinal apoptosis and its involved signaling pathways and consecutively reduces neuronal tissue damage.

Methods

IRI was performed in rat’s retinae for 1 hour. The water-soluble CORM ALF-186 (10 mg/kg) was administered intravenously via a tail vein after reperfusion. After 24 and 48 hours, retinal tissue was harvested to analyze mRNA and protein expression of Bcl-2, Bax, Caspase-3, ERK1/2, p38 and JNK. Densities of fluorogold pre-labeled retinal ganglion cells (RGC) were analyzed 7 days after IRI. Immunohistochemistry was performed on retinal cross sections.

Results

ALF-186 significantly reduced IRI mediated loss of RGC. ALF-186 treatment differentially affected mitogen-activated protein kinases (MAPK) phosphorylation: ALF-186 activated p38 and suppressed ERK1/2 phosphorylation, while JNK remained unchanged. Furthermore, ALF-186 treatment affected mitochondrial apoptosis, decreasing pro-apoptotic Bax...
and Caspase-3-cleavage, but increasing anti-apoptotic Bcl-2. Inhibition of p38-MAPK using SB203580 reduced ALF-186 mediated anti-apoptotic effects.

Conclusion
In this study, ALF-186 mediated substantial neuroprotection, affecting intracellular apoptotic signaling, mainly via MAPK p38. CORMs may thus represent a promising therapeutic alternative treating neuronal IRI.

Introduction
As an organ with high demand of nutrients and oxygen, the human brain is very sensitive to hypoperfusion. In patients suffering from stroke, irreversible neuronal cell death may occur within minutes leading to substantial neuronal deficit and increased mortality.[1–4]

Similarly, retinal ischemia and reperfusion injury leads to neuronal cell death and plays an important role in the pathophysiology of several eye diseases such as diabetic retinopathy and glaucoma.[5, 6]

In the last years a multitude of substances have been tested pre-clinically regarding their neuroprotective potential.[7] However, all of them showed either a weak effect only, or failed to prove relevance in clinical practice at all.

In human organism, carbon monoxide (CO) is produced endogenously as one elimination product of heme oxygenase (HO), an enzyme that facilitates heme catabolism and catalyzes the degradation of heme to biliverdin, CO and ferrous ions.[8, 9] It has been shown that even exogenously applied CO may mediate protection in the context of ischemia and reperfusion. Although poisonous at high concentration, inhaled carbon monoxide has attracted attention as a potential therapeutic agent. Being administered after ischemia-reperfusion injury (IRI) and transplantation, CO appears to protect vital organs including heart, lung, kidney and liver.[10–13] To do so, CO interacts with a variety of physiological processes, among which some of them are responsible for apoptosis. As far as known, targets of CO action are the binding to soluble guanylate cyclase (sGC), stimulation of cGMP production, activation of calcium-dependent potassium channels and stimulation of mitogen-activated protein kinases.[14]

Due to the fact of difficulties in handling a gaseous–and dose-dependently toxic–agent and the negative effects of inhaled CO on oxygen transport, molecules containing a heavy-metal framework were created as carriers of covalently-bound CO, defined as carbon monoxide releasing molecules (CORM). These substances, administered either orally or intravenously ensure a constant liberation of CO and delivered a predictable and controllable amount of CO to organs and tissue.[15] The carbon monoxide releasing molecule ALF-186 presents a special group of CORMs, which is water-soluble and releases its bound CO molecules in a slower manner than other CORMs do.[16]

Although described in some organ systems, therapeutic effects of CORMs on neuronal structures following neuronal damage have not been investigated so far. In this in-vivo study we analyzed the impact of the CORM ALF-186 after neuronal IRI and the underlying mechanism. We hypothesized that ALF-186 treatment mediates anti-apoptotic effects and protects retinal ganglion cells after IRI via the p38 MAPK. In future, CO might be a promising therapeutic option reducing neuronal damage.
Materials and Methods

Animals

Adult male and female Sprague-Dawley rats (1:1, 280-350g bodyweight, Charles River, Sulzfeld, Germany) were used in these experiments. Animals were fed with a standard diet ad libitum, being kept on a 12-h light/12-h dark cycle. All procedures involving the animals concurred with the statement of The Association for Research in Vision and Ophthalmology for the use of animals in research and were approved a priori by the Committee of Animal Care of the University of Freiburg in accordance with the ARRIVE guidelines (Permit No: 35-9185.81/G-11/81). All types of surgery and manipulations were performed under general anesthesia with isoflurane/O₂ for retrograde labeling with fluorogold (FG) or with a mixture of intraperitoneally administered ketamine 50 mg/kg (Ceva-Sanofi, Duesseldorf, Germany) and xylazine 2 mg/kg (Ceva-Sanofi) for the ischemia-reperfusion experiment. Body temperature was maintained at 37°C ± 0.5°C with a heating pad. After surgery, buprenorphine (Temgesic 0.5 mg/kg; Essex Pharma, Solingen, Germany) was applied intraperitoneally to prevent pain. During recovery from anesthesia, the animals were placed in separate cages. The number of animals used for RGC quantification and molecular analysis was n = 8 per group. For analysis of mRNA- and protein-expression retinal tissue was harvested at t = 24 h and 48 h after reperfusion. (see flow-chart in Fig 1).

Retrograde labeling of RGC

Sprague Dawley rats were anesthetized, placed in a stereotactic apparatus (Stoelting, Kiel, Germany) and retrograde RGC-labeling was done as described previously, briefly summarized as follows: The skin overlying the skull was cut open and retracted. The lambda and bregma sutures served as landmarks for drilling three holes on each site of the bregma sutures. A total amount of 7.8 μl fluorogold (FG) (Fluorochrome, Denver, CO, USA) dissolved in DMSO/PBS was injected into both superior colliculi through the drilling holes. To ensure adequate retrograde transport of FG into the RGC’s perikarya, further experimental interventions were done 7 days after retrograde labeling.

Retinal ischemia/reperfusion injury and treatment with ALF-186

Following randomization, rats were sedated as previously described [17–19] and the anterior chamber of the left eye was cannulated with a 30-gauge needle connected to a reservoir containing 0.9% NaCl. Intraocular pressure was increased to 120 mmHg for 60 minutes and ocular ischemia was confirmed microscopically by interruption of the retinal circulation. Reperfusion was initiated by removing the needle tip promptly. Rats without immediate recovery of retinal perfusion at the end of the ischemic period or those with lens injuries were excluded from the study, since the latter prevents RGC death and promotes axonal regeneration. To evaluate possible neuroprotective effects of the treatment with carbon monoxide, released from ALF-186, animals were randomly assigned to receive either treatment with ALF-186 (10 mg/kg body weight i.v., dissolved in PBS) or PBS (vehicle control) alone. Both treatment options were injected intravenously, either immediately or with a delay of 3 hours after IRI. To exclude that the molybdenum-containing metal backbone of ALF-186 exerts protective or crucial effects itself, inactivated ALF-186 (iALF-186) was injected 24 hours after dissolving. In a later set of experiments, rats received the p38 MAPK inhibitor SB203580 (1 mg/kg bodyweight i.v., Sigma-Aldrich, #S8307) sixty minutes prior to IRI and ALF-186 treatment.
RGC quantification

Animals were sacrificed by CO$_2$-inhalation 7 days after ischemia. Retinal tissue was immediately harvested, placed in ice-cold Hank’s balanced salt solution and further processed for whole mount preparation. Retinae were carefully placed on a nitrocellulose membrane with the ganglion cell layer (GCL) on top. After removing the vitreous body, retinae were fixed in 4% paraformaldehyde for 1 h and then embedded in mounting media (Vectorshied; Axxora, Loerrach, Germany). The densities of FG-positive RGC were determined in blinded fashion using a fluorescence microscope (AxioImager; Carl Zeiss, Jena, Germany) and the appropriate bandpass emission filter (FG: excitation/emission, 331/418 nm), as previously described. Briefly, we photographed 3 standard rectangular areas (0.200 mm x 0.200 mm = 0.04 mm$^2$) at 1, 2 and 3 mm from the optic disc in the central regions of each retinal quadrant. Hence, we evaluated an area of 0.48 mm$^2$ per retina. To calculate the average RGC density in cells/mm$^2$, we multiplied the number of analyzed cells/0.04 mm$^2$ with 25. Secondary FG stained activated microglia cells (AMC) after RGC phagocytosis were identified by morphologic criteria and excluded from calculation. All data are presented as mean RGC densities [cells/mm$^2$] ± SD.
Immunohistochemical staining

Immunohistochemistry was performed to identify p38 expressing cells in the retina 24 h after intervention. Rat eyes (n = 2 per group) were enucleated and immediately fixed in 4% paraformaldehyde overnight at 4°C. After washing in Dulbecco’s phosphate buffered saline before and after post-fixation in 30% sucrose for 5–6 h at 22°C, eyes were embedded in compound (Tissue-Tek; Sakura-Finetek, Torrance, CA, USA), and frozen in liquid nitrogen. Frozen sections (10 μm) were cut through the middle third of the eye and collected on gelatinized slides. Double immunolabeling was performed according to standardized protocols with monoclonal antibodies brain-specific homeobox/POU domain protein 3a Brn-3a (sc-390078; dilution 1:200; Santa Cruz Biotechnology; Dallas, TX, USA) with p38 (ab7952, dilution 1:200; Abcam, Cambridge, UK). Primary antibodies were then conjugated with their corresponding secondary antibody (red fluorescence: Alexa Fluor 647, mouse anti-rabbit, dilution 1:1000, Jackson ImmunoResearch Europe, 211-605-109; green fluorescence: Cyanine Cy™2, donkey anti-goat, dilution 1:1000, Jackson ImmunoResearch Europe, 705-225-147). The nuclei of cells in the retina were stained with 4´,6-diamino-2-phenylindole dihydrochloride hydrate (DAPI, Sigma, Taufkirchen, Germany) added to the embedding medium (Mowiol; Calbiochem, San Diego, CA, USA). Slides were examined under a fluorescence microscope (Axiophot; Carl Zeiss, Jena, Germany).

Western blot analysis

24 h and 48 h after ischemia retinal tissue for analysis of protein expression was harvested. Total protein from ¼ of retina was extracted and processed for Western Blot as described previously. The membranes were blocked with 5% skim milk in Tween20/PBS and incubated in the recommended dilution of protein specific antibody (p-ERK1/2 #4370, p-p38 #9211, p-JNK #9251, cleaved Caspase-3 #9664, Bax #2722, Bcl-2 #2876, Cell Signaling Technology, Danvers, MA, USA) overnight at 4°C. After incubation with a horseradish peroxidase-conjugated anti-rabbit secondary antibody (GE Healthcare, Freiburg, Germany), proteins were visualized using the ECL plus Chemiluminescence Kit (GE Healthcare). For normalization, blots were re-probed with ERK1/2 (#4695), p38 (#9212), JNK (#9258), Caspase-3 (#9665) and β-Actin (#4967S, all Cell Signaling Technology, Danvers, MA, USA). Relative changes in protein expression in IR injured retinas either with or without ALF-186 were calculated in relation to the corresponding non-ischemic retinas.

Real time polymerase chain reaction (RT-PCR)

From retinal tissue harvested 24 h and 48 h after ischemia, total RNA from ¼ of retina was extracted using a column-purification based kit (RNeasy Micro Kit, Qiagen, Hilden, Germany) according to the manufacturer’s instructions. Reverse transcription was performed with 50 ng of total RNA using random primers (High Capacity cDNA Reverse Transcription Kit, Applied Biosystems, Darmstadt, Germany). Real time polymerase chain reactions (RT-PCR) were done with TaqMan® probe-based detection kit (TaqMan® PCR universal mastermix, Applied Biosystems, Darmstadt, Germany). Following primers were used: Caspase-3 #Rn00563902_m1, Bax #Rn02532082_g1 and Bcl-2 #Rn99999125_m1 (all from Applied Biosystems, Darmstadt, Germany). The PCR assays were then performed on a RT-PCR System (ABI Prism 7000, Applied Biosystems, Darmstadt, Germany) with the following cycling conditions: 95°C for 10 min, 40 cycles of 95°C for 10 sec and 60°C for 1 min. Reaction specificity was confirmed by running appropriate negative controls. Cycle threshold (CT) values for each gene of interest were normalized to the corresponding CT values for GAPDH (∆CT). Relative gene expression in IR injured retinal tissue either with injection of ALF-186 or PBS was calculated in relation to the corresponding gene expression in the non-injured retinal tissue of each individual animal (∆∆CT).
Statistical analysis
Data was analyzed using a computerized statistical program (SigmaPlot Version 11.0, Systat Software Inc., San Jose, CA, USA). The results are presented as means (±SD) after normal distribution of data had been verified. One-way ANOVA for repeated measurements was used for between-group comparisons with post hoc Holm-Sidak test and Kruskal Wallis test for data with lack of normal distribution. A p-value < 0.05 was considered statistically significant.

Results
ALF-186 treatment protects retinal ganglion cells against ischemia reperfusion injury (IRI) in a time-dependent manner
To answer the question of the influence of ALF-186 regarding retinal ganglion cell survival, the amount of vital neuronal cells was quantified. Retinal IRI damage caused a loss of retinal ganglion cells of approximately 40% (Fig 2A respective image and Fig 2B: columns 1 and 2: IRI 1808±262 vs. untreated 2503±295; *** = p<0.001). Immediate (0 h) treatment with ALF-186 almost completely neutralized this effect (Fig 2A and 2B, column 3: IRI 1808±262 vs. IRI +ALF-186 0 h 2499±377; *** = p<0.001). Even delayed ALF-186 application (3 h) reduced...
RGC loss significantly (IRI 1808±262 vs. IRI±ALF-186 3h 2202±239; p*** = <0.001). In contrast, administration of inactivated ALF-186 (iALF) did not result in a change of RGC compared to untreated animals. The p38 inhibitor SB203580, injected prior to retinal IRI, abolished ALF’s effect in part (Fig 2A and 2B, column 7: IRI+ALF-186 2499±377 vs. SB203580 +IRI+ALF-186 2133±101; * = p<0.05). SB203580 alone showed no effect on ganglion cell loss (IRI 1808±262 vs. IRI+SB203580 1816±162).

**ALF-186 suppresses phosphorylation of ERK1/2, while inducing phosphorylation of p38 after IRI**

To answer the question if intracellular signaling mechanisms of apoptosis are involved, we analyzed the effect of ALF-186 on the phosphorylation of MAPK. After IRI, treatment with ALF-186 decreased IRI-induced ERK1/2 phosphorylation. The extent depended on the time-point of enucleation, with a stronger effect at 24 than 48 hours after IRI (Fig 3A and 3B; IRI 1.23±0.14 vs. IRI+ALF-186 0.87±0.21 fold change at 24 h; ** = p<0.01; IRI 1.16±0.16 vs. IRI+ALF-186 0.95±0.17 fold change at 48 h; * = p<0.05). In contrast, no ALF-186 mediated effect was detectable for JNK-phosphorylation (Fig 3C and 3D; IRI 1.14±0.23 vs. IRI+ALF-186 1.03±0.28 fold change at 24 h; IRI 0.94±0.09 vs. IRI+ALF-186 0.90±0.12 fold change at 48 h).

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**Fig 3. Effect of ALF-186 treatment on the phosphorylation of the mitogen-activated protein (MAP) kinases ERK1/2 and JNK after IRI.** (A) Representative western blot image (n = 8) showing the suppression of the phosphorylated MAPK ERK1/2 compared to total ERK1/2 after immediate ALF-186 treatment. Enucleation was either performed 24 hours or 48 hours after IRI. (B) Densitometric analysis (n = 8) of western blots for phosphorylated ERK1/2 compared to its total protein formation 24 and 48 hours after IRI+ALF-186 (data are means±SD; IRI vs. IRI+ALF-186 24h, ** = p<0.01 and IRI vs. IRI+ALF-186 48h, * = p<0.05). (C) Representative western blot image (n = 8) showing no effect of the phosphorylated MAPK JNK compared to total JNK after immediate ALF-186 treatment. Enucleation was either performed 24 hours or 48 hours after IRI. (D) Densitometric analysis (n = 8) of western blots for phosphorylated JNK compared to its total protein formation 24 and 48 hours after IRI+ALF-186.

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Instead, ALF-186 significantly induced p38-phosphorylation. Again, we observed a more pronounced effect at 24 h after IRI (Fig 4A and 4B; IRI 1.03±0.12 vs. IRI+ALF-186 1.42±0.15 fold change at 24 h; *** = p<0.001; IRI 1.06±0.13 vs. IRI+ALF-186+1.20±0.08 fold change at 48 h; * = p<0.05). Inactivated ALF-186 did not affect p38 phosphorylation (Fig 4C and 4D; IRI 1.05±0.09 vs. IRI+ALF-186 0.95±0.13). To confirm ALF’s mechanism of action regarding the involvement of p38, we next injected the p38 inhibitor SB203580 intravenously prior to IRI. While SB203580 had no influence on p38-phosphorylation after IRI, its presence significantly reduced phosphorylation of p38 in the context of ALF-186. This effect was quantified densitometrically, showing that SB203580 abolished the ALF-186 mediated increase of p38-phosphorylation completely (Fig 4C and 4D; IRI 0.90±0.35 vs. SB203580+IRI+ALF-186 0.90±0.32).

ALF-186 inhibits IRI-induced Caspase-3 mRNA expression and Cleavage

To study the extent to which ALF-186 influences apoptosis, we next analyzed Caspase-3 mRNA expression and cleavage.
ALF-186 treatment decreased IRI-induced Caspase-3 mRNA expression significantly (Fig 5A; IRI 1.80±0.14 vs. IRI+ALF-186 1.33±0.14; ** = p<0.01). P38 inhibition prior to IRI and ALF-186 treatment abrogated ALF-186 mediated reduction of Caspase-3 mRNA (Fig 5A; IRI+ALF-186 1.33±0.14 vs. IRI+SB203580+ALF-186 1.86±0.14; * = p<0.05), while the inhibitor alone had no effect.

ALF-186 affects Caspase-3 cleavage in a similar way. Animals treated with ALF-186 showed reduced Caspase-3 cleavage after IRI (Fig 5B and 5C; IRI 1.39±0.11 vs. IRI+ALF-186 1.06±0.17; * = p<0.05 and IRI+ALF-186 vs. IRI+SB203580+ALF-186, * = p<0.05). P38 inhibition prior to IRI and ALF-186 treatment abrogated ALF-186 mediated reduction of Caspase-3 cleavage (Fig 5B and 5C; IRI+ALF-186 1.06±0.17 vs. IRI+SB203580+ALF-186 1.33±0.14; ** = p<0.01).
1.07±0.16; * = p<0.05). SB203580 significantly reduced ALF’s effect (Fig 5B, 1.07±0.16 vs. 1.39±0.15; * = p<0.05).

**ALF-186 reduces mRNA and protein expression of Bax**

To further analyze the role of ALF-186 in apoptosis, we evaluated the effect of ALF-186 on Bax, a protein enhancing apoptosis. Treatment with ALF-186 significantly decreased IRI induced Bax mRNA (Fig 5D; IRI 1.54±0.17 vs. IRI+ALF-186 1.14±0.18; * = p<0.05) as well as protein expression (Fig 5E; IRI 1.50±0.38 vs. IRI+ALF-186 0.89±0.06; * = p<0.05). P38 inhibition prior to IRI and ALF-186 treatment significantly increased Bax mRNA and protein expression again (Fig 5D; IRI+ALF-186 0.89±0.06 vs. SB203580+IRI+ALF-186 1.43±0.16; * = p<0.05; Fig 5F; IRI+ALF-186 0.89±0.06 vs. SB203580+IRI+ALF-186 1.42±0.20; * = p<0.05), while the inhibitor alone had no effect.

**ALF-186 increases mRNA and protein expression of Bcl-2**

Moreover, Bcl-2, a protein known to inhibit apoptosis, was analyzed to answer the question of ALF-186 influence on apoptosis. ALF-186 treatment significantly increased Bcl-2 mRNA (Fig 6A; IRI 1.07±0.16 vs. IRI+ALF-186 1.91±0.18; ** = p<0.001) and protein expression (Fig 6B and 6C; n = 6; IRI 0.79±0.14 vs. IRI+ALF-186 1.16±0.16; ** = p<0.01) after IRI. This effect was reduced significantly by p38 inhibition (Fig 6A; IRI+ALF-186 1.91±0.18 vs. SB203085+IRI+ALF-186 1.29±0.14, ** = p<0.001; Fig 6B and 6C; IRI+ALF-186 1.16±0.16 vs. SB203085+IRI+ALF-186 0.84±0.06; * = p<0.05), while the inhibitor alone had no effect.

**ALF-186 induced p38 expression in the ganglion cell layer and inner nuclear layer**

Retinal cross sections were evaluated for p38 immunoreactivity to answer the question, which cells were responsible for p38 upregulation after IRI+ALF. In untreated eyes and after IRI, there was only a weak p38 expression pronounced in the area of the ganglion cell layer (GCL) (Fig 7A, 7D and 7G). Co-staining with Brn3a (B, E, H) revealed a low degree of double-immunoreactivity with RGCs, the p38 signal was seen in a low level in the area of the nerve fibers on top of the GCL (C, F, I). ALF treatment directly after IRI induced upregulation of p38 (J) in the GCL and inner nuclear layer (INL). Double-immunoreactivity with Brn3a (L and Fig 8) was partially seen. Scale bar in L is 100 μm.

**Discussion**

The main findings of this *in vivo* study can be summarized as follows: (1) Treatment with the carbon monoxide releasing molecule (CORM) ALF-186 alleviated loss of retinal ganglion cells due to ischemia reperfusion injury (IRI). (2) ALF-186 mediates anti-apoptotic signaling via p38, but not extracellular regulated kinases (ERK1/2) and c-Jun N-terminal kinases (JNK). (3) ALF-186 mediated anti-apoptotic signaling, decreasing caspase-3 and the pro-apoptotic protein Bax, while increasing anti-apoptotic proteins like Bcl-2. (4) Inhibition of p38 MAPK counteracted ALF’s protective effect in retinal IRI. (5) ALF’s effect on molecular mechanisms predominantly proceeded within the first 24h showing a marked fading thereafter.

Carbon monoxide (CO) exerts a variety of pharmacological effects, mediating among others anti-apoptotic effects in the cardiovascular, immune and nervous systems.[20, 21] Due to difficulty handling in treatment with the potentially toxic volatile agent CO, carbon monoxide releasing molecules have attracted attention. These molecules contain metal carbonyls like manganese, ruthenium, boron and iron as CO carrier leading to a constant and reliable release.
Fig 6. Effect of ALF-186 treatment on retinal protein and mRNA expression of Bcl-2. (A) Fold induction of Bcl-2 mRNA after IRI+ALF-186 and p38 inhibition in ischemic retinal tissue compared to GAPDH in relation to the corresponding non-ischemic retinae analyzed by RT-PCR (n = 8; data are mean±SD; IRI vs. IRI+ALF-186, *** = p<0.001 and IRI+ALF-186 vs. SB203580+IRI+ALF-186, *** = p<0.001). (E) Representative western blot image showing no change of Bcl-2 due to IRI (lane 2) but its strong induction (lane 4) compared to β-Actin after ALF-186 treatment. This effect was abrogated by p38 inhibition using SB203580 (lane 6). (F) Densitometric analysis of n = 8 western blots for Bax expression (data are mean±SD; IRI vs. IRI+ALF-186, ** = p<0.01 and IRI+ALF-186 vs. SB203580+IRI+ALF-186, * = p<0.05).

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Fig 7. ALF-186 induced p38 MAPK in retinal ganglion cell layer. Retinal cross sections were evaluated for p38 immunoreactivity to answer the question, which cells were responsible for p38 upregulation after IRI+ALF. In controls and after IRI+PBS, there was only a weak p38 expression, pronounced in the area of the ganglion cell layer (GCL) (Fig 5, right side, A, D, G). Co-staining with Brn3a (B, E, H) revealed a low degree of double-immunoreactivity, the p38 signal was seen in a low level in the area of the nerve fibers on top of the GCL (C, F, I). ALF treatment directly after IRI induced upregulation of p38 (J) in the GCL and inner nuclear layer (INL), as expected from the results of the p38 mRNA expression. Double-immunoreactivity with Brn3a (K+L) was more frequent and more intense but not exclusively expressed in RGCs. Scale bar in L is 100 μm.

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Since water-soluble substances are available, oral or intravenous application simplified CO treatment avoiding environmental impact and potential poisoning.

In this in-vivo study, we have examined the potential protective effect of ALF-186 on neuronal tissue. Hence, we chose a well-established model of retinal IRI in rats.[17–19] We found that treatment with ALF-186 reduced IRI mediated loss of retinal ganglion cells and therefore mediated protection. While protective effects of carbon monoxide are well studied in a variety of damage models of non-neuronal organs, studies describing the protective effect of CO and CORM on neuronal tissue are limited.[22–26] Zeynalov and Doré demonstrated that inhalative carbon monoxide reduced infarct size after focal transient brain ischemia in a mouse model.[27] Wang et al. also exposed mice to carbon monoxide after middle cerebral artery occlusion and described a reduced infarct volume.[28] The neuroprotective effects in this study are comparable to our previous findings.[17]

Intracellular signaling via MAPK ERK1/2, p38 and JNK plays a significant role in the regulation of neuronal [29–31] and retinal [32–34] apoptosis. However, the effects of ERK1/2 phosphorylation are considered to be contradictory: both survival and death signals can activate ERK1/2 and therefore activation itself may result in neuronal death or survival.[29] Cellular stress triggers MAPK p38 and JNK activation, for example in the context of cerebral ischemia. [35] It is assumed that p38 and JNK mediated apoptosis is transmitted by mitochondrial pro-apoptotic proteins like Bax and Bcl-2.[35]

In our study we have found that ALF 186 modulated MAPK differently. ALF treatment resulted in a decreased ERK1/2 phosphorylation and, on contrary, increased p38 activation while JNK phosphorylation did not change. Following cell and tissue damage, the influence of carbon monoxide on MAPK is a topic of controversy in the literature whereby it is important to note that the type of injury differently affected the MAPK. Our findings are in accordance to the results of Sethi et al., who exposed primary pulmonary artery endothelial cells of the rat to carbon monoxide after TNFα stimulation. Injured cells presented increased ERK1/2 and JNK activation in contrast to p38 suppression. Application of CO reversed TNFα mediated effects on ERK1/2 and p38 while CO had no influence on JNK.[36] Li et al. describe in their study the p38 MAPK pathway as a key signaling mechanism of CO´s protective effect. In primary hepatocytes of rats´ ethanol increased p38 phosphorylation while additional CORM further stimulated p38 phosphorylation. JNK and ERK1/2 were not affected.[37] The cytoprotective effect of ERK1/2 downregulation was described by Choi et al. Glucose deprivation led to an increase of
ERK1/2 and p38 phosphorylation in BNL CL.2 cells whereas JNK was not affected. In contrast to our findings, the presence of CORM only suppressed ERK1/2 phosphorylation.[38] Ning et al. showed in their in-vitro study that stimulating lung cells, with IL16 only affected MAPK ERK1/2, but neither p38 nor JNK, and increased levels of phosphorylated ERK1/2. Exposure to CO (250 ppm) mitigated this effect.[39] Similarly, CO treatment suppressed elevated ERK1/2 phosphorylation in liver grafts after cold IRI due to orthotopic liver transplantation in rats. Although CO also decreased p38 phosphorylation, the influence of CO was identified as not significant.[13] Our research group showed that CORM increased p38 phosphorylation in staurosporine treated EBL cells, but not ERK1/2 nor JNK.[40] In another study, postconditioning with inhalative CO (250 ppm) showed similar effects on retinal ganglion cells with an increase of cell count after IRI. Interestingly, an opposite effect on MAPK was demonstrated. [19] Inhalative CO reduced p38 phosphorylation and induced ERK1/2 activation while JNK was not affected. The reason for this might be the different way of administration.

Due to neuronal injury, neurons release molecules interacting cell survival and cell death signaling. In particular, Bcl-2 family proteins like Bax and Bcl-2 are involved in IRL.[41]
Thereby, Bcl-2 promotes neuronal survival modulating Ca$^{2+}$ signaling and Bax operates as an apoptotic stimulus. Shifting this balanced Bax/Bcl-2 ratio towards Bax causes mitochondrial damage that activates the caspase cascade.[42]

Our study demonstrated, that ALF-186 reduced IRI induced apoptosis by inhibition of Bax expression, activation of Bcl-2 and, consecutively, by suppression of caspase-3 cleavage. In part, these results are in agreement with Schallner et al.: Inhalation of CO prior to IRI caused a decrease of Bax and caspase-3 cleavage while Bcl-2 was not affected.[19] Cheng and Levy also reported a beneficial effect of carbon monoxide on neuronal tissue. In their study, subclinical CO decreased caspase-3 activation in mice pulps following isoflurane exposition.[43] In a study with endothelial lung cells Wang et al. described that CO inhibited hyperoxia induced cell death by decreasing pro-apoptotic Bax.[44] Nakao et al. examined the effect of CO treatment on intestinal apoptosis due to IRI in rats. CO inhalation reduced Bax and upregulated Bcl-2.[45]

Pretreatment with p38 inhibitor SB203580 reduced the impact of ALF-186 on retinal apoptosis. We have found that p38 inhibition resulted in an increase of caspase-3 cleavage and Bax expression and also in a decrease of Bcl-2 expression. Furthermore, p38 inhibition reversed ALF-186 mediated p38 phosphorylation, reduced neuronal ganglion cell count and cancelled neuroprotection. We focused on this proposed mechanism as part of the carbon monoxide mediated neuroprotective effect in Fig 9. Although neuroprotective effects are described [46], in our study SB203580 alone did not affect retinal apoptosis. This is also confirmed by Mori et al., who did not notice protection in traumatic brain injury after treatment with SB203580.[47]

In summary, we demonstrate that the treatment with ALF-186 as a therapeutic option reduced retinal apoptosis after IRI in-vivo mainly in the first 24 hours after neuronal damage. ALF186 affected and modulated intracellular MAPK and apoptosis signaling and reduced loss of retinal ganglion cells, therefore representing a novel and promising alternative in treating ischemia-related injury of neuronal organs.

Author Contributions

Conceptualization: JB UG.

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