Review

Therapeutic Potential of Hemoglobin Derived from the Marine Worm Arenicola marina (M101): A Literature Review of a Breakthrough Innovation

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Abstract: Oxygen (O2) is indispensable for aerobic respiration and cellular metabolism. In case of injury, reactive oxygen species are produced, causing oxidative stress, which triggers cell damaging chemical mediators leading to ischemic reperfusion injuries (IRI). Sufficient tissue oxygenation is necessary for optimal wound healing. In this context, several hemoglobin-based oxygen carriers have been developed and tested, especially as graft preservatives for transplant procedures. However, most of the commercially available O2 carriers increase oxidative stress and show some adverse effects. Interestingly, the hemoglobin derived from the marine lugworm Arenicola marina (M101) has been presented as an efficient therapeutic O2 carrier with potential anti-inflammatory, antibacterial, and antioxidant properties. Furthermore, it has demonstrated promise as a supplement to conventional organ preservatives by reducing IRI. This review summarizes the properties and various applications of M101. M101 is an innovative oxygen carrier with several beneficial therapeutic properties, and further research must be carried out to determine its efficacy in the management of different pathologies.

Keywords: Arenicola marina; hemoglobin-based oxygen carriers; therapeutic oxygen carriers; M101

1. Introduction

The existence and maintenance of human life is dependent on oxygen (O2) as it is vital for life-sustaining aerobic respiration in humans [1]. At the cellular level, O2 acts as the terminal electron acceptor at the end of the electron transport chain within the mitochondrial inner membrane, whereby oxidative phosphorylation results in the synthesis of adenosine triphosphate (ATP), which is necessary for all active metabolic processes [2]. O2 is also vital for protein synthesis, maturation of extracellular matrices such as collagen and nitric oxide synthesis, eventually, playing a major role in the regulation of vascular tone and angiogenesis [3]. However, in the presence of O2, normal metabolism can also generate reactive oxygen species (ROS). In a wound microenvironment, large amounts of molecular O2 are partially reduced to form ROS [3]. ROS are small and highly reactive molecules including superoxide ions and peroxides [4] that influence cell migration, proliferation and angiogenesis [3]. Exacerbation of hypoxic injury after restoration of oxygenation (reoxygenation) is an important mechanism of cellular injury, especially, in transplantation procedures. Cellular hypoxia and reoxygenation are two essential elements of ischemia-reperfusion injury (IRI) [5].
The imbalance between ROS production and the ability of cells and tissue to detoxify these reactive products causes oxidative stress, triggering the production of cell damaging mediators and consequent IRI [6–8]. Herein, the excess production or decreased scavenging of ROS has the potential to cause cell injury contributing to human aging and to the pathogenesis of several diseases such as neurodegeneration, diabetes, cancer and atherosclerosis [4]. Oxidative stress is considered to be the major culprit in orchestrating cellular damage in hypoxia-reoxygenation injury and eventual IRI [9,10].

Physiologically, $O_2$ is bound reversibly to hemoglobin (Hb), a metalloprotein containing an iron atom at the center of a tetrapyrilic group or porphyrin, contained in red blood cells (RBCs). Hb is a molecule that is responsible for carrying almost all of the $O_2$ in the blood, and in humans, it is composed of four subunits (2 alpha and 2 beta globin chains), each of them contains a heme that is able to reversibly bind the $O_2$. In the absence of protective enzymes such as superoxide dismutase (SOD) and catalase, Hb is oxidized to methemoglobin (MetHb), thus, compromising its ability to deliver $O_2$ [11]. Hb oxidation can compromise efficient $O_2$ binding to Hb through the formation of toxic heme degradation products, thereby, hampering optimal tissue perfusion [12]. Sufficient $O_2$ supply to tissues is crucial for optimal organ function, the complete failure of which can require an organ transplant. Good tissue oxygenation is mandatory during organ preservation or perioperatively during different major surgical procedures [13,14]. Rigorous research has been carried out over the years in order to find solutions to reach the optimal oxygen requirement during surgery or organ preservation. No single strategy has shown complete success in this context. Over the past decades, artificial $O_2$ carriers have been proposed to improve $O_2$ supply to tissues [15], including Hb-based $O_2$ carriers (HBOCs) and perfluorocarbon-based $O_2$ carriers [16,17]. Interestingly, some HBOCs have shown therapeutic promise in maintaining graft/organ vitality and function, and in minimizing the risk of reoxygenation injuries or IRI [18,19]. Also, considering their additional benefits as potential blood substitutes without the inconvenience of crossmatching, their application in maintaining optimum tissue oxygenation is deemed highly appropriate and is under further investigation [20].

Indeed, several mammalian source Hbs have also been tested for their potential as blood substitutes and therapeutic $O_2$ carriers [21–25]. Interestingly, Hemopure (HBOC-201) has been approved by the South African drug council for the treatment of anemia. Moreover, this product is allowed to be used in United States in qualified patients to treat severe life-threatening anemia under the FDA’s expanded compassionate use access program [20,26–29]. Additionally, one randomized, multicenter trial investigation of HBOC-201 in non-cardiac surgery patients indicated that its use reduced RBC transfusions in $43\%$ of patients without notable differences in mortality and serious adverse events, though there was a notable increase in the associated non-serious events (i.e., hypertension and fever) [26]. Unfortunately, while many of these HBOCs showed promise in early clinical trials [15,28,30,31], they eventually failed Phase III clinical trials due to severe adverse effects, such as hypertension, myocardial infarction, stroke, renal damage and tissue toxicity, which are caused by rapid Hb oxidation and scavenging of the vasodilator nitric oxide by Hb [28,32–34]. These side effects can be attributed to the removal of intracellular mammalian Hb from the protective environment of the RBC and its release, even the modified forms, into the harsh extracellular environment of the bloodstream [12].

