A Complete Review of Preclinical and Clinical Uses of the Noble Gas Argon: Evidence of Safety and Protection

Abstract
The noble gas argon (Ar) is a “biologically” active element and has been extensively studied preclinically for its organ protection properties. This work reviews all preclinical studies employing Ar and describes the clinical uses reported in literature, analyzing 55 pertinent articles found by means of a search on PubMed and Embase. Ventilation with Ar has been tested in different models of acute disease at concentrations ranging from 20% to 80% and for durations between a few minutes up to days. Overall, lesser cell death, smaller infarct size, and better functional recovery after ischemia have been repeatedly observed. Modulation of the molecular pathways involved in cell survival, with resulting anti-apoptotic and pro-survival effects, appeared as the determinant mechanism by which Ar fulfills its protective role. These beneficial effects have been reported regardless of onset and duration of Ar exposure, especially after cardiac arrest. In addition, ventilation with Ar was safe both in animals and humans. Thus, preclinical and clinical data support future clinical studies on the role of inhalatory Ar as an organ protector.

Keywords: Argon, cardiac arrest, ischemia-reperfusion injury, neuroprotection

Introduction
Argon (Ar) is a noble gas discovered in 1894.[i] Ar is a colorless, tasteless, odorless, noncorrosive, nonflammable, and nontoxic gas.[ii] It is the most abundant noble gas, and the third major component of the air, with a concentration of 0.93%. Ar’s density is 38% higher than air, and the solubility in water and plasma is 24-fold lower than that of carbon dioxide (CO₂).[iii]

Ar has a full electron valence shell which avoids covalent binding with other elements; therefore, it is usually considered a nonreactive chemical gas, as known by its Greek name “αργός,” meaning “inert.”[iv] Although labeled as a “biologically” inert gas, recent evidence suggests that the drug can have significant effects. Narcotic effects have been described in divers since 1939 and further confirmed by other studies.[v‑10] Besides narcosis, evidence of neuroprotection has been rising since the first report in a rodent model of hypoxic brain injury, in 1998.[xi] In more recent years, the protective effects of Ar after ischemia became more evident preclinically.

The aim of this work was to review all the preclinical studies employing Ar, both in vivo and in vitro, focused on organ protection, and its clinical uses.

Materials and Methods
A concurrent PubMed and Embase search was carried out and updated till October 2017. “Ar” was used as MeSh and Emtree term for PubMed and Embase search, respectively. The following keywords were then used to narrow and focus the search: “neuroprotection,” “cardioprotection,” “organ protection,” “cellular protection,” “organ transplantation,” “ischemia-reperfusion injury,” “hypoxic-ischemic injury,” “perinatal asphyxia,” “cardiac arrest (CA),” “traumatic brain injury,” “pneumoperitoneum,” “lung volume measurement,” “cardiac output measurement,” “narcosis,” “diving,” and “hyperbarism.” During the selection, only English-written articles with full text available were considered. Both in vitro and in vivo studies and human application of Ar were included in this review. Articles concerning the surgical use of Ar, i.e., “Ar laser,” “Ar beam,” and “Ar coagulation” were excluded. Articles selection was performed independently.

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by two authors (FN, SR). Fifty-five pertinent articles were found and included. Original results from a study performed by our group on a rat model of acute myocardial infarction (MI) were added.

**Preclinical uses**

Forty-four preclinical studies were found. The majority ($n = 25$) of in vitro and in vivo studies with Ar focused on neuroprotection after a variety of brain damages, in which different neurological and behavioral tests were combined with histological and biochemical assays. Four studies evaluated the presence of cardioprotective effect after myocardial ischemia/reperfusion (I/R). Nine studies focused on the preservation of other organs and tissues, such as ex vivo perfused kidney and lung, and culture of human airway epithelium, kidney tubular, and osteosarcoma cells. A single study evaluated the influence of Ar inhalation on liver regeneration after partial heptectomy. Three in vivo studies investigated the effects and the mechanisms of action of Ar under hyperbaric condition. Five in vivo studies specifically focused on the safety: In 2, Ar was administered as respiratory mixture to pigs, while, in 3 studies, Ar was used to induce pneumoperitoneum for surgical laparoscopic procedures in place of CO$_2$. The area of interest is summarized in Figure 1a.

Overall, both in vitro and in vivo studies provided evidence in support of a protective role of Ar, independently from the disease model and the duration and onset of exposure to the treatment (i.e., pre- or post-acute event) [Figure 1b].

**In vitro studies**

Seventeen articles investigated the protective effects of Ar on cell damage [Table 1]. In details, two studies were performed in rodent cardiomyocytes, and one of these included also additional investigations in human right atrial appendages obtained from patients undergoing coronary artery bypass or valve replacement surgery. Nine in rat neurons or hippocampal slices in human cells (neuroblastoma, kidney, airway epithelium, osteosarcoma) and one in whole blood from rodents. One in vitro study investigated whether Ar interacted with tissue plasminogen activator (tPA). The duration of the imposed insult varied from 30 min of hypoxia in cardiomyocytes to several hours under oxygen glucose deprivation condition, up to 3 h in the drug-induced toxicity studies.

In these studies, cells and/or tissues were incubated in chambers containing Ar atmosphere in a concentration varying from 25% to 75% in air and/or O$_2$. The average Ar exposure was 90 min, ranging from a minimum of 5 min up to 3 h later. In the majority of the studies, Ar was administered either immediately after the onset of the insult or delayed, up to 3 h later. In 3 studies, Ar was administered as a preconditioning agent.

Overall, the in vitro studies showed improved cell survival and viability when Ar was used both pre and after the insult. More specifically, Ar reduced apoptosis in neuronal, tubular kidney, and airway epithelial cells. As for cardioprotection after I/R, besides increasing cell viability, Ar also diminished early depolarizations in the ventricle tissue, limiting the onset of ischemia-triggered arrhythmias. When Ar was applied on the human right atrial appendages after hypoxia/reoxygenation injury, the recovery of contractile activity was increased.

**In vivo studies**

Twenty-four studies employed Ar as a potential protective treatment for brain, heart, and other organ preservation after different ischemic insults. Six studies used swine models and 18 rodent ones, including rats, guinea-pigs, and rabbits. In addition, three studies investigated the effects of Ar under hyperbaric conditions. A single study evaluated the effects on the initiation of liver regeneration in rats.
### Table 1: In vitro studies employing argon for cell and tissue preservation

| Article | Cell/tissue | Model | Ar treatment | Histopathology/functionality | Biochemistry |
|---------|-------------|-------|--------------|------------------------------|--------------|
| Lemoine et al., 2017[17] | 1. Cardiomyocytes  
2. Human right atrial appendage | H/R | Ar 70%  
Duration: 5 min | EAD: ↑  
Conduction blocks: ↑  
Reentries: ↑  
Contractile force: ↑ | NA |
| Mayer et al., 2016[18] | Cardiomyocytes | Hypoxia + Ar preconditioning | Ar 50%  
Duration: 1 h | Cell survival: ↑ | HSP27: +  
SOD2: +  
VEGF: +  
iNOS: +  
COX2: ≈  
STAT3: ≈  
HSPA4: ≈  
HIF1α: ≈  
ERK1-2 phosphorylation: ≈  
Akt: ≈ |
| Hafner et al., 2016[19] | Airway epithelial cells | N₂O₂ oxidative stress + Ar preconditioning | Ar 30-50%  
Duration: 30-180 min | Apoptosis: ↑  
Cell viability: ↑ | JNK: +  
p38: +  
ERK 1-2: +  
Akt: ≈ |
| Ulbrich et al., 2015[20] | Human neuroblastoma cells | Drug-induced cell injury (rotenone) | Ar 25-75%  
Duration: 2-4 h | Apoptosis: ↑ | TLR 2 - 4: -  
IRAK 4: -  
ERK1 - 2: +  
Caspase-3: -  
p-AKT: ≈  
HIF1α: ≈  
Bel 2: ≈  
Mitocondrial ΔΨ: ↑  
ROS: -  
NF-kB: -  
STAT3: -  
IL-8: -  
Caspase 3: -  
Cytochrome c: -  
Mitocondrial ΔΨ: ↑  
Phospho-ERK 1-2: ↑  
IL 1β: ≈  
TNFα: ≈  
IL 6: ≈  
Cell reducing ability: ↑ |
| Rizvi et al., 2010[21] | Human tubular kidney cells | OGD + Ar preconditioning | Ar 75%  
Duration: 3 h | Cell viability: ≈ | NA |
| Ulbrich et al., 2016[22] | Human neuroblastoma cells | Drug-induced cell injury (rotenone) | Ar 75%  
Duration: 2 h | NA | NA |
| Spaggiari et al., 2013[23] | Human osteosarcoma cells | Drug-induced cell injury (STS, MTX, rotenone, antimycin A, menadione) | Ar 75%  
Duration: 6-16 h | NA | NA |
| Fahlenkamp et al., 2012[24] | Neuronal, astroglial cells, and microglial cells | Selective influence of Ar on ERK 1-2 | Ar 50%  
Duration: 15-120 min | NA | NA |
| Jawad et al., 2009[25] | Neuronal cells | OGD+hypoxia | Ar 75%  
Duration: 24 h  
Ar 25-74%  
Duration: 72 h  
Ar 25-75%  
Duration: 3 h  
Ar 70% + MTH  
Duration: 2h | Cell viability: ↑  
OGD and TBI tissue injury: ↑  
Neuronal death: ↑  
HO-1: +  
p-Akt: + | NA |
| Loetscher et al., 2009[26] | Hippocampal slices | 1. OGD  
2. TBI above CA1 | OGD | LDH release: - | NA |
| David et al., 2012[27] | Brain coronal slices | OGD | OGD | NA |
| Zhao et al., 2016[28] | Rat cortical neuronal cultures | OGD | NA |

Contd...
In the majority of the studies, Ar was applied as a potential treatment and thus administered either immediately after the insult, i.e., after resuscitation from CA,\[13\] or delayed up to 3 h later.\[14\] Only in 3 studies specifically focused on myocardial I/R, EVLP, and liver regeneration, Ar was employed as a preconditioning agent, before induction of the insult.\[45,52,53\]

Regarding the use of Ar as neuroprotective agent, only 4 studies\[27,34,35,38\] evaluated the postinsult neurological recovery, and in half of them\[35,37\] there was evidence for a better outcome after Ar inhalation when compared to the control.

All the investigations, except one focusing on the safety of Ar,\[36\] included histological evaluations, in different models and tissues.

**Table 1: Contd...**

| Article, Year | Cell/tissue, Model | Ar treatment | Histopathology/functionality | Biochemistry |
|--------------|-------------------|-------------|-------------------------------|--------------|
| Zhao et al., 2016\[29\] | Rat cortical neuronal cultures, OGD | Ar 70% | Cell viability: ↑ | PI-3K: + |
| | | Duration: 2 h | Erk 1/2: + | |
| | | | mTOR: + | |
| | | | Nrf2: + | |
| | | | ROS: - | |
| Harris et al., 2013\[30\] | Mouse hippocampal slices, Focal mechanical trauma | Ar 33% | Cell injury (PI): ↑ | NA |
| | | Duration: 30 mins-24 h | Number of OHC: ↑ | |
| | | | Number of IHC: ↑ | NA |
| Yarin et al., 2005\[31\] | Rat organ of Corti cultures, 1. Hypoxia, 2. Drug-induced cell injury (cisplatin), 3. Drug-induced cell injury (gentamycin) | Ar 95% | 1. Ar 95% | NA |
| | | Duration: 30 h | Number of OHC: ↑ | |
| | | | Number of IHC: ↑ | |
| | | 2. and 3. Ar 74% | 2. Number of OHC: ↑ | |
| | | Duration: 48 h | Number of IHC: ≈ | |
| | | | 3. Number of OHC: ↑ | |
| | | | Number of IHC: ↑ | |
| David et al., 2013\[32\] | Enzymatic and thrombolytic efficiency of tPA, 1. Catalytic efficiency: tPA + specific chromogenic substrate + distilled water saturated solution, 2. Thrombolytic efficiency | Ar 25-75% | NA | 1. Catalytic efficacy |
| | | Duration: 120 min | Ar 75%: + | |
| | | | Ar 50: ≈ | |
| | | | Ar 25%: - | |
| | | | 2. Thrombolytic efficacy | Ar 75%: + |
| | | | Ar 50: ≈ | |
| | | | Ar 30%: - | |
| Grüßer et al., 2017\[33\] | Hippocampal slices, TBI | Ar 50% | Cell survival≈ | NA |
| | | Duration: 2 h | 1. Catalytic efficacy | |
| | | | Ar 75%: + | |
| | | | Ar 50: ≈ | |
| | | | Ar 25%: - | |

↑: Treatment with Ar was significantly cell protective compared to control; ≈: Treatment with Ar was equivalent to control; +: Increased with Ar treatment compared to control; -: Decreased with Ar treatment compared to control, Ar: Argon, O\(_2\): Oxygen, CO\(_2\): Carbon dioxide, N\(_2\): Nitrogen, He: Helium, Kr Krypton, Xe Xenon, Ne: Neon, NA: Not available, H/R: Hypoxia-reoxygenation, EAD: Early after depolarization, HSP: Heat shock protein, SOD: Superoxide dismutase, VEGF: Vascular endothelial growth factor, iNOS: Inducible nitric oxide synthase, COX: Cyclooxygenase, STAT: Signal transducers and activator of transcription, HIF: Hypoxia-inducible factor, ERK: Extracellular signal-regulated protein kinases, p-Akt (also known as PKB): Phospho protein kinase B, N\(_2\)O\(_2\): Nitrogen peroxide, JNK: c-Jun N-terminal kinase, p38: Protein 38, TLR: Toll-like receptor, IRAK: IL-1 Receptor-associated kinase, Bcl: B-cell lymphoma, OGD: Oxygen-glucose deprivation, ∆Ψ: Membrane potential, ROS: Reactive oxygen species, NFκB: Nuclear factor-kB, IL: Interleukin, MTX: Mitoxantrone, STS: Broad-spectrum kinase inhibitor staurosporine, LPS: Lipopolysaccharide, TNF: Tumor necrosis factor, TBI: Traumatic brain injury, LDH: Lactate dehydrogenase, MTH: Mild therapeutic hypothermia, HO-1: Heme oxygenase-1, PI-3K: Phosphatidylinositol-3-kinases, mTOR: Mechanistic target of rapamycin, Nrf2: Nuclear factor (erythroid-derived 2) factor 2, Pl: Propidium iodide, OHC: Outer hair cells, IHC: Inner hair cells, tPA: Tissue plasminogen activator

These studies reproduced the following human conditions: ischemic stroke, neonatal hypoxic-ischemic encephalopathy (HIE), subarachnoid hemorrhage, MI; CA, organ (kidney and lung) transplantation, exposure to hyperbaric condition, and liver surgery [Tables 2-4].

The minimum duration of the ischemic insult was 7 min in a model of CA in rats,\[15\] while the maximum duration was 18 h in the *ex vivo* lung perfusion (EVLP).\[52\] In these studies, Ar was always administered as inhalatory mixture, with a concentration varying from 25% up to 88% either in air or in a mixture of nitrogen (N\(_2\)) and O\(_2\). The average Ar exposure was approximately 1–4 h,\[27,29,34\] ranging from a minimum of 3 5-min cycles in the MI studies,\[45\] up to 30 h in the organ transplantation models.\[48\]
Table 2: *In vivo* studies employing argon for protection in models of acute cerebral and myocardial injury, for narcosis and for liver regeneration

| Article                      | Disease          | Model                                                                 | Comparison and duration of treatment | Functional assessment^1| Histopathology                          | Biochemistry                       |
|------------------------------|------------------|----------------------------------------------------------------------|--------------------------------------|------------------------|------------------------------------------|------------------------------------|
| **Models of acute cerebral injury** |                  |                                                                      |                                      |                        |                                         |                                    |
| Höllig *et al.*, 2016[^34^]  | SAH              | Endovascular perforation technique                                    | Ar/O₂ 50/50                          | ≈                      | Hippocampus DG: ↑                        | HIF1α: +                           |
|                              |                  |                                                                      | Duration: 1 h                         |                        | Hippocampus CA1: ≈                       | HO-1: +                            |
|                              |                  |                                                                      |                                      |                        | Hippocampus CA3/4: ≈                    | Bcl-2: +                            |
| Zhuang *et al.*, 2012[^35^]   | Perinatal HIE    | Right common carotid artery ligation + hypoxia                        | Ar/O₂ 70/30                          | ≈↑                     | Hippocampus CA1: ↑                       |                                    |
|                              |                  |                                                                      | Duration: 90 min                      |                        | Infarct size: ↑                          |                                    |
| Alderliesten *et al.*, 2014[^36^] | Perinatal HIE   | Hypoxia                                                              | Ar/room air30-80/70-20               | NA                     | NA                                       | NA                                 |
|                              |                  |                                                                      | Ar/room air 50/50                    |                        | NA                                       | NA                                 |
|                              |                  |                                                                      | Ar/room air 50/50 + MTH              |                        | NA                                       | NA                                 |
|                              |                  |                                                                      | Duration: 1 h × 3                    |                        | NA                                       |                                   |
|                              |                  |                                                                      | Ar/O₂/ N₂ 45-50/30-21/25-29          | aEEG ↑                  | Neocortex: ↑                            |                                    |
|                              |                  |                                                                      | + MTH                                |                        | Subcortical: ↑                           |                                    |
|                              |                  |                                                                      | Duration: 2-26 h                     |                        |                                          |                                    |
| Zhao *et al.*, 2016[^29^]     | Perinatal HIE    | Right common carotid ligation + hypoxia                              | Ar/O₂ 70/30                          | NA                     | Infarct size: ↑                          |                                    |
|                              |                  |                                                                      | Duration: 2 h                        |                        |                                          |                                    |
| Zhao *et al.*, 2016[^28^]     | Perinatal HIE    | Right common carotid ligation + hypoxia                              | Ar/O₂ 70/30 + MTH                  | NA                     | Infarct size: ↑                          |                                    |
|                              |                  |                                                                      | Ar/O₂ 70/30                          |                        |                                          |                                    |
|                              |                  |                                                                      | Duration: 2 h                        |                        |                                          |                                    |
| Ryang *et al.*, 2011[^38^]    | Ischemic stroke  | Transient middle cerebral artery occlusion                           | Ar/O₂ 50/50                          | ↑                      | Neocortex: ↑                            |                                    |
|                              |                  |                                                                      | Duration: 1 h                        |                        | Basal ganglia: ↑                         |                                    |
|                              |                  |                                                                      |                                      |                        | Infarct size: ↓                          |                                    |
| David *et al.*, 2012[^27^]    | 1. Ischemic stroke 2. Excitotoxicity | 1. Middle cerebral artery occlusion 2. Intrastriatal injection of NMDA | 1. Ar/O₂ 50/50 2. Ar/N₂/O₂ 15-75 | 1. ↑ 2. ↓   | 1. ≈                                     |                                    |
|                              |                  |                                                                      | Duration: 3 h                        |                        | 2. ↑                                     |                                    |
| Fahlenkamp *et al.*, 2014[^39^] | Ischemic stroke | Transient middle cerebral artery occlusion                           | Ar/O₂ 50/50                          | NA                     | Ischemic area: ≈↑                         | IL-1β, IL-6: +                      |
|                              |                  |                                                                      | Duration: 1 h                        |                        |                                          | + iNOS, TGF-β, NGF:                  |

Contd...
| Article                         | Disease          | Model                     | Comparison and duration of treatment | Functional assessment | Histopathology | Biochemistry          |
|--------------------------------|------------------|---------------------------|--------------------------------------|-----------------------|----------------|-----------------------|
| **Models of acute cerebral injury** |                  |                           |                                      |                       |                |                       |
| Ulbrich *et al*., 2014[40]     | Retinal I/R      | Anterior chamber hypertension | Ar/O₂/N₂ 25-75/21/4-54 Duration: 1 h | NA                    | Ganglion cell: ↑ | Caspase 3: - Bax: -  |
|                               |                  |                           |                                      |                       |                | Bcl-2: - NF-kB: -    |
|                               |                  |                           |                                      |                       |                | Blood WC count: -    |
|                               |                  |                           |                                      |                       |                | HSP: -                |
|                               |                  |                           |                                      |                       |                | ERK 1-2: +           |
|                               |                  |                           |                                      |                       |                | HO-1: +              |
|                               |                  |                           |                                      |                       |                | IL-8: -               |
|                               |                  |                           |                                      |                       |                | MitocondrialΔΨ: +    |
|                               |                  |                           |                                      |                       |                | ROS: -                |
|                               |                  |                           |                                      |                       |                | NF-kB, STAT3: -      |
| Ulbrich *et al*., 2015[41]     | Retinal I/R      | Anterior chamber hypertension | Ar/O₂/N₂ 75/21/4 Duration: 1 h | NA                    | Ganglion cells: ↑ |                       |
| Ulbrich *et al*., 2016[22]     | Retinal I/R      | Anterior chamber hypertension | Ar/O₂/N₂ 75/21/4 Duration: 1 h | NA                    | ↑              |                       |
| **Models of acute myocardial injury** |                  |                           |                                      |                       |                |                       |
| Pagel *et al*., 2007[45]       | Infarction       | LAD occlusion             | Ar/O₂ 70/30 Duration: 5 mins x 3* | NA                    | Infarct size: ↑ | NA                    |
| Ristagno *et al*., 2014[13]    | Cardiac arrest   | LAD occlusion + VF        | Ar/O₂ 70/30 Duration: 4 h           | cUS: Systolic function (EF) ≈↑ | Infarct size: ≈↑ | hs-cTnT: ≈↑          |
| Lemoine *et al*., 2017[17]     | Infarction       | LAD ligation              | Ar/O₂ 80/20 Duration: 20 min        | cMR: systolic function (LV volumes, EF) ↑ | NA            | NA                    |
| **Models of narcosis**         |                  |                           |                                      |                       |                |                       |
| Balon *et al*., 2002[42]       | Ar narcosis      | Hyperbaric chamber        | Ar 2 MPa + O₂ 0,04 MPa Duration: 2h | NA                    | NA              | Striatal DA release: - |
| Balon *et al*., 2003[43]       | Ar narcosis      | Hyperbaric chamber        | Ar 2 MPa + O₂ 0,04 MPa Duration: 100 min | First phase of compression→ hyperactivity, then hypoactivity | NA              | Striatal DA release: - |
| Abraini *et al*., 2003[44]     | Ar narcosis      | Hyperbaric chamber        | Ar + O₂ 0,03 MPa Compression rate: 0,1 MPa/min | Increased threshold pressure for loss of righting reflex | NA              | NA                    |
| **Models of liver regeneration** |                  |                           |                                      |                       |                |                       |
| Ulmer *et al*., 2017[13]       | Liver regeneration | Partial Hepatectomy     | Ar/O₂ 50/50 Weight of residual liver: ≈ | Proliferation↓Apoptosis≈ | AST, ALT, bilirubin:≈ | BrdU↓ Ki-67↓ UnEL≈ HGF↓ IL-6↓ |
Inhalatory argon and organ protection

Table 2: Contd...

| Article | Disease | Model | Comparison and duration of treatment | Functional assessment | Histopathology | Biochemistry |
|---------|---------|-------|--------------------------------------|----------------------|---------------|-------------|
| Models of liver regeneration | | | | | | |
| Duration: 1 h* | Liver-body weight ratio: ≈ | TNF≈ | ERK 1/2 ≈ |

*In this study, Ar was administered as pretreatment before MI, †In this study, Ar was administered before injury (preconditioning). ↑: Treatment with Ar was significantly protective compared to control; ≈↑: Treatment with Ar showed a trend toward protection compared to control; ≈: Increased with Ar treatment compared to control; ‑: Decreased with Ar treatment compared to control; ↓: Treatment with Ar was significantly detrimental. | Functional assessment included: neurological tests for model of acute cerebral injury; myocardial function after acute ischemic injury; and motor assessment under hyperbaric condition. Ar: Argon, O2: Oxygen, N2: Nitrogen, He: Helium, Xe: Xenon, NO: Nitrosous oxide, Ne: Neon, NA: Not available, SAH: Subarachnoid hemorrhage, DG: Dentate gyrus, HIF: hypoxia inducible factor, HO-1: heme oxygenase-1, HIE: hypoxic ischemic encephalopathy, Bcl-2: B-cell lymphoma 2, MTH: Mild therapeutic hypothermia; aEEG: amplitude-integrated electro-encephalogram, PCr/pi: Ratio of phosphocreatine to inorganic phosphate, NTP: Nucleoside triphosphate, epp: Exchangeable phosphate pool, Lac: Lactate, NAA: N-acetylaspartate, MDA: Malondialdehyde, GSH: Glutathione, GSSG: Glutathione disulfide, TNFα: Tumor necrosis factors, IL: interleukin, SOD: Superoxide dismutase, NQO1: nicotinamide adenine dinucleotide phosphate (NADP) dehydrogenase quinone 1, NF-kB: Nuclear factor-KB, GFAP: Glial fibrillary acid protein, NMDA: N-Methyl-D-aspartic acid, INOS: inducible nitric oxide synthase, TGF-β: Transforming growth factorβ, NGF: Nerve growth factor, I/R: Ischemia/reperfusion, Bax: Bcl-2-associated X protein, WC: White cells, HSP: Heat shock protein, ERK 1/2: Extracellular signal-regulated kinase, AJP: Membrane potential, ROS: Reactive oxygen species, STAT3: Signal transducer and activator of transcription 3, LAD: Left anterior descending coronary artery, VF: Ventricular fibrillation, cUS: Cardiac ultrasound, EF: Ejection fraction, hs-cTnT: High-sensitivity cardiac troponin T, cMR: Cardiac magnetic resonance, LV: Left ventricle, WMS: Wall motion score, MPAs: Megapascal, DA: Dopamine, AST: Aspartate aminotransferase, ALT: Alanine aminotransferase, BrDU: Bromodeoxyuridine, HGF: Hepatocyte growth factor

brain areas. Overall, there was evidence for a mitigation of the brain tissue injury. In one study on a rat model of middle cerebral arterial occlusion,[27] Ar had contrasting effects, with a reduced infarct size in the cortical area, but an increased injury in the subcortical area, compared to sham animals. However, in this study, animals treated with Ar had a significantly higher body temperature (average of 38.5°C), which could explain the greater brain damage compared to the controls.[27] In studies focused selectively on I/R injury in the retina, Ar confirmed its protective effects in a dose- and time-dependent manner[60] [Table 2].

In a model of perinatal HIE, Ar inhalation had no effect on heart rate, arterial blood pressure and blood gases, cerebral O2 saturation, and electrocortical brain activity.[36]

Studies focused on neurological injury following resuscitation from CA are described in a dedicated section of this review.

Regarding the use of Ar as cardioprotective treatment, 3 studies were individuated,[13,17,45] one of which investigated the cardioprotection within the broader post-CA syndrome.[13] Overall, the studies provided evidence for a reduced infarct size and an improved left ventricle systolic function, evaluated either by echocardiography or magnetic resonance [Table 2].

Additional data come from a preliminary study in a rat model of myocardial I/R injury recently performed by our group (published in this review as original data). The left anterior descending was occluded for 30 min, and animals were randomized to receive 1-h ventilation with Ar 70% versus N2 70% in O2, starting 5 min before reperfusion. One hour after reperfusion, rats were weaned from mechanical ventilation and returned to their cages. Six h after reperfusion plasma samples were collected for high-sensitive cardiac troponin T (hs-cTnT) assay and animals were sacrificed. Myocardial infarct size was assessed by tetrazolium chloride staining, and inflammatory response was evaluated in terms of neutrophil infiltration by naphthol staining. Rats treated with Ar showed a lower plasma level of hs-cTnT compared to controls (median [interquartile range]: 3128 ng/L [1832–7053] vs. 7432 ng/L [4965–10000]; P < 0.05) together with a trend toward a reduction in neutrophil infiltration (mean ± Standard error of mean: 77 ± 7 cells/mm2 vs. 92 ± 8.5 cells/mm2; P = not significant). Nevertheless, there was no reduction in infarct size.

In vivo studies on narcosis

Ar showed narcotic effects only under hyperbaric conditions.[42,55] Different theories have been proposed to explain this property.[9] Most likely, the increase of the inert gas pressure induces modification on cytoplasmic membrane and/or receptor condition, subsequently modifying the neurotransmission. Indeed, when rats were exposed to Ar at ≈ 19 atm for 2 h, dopamine release from striatum decreased by approximately 10% compared to animals that were not exposed to the gas.[41] Since the striatum coordinates different aspects of motor planning and motivation, a subsequent study investigated the relationship between motor activity and dopamine release in rats exposed to Ar in a pressurized chamber for 100 min. During the compression period, a transient phase of hyperactivity was observed, followed by a decrease in...
the motor activity. More specifically, exposure to Ar at the pressure of ≈ 10 atm, showed motor hyperactivity, probably related both to the higher pressure and to the narcotic potency of the gas, dependent on its lipid solubility. When the pressure was raised to ≈ 20 atm, loss of righting reflex was observed, indicating the onset of the anesthetic effect. This subsequent decrease of behavioral activity could be ascribed to the decrease of dopamine release from the striatum that occurred when the 20 atm pressure was achieved and was kept constant for at least 10 min.\(^{[43]}\)

Another study concerning Ar anesthetic properties analyzed the contribution of gamma-aminobutyric acid (GABA) transmission as a possible mechanism of action for narcosis. Rats were treated with different GABA receptor antagonists (gabazine and flumazenil, as GABA\(_A\) antagonists, and 2-idrossisaclofene, as GABA\(_B\) antagonist) and then exposed to Ar in a pressurized chamber. Pretreatment with GABA\(_A\) inhibitors increased significantly the Ar threshold pressure for the loss of righting reflex. Instead, pretreatment with GABA\(_B\) inhibitor showed no significant effects on narcosis\(^{[44]}\) [Table 2].

**Ex vivo studies**

Regarding the use of Ar for organ preservation, kidneys and lungs were the preferred targets.\(^{[46‑48,52,54]}\) Indeed, reduced acute tubular necrosis and inflammatory lesions accompanied by better renal function and faster recovery were observed after heterotopic autotransplantation in which the removed kidney was preserved in a cold storage solution

| Article                  | Organ          | Model                        | Animal | Comparison and duration of treatment | Histopathology                                      | Functional assessment | Biochemistry                  |
|--------------------------|----------------|------------------------------|--------|--------------------------------------|-----------------------------------------------------|------------------------|------------------------------|
| Irani *et al.*, 2011[46] | Kidney         | I/R + Renal transplantation  | Rat    | Cold preservation solution saturated with Ar 100% | Organ structural architecture: ↑ Necrosis: ↑ | CrCl: ↑ Urinary albumin: ↑ | CD10: + Caspase-3: - |
| Faure *et al.*, 2016[48] | Kidney         | I/R + Renal transplantation  | Pig    | Preservation solution saturated with Ar 100% | Tubular necrosis: ↑ Fibrosis: ↑ Apoptosis: ↑ Inflammation: ↑ | Diuresis: ↑ CrCl: ↑ EFNa: ↑ RAA/TBARS: + HSP 27: + IL1, IL6, TNFα: - | Urinary urea/plasma urea ratio: ↑ Septal thickness: ≈ CONSOLIDATION: ≈ |
| Martens *et al.*, 2016[47] | Lung           | 2-h warm ischemia            | Pig    | EVLP: Ar 70% Duration: 6 h | BAL: ≈ W/D: ≈ Necrosis: ≈ Inflammation: ≈ | PVR: ≈ Ppeak: ≈ PaO2/FIO2: ≈ | CT: ground glass opacities: ≈ |
| Martens *et al.*, 2017[52] | Lung           | 18-h cold ischemia           | Pigs   | Pre-conditioning + EVLP Ar79-88% Duration: 28 h | BAL: ≈ W/D: ≈ Necrosis: ≈ Inflammation: ≈ Congestion: ≈ | PVR: ≈ Ppeak: ≈ PaO2/FIO2: ≈ | CT: inflammatory cytokines: ≈ |
| Smith *et al.*, 2017[54] | Kidney         | I/R                          | Pig    | EVNP: Ar 70% Duration: 1 h | Tubular dilatation: ≈ Epithelial shedding: ≈ Epithelial flattening: ≈ Tubular debris: ≈ Vacuolation: ≈ Condensed tubular nuclei: ≈ | Renal blood flow: + Urine Output ≈CrCl ≈Fractional excretion of sodium: ≈ | Urinary HIF-1α: ≈ Urinary IL-6: ≈ Urinary IL-8: ≈ Urinary TNF-α: ≈ Cytoplasmic HIF-1α: ≈ Nuclear HIF-1 α: ≈ |
Table 4: *In vivo* studies in models of cardiac arrest employing argon for neuroprotection

| Article             | Model of CA                  | Animal | No-flow time (mins) | CPR duration (mins) | Comparison                          | Treatment onset | Neurobehavioral tests | Histopathology | Biochemistry |
|---------------------|-------------------------------|--------|---------------------|---------------------|-------------------------------------|----------------|-----------------------|----------------|--------------|
| Brücken *et al.*, 2017♂[9] | Transesophageal electrical stimulation | Rat    | 9                   | 3                   | Ar/O₂ 70/30 + MTH Duration: 1 h after ROSC | 1 h after ROSC | ↓ NA NA NA ↓ NA ↓ NA ↓ NA | NA NA NA | NA NA NA |
| Zuercher *et al.*, 2016♂[10] | KCl + esmolol | Rat    | 8                   | 8                   | Ar/O₂ 50/50 Duration: 24 h after ROSC | 15 min after ROSC | ≈ ≈ NA ≈ ≈ NA NA ≈↑ | NA NA NA | NA NA NA |
| Brücken *et al.*, 2015♂[14] | Transesophageal electrical stimulation | Rat    | 7                   | 3                   | Ar/O₂ 70/30 Duration: 1 h after ROSC | 1-3 h after ROSC | ↑ ↑ NA NA NA NA NA ↑ ↑ ↑ | ↑ ↑ ↑ ↑ | NA NA NA |
| Brücken *et al.*, 2014♂[12] | Transesophageal electrical stimulation | Rat    | 7                   | 3                   | Ar/O₂ 40/30/30 Ar/O₂ 70/30 Duration: 5 h after ROSC | Immediately after ROSC | ↑ ↑ NA NA NA NA NA ↑ ↑ NA | ↑ ↑ NA | ↑ ↑ NA |
| Ristagno *et al.*, 2014♂[13] | LAD occlusion | Pig    | 8                   | 5                   | Ar/O₂ 70/30 Duration: 4 h | NA NA NA NA ↑ ↑ ↑ ↑ | ↑ ↑ NA NA NA | NSE: ↑ ↑ | ↑ ↑ ↑ ↑ |

↑: Treatment with argon was significantly neuroprotective compared to control, ≈↑: Treatment with argon showed a trend towards neuroprotection compared to control, ≈: Treatment with argon was equivalent to control, ↓: Treatment with argon was significantly worse than control. Ar: Argon, O₂: Oxygen, N₂: Nitrogen; He: helium; NA: Not available; CA: Cardiac arrest; CPR: Cardiopulmonary resuscitation, NDS: Neurological deficit score, NAS: Neurological alertness score, OFT: Open field test, MWT: Morris water maze, VPT: Vertical pole test; TRT: Tape removal test, OPC: Overall performance category, RTD: Rotarod, MTH: Mild therapeutic hypothermia, ROSC: Return of spontaneous circulation, LAD: Left anterior descending coronary artery, NSE: Sieric neuron-specific enolase, BDNF: Brain-derived neurotrophic factor.
saturated with pure Ar. Smith et al. studied the effects of ex vivo normothermic perfusion with a gas composition of 70% Ar for 1 h, in a porcine model of kidney ischemia-reperfusion injury. The authors did not observe any organoprotective effect as far as hemodynamics, renal function, functional parameters, inflammatory markers, and histological analysis were concerned.\footnote{14} In EVLP, following a period of warm ischemic injury in the swine, 6-h ventilation with a respiratory mixture containing 70% Ar in O\textsubscript{2} did not show any benefit in graft function.\footnote{47} Indeed, no significant differences in ventilator parameters, such as pulmonary vascular resistance, peak airway pressure, and blood gases, nor in the histopathology, were observed, compared to a control ventilation with N\textsubscript{2}/O\textsubscript{2}. In 2017, the same group confirmed the above results in a model of cold I/R injury during EVLP.\footnote{51} Overall lungs were exposed to Ar at a different concentration ranging from 79% to 88% in O\textsubscript{2} for 28 h. At the end of the exposure, no beneficial effects on graft quality were observed [Table 3].

**Use of argon after cardiac arrest**

The use of Ar ventilation for neuroprotection after CA was explored in 6 studies,\footnote{12,16,49} [Table 4]. All these studies reported both neurological recovery, evaluated with various neurobehavioral tests, and tissue damage, assessed by histopathology and biochemistry. Of the 5 studies using the rat model of CA, 4 were from the same research group\footnote{12,14,15,40} that investigated the effect of 1 h of ventilation with a mixture of 70% Ar in O\textsubscript{2} initiated 1 h after resuscitation. After a no-flow of 7 min, postresuscitation ventilation with Ar led to a better and faster neurological recovery during the following 7 days compared to the control ventilation. The results were assessed by a battery of tests, including the rodent neurological deficit score, the open field, and the Morris water maze tests.\footnote{15} This good functional recovery was paralleled by a significantly lower brain damage in the neocortex and the hippocampal CA 3 and 4 regions. In a subsequent study employing the same model of CA and cardiopulmonary resuscitation, it was confirmed that such beneficial effects of Ar treatment were dose-dependent.\footnote{12} Indeed, both the neurological outcome and the histopathological brain damage were confirmed to be improved compared to controls, after exposure to Ar at a concentration of either 40% or 70% after resuscitation; however, the neuroprotective effects were significantly more pronounced when the inhalatory mixture contained 70% Ar in place of 40%.\footnote{12} A third study introduced further data showing that the 70% Ar-induced neuroprotection persisted even when the onset of the treatment was delayed up to 3 h after resuscitation.\footnote{14} More recently, neuroprotection of Ar was investigated in conjunction with mild therapeutic hypothermia (MTH), based on the hypothesis that the combination of these 2 interventions could boost protection, further improving neurological outcome after CA.\footnote{49} Thus, in the same model, rats were treated with MTH at 33°C (6 h) and 1 h ventilation with 70% Ar. Surprisingly, the results showed that the combined treatment, similarly to a control ventilation with air/O\textsubscript{2}, generated worse neurological recovery together with greater neuronal degeneration in hippocampal CA1 region when compared to MTH alone. Unfortunately, in this study the treatment with Ar alone, as a reference, was absent and thus no further considerations on the effect of Ar could be made.

In a model of CA with an underlying acute MI in swine, a 4-h ventilation with a mixture of 70% Ar–30% O\textsubscript{2} initiated immediately after resuscitation was compared to a control ventilation with an equivalent percentage of N\textsubscript{2}/O\textsubscript{2}. This study confirmed the Ar-induced neuroprotective effects previously shown in rats. More specifically, animals that received Ar achieved a faster and more complete neurological recovery, already 24 h after CA, in contrast to controls, in which a neurological impairment persisted up to 72 h after resuscitation.\footnote{113} The improvement in neurological recovery was confirmed by different neurological tests and by the circulating level of serum neuron-specific enolase, a biomarker of neuronal damage. Histopathology supported this benefit on neurological recovery, showing lesser brain injury in animals that were treated with Ar compared to controls. In addition, a trend toward a smaller myocardial infarct size was also observed after inhalation of Ar.\footnote{113} The same group has recently replicated these results showing a beneficial effects of Ar ventilation on neuro- and cardioprotection in a more severe model of CA (abstract only available).\footnote{56}

In a model of CA in rodents induced with KCl and esmolol, 24-h ventilation with a mixture of 50% Ar-50% O\textsubscript{2} was conducted, starting 15 min after resuscitation. As the primary outcome, neuronal damage was assessed by histopathology, while as secondary outcome, neurologic tests were performed. A trend toward a lesser neuronal damage after Ar treatment was reported.\footnote{116}

**Mechanism of organ protection**

The description of the potential mechanisms of action involved in the Ar protection derives mainly from in vitro studies and have been further confirmed in vivo. Ar appeared to possess O\textsubscript{2}-like properties, which could explain its neuroprotective effects by partially restoring mitochondrial respiratory enzyme activity and reducing N-methyl-D-aspartic acid-induced neuronal death.\footnote{27} Moreover, Ar plays anti-apoptotic effects modulating the molecular pathways involved in cell survival. In details, it increases extracellular signal-regulated kinase (ERK) 1/2 phosphorylation, already after 30 min of exposure; it blocks the apoptosis cascade;\footnote{24,29,41} it upregulates the expression of the anti-apoptotic protein B-cell lymphoma-2;\footnote{28,35} it activates the toll-like receptor 2 and 4, which reduce caspase-3 activity;\footnote{28,22,28} and mediate the intracellular signaling involved in the production of pro-inflammatory cytokines, growth factors, and cell survival.\footnote{8} [Table 1].
Reperfusion can also contribute to cell death, and particularly on cardiomyocyte, it can induce ventricular arrhythmias and contractile dysfunction subsequent to the opening of the mitochondrial permeability transition pore (MPTP). MPTP is a nonselective channel of the inner mitochondrial membrane that opens at reperfusion, causing mitochondrial calcium-phosphate overload, and dissipates mitochondrial membrane potential. Phosphatidylinositol-3-kinase, ERK 1–2 and 70 kDa ribosomal protein s6 kinase inhibit MPTP opening by their actions on several downstream signaling molecules that modulate the transition state of the pore either directly (via endothelial nitric oxide synthase, p53, glycogen synthase kinase) or indirectly by affecting the pro- and anti-apoptotic B-cell lymphoma protein. Thus, Ar by acting on the same pathways, directly or indirectly, through reperfusion injury salvage kinase, prevents the MPTP opening.\[^{17,45}\]

**Safety of argon use**

Even though safety was not the primary endpoint of all the above described *in vivo* studies, none of them has ever reported safety concerns related to the inhalatory exposure to Ar. In 12 *in vivo* studies,\[^{12-16,27-36,38,45,49,52}\] continuous hemodynamic parameters and repeated blood gas analyses were performed during Ar ventilation. Ar did not affect neither hemodynamics nor ventilation, in terms of respiratory gas exchange. Three studies have also reported body weight as an indicator of animal’s wellbeing during the days after Ar ventilation, and none of them reported significant changes.\[^{16,34,35}\]

Two studies aimed to specifically evaluate the safety of Ar ventilation in pigs.\[^{36,52}\] In 1 study, a 6-h ventilation with Ar 79% in pigs showed no toxic effects, as demonstrated by serum biomarkers and assessment of liver and kidney function and structure.\[^{52}\] In the other study, neonatal piglets were ventilated with Ar at different concentrations, ranging from 30% to 80% in O$_2$, for a total of 3 h. Effects on hemodynamics and blood gases were evaluated, and prolonged ventilation with Ar was confirmed to be safe at each concentration. A similar study observed an augmented hypothermic protection at 48 h after hypoxia-ischemia with Ar from 2 to 26 h, shown by improved brain energy metabolism, faster electrical brain activity recovery and reduced cell death.\[^{37}\]

In 3 studies, Ar was used as a substitute of CO$_2$ in inducing pneumoperitoneum during laparoscopic surgery to evaluate its safety profile.\[^{3,50,51}\] Ar insufflation into the abdominal cavity produced hemodynamic modifications, mainly related to increases in abdominal pressure and perhaps to a potential effect on systemic vascular resistances (effect observed in 1 study\[^{50}\] but not replicated in the other\[^{51}\]). No significant changes in respiratory functions were observed. Since Ar is lesser soluble than CO$_2$, an increased risk of hemodynamic instability exits in the instance of accidental gas embolism.\[^{31}\]

Finally, because Ar is denser than air, it could be argued that ventilation with such a gas could increase respiratory resistance.\[^{31}\] However, none of the preclinical studies reported such a condition after prolonged exposure to Ar. Moreover, a study with a model of vigorous ventilation in a respiratory resistance machine showed no changes in respiratory resistance at atmospheric pressure, and up to 4 atm, when Ar 80% was compared to air.\[^{31}\]

**Clinical uses of argon**

Since the sixties, Ar was applied for the measurement of lung volumes,\[^{57-59}\] with the rebreathing/gas-dilution technique. Ar 6%–7% has been also used to calculate the cardiac output CO,\[^{60,61}\] and for the estimation of global myocardial blood flow.\[^{62}\]

In these studies, subjects were exposed to a single or a few breaths of an isobaric mixture of Ar and no conclusions on safety issues may be extrapolated.

The feasibility of Ar use was addressed by studies involving divers. In these studies, divers breathed Ar mixtures for long periods in hyperbaric conditions. In 1939, motor and cognitive impairment, such as slowed mental activity, inability to perform efficient manual work and emotional disturbances, which might lead to loss of consciousness, were described in divers breathing a mixture of 69% Ar, 20% O$_2$ and 11% N$_2$ from 1 to 10 atm.\[^{51}\] In another study, 10 subjects performed mental arithmetic tasks while exposed to Ar 80% in O$_2$ at 1, 4, and 7 atm. Again, Ar under hyperbaric condition showed more narcotic effects.\[^{58}\]

The longest exposure to Ar under hyperbaric condition was in 1998.\[^{63}\] Four male volunteers participated in a simulated diving at 10 m for 7 days with following gas mixture: O$_2$ 0.2 ± 0.005 kg/cm$^2$, N$_2$ 0.8 ± 0.01 kg/cm$^2$ and Ar 1.0 ± 0.01 kg/cm$^2$. No effects on central nervous system electrophysiology and functional test, cardiopulmonary system, i.e., electrocardiogram, oxihemometry, and biochemical analysis of urine, were detected. All measured data did not exceed physiological range. Under hyperbaric pressure, a workload of 100 Wt was performed 62% higher when breathing Ar 15% in O$_2$ than N$_2$ 15% in O$_2$. Moreover, the exposure to a hypoxic Ar-N$_2$ mixture containing O$_2$ 0.15 ± 0.005 kg/cm$^2$, instead of O$_2$ 0.2 ± 0.005 kg/cm$^2$, determined an increase in the performed work volume, showing that Ar causes a positive effect on organism adaptation to hypoxia.

Another study performed on 8 human volunteers investigated the possible adverse effects from breathing isobaric mixture of 80% Ar and 20% O$_2$ for 30 min. The authors did not observe narcosis nor coagulation nor fibrinolysis abnormalities, which would have been present in case of gas embolism.\[^{64}\]

More recently, Ar inhalation has been proposed to measure the global cerebral blood flow (CBF), by a
Inhalatory argon and organ protection

Nespoli, et al.: Inhalatory argon and organ protection

...respectively. In vitro 2-fold Argon mediates anti-apoptotic signaling and neuroprotection via...210-fold lower than xenon.

Summary of evidence and future perspectives
Ar might diminish the neurological and myocardial damage after a hypoxic-ischemic insult. Indeed, Ar has been tested in different models of ischemic insult, at concentrations ranging from 20% up to 80%. Overall, Ar emerged as a protective agent on cells, tissues, and organs, showing less cell death, reduced infarct size and faster functional recovery. More specifically, encouraging data has been reported in animal studies on CA in which a better and faster neurological recovery was achieved when Ar was used in the postresuscitation ventilation. More importantly, these benefits have been replicated in different studies, enrolling both small and large animals. Ventilation with Ar in O₂ has been demonstrated to be safe both in animals and humans.

Based on all the evidence described in this article, the protective effects of Ar should be evaluated in a clinical scenario, in particular, those concerning the neuroprotective properties after CA which are strongly supported by preclinical evidence. Finally, Ar can be simply administered through a ventilator which is modified to be compatible with the gas mixture Ar/O₂/air, in variable proportions, provided in dedicated cylinders. Indeed, Ar has a relatively low cost, approximately 9 Euro cent per liter,[13] 10-fold lower than anesthesia agents such as sevoflurane and isoflurane and 100-fold lower than xenon.

Conclusions
Ar use has been described in a variety of preclinical and human studies. Inhalatory Ar seems to be potentially beneficial in the context of organ protection after different types of insults. Safety data show nontoxic effects. Thus, preclinical and clinical data support future clinical studies on the inhalatory Ar therapy.

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Conflicts of interest
There are no conflicts of interest.

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Inhalatory argon and organ protection

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