Conservation of Binding of Llama and Other Animals’ Hemoglobins to Haptoglobins Across Species

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Abstract: The plasma protein haptoglobin binds hemoglobin released from lysed erythrocytes. It causes removal of the free hemoglobin, thus preventing pathological oxidation of cells. Hemoglobin higher level structures are well conserved across animal species but primary structures can be as little as 50% homologous. Because of these differences, the question arose as to what extent hemoglobins can bind to haptoglobins from different species. The charge properties and binding to three genetic variants of human haptoglobin were compared by non-denaturing agarose gel electrophoresis, with llama, human, dog, pig, horse and goat hemoglobins. In this study, it was reported for the first time that llama and alpaca hemoglobins differed in electrophoretic mobility from hemoglobins from several animal species and humans. Llama hemoglobin was more positively charged than the other mammalian hemoglobins. Electrophoretic mobility changes of the animal hemoglobins in the presence of human plasma and two different purified human haptoglobin genetic variants suggested that hemoglobins from all animals in this study could bind all three genetic variants of human haptoglobin. In all cases, the llama hemoglobin-haptoglobin samples had lesser mobility than those of the other mammals. This study showed that the binding sites on hemoglobin and haptoglobin for each other have been evolutionarily conserved despite differences in primary structure and marked difference in the charge of llama hemoglobin from the other animal species and humans.

Keywords: Blood, Erythrocytes, Haptoglobin, Hemoglobin, Llama, Proteins, Polymorphism

Introduction

Hemoglobin is essential in vertebrates as an oxygen transport protein but when released from lysed erythrocytes can damage cells and tissues and especially kidneys by oxidative processes. A major mechanism for prevention of hemoglobin-mediated injury involves the plasma protein haptoglobin which is synthesized in the liver (Quaye, 2008; Shaer et al., 2014) (Fig. 1). Haptoglobin has an extremely high affinity (Ka ~ 10^{15}) for hemoglobin and its binding leads to decreases in hemoglobin mediated damage to kidneys (Quaye, 2008). The hemoglobin-haptoglobin complex is removed from blood by binding to the macrophage cell surface protein CD163 followed by endocytosis (Quaye, 2008; Shaer et al., 2014) (Fig. 1).

Hemoglobin varies among animal species in its sequence, its oxygen-binding and its regulation by small molecules (Perutz, 1983). The structural variation in hemoglobins and increased affinity for oxygen have been correlated with such factors as species habitats and lifestyles such as high altitude environments for llama, guinea pigs and other Andean species (Perutz, 1983; Jessen et al., 1991; Storz, 2007; Storz and Moriyama, 2008; Painet and Jaenicke, 2010). Increased affinity for oxygen for some hemoglobins is also correlated with high altitude flight for vultures and black headed gull, bar headed goose and Andean geese (Hiebl et al., 1988; Godovac-Zimmermann et al., 1988; Jessen et al., 1991). In other cases, lowered affinity of hemoglobin for oxygen is correlated with certain amino acid sequence substitutions and cold habitats such as for Antarctic penguins and fish (Notothenioide) and woolly mammoths (Stam et al., 1997; diPrisco et al., 2000; Campbell et al., 2010).
Diving mammals such as seals, sea lions and walruses also express hemoglobins with sequence differences in comparison to other mammals (Watanabe et al., 1986; Lin et al., 1989; Ikehara et al., 1996). However, in an extreme example, certain Antarctic fish genetically do not express hemoglobin which is a condition not described in other vertebrates (Cocca et al., 1995).

In the case of the llama, its hemoglobin differs biochemically from those of many other mammals living at lower altitudes. Llama hemoglobin is able to bind oxygen more efficiently at low oxygen concentrations and responds differently to 2-diphosphoglycerate than some other species such as human (Piccinini et al., 1990; Storz, 2007).

It has been repeatedly found that hemoglobins from phylogenetically separated species or groups generally differ more in amino acid sequence than do hemoglobins from phylogenetically similar species which are more similar in amino acid sequence to each other. Therefore, based on extensive studies, there is good evidence for interspecies differences in hemoglobin amino acid sequences and how these affect the properties of hemoglobin. The interspecies differences in hemoglobin amino acid sequence suggest the question of whether hemoglobin interactions with other proteins vary among animal species.

Haptoglobin consists of two polypeptide chains which are a smaller alpha subunit and a larger beta subunit. These polypeptides are usually linked by disulfide bonds in a basic tetrameric (alpha chain -beta chain)2 arrangement (Fig. 2). In most species, the alpha-beta units are usually but not always linked by disulfide bonds. Haptoglobin also shows interspecies biochemical differences. Human haptoglobin has a genetic polymorphism that is absent from other mammals including primates (Quaye, 2008; Wobeto et al., 2008; Campbell et al., 2010; Shaer et al., 2014). It exists as Hp 1:1, 2:1 and 2:2 variants (Fig. 2). Ruminant haptoglobins such as those from cattle, goat, sheep, deer and water buffalo differ from those of other mammals by existing in a multimeric form. The haptoglobins of these species are different from those of other mammals but similar to the human Hp 2:2 allele in having a partial tandem duplication of sequence in the haptoglobin alpha chain (Busby and Travis, 1978; Lai et al., 2007; Wicher and Fries, 2007; Lai et al., 2008; Wicher and Fries, 2010; Niranjan et al., 2015). Haptoglobins from carnivores have their own structural distinctions from other mammalian haptoglobins. The haptoglobin alpha-beta subunit pairs of dog, bear and cat are linked non-covalently into tetramers. There are no disulfide bonds between the alpha subunits in the tetramer unlike in haptoglobins of other species. The dog alpha chain also has N-linked oligosaccharides unlike those of other mammals (Kumazaki et al., 1992; Mominoki et al., 1995). The amino acid sequence of dog haptoglobin alpha chain is 68% identical to human haptoglobin alpha chain and the respective beta subunits are 85% identical in amino acid sequence (Kumazaki et al., 1992).

The entries in the UniProt sequence database provide much evidence of sequence differences among haptoglobins that often but not always group according to phylogenetic classifications. For example, the Sambar deer (Rusa unicolor swinhoei) haptoglobin is 98.2% identical in sequence to another deer Cervus elaphus, 91.4% identical to cattle and water buffalo haptoglobins but shows lower sequence identity to haptoglobins from human (75.5%), gorilla (72.95), mouse (73.2%) and rat (72.6%). Haptoglobins also differ among species in that these proteins have been found in fish and mammals and ostriches but are absent from chicken and the frog Xenopus. In chickens another hemoglobin binding protein PIT54 is present in blood. Ostrich blood contains both PIT54 and haptoglobin (Wicher and Fries, 2006).
Fig. 2: Human haptoglobin exists as three genetic variants that differ in oligomeric arrangement: Human haptoglobin variants are depicted. Haptoglobin 1:1 (Hp 1:1) consists of two β chains and two smaller α chains and is a tetramer (α1β2). Haptoglobin 2:1 (Hp2:1) expresses the α1 polypeptide and the α2 polypeptide and the β chain. Haptoglobin 2:1 consists of the tetramer found in Hp 1:1 (α1β2) and also higher order linear oligomers (α2β)n. Only one example is diagrammed. Haptoglobin 2:2 consists only of higher order oligomers of (α2β) but these are more compactly or even circularly arranged in contrast to Hp 2:1.

Differences in immunochemical reactivities provide other evidence for similarities and differences in haptoglobin structure among species. A mouse monoclonal antibody to human haptoglobin recognized goat, sheep and bovine haptoglobins and to a lesser extent horse and rabbit haptoglobins. This antibody did not react with haptoglobins from dog, fox, cat and pig (Katnik et al., 1998). Another study found that an antibody preparation reacted with goat, sheep and bovine haptoglobins but not with pig haptoglobin (Busby and Travis, 1978). A commercial immunoturbidometric assay for human haptoglobin reacted with the dog protein and another polyclonal antibody to human haptoglobin reacted with the dog protein in ELISA and immunodiffusion assays (Tecles et al., 2007). The immunochemical results indicate that haptoglobins vary in structural details among different animal species.

Because both hemoglobin and haptoglobin from different species differ structurally, the question arises as to the extent to which hemoglobins and haptoglobins from different species can bind each other. Two opposing hypotheses can be made. (1) Haptoglobin binding to hemoglobin is of fundamental physiological importance and once evolved, the protein structural elements involved have been conserved among vertebrate hemoglobins and haptoglobins although other sites on these molecules have changed during the evolution of different species.

(2) The evolutionary changes in haptoglobin and hemoglobin sequences have affected the abilities of haptoglobins from some species to bind to hemoglobins from other species.

The objectives of this study were to examine: (1) charge differences of different animal hemoglobins by electrophoresis (2) binding of hemoglobins from different species to the plasma protein haptoglobin in human and horse plasmas (3) binding of hemoglobins from different animal species to human haptoglobin genetic variants Hp1:1 and Hp2:2 (4) isolation of haptoglobin from plasmas of different animal species by affinity chromatography with llama hemoglobin-Sepharose conjugate.

Materials and Methods

Materials

Llama, canine, bovine, goat, equine and porcine whole blood samples were obtained from Lampire Biological Laboratories, Pipersville, PA, USA. Llama, feline and anteater blood samples were provided by Dr. Woldemeskel, University of Georgia, Veterinary Diagnostic and Investigational Laboratory, Tifton, GA, USA. Whole human blood samples were obtained from Zen Bio Inc., Research Triangle Park, NC, USA. Ficoll-Paque Plus (Ficoll) was purchased from GE Healthcare Bio-Sciences, Pittsburgh, PA, USA. Cyanogen bromide-activated Sepharose CL-4B and other biochemical reagents were obtained from Sigma Chemicals, St. Louis MO, USA. Agarose, Tris-Glycine pH 8.3 electrophoresis buffer, Kaleidoscope Precision Plus pre-stained molecular weight standards, nitrocellulose and other electrophoresis materials were obtained from Biorad Laboratories, Hercules, CA, USA. The Bicinchoninic Acid (BCA) protein concentration determination kit was
Agarose gel Electrophoresis

Isolation of Erythrocytes and Plasma from Whole Blood

In order to isolate fractions containing plasma and erythrocytes, acid citrate dextrose anti-coagulated whole blood was processed by centrifugation through layers of Ficoll by adapting the manufacturer’s instructions (GE Healthcare manual: Instructions 71-7167-00AG). Volumes of 2 to 4 mL of whole blood were layered onto 3 mL of Ficoll in 15 mL conical centrifuge tubes and centrifuged in a bench centrifuge for 8 min. The layer above the Ficoll constituted plasma, and the erythrocytes were found as a pellet below the Ficoll.

The plasma layer was removed by aspiration with a disposable transfer pipette and centrifuged at 14,000 rpm in a Hettich microcentrifuge to sediment platelets and the supernatant was removed and retained as a preparation of plasma. The erythrocytes were retrieved after removal of the Ficoll, resuspended in 10 mL of Tris Buffered Saline (TBS), sedimented by centrifugation and washed twice with 10 mL volumes of TBS. The final erythrocyte pellets of 12 mL settled volume of cells were resuspended to 30 mL in TBS and stored at -20°C.

Isolation of Hemoglobin

Isolation of hemoglobins was done by osmotic or freeze thaw lysis of erythrocytes previously washed with Tris-buffered saline. Erythrocytes in 30 mL of TBS (21 mL settled volume) were thawed from minus 20°C storage. This treatment lysed the cells and released their contents. After centrifugation for 20 min at 14,000 rpm (> 13,000g) in a Hettich Mikro 120 (in 1.5 mL volumes), the supernatants were retained and combined as a hemoglobin preparation.

Agarose gel Electrophoresis

Agarose gel electrophoresis was done by using 2-2.5 cm thick 1% agarose gels in 25 mM Tris 250 mM Glycine pH 8.3 buffer. A Biorad Midi horizontal gel electrophoresis chamber was used. The thickness of the gels accommodated larger sample volumes to enhance visualization of the red color of hemoglobin and its complexes. Sample buffer consisted of 200 µL of 250 mM Tris, 25 mM glycine pH 8.3 mixed with 800 microliters of water and 1 mL of glycerol.

Sample preparation for electrophoresis was done for samples with and without addition of plasma or purified haptoglobin. To 50 µL of hemoglobin was added 20 µL of sample buffer or 50 µL of plasma or for negative controls - 50 µL of Tris Glycine buffer. To assess interaction of hemoglobins from different animal species with human haptoglobin variants Hp1:1 and Hp2:2, 50 µL of 2 mgmL of haptoglobin were mixed with 50 µL of hemoglobin and 20 µL of sample buffer. Fifty µL to seventy µL of samples of different hemoglobin from different animal species with or without plasma were loaded in the sample wells of gels which were run at 80-100 V at constant voltage until a bromophenol blue marker in one sample well was about one centimeter from the end of the gel.

In the case of experiments to assess the interaction of hemoglobins from different animal species with horse plasma, 10 µL of hemoglobin were mixed with 90 µL of horse plasma and 10 µL of sample buffer and 40 µL of sample was loaded per sample well. In certain cases, the electrophoresis was photographically documented after the samples had moved 2 to 3 cm from the point of application to better visualize bands since they became fainter due to diