Hepatoprotective Effects of Poly-[hemoglobin-superoxide Dismutase-catalase-carbonic anhydrase] on alcohol damaged primary rat hepatocyte culture in vitro

Wenhua Jiang, Yuzhu Bian, Zhenghui Wang, and Thomas Ming Swi Chang
Artificial Cells and Organs Research Centre, Departments of Physiology, Medicine and Biomedical Engineering, Faculty of Medicine, McGill University, Montreal, Quebec, Canada

Abstract: We have prepared a novel nanobiotherapeutric, Poly-[hemoglobin-superoxide dismutase-catalase-carbonic anhydrase], which not only transports both oxygen and carbon dioxide but is also a therapeutic antioxidant. Our earlier study in a severe sustained 90 minutes hemorrhagic shock rat model shows that it has a hepatoprotective effect. We investigate its hepatoprotective effect further in this present report using an alcohol damaged primary hepatocyte culture model. Results show that it significantly reduced ethanol-induced AST release, lipid peroxidation and ROS production in rat primary hepatocytes culture. It also significantly enhanced the viability of ethanol-treated hepatocytes. Thus, the result shows that Poly-[hemoglobin-superoxide Dismutase-catalase-carbonic anhydrase] also has some hepatoprotective effects against alcohol-induced jury in in-vitro rat primary hepatocytes cell culture. This collaborate our earlier observation of its hepatoprotective effect in a severe sustained 90 minutes hemorrhagic shock rat model.

Key Words: PolyHb, Polyhemoglobin-superoxide Dismutase-catalase-carbonic anhydrase, primary rat hepatocytes, alcohol, reactive oxygen species, malondialdehyde, aspartate aminotransferase, hepatoprotective.

Corresponding author: Professor TMS Chang, Director, Artificial Cells and Organs Research Centre, Faculty of Medicine, McGill University, Montreal, QC Canada, H3G1Y6 artcell.med@mcgill.ca

INTRODUCTION
Red blood cells have 3 major functions: transport oxygen, transport carbon dioxide and remove oxygen radicals. Poly-[Hemoglobin-Superoxide dismutase-catalase] (Poly-[Hb-SOD-CAT]) was prepared to carry oxygen and removal oxygen radicals (D'Agnillo D & Chang, 1998). It could protect the brain from ischemia-reperfusion injuries in a severe hemorrhagic shock-cerebral ischemia rat model (Powanda & Chang, 2002). Others showed that it could also prevent ischemia-reperfusion injury in liver and kidney transplantation in rats (Chang, E.J., et al.; 2004).

Recent study shows that increase in tissue CO$_2$ may also be related to organ ischemia (Tronstad C. 2010). Thus we have more recently devised a novel nanobiotherapeutic with enhancement of all 3 rbc functions in the form of Poly-[Hemoglobin-Superoxide dismutase-catalase-carbonic anhydrase] (Poly-[Hb-SOD-CAT-CA]) (Bian, Rong & Chang, 2011). Our study in a severe 90 minutes sustained hemorrhagic shock rat model shows that it is more effective than whole blood in the protective effect for the heart, liver, kidney and intestine (Bian & Chang 2015). The purpose of the present study is to further investigate the hepatoprotective effect of
Poly-[Hb-SOD-CAT-CA] using an alcohol damaged primary rat hepatocytes culture model. Alcohol-dependent liver disease (ALD) ranges in severity from simple steatosis (fatty liver) to steatohepatitis, cirrhosis and hepatocellular carcinoma (Tome & Lucey 2004). Although the exact mechanism of alcohol-induced hepatotoxicity is not clear yet, this adverse effect has been closely related to reactive oxygen species (ROS) generation during ethanol metabolism (Ishii, Kurose & Kato, 1977).

MATERIALS AND METHODS

Materials

The stroma-free hemoglobin (SFHb) used in our experiments was extracted from bovine blood purchased from the McGill University McDonald Campus Cattle Complex (Sainte-Annee-Bellevue, Canada). Glutaraldehyde (25%) was purchased from Poly-science (Warrington, PA). Drabkin’s Reagent, L-lysine (monohydrochloride, Sigma Ultra >99%), SOD, CAT, and bovine CA were purchased from Sigma-Aldrich (Ontario, Canada). All other chemicals or reagents of analytical grade were purchased from Sigma-Aldrich.

Preparation of PolySFHb, Poly-[Hb-SOD-CAT-CA]

PolySFHb, Poly-[Hb-SOD-CAT-CA] were prepared using the glutaraldehyde methods described (Chang 2007, Bian & Chang 2015). For the preparation of Poly-[Hb-SOD-CAT-CA], SOD (1050 units/mL), catalase (21,000 units/mL), and carbonic anhydrase (1070 units/mL) were added to the 20 mL solution containing stroma-free hemoglobin (7 g/dl) in 50mM sodium phosphate buffer (pH 7.4). For the preparation of PolySFHb, an equivalent volume of buffer containing no enzymes was added. The Hb concentration in the PolySFHb and Poly-[Hb-SOD-CAT-CA] preparations was colorimetrically determined by reacting the samples with Drabkin’s reagent (Sigma-Aldrich).

Determination of CAT and SOD Activity:

The method for catalase measurement was according to that of D’Agnillo & Chang (1989) and Chang (2007). The catalase activity was expressed in units per grams of hemoglobin. The measurement for SOD activity was based on the reduction of cytochrome c by superoxide.

Rat primary hepatocytes culture and treatments.

Primary Hepatocytes were isolated from rats according to the two-step collagenase perfusion technique described by Seglen (1976) with minor modifications as follows. After perfusion with a chelating agent to allow the cleavage of the hepatic desmosomes, hepatic collagen was hydrolyzed by ex situ perfusion with a collagenase solution supplemented with dispase at a rate of 35ml/minute at 37°C for 10-12 minutes. The liver was excised and minced in cold William’s E medium (GIBCO) to free the hepatocytes. The hepatocytes were collected and filtered through a 100 um nylon mesh on ice. Then sedimented by centrifugation, resuspended and washed 2-3 times at 50g at 4°C for 5 minutes, the hepatocytes were purified by 90% Percoll solution (Sigma). The viability of hepatocytes measured by trypan blue exclusion test was consistently within the range of 88-94%.

The primary hepatocytes were cultured with complete culture medium (William’s E medium supplemented with 5% FCS, 1% penicillin/streptomycin,4ug/mL insulin, 2mM glutamate,15mM HEPES) in previously collagen-coated culture wells at a density of 1x10⁵ cells/mL. Four hours after plating, the medium was changed with fresh medium to remove the
non-viable cells and cell particles. After 24h of incubation at 37°C in a humidified atmosphere composed of 5% CO₂ in air, the cultured cells were seeded in 96-well plates at a density of 4.8x10⁴ cells/mL. Cell viability evaluation was based on 12-well plates containing 200 μL per well at a density of 2.4x10⁵ cells/mL. MDA content and AST leakage analysis was based on 6-well plates containing 600 μL per well at a density of 2.4x10⁵ cells/mL. Reactive oxygen species (ROS) in live cells analysis was carried out in 6-well plates containing 1000 μL per well at a density of 2.4x10⁵ cells/mL. The following groups of study were carried out each of the analysis:
1: Add alcohol 200 mmol/L to the hepatocyte culture and incubate for 12h
2: 3 groups pre-incubated at 37°C for 2 h, each group with respectively 5, 10 or 15ul Poly-[Hb-SOD-CAT-CA]. After this, Alcohol (200 mmol/L) was added and incubate for 12 hours
3: 3 groups pre-incubated at 37°C for 2 h each group with respectively 5, 10 or 15ul PolySFHb. After this Alcohol (200 mmol/L) was added and incubate for 12 hours

Measurement of aspartate aminotransferase (AST) activity
AST leakage of the hepatocytes was used to study membrane integrity of the cells. The culture medium was centrifuged at 1600g for 10 min and the supernatant was collected. AST was measured using commercially available assay kits (Cat.NO.MAK 055, Sigma, USA). This was based on the mechanism that AST facilitates the conversion of aspartate and alpha-ketoglutarate to oxaloacetate and glutamate. This AST activity assay is based on the transfer of an amino group from aspartate to alpha-ketoglutarate resulting in the generation of glutamate, and the production of a colorimetric product proportional to the AST enzymatic activity present. After incubation, culture supernatant was removed and the AST level was measured using a microplate reader at a wavelength of 40nm according to the supplier instructions, the results were expressed as units/liter (U/L).

Assaying of malondialdehyde (MDA) content
MDA content of the cells was measured by a OXI-TEK TBARS Assay kit (enzolife sciences, Cat.NO.ALX-850-287-K101,CA). This was based on the mechanism that the MDA can react with thiobarbituric acid (TBA) to generate a color compound with a maximal absorbance at 532 nm. In brief, 2x10⁵ hepatocytes in 500 ul of PBS were sonicated for 5 seconda intervals at 40V setting over ice to collect the whole homogenates in the assay. 0.1 mL of the whole homogenates was mixed with 2.5 mL TBA/Buffer Reagent and incubated in a water bath for 60 min. The absorbance was recorded on the microplate reader at a wavelength of 532 nm. MDA content was expressed as nmol/ml to reflect lipid oxidation.

Measurement of oxidative stress
The measurement of ROS production in hepatocytes was quantified by CellRox Green (Invitrogen, USA,CAT.NO.C10444) staining using fluorescence microscopy (Sollinger et al. 2014). In brief, after incubation, hepatocytes were washed with phosphate-buffered saline and stained with 5 mmol/L CellRox Green and 10ug/mL of Hoechst 33342 (Life Technologies, USA) for 30 minutes. Then placed on fluorescence microscope (Carl Zeiss, Oberkochen, Germany) with a 10-fold magnification objective. Excitation/Emission was recorded at 485/520 nm, respectively. Total nuclei and CellRox Green positive nuclei were determined simultaneously and an overlap of staining of nuclei was performed. CellRox-positive nuclei were only counted in the case of a positive overlap.
Analysis of cell viability

Cell viability was measured by the MTT assay that is based on the production of a blue product, formazan, by the living cells (Kang et al 2005). After the cells were treated as described previously, 20uL MTT (Sigma, USA) solution (5 g/L) was added into each well and incubated 4 h at 37°C. Then, 150 uL DMSO were added into each well to dissolve the formazan generated. The absorbance of each well was recorded on a microplate reader (Biotek) at a wavelength of 490 nm. The data was recorded using the software package Gen 5™. The viability was calculated as follows:

\[
\text{Viability} = \frac{\text{Average of test wells OD} - \text{Average of blank wells OD}}{\text{OD}} \times 100
\]

where OD is optical density.

Statistical analysis

Values were expressed as mean±SD. To evaluate the differences between the groups studied, one way analysis of variance (ANOVA) was used. Differences were considered statistically significant when P<0.05.

RESULT

Molecular Weight Distribution of PolyHb-SOD-CAT-CA and Enzyme Activities

Sepacryl S-300 gel column chromatography was used to analyze the molecular weight distribution of PolyHb-SOD-CAT-CA. The molecular weight distributions for Poly-[Hb-SOD-CAT-CA] show three molecular components: (1) low (<100kDa); (2) intermediate (100-450kDa); and (3) high molecular weight (>450kDa). The sample contains about 86% components with molecular weight higher than 100kDa. The low molecular weight fraction(<100kDa) was discarded before use I the present study. Most of the Hb, SOD, CAT, and CA activity were in the Poly-[Hb-SOD-CAT-CA] fraction with a molecular weight of >450 kDa. SOD activity was 51,608.33 and 89,678.66(U/g) in Poly-[Hb-SOD-CAT-CA] and PolySFHb respectively, CAT activity was 12,301,682.7 and 275,983.0(U/g) in Poly-[Hb-SOD-CAT-CA] and PolySFHb respectively.

Effects of PolySFHb and PolyHb-SOD-CAT-CA on the levels of hepatic transaminase (AST) released into the culture medium.

Incubation of rat primary hepatocytes with ethanol for 12 h, resulted in a significant increase in AST leakage compared to control cells. Pre-incubation with 5,10 or 15 ul of PolySFHb did not have any reduction of AST release. Poly-[Hb-SOD-CAT-CA] pre-incubation significantly reduced the release of AST only at the higher dosage of 15 ul. There was no significant effect at the lower dosage of 5 and 10 ul (Fig.1).
Figure 1. Effects of Poly-[Hb-SOD-CAT-CA] (pe) on ethanol-induced AST release in rat primary hepatocytes. Rat primary hepatocytes were pre-incubated with 5ul, 10ul or 15ul of Poly-[Hb-SOD-CAT-CA] (pe) or PolyHb (p) for 2h before treated with 200mmol/L alcohol (a) for 12h. *P<0.05 versus 200mm alcohol group.

Effects of PolySFHb and PolyHb-SOD-CAT-CA on ethanol-induced lipid peroxidation
The effects of Poly-[Hb-SOD-CAT-CA] on ethanol-induced lipid peroxidation were measured by following the MDA level in culture lysates. Ethanol-exposure for 12 h significantly (P < 0.01) increased intracellular MDA level compared to control in rat primary hepatocytes. Pre-incubation with 5, 10 or 15 ul PolySFHb did not have any reduction in MDA formation of rat primary hepatocytes. However, 15 ul of PolyHb-SOD-CAT-CA resulted in a significant decrease in the MDA levels.
Figure 2. Poly-[Hb-SOD-CAT-CA] down-regulates ethanol-induced MDA in rat primary hepatocytes. Rat primary hepatocytes were pre-incubated with Poly-[Hb-SOD-CAT-CA] (pe) PolySFHb (p) for 2h before exposure to 200mmol/L alcohol (a) for 12h. *P<0.05 versus 200mm alcohol group.

Poly-[Hb-SOD-CAT-CA] prohibits ethanol-induced ROS production in rat primary hepatocytes

The effects of Poly-[Hb-SOD-CAT-CA] on ethanol-induced ROS production were measured by the CellRox green reagent. Ethanol-exposure for 12 h significantly (P < 0.01) increased ROS production compared to control in rat primary hepatocytes. Pre-incubation with 5, 10 or 15 ul PolySFHb did not have any reduction on ROS production of rat primary hepatocytes (Fig. 3, Fig.4 G, H and I). However, ROS production were significantly reduced when pre-incubated with 10ul and 15ul of Poly-[Hb-SOD-CAT-CA] (Fig. 3, Fig.4 E and F).
Figure 3. Effects of Poly-[Hb-SOD-CAT-CA] on ROS generation in rat primary hepatocytes injured by alcohol (a). Rat primary hepatocytes were pre-incubated with 5ul, 10ul or 15ul of Poly-[Hb-SOD-CAT-CA] (pe) and PolyHb (p) for 2h before treated with 200mmol/L alcohol (a) for 12h. *P<0.05 versus 200mm alcohol group.

Figure 4. ROS production was determined by CellRox Green and Hoechst33342 staining in rat primary hepatocytes. Rat primary hepatocytes were pre-incubated with Poly-[Hb-SOD-CAT-CA] or PolySFHb for 2h before treated with 200mmol/L alcohol for 12h. B: control, C: 200mm alcohol, D: 5ul PolyHb-SOD-CAT-CA +200mm alcohol, E: 10ul PolyHb-SOD-CAT-CA +200mm alcohol , F: 15ul PolyHb-SOD-CAT-CA +200mm alcohol, G: 5ul PolyHb+200mm alcohol , H: 10ul PolyHb+200mm alcohol, I: 15ul PolyHb+200mm alcohol. (Arrow points an example of overlap of staining).
Effects of PolyHb-SOD-CAT-CA on the viability of rat primary hepatocytes

The results of MTT assay (Fig.5) showed that 5, 10, 15 ul PolySFHb did not improve the viability of rat primary hepatocytes when compared with the group exposed to 200mm alcohol for 12h without pre-incubation with PolySFHb. On the other hand, Poly-[Hb-SOD-CAT-CA] significantly increased the viability of hepatocytes compared with the 200mm alcohol group, but only at the higher dose of 15ul.

![Figure 5](image.png)

**Figure 5.** Effects of PolyHb-SOD-CAT-CA on the viability of rat primary hepatocytes. Rat primary hepatocytes were preincubated with 5ml, 10ml or 15ml of PolyHb-SOD-CAT-CA (pe) and PolyHb (p) for 2h before treated with 200mmol/L alcohol (a) for 12h. *P<0.05 versus 200mm alcohol group.

**DISCUSSION**

In the present study, the decrease of cell viability and the significant elevated activities of AST in the supernatants of the alcohol-induced hepatocytes indicated the cellular leakage and hepatocyte damage. Pretreatment of the hepatocytes with Poly-[Hb-SOD-CAT-CA] (15 µl group) significantly increased the cell viability and decreased the release of AST, indicating that Poly-[Hb-SOD-CAT-CA] could maintain the functional integrity of the hepatocyte membrane, and protect the hepatocytes against alcohol-mediated toxicity. Moreover, alcohol-mediated free radical formation has a great potential to react with lipid molecules, which leads to lipid peroxidation. During alcohol metabolism, alcohol oxidized into aldehydes especially acetaldehydes and malonaldehydes (MDA). MDA is a decomposition product of lipid hydroperoxides (Gutteride 1995). It is widely used as a marker of lipid
peroxidation (Mansour, 2000) and its elevated level could reflect the degree of lipid peroxidation injury in hepatocytes (Hu et al 2001). In our present study, MDA levels in the supernatant of alcohol treated hepatocytes were significantly elevated, pretreatment of the hepatocytes with Poly-[Hb-SOD-CAT-CA] (15 µl group) significantly suppressed the elevation of MDA caused by alcohol, indicating the anti-lipid peroxidation effect of Poly-[Hb-SOD-CAT-CA].

ROS generated by alcohol metabolism is a putative mechanism underlying alcoholic liver diseases. In this study, ROS level of alcohol treated hepatocytes was significantly elevated. Pretreatment of the hepatocytes with Poly-[Hb-SOD-CAT-CA] (10,15 µl groups) significantly reduced the elevation of ROS caused by alcohol. Thus, Poly-[Hb-SOD-CAT-CA] treatment significantly prevented ethanol-induced intracellular ROS accumulation alcohol treated hepatocytes.

In summary, the present findings demonstrated the hepatoprotective effect of Poly-[Hb-SOD-CAT-CA] against rat hepatocytes damage induced by alcohol. The hepatoprotective activity of Poly-[Hb-SOD-CAT-CA] may be attributed to the enhancement of the hepatic antioxidant system, such as down-regulating hepatic transaminase enzymes, MDA and intracellular ROS generation. This result study supports and explains the hepatoprotective effect of Poly-[Hb-SOD-CAT-CA] in our earlier study in a severe 90 minutes sustained hemorrhagic shock rat model (Bian & Chang 2015).

**Disclosure statement**

This laboratory is not connected to any commercial organizations. The research support for the study reported here comes from the Canadian Blood Service/Canadian Institutes of Health Research joint program that requires the authors to state that: "The opinion expressed in this article is the opinions of the investigators of this paper and not neccessary those of the Canadian Blood Service nor of the Canadian Government ". Dr W Jiang is a China Scholarship Council visiting scholar from the Department of Histology and Embryology, Norman Bethune Medical College of Jilin University, Changchun 130021, Jilin Province, China

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