In this context, natural extracellular Hb found in annelids may prove to be advantageous over mammalian Hbs [12,35]. For instance, Arenicola marina-derived natural extracellular Hb has been studied in the recent past [35–37]. Arenicola marina’s Hb (M101, commercially manufactured as HEMO2life® by Hemarina (Morlaix, France) has been shown to effectively deliver $O_2$ in vivo [38,39] without any signs of oxidation, vasoconstriction, or hypertension [35,40]. Large heme pockets of Arenicola marina Hb allow $O_2$ to easily escape in passive way [41–43]. It is notable that Hb of Arenicola marina has a significantly higher $O_2$ affinity as reflected by its lower p50 value ($7.05 \pm 0.93$ mmHg) compared to that of human blood, targeting its $O_2$ delivery action to a more hypoxic environment [37]
(Table 1). However, the p50 of M101 is closely related to the p50 of human HbA inside the red blood cell and its cooperativity is also similar [37].

Table 1. Properties of Arenicola marina’s Hb.

| Properties                      | Hb of Arenicola marina | References |
|---------------------------------|------------------------|------------|
| Molecular weight                | 3600 kDa               | [35,44,45] |
| Size                            | 15 by 25 nm            | [44,46]    |
| Shape of central piece          | Ellipsoid (hexagonal bilayer) | [46] |
| O₂ binding sites                | 156                    | [35,44]    |
| Heme pockets                    | large                  | [36,47]    |
| Auto-oxidation rate kₐox (h⁻¹)  | 0.014                  | [35,48]    |
| Redox potential Eₒ (mV)         | −50                    | [48]       |
| p50                             | 7.05 ± 0.93 mmHg       | [40,49]    |
| n50                             | 2.54 ± 0.23            | [29]       |

Arenicola marina inhabits the intertidal area in a sulfide-rich environment and is exposed to pronounced fluctuations in environmental conditions [50]. The manufacturing of the product HEMO₂life® is done under GMP and starts by freezing the worms to create a hemorrhagic shock and to release its extracellular Hb (M101). After successive steps of solid/liquid extraction, purification, filtration and gamma irradiation, the final result is a class III medical device containing M101 [51]. Preclinical studies in rats and hamsters using M101 have shown reduced microvascular vasoconstriction and no significant impact on mean arterial blood pressure compared with other HBOCs that contain bovine or human Hb [40]. Moreover, the biocompatibility and absence of toxicity of HEMO₂life® has also been evaluated according to international standards (ISO 10993).

In view of the beneficial properties exhibited by M101, the aim of this review is to summarize the outcomes of M101 application to several in vitro, in vivo and human trials and to assess its potential therapeutic and clinical relevance. To perform this review, only studies published in the English language were included and the last search was carried out in August 2020. The following keywords were used for the search: (“M101” OR “HEMO₂life®” OR “Arenicola marina” OR “extracellular hemoglobin” OR “HEMOXYCarrier”) and a systematic literature search was performed in the Pubmed/Medline database. One hundred ninety-one articles were found and after they were reviewed for relevance and repetition, only 16 articles were included in this review.

2. Structure of M101

The description of the quaternary structure of M101 and an assessment of its heme content was first carried out in 1996. Multi-angle laser-light scattering electrospray-ionization mass spectrometry [50] gave a molecular mass of 3648 ± 24 kDa and a gyration radius of 11.3 ± 1.7 nm. According to Zal et al., in the reduced condition, the Hb is composed of eight globin chains of molecular masses 15952.5 Da (a1), 15974.8 Da (a2), 15920.9 Da (b1), 16020.1 Da (b2), 16036.2 Da (b3), 16664.8 Da (c), 16983.2 Da (d1), 17033.1 Da (d2). In the native Hb, chains b, c, d occur as five disulfide-bonded trimer subunits T with masses of 49560.4 Da (T1), 49613.9 Da (T2), 49658.6 Da (T3), 49706.8 Da (T4), 49724.5 Da (T5). There are two different linker chains, L1 and L2, involved in homo- and heterodimer formation, with a mass of 25174.1 and 26829.7 Da, respectively [45].

Thus, the Arenicola marina Hb would be composed of 198 polypeptide chains with 156 globin chains and 42 linker chains, each twelfth being in contact with 3.5 linker subunits, providing a total mass of 3682 kDa [50]. The phylogenetic analyses of the Arenicola marina globin chains confirmed a more intricate structural organization of this extracellular globin compared to the previous description [50] and have allowed DNA sequencing of Arenicola
marina globin genes, which assigned the five Arenicola marina sequences to the different globin sub-families. The cDNA-derived amino acid sequence exhibits 12 cysteine residues, which is in agreement with previous studies on Arenicola marina’s Hb [52]. However, two of the globins were found to be A2 globin chains lacking the cysteine residues proposed to be involved in the binding of hydrogen sulfide by such Hb, which has been linked to the evolution of the Hb sulfide binding function in annelids inhabiting sulfide-rich environments [45,53]. In summary, M101’s quaternary structure comprises two overlapping hexagons, which are approximately 25 nm between parallel sides (face view) and 15 nm thick (profile view) (Figure 1).

Figure 1. Diagrammatic representation of the structure of Arenicola marina’s hemoglobin (M101) molecule.

3. Properties of M101

The Arenicola marina Hb M101 can carry up to 156 O₂ molecules (versus 4 for human hemoglobin) in a saturated state, hence, it exhibits a greater O₂-binding capacity and antioxidative properties. M101 releases O₂ according to a simple gradient [35] and without co-factors. M101 can completely inhibit reduction of nitroblue tetrazolium in the presence of superoxide radicals, thereby, demonstrating intrinsic SOD-like activity linked to Cu/Zn metals and an ability to ameliorate oxidative stress [54].

Sulfide concentration in the pore water of the sediment (lugworm burrow) can reach several hundred µmol/L and is considered an important environmental factor for a sediment-dwelling animal such as Arenicola marina [55,56]. In the presence of sulfide, vertebrate oxyhemoglobin forms sulfhemoglobin. The inability of Arenicola marina’s Hb to form this compound might indicate a biochemical adaptation that is necessary for Arenicola marina during exposure to high concentrations of sulfide. The free cysteine on the polypeptide d chains plays a role in this mechanism by inhibiting the formation of sulfhemoglobin and allowing sulfide oxidation. The oxidation product, defined as the brown pigment, and which possesses higher sulfide oxidation activity than hexagonal bilayer Hb, could be due to some modification or breakage of the prosthetic-group-protein linkage during the reaction leading to its formation. However, for Arenicola marina, this would be a detoxification mechanism (i.e., sulfide immobilization by the Hb), and not an adaptation to symbiotic life as is the binding of sulfide by vestimentiferan Hb [50]. Interestingly, M101 is able to work in a broad range of temperatures (4 ℃ to 37 ℃) as opposed to other therapeutic oxygen carriers manufactured using vertebrate Hb, which only work at 21 ℃ to 37 ℃ [19,57]. In vivo, studies carried out on mice and rats showed that the half-life of M101 molecule in circulation is 2.5 days [38]. The important properties of M101 have been listed below (Table 2).
Table 2. Comparison between the properties of *Arenicola marina’s* Hb and human Hb under human physiological conditions (pH 7–8).

| Properties                  | *Arenicola marina’s* Hb | Human Hb | References |
|-----------------------------|-------------------------|----------|------------|
| Colloid oncotic pressure    | 1 mm Hg                 |          | [35,40]    |
| Size/weight                 | 3600 kDa                | 64,458 Da| [35]       |
| Half life                   | 50 h                    | few seconds | [35,40] |
| Working temperature range   | Broad (4 °C to 37 °C)   | 37 °C    | [35,40]    |
| p50 (mmHg)                  | 7.05 ± 0.93             | 8–18     | [35,40,58] |
| n50                         | 2.54 ± 0.23             | 2.3–3.0  | [35,40,59] |
| Bohr coefficient            | −0.5                    | −0.48    | [35,40,60] |
| ΔH (kJ mol⁻¹)               | −19                     | −50.7    | [35,40,61] |
| COP (mmHg)                  | 1.0                     | nd       | [40]       |
| Viscosity (cP)              | 1.23                    | nd       | [40]       |
| SOD activity (U/mg Hb)      | 3.53 ± 0.02             | 0        | [40]       |
| Cyanide (CN) inhibition     | 100%                    | 100%     | [40]       |
| Fe (atom/molecule)          | 156                     | 4        | [40]       |
| Cu (atom/molecule)          | 3.58 ± 1.17             | 0        | [40]       |
| Zn (atom/molecule)          | 5.13 ± 0.75             | 0        | [40]       |

4. Applications of M101

M101 has already demonstrated its efficacy as a supplement to the solution for graft/organ preservation under static and perfusion conditions during organ transplant. Several in vivo preclinical trials involving different vital organs, for instance, heart [62], kidneys [58,63], liver [64], pancreas [65], lungs [66] have exhibited its ability to maintain and improve graft viability and function. Furthermore, its safety and efficiency has also been established in the human trial OXYOP (Clinical Trial Registry No. NCT 02652520) involving a kidney transplant procedure where HEMO2life® as the class III medical device containing M101 (1 g/L) was added to a preservation solution [51]. Interestingly, no immunological, allergic or prothrombotic effects were observed with M101 application. Furthermore, addition of M101 decreased the delayed graft function (DGF) and improved the renal function [51]. This confirmed the safety and efficacy of M101 for its potential clinical applications. M101 has also been proposed as an instant blood substitute for wounded US army soldiers [67].

5. M101 and Conventional Preservative Techniques

Hypothermic cold storage (CS) is a conventional strategy based on the principle of metabolism reduction with temperature to minimize ischemic injuries during organ preservation [68], however, even slow metabolism requires O₂ [69,70]. Furthermore, the delayed O₂ delivery to ischemic tissue during CS could also aggravate oxidative stress if the cells fail to restore oxidative respiration [71]. Therefore, the use of an O₂ carrier during the entire preservation procedure could be protective, allowing cold-preserved cells to maintain their ATP reserve, thus, establishing a balanced energy metabolism at reperfusion and coping with the sudden influx of O₂ without causing oxidative stress.

In vitro, M101 supplementation to a range of solutions used in the clinic has demonstrated utility in cold-stored cultured cells. Furthermore, in vivo, M101 addition to University of Wisconsin (UW) and histidine-tryptophan-ketoglutarate (HTK) solution [72] has shown an improvement of graft function and reduction in DGF [57,63].

However, the tissue oxygenation ability of M101 is in accordance with the physiological needs as O₂ binding and release occurs passively in a simple O₂ gradient. The intrinsic Cu/Zn-SOD activity of M101 is crucial for the prevention and management of
IRI [58,63,73]. Thus, the simple addition of M101 to CS presents excellent potential for clinical translation. Besides static CS, machine preservation (MP) of grafts has attracted mounting interest as it exhibits improvement in graft quality and function [74–76]. Its ability to get rid of metabolites and cellular waste produced during ischemia has been proposed as the reason for its beneficial effects. However, the presence of these products at reperfusion is most likely associated with the intense activation of the innate immune pathway [77,78]. In this context, M101 supplementation to MP for uniform and efficient O$_2$ delivery to the entire graft could promote organ protection because of the synergistic beneficial effects of M101 and MP [79,80].

6. M101 as an Extracellular O$_2$ Carrier

Conventional preservative techniques provide O$_2$ in excess, which could induce oxidative stress. On the contrary, M101 has the unique property of providing O$_2$ against a gradient, according to the physiological needs of the cell, thus, eliminating the risk of hyperoxia and oxidative stress [19,35,38]. M101 supplementation increases cellular ATP production, thus, it efficiently maintains the metabolic processes, and protects mitochondria by decreasing the need to switch from mitochondrial respiration to anaerobic glycolysis. The ability of M101 to maintain high ATP levels during the preservation process may also benefit the restoration of energy homeostasis upon reperfusion due to less metabolic stress on oxidative pathways [57]. M101 remains stable in different organ preservation solutions of varying ionic compositions and osmolarities and demonstrates its O$_2$ carrying and antioxidant (SOD) properties [35]. So far, M101 application in vitro, in vivo and clinically has shown great promise (Tables 3 and 4).

Hence, M101 improves graft vitality by promoting tissue oxygenation, maintaining tissue integrity, reducing IRI, inflammation and fibrosis (Figure 2).

![Figure 2](image)

Figure 2. The use of Hb derived from *Arenicola marina* (M101) as a graft preservative can reduce ischemic reperfusion injury and delayed graft function, thus, improving graft vitality and a successful transplant outcome.
### Table 3. In vitro studies involving M101 application.

| Application                                                                 | Concentration & Study Type | Results                                                                                                                                   | Conclusion                                                                                           | Reference |
|----------------------------------------------------------------------------|----------------------------|------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------|-----------|
| Human oral epithelial cells (EC), human oral fibroblasts, human osteoblasts, BALB/3T3 cells P. gingivalis | 1 g/L. Cells were exposed to M101 for up to 24 h and cytocompatibility was determined. EC exposed to lipopolysaccharide of P. gingivalis (Pg-LPS) or infected with P. gingivalis to mimic inflammatory state were treated with M101 and inflammatory markers were studied. P. gingivalis biofilms grown on glass surface were treated with M101. | No biologically significant reduction in cellular viability and metabolic activity in all the four cell types tested. → pro-inflammatory markers’ release (TNF-α, NF-κB, RANKL, IL-1β, IL-8, RANTES, IP-10). → anti-inflammatory and pro-resolution mediators (IL-2, IL-4, IL-10, IL-11, IL-15, PDGF-BB, TGF-β, Resolvin-E1 receptor) (p < 0.005). → extracellular matrix and immune modulators (TIMP-2, M-CSF and ICAM-1) (p < 0.005). → P. gingivalis growth. | M101 decreases P. gingivalis-induced inflammation. | [81]      |
| Cold stored porcine proximal tubular cells line (kidney epithelial cells (LLC-PK1 cells)) | M101 was supplemented (0–10 g/L) to University of Wisconsin (UW), histidine-tryptophane-ketoglutarate (HTK), Institut Georges Lopez (IGL-1), Celsior, RingerLactate (RL) and Perfadex. LLC-PK1 cells or N2 gas was used to deoxygenate the solution. M101 SOD activity was evaluated by nitroblue tetrazolium assay (NBT). | Commercial preservative solutions: → cell viability (LDH release); → metabolic activity (MTT) and ATP content. M101 supplementation in commercial preservative solutions: → cell integrity; → cell functionality. Total inhibition of NBT formation with M101 (93.5 ± 1.1%). | Commercial preservative solutions alone were deleterious to cells. M101 protects cells in vitro against cold preservation lesions. M101 is an effective antioxidant. | [57]      |
| Human primary aortic endothelium cells (HAECs)                             | In vitro model of cold hypoxia and reoxygenation simulating reperfusion on cold-preserved cells. HAECs were subjected to 24 h hypoxia at 4 °C in UW solution mimicking cold ischemia during organ preservation. M101 (0, 1, 2.5, 5, 10 g/L), was added to UW solution. UW was removed and cells were re-cultured at 37 °C to mimic organ reperfusion. Necrosis assay, lactate dehydrogenase (LDH) release. | Cells after 24 h cold hypoxia: → LDH released from necrotic cells (32%); → intracellular ATP (47%); → mitochondrial respiratory chain complex II activity. Cells M101 + UW: → LDH, → intracellular ATP; → mitochondrial respiratory chain complex II. Cells in UW alone: → cell organelles altered and damaged (stress). Cells M101 + UW: → necrotic bodies; → organelles integrity presence of autophagosomes rather than necrotic bodies. Cells after 24 h simulated reperfusion: UW alone: → Cell injuries; → LDH release (51%); → intracellular ATP (24%); → respiratory mitochondrial complex II activity (33%). Cells M101 + UW: → LDH release (13%); → intracellular ATP (86%); → metabolic activity (XTT) for the 5 g/L dose (62%) (p ≤ 0.05 vs. 0 g/L). | M101 imparts dose dependent protection against cold hypoxia. M101 conserves cell integrity during preservation. M101 provides dose-dependent protection against IRI. | [63]      |
### Table 3. Cont.

| Application | Concentration & Study Type | Results | Conclusion | Reference |
|-------------|-----------------------------|---------|------------|-----------|
| In vitro Measurement of NO and CO Binding rates | Nitric oxide (NO) and carbon monoxide (CO) reaction kinetics measurements. Briefly, 3 mL of a deoxygenated PBS solution containing HbV, human HbA, Oxyglobin (Biopure) and M101 at [heme] = 3 µM was rapidly mixed with NO or CO containing PBS solution. Changes were recorded at 10, 30 and 60 min post-infusion. | M101 showed a lower binding rates of NO and CO than human Hb and polymerized bovine Hb. | M101 has different binding rates for gases as compared to human Hb. | [40] |

**Note:** ↘ decrease; ↗ increase.

### Table 4. Ex vivo, in vivo and clinical studies involving M101 use.

| Application | Concentration & Study Type | Results | Conclusion | Reference |
|-------------|-----------------------------|---------|------------|-----------|
| Isolated Langendorf-perfused rat hearts (n = 12/group) | Evaluation of M101 (1 g/L) as a protective additive to Celsior solution for static storage of donor hearts prior to transplantation. Heart function in Celsior solution, either alone (control) or with the addition of M101 was measured by intra-ventricular balloon before arrest with cold (7.5 °C). Cold storage (CS) lasted 8 h prior to reperfusion (60 min). Hearts (minced and homogenized) were also assessed by 2,3,5-triphenyltetrazoliumchloride (TTC) staining as a measure of viability and infarct size. | M101: ↗ recovery of left ventricular developed pressure. Recovered heart rate to pre-ischemic value (final recovery between 84 and 89% pre-ischemic value). ↗ coronary flow (7.5 ± 0.7 mL/min) compared to control (5.4 ± 0.4 mL/min) (p < 0.05). Viability and infarct size measurements were similar between groups. | The addition of M101 to Celsior preservation solution significantly improved post-ischemic recovery of heart function. | [62] |
| Rat pancreas preservation | Preservation solution with or without M101 at 2 g/L was injected into the pancreas via the pancreatic duct. Pancreas was removed and placed in the preservation solution at 4 °C, and cold ischemia kinetics were then determined. Samples were taken at 0, 2, 4, 6, 8, 12 or 18 h (n = 4–6). Metabolite extraction was performed on fresh tissue, while protein extraction and OCT slide analysis were performed on snap-frozen tissue at each time-point. Islet isolation was performed after cold ischemia (30 min, 4, 6, 8, 12 or 18 h). For the experiments with M101 perfusion, 2 mL of preservation solution with or without M101 at 2 g/L was injected into the pancreas via the pancreatic duct. Pancreases were preserved for 6 h at 4 °C in the presence of M101, before islet isolation process. 12 h of rat pancreas preservation in the presence of M101. | M101: ↗ maintenance of mitochondrial complex 1 pancreas activity throughout ischemia kinetics compared to controls; ↘ Lactate levels compared to the control; ↘ variability of active caspase-3 levels; ↘ transient phosphorylation of p38, which is observed in the control, after 4 h (p = 0.002) and 6 h. (p = 0.008) of ischemia; ↘ oxidative stress ROS (25%) compared to control; ↘ p38 activity in islets; ↘ necrosis (HMGB1); ↘ cellular stress pathway (p38 MAPK) activity. | Safe, efficient, anti-oxidant. Freshly isolated islets had improved function when M101 injected in the pancreas. | [65] |
Table 4. Cont.

| Application | Concentration & Study Type | Results | Conclusion | Reference |
|-------------|----------------------------|---------|------------|-----------|
| Human pancreas | Evaluation of M101 effects during cold preservation. Human pancreases (n=3) underwent 6 h of cold ischemia between removal from donors and arrival in our laboratory. M101 (2 g/L) was added to Belzer® solution (UW solution, Bridge to Life) and then injected into the pancreas for 3 h. Protein extraction was performed on snap-frozen pancreatic tissue after 1 or 3 h of M101 exposure at 4 °C. Islets were isolated after 9 h of cold ischemia and counted after 12 h culture post-isolation in CMRL medium (Sigma-Aldrich) at 37 °C. Functionality of islets was assessed by a glucose-stimulated insulin secretion test. A Spearman’s correlation was performed to check the influence of age on IEQ or insulin secretion. | M101: ▶ HIF1-α expression in pancreas after 1 h of exposure, but not after 3 h; ▶ AKT phosphorylation in tissue after 3 h of exposure to M101 (p = 0.08); ▶ increased complex 1 activity; ▶ cleaved caspase-3 in tissue; ▶ islet yield and function of pancreases relative to the control. ▶ insulin secretion from islets (both in basal and in stimulated conditions); ▶ overall insulin content; ▶ complex 1 mitochondrial activity; ▶ activation of AKT activity (a cell survival marker); ▶ variability of caspase 3; ▶ oxidative stress (ROS); ▶ necrosis (HMGB1); ▶ cellular stress pathway (p38 MAPK) activity; ▶ islet yield and function. Despite the absence of M101 during the first period of ischemia, positive effects were observed in human pancreas during preservation. | | [65] |
| Pig pulmonary preservation and post-transplant lung function | 36 h cold preservation with 1 g/L of M101. Normothermic ex vivo lung perfusion (EVLP) followed by lung transplant (4 h reperfusion). | M101-treated lungs improved physiologic parameters: ▶ oxygenation than in the controls significantly; ▶ edema formation significantly; ▶ circulating IL-6 within recipient plasma after transplantation. M101 during an extended pulmonary preservation period led to significantly superior early post-transplant lung function. | | [82] |
| Inflammatory calvarial abscess model in mice | M101 (1 g/L). P. gingivalis (5 × 10^8 CFU) was injected subcutaneously on the calvaria to induce inflammatory abscess which was treated with M101 (1 g/L) subcutaneous injection. The healing response was followed for 5 days. | M101: ▶ clinical abscess size significantly; ▶ inflammatory score significantly; ▶ osteoclastic activity. M101 decreased P. gingivalis elicited inflammation and tissue destruction. | | [81] |
| Traumatic brain injury (TBI) rat | M101 (12.5 mL/kg i.v. over 2 h = 625 mg/kg cumulated doses); M101 (12 mL/kg i.v. over 1 h = 600 mg/kg). | M101 (12.5 mL/kg i.v. over 2 h): ▶ MAP (mild 9 mm Hg) of healthy rats without constriction of cerebral pial arterioles. M101 (12 mL/kg i.v. over 1 h): ▶ MAP (modest 27 mm Hg) (peak, 123 ± 5 mm Hg [mean ± standard error of the mean]) of CCI-TBI rats; ▶ PbtO2 brain tissue oxygen (peak, 25 ± 5 mm Hg), restored brain tissue oxygen to 86% of pre-injury level. Proof of concept that M101 may be beneficial in TBI treatment | | [39] |
| Mice: in vivo oxygenation potency of M101 toward HT29 human colorectal adenocarcinoma subcutaneous tumors | M101 I.V. injection at 600 mg/kg and 1200 mg/kg. Single I.V. injection of M101 in mice bearing human-derived subcutaneous tumors. Expression of hyperoxia marker anti-glucose transporter Glut-1 (immunohistochemistry). Rapid diffusion of M101 in brain, liver, lungs and ovaries: ▶ size of carcinomatous areas; ▶ dissociation of tumor; ▶ intensity of Glut-1 staining ∼20% after 1 h when M101 was injected at 1200 mg/kg; ▶ tumor hypoxia of ∼23% (even 5 h after the same treatment); ▶ hypoxia in poorly vascularized tissues. Lower M101 doses (60 or 600 mg/kg), no or intermediate hypoxia reductions. No side effects. Potential oxygen carrying therapeutic product. Ability of M101 to diffuse within poorly vascularized tissues and to behave as a potent O2 carrier towards vertebrate tissues, without inducing obvious side effects. | Rapid diffusion of M101 in brain, liver, lungs and ovaries: ▶ size of carcinomatous areas; ▶ dissociation of tumor; ▶ intensity of Glut-1 staining ∼20% after 1 h when M101 was injected at 1200 mg/kg; ▶ tumor hypoxia of ∼23% (even 5 h after the same treatment); ▶ hypoxia in poorly vascularized tissues. Lower M101 doses (60 or 600 mg/kg), no or intermediate hypoxia reductions. No side effects. Potential oxygen carrying therapeutic product. Ability of M101 to diffuse within poorly vascularized tissues and to behave as a potent O2 carrier towards vertebrate tissues, without inducing obvious side effects. | | [38] |
Table 4. Cont.

| Static lung transplantation pigs | HEMO<sub>2</sub> life added to preservative solution (Perfadex<sup>®</sup>). 24 h of hypothermic preservation followed by lung transplantation. Five hours of lung reperfusion. | HEMO<sub>2</sub> life group:  
  - ↓ graft vascular resistance (<i>p < 0.05</i>);  
  - ↓ graft oxygenation ratio (<i>p < 0.05</i>);  
  - ↓ serum HMG B1. | HEMO<sub>2</sub> life improved early graft function after prolonged cold ischemia. [66] |
| Mice: testing immunogenicity of M101 | Antibody response after 1 or 2 I.V. administrations of M101 in hyper-responsive strain (BP/2 mice) that easily produce antibodies. Plasma levels of IgE and IgG2a measured by ELISA. Comparison with a negative control groups (M101 buffer) and Ovalbumin as a positive control. | M101 did not cause any immunogenic reaction. [35] |
| Pig: kidney autotransplantation model | Four groups were studied for organ preservation and subsequent in vivo transplantation:  
  1. UW;  
  2. UW + M101 (5 g/L);  
  3. HTK;  
  4. HTK + M101 (5 g/L). Controls were sham-operated animals. Animals were followed-up until the sacrifice at 3 months. | In M01 kidney grafts:  
  - ↑ resumption of urine production;  
  - ↑ kidney graft function recovery;  
  - ↑ preservation of tissue integrity;  
  - ↓ inflammation;  
  - ↓ invasion of both innate (ED1+) and adaptative (CD3+) immune cells;  
  - ↓ chronic fibrosis;  
  - ↓ tubular atrophy. | Beneficial use of M101 in 2 of the most used preservation solutions (better short-term function recovery and reduced development of fibrosis, the main cause of graft loss). [57] |
| Pig: kidney autotransplantation model | Kidney was harvested, cold flushed and preserved for 24 h at 4 °C before transplantation. Dose-ranging study: 1, 2, and 5 g/L M101 supplementing UW solution. Pigs placed in metabolic cages for diuresis and serum creatinine measurements. Biopsies for conventional histology and fibrosis evaluated on sacrifice by Sirius red staining. | ↓ intensity of IRI;  
  - ↓ interstitial fibrosis;  
  - ↑ graft outcome. Area under the curve of serum creatinine for the first 2 weeks after transplantation showed that the 1 g/L and 2 g/L groups were efficient and nearly identical. | HEMO2Life improved kidney graft function. Beneficial use of M101 at lower dosage. [63] |
| Kidney machine preservation in a porcine transplantation model | Kidneys were submitted to 1 h-warm ischemia, followed by 23 h hypothermic preservation in Waves machine perfusion (MP) before auto-transplantation. Four groups were studied:  
  1. W (MP without 100%-O<sub>2</sub>);  
  2. W-O<sub>2</sub> (MP with 100%-O<sub>2</sub>; also called hyperoxia);  
  3. W-M101 (MP without 100%-O<sub>2</sub> + M101 2 g/L);  
  4. W-O<sub>2</sub> + M101 (MP with 100%-O<sub>2</sub> + M101 2 g/L). | First week post-transplantation, W-M101 group:  
  - ↑ renal resistance;  
  - ↑ blood creatinine;  
  - ↑ glomerular filtration rate;  
  - ↑ KIM-1;  
  - ↑ IL-18 blood levels;  
  - ↑ kidney fibrosis. | Supplementation with M101 associated with or without 100% O2 improved the Waves MP effect upon kidney recovery and late graft outcome. [80] |
**Table 4. Cont.**

| Cold preservation in a preclinical porcine model of kidney donation after cardiac death | Comparison of addition of M101 in CS and MP (LifePort). |
| CS + M101: | In the CS arm, M101 improved long-term function, normalizing creatininemia. |
| ↗ | In the MP arm, M101 improved both short- and long-term functional outcomes, as well as tissue integrity. |
| ↓ | M101 is a viable strategy to improve current organ preservation method in marginal organ transplantation. |

| Pig liver orthotopic allotransplantation model | HOPE or SCS+M101 compared with SCS alone: |
| Addition of M101 to static cold storage (SCS) on the quality of pig liver graft preservation. Pig liver grafts were preserved using: 1. SCS; 2. Hypothermic oxygenated perfusion machine (HOPE); 3. SCS + M101 (1 g/L). | After 9 h of ex vivo preservation, liver functions: |
| HOPE ≥ SCS + M101 > SCS | When added to SCS, M101 effectively oxygenated liver grafts during preservation, preventing post-transplant injury without reaching the level of HOPE. This needs to be weighed against the cumbersomeness and cost of infusion machines, which require continuous presence of highly-skilled operators, and are associated with a risk of vascular trauma in organ grafts that might compromise the outcome of transplantation. |
| ↗ | Mitochondrial function; |
| ↗ | ATP synthesis; |
| ↗ | Anti-oxidant capacities; |
| ↗ | Hepatocyte architecture preservation; |
| ↗ | Free radical production; |
| ↗ | Inflammatory mediators. |
| ↓ | Blood aspartate; |
| ↓ | Blood alanine aminotransferases; |
| ↓ | Blood LDH at day one post-transplant. |
| At days 1 and 3: | At day 7: |
| ▲ TNF-α levels. | Liver cell necrosis; |
| HOPE ≥ SCS + M101 > SCS | Liver inflammation; |

| Hamster & rat cardio-vascular study | In hamster: absence of microvascular vasoconstriction and no significant effect on mean arterial blood pressure. |
| Hamster study: dorsal skinfold window chamber model to evaluate microcirculation | In rat: minor effects on mean arterial pressure (differences not statistically significant compared to control), heart rate and myocardial contractility. |
| Changes were recorded 60 min after M101 infusion. Rat study: to evaluate cardiovascular effects. M101 (600 mg/kg) or saline solution 0.9% (control). Changes were recorded 45 min after M101 infusion. | M101 appears to have no vasoactivity at the microvascular level. |

| Human safety and proof-of-principle study | First-in-human use of M101 (1 g/L) for organ preservation. Open-label study investigating the safety of M101 used ex vivo as an additive to the preservation solution in kidney transplantation (Clinical Trial Registry No. NCT 02652520). Grafts were preserved either in cold storage (standard donor) or on machine perfusion (extended criteria donor). |
| 60 graft kidneys from 60 deceased donors were preserved with M101. M101 is safe for the graft and for the recipient. No allergic or hypersensitivity reactions or infections related to the product were reported. Less DGF (at least one dialysis) in the M101 group was reported, and the difference between the two groups (23% vs. 33%) was clinically relevant but not statistically significant. When a more stringent definition (more than one dialysis) is used, the difference (7% vs. 26%) was statistically significant (p = 0.038). |
| This study demonstrated that the addition of the oxygen carrier M101 to preservation solution is safe. Although the study was not designed to show the superiority of M101, the analysis of the secondary efficacy end points is highly promising, with better renal function in recipients of the kidneys preserved with M101. |
7. Perspectives

Owing to its interesting \(O_2\) carrying capacity, the potential of M101 as an \(O_2\) transporter for numerous pathologies involving hypoxia is worth exploring. Furthermore, its prospects as a therapeutic \(O_2\) carrier for conditions involving massive hemorrhage, for instance, accidents, terrorist attacks and war injuries should be investigated. M101’s potential application as a temporary or intermittent use alternative for reducing the adverse side-effects associated to pathologies requiring frequent blood transfusions should be further investigated. Additionally, with the recent evolution of its anti-inflammatory and anti-bacterial properties, further research is essential to examine its therapeutic potential for the treatment of inflammatory and infectious diseases. Besides, its incorporation into scaffolds should be tested to establish its feasibility in clinical applications.

Interestingly, the product HEMO\(_2\)life\(^\circ\) has been used to preserve a face transplant during a full-face re-transplantation. The patient was transplanted in January 2018 with a graft preserved with HEMO\(_2\)life\(^\circ\). The surgery was a success and 30 months post-surgery, the patient is safe and the graft has not been rejected \(^{[83]}\). Recently, M101 has also been proposed as a “molecular respirator”, a potential tool to help COVID-19 patients in their struggle with hypoxemia \(^{[84]}\). Testing this hypothesis further could be instrumental in developing a potential therapeutic strategy to combat the COVID-19 crisis.

8. Conclusions

Therapeutic \(O_2\) carriers could be used as graft/organ preservatives and could also present an alternative in cases requiring frequent blood transfusion. However, extensive investigation is warranted to establish their safety and efficacy, especially for their potential clinical application. Moreover, M101, being an efficient \(O_2\) transporter, may be a promising medical product with therapeutic potential for the management of several pathologies.

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References

1. Briehl, M.M. Oxygen in human health from life to death—An approach to teaching redox biology and signaling to graduate and medical students. *Redox Biol.* 2015, 5, 124–139. [CrossRef]
2. Dunn, J.-O.; Mythen, M.G.; Grocott, M.P. Physiology of oxygen transport. *BJA Educ.* 2016, 16, 341–348. [CrossRef]
3. Sen, C.K. Wound healing essentials: Let there be oxygen. *Wound Repair Regen.* 2009, 17, 1–18. [CrossRef]
4. Morrell, C.N. Reactive oxygen species: Finding the right balance. *Circ. Res.* 2008, 103, 571–572. [CrossRef]
5. Li, C.; Jackson, R.M. Reactive species mechanisms of cellular hypoxia-reoxygenation injury. *Am. J. Physiol. Physiol. Cell Physiol.* 2002, 282, C227–C241. [CrossRef]
6. Van Golen, R.F.; Reiniers, M.J.; Vrisekoop, N.; Zuurbier, C.J.; Olthof, P.B.; van Rheenen, J.; van Gulik, T.M.; Parsons, B.J.; Heger, M. The Mechanisms and Physiological Relevance of Glycocalyx Degradation in Hepatic Ischemia/Reperfusion Injury. *Antioxid. Redox Signal.* 2014, 21, 1098–1118. [CrossRef]
7. Ivanovic, Z. Stem cell evolutionary paradigm and cell engineering. *Transfus. Clin. Biol.* 2017, 24, 251–255. [CrossRef]
8. Wong, H.-S.; Dighe, P.A.; Mezera, V.; Monternier, P.-A.; Brand, M.D. Production of superoxide and hydrogen peroxide from specific mitochondrial sites under different bioenergetic conditions. *J. Biol. Chem.* 2017, 292, 16804–16809. [CrossRef]
9. Liaudet, L.; Szabó, G.; Szabó, C. Oxidative stress and regional ischemia-reperfusion injury: The peroxynitrite-poly(ADP-ribose) polymerase connection. *Coron. Artery Dis.* 2003, 14, 115–122. [CrossRef]
10. Szabó, G.; Soós, P.; Bährle, S.; Zsengellér, Z.; Flechtenmacher, C.; Hagl, S.; Szabó, C. Role of poly(ADP-ribose) polymerase activation in the pathogenesis of cardiopulmonary dysfunction in a canine model of cardiopulmonary bypass. *Eur. J. Cardio-Thoracic Surg.* 2004, 25, 825–832. [CrossRef]
11. Baidukova, O.; Wang, Q.; Chaivaree, S.; Freyer, D.; Prapan, A.; Georgieva, R.; Zhao, L.; Báumlér, H. Antioxidative protection of haemoglobin microparticles (HbMPs) by PolyDopamine. *Artif. Cells Nanomed. Biotechnol.* 2018, 46, S693–S701. [CrossRef]
65. Thuillier, R.; Dutheil, D.; Trieu, M.T.N.; Mallet, V.; Allain, G.; Rousselot, M.; Denizot, M.; Goujon, J.-M.; Zal, F.; Hauet, T. Supplementation With a New Therapeutic Oxygen Carrier Reduces Chronic Fibrosis and Organ Dysfunction in Kidney Static Preservation. Am. J. Transplant. 2011, 11, 1845–1860. [CrossRef]  
66. Glorion, M.; Polard, V.; Faveureau, F.; Hauet, T.; Zal, F.; Fadel, E.; Sage, E. Prevention of ischemia-reperfusion lung injury during static cold preservation by supplementation of standard preservation solution with HEMO²Life® in pig lung transplantation model. Artif. Cells Nanomed. Biotechnol. 2018, 46, 1773–1780. [CrossRef]  
67. Gupta, A.S. 2017 Military Supplement: Hemoglobin-based Oxygen Carriers: Current State-of-the-Art and Novel Molecules. Shock 2017. [CrossRef]  
68. Salahudeen, A.K. Cold ischemic injury of transplanted kidneys: New insights from experimental studies. Am. J. Physiol. Ren. Physiol. 2004, 287, F181–F187. [CrossRef]  
69. Bernhardt, W.M.; Gottmann, U.; Doyon, F.; Buchholz, B.; Campean, V.; Schödel, J.; Reisenbuechler, A.; Klaus, S.; Arend, M.; Flippin, L.; et al. Donor treatment with a PHD-inhibitor activating HIFs prevents graft injury and prolongs survival in an allogenic kidney transplant model. Proc. Natl. Acad. Sci. USA 2009, 106, 21276–21281. [CrossRef]  
70. Bos, E.M.; Leuvenink, H.G.D.; Snijder, P.M.; Kloosterhuis, N.J.; Hillebrands, J.-L.; Leemans, J.C.; Florquin, S.; van Goor, H. Hydrogen Sulfide-Induced Hypometabolism Prevents Renal Ischemia/Reperfusion Injury. J. Am. Soc. Nephrol. 2009, 20, 1901–1905. [CrossRef]  
71. ‘T Hart, N.A.; van der Plaats, A.; Faber, A.; Leuvenink, H.G.D.; Olinga, P.; Wiersema-Buist, J.; Verkerke, G.J.; Rakhorst, G.; Ploeg, R.J. Oxygenation during hypothermic rat liver preservation: An in vitro slice study to demonstrate beneficial or toxic oxygenation effects. Liver Transpl. 2005, 11, 1403–1411. [CrossRef] [PubMed]  
72. Maathuis, M.-H.J.; Leuvenink, H.G.D.; Ploeg, R.J. Perspectives in organ preservation. Transplantation 2007, 83, 1289–1298. [CrossRef] [PubMed]  
73. Ahlenstiel, T.; Burkhardt, G.; Köhler, H.; Kuhlmann, M.K. Improved cold preservation of kidney tubular cells by means of adding flavonoids to organ preservation solutions. Transplantation 2006, 81, 231–239. [CrossRef] [PubMed]  
74. Joehmans, I.; Moers, C.; Smits, J.M.; Leuvenink, H.G.D.; Treckmann, J.; Paul, A.; Rahmel, A.; Squifflet, J.-P.; van Heurn, E.; Monbaliu, D.; et al. Machine perfusion versus cold storage for the preservation of kidneys donated after cardiac death: A multicenter, randomized, controlled trial. Ann. Surg. 2010, 252, 756–764. [CrossRef]  
75. Moers, C.; Smits, J.M.; Maathuis, M.-H.J.; Treckmann, J.; van Gelder, F.; Napieralski, B.P.; van Kasterop-Kutz, M.; van der Heide, J.J.H.; Squifflet, J.-P.; van Heurn, E.; et al. Machine perfusion or cold storage in deceased-donor kidney transplantation. N. Engl. J. Med. 2009, 360, 7–19. [CrossRef] [PubMed]  
76. Vaziri, N.; Thuillier, R.; Favreau, F.D.; Eugene, M.; Milin, S.; Chatauret, N.P.; Hauet, T.; Barrou, B. Analysis of machine perfusion benefits in kidney grafts: A preclinical study. J. Transl. Med. 2011, 9, 15. [CrossRef] [PubMed]  
77. Eleftheriadis, T.; Lawson, B.R. Toll-like receptors and kidney diseases. Inflamm. Allergy Drug Targets 2009, 8, 191–201. [CrossRef]  
78. Evankovich, J.; Billiar, T.; Tsung, A. Toll-like Receptors in Hepatic Ischemia/Reperfusion and Transplantation. Available online: https://www.hindawi.com/journals/grp/2010/537263/ (accessed on 1 September 2020).  
79. Kaminski, J.; Hannaert, P.; Kasil, A.; Thuillier, R.; Leize, E.; Delpy, E.; Steichen, C.; Goujon, J.M.; Zal, F.; Hauet, T. Efficacy of the natural oxygen transporter HEMO²Life® in cold preservation in a preclinical porcine model of donation after cardiac death. Transpl. Int. 2019, 32, 985–996. [CrossRef]  
80. Kasil, A.; Giraud, S.; Couturier, P.; Amiri, A.; Danion, J.; Donatini, G.; Matillon, X.; Hauet, T.; Badet, L. Individual and Combined Impact of Oxygen and Oxygen Transporter Supplementation during Kidney Machine Preservation in a Porcine Preclinical Kidney Transplantation Model. Int. J. Mol. Sci. 2019, 20, 1992. [CrossRef]  
81. Batool, F.; Stutz, C.; Petit, C.; Benkirane-Jessel, N.; Delpy, E.; Zal, F.; Leize-Zal, E.; Huck, O. A therapeutic oxygen carrier isolated from Arenicola marina decreased P. gingivalis induced inflammation and tissue destruction. Sci. Rep. 2020, 10, 14745. [CrossRef]  
82. Ali, A.; Watanabe, Y.; Galasso, M.; Watanabe, T.; Chen, M.; Fan, E.; Broach, L.; Ramadan, K.; Ribeiro, R.V.P.; Stansfield, W.; et al. An extracellular oxygen carrier during prolonged pulmonary preservation improves post-transplant lung function. J. Hear. Lung Transplant. 2020, 39, 593–595. [CrossRef] [PubMed]  
83. Lantieri, L.; Cholley, B.; Lemogne, C.; Guillemain, R.; Ortonne, N.; Grimbert, P.; Thervet, E.; Lellouch, A.G. First human facial retransplantation: 30-month follow-up. Lancet 2020, 396, 1758–1765. [CrossRef]  
84. Lupon, E.; Lellouch, A.G.; Zal, F.; Cetrulo, C.L.; Lantieri, L.A. Combating hypoxemia in COVID-19 patients with a natural oxygen carrier, HEMO²Life® (M101). Med. Hypotheses 2021, 146, 110421. [CrossRef] [PubMed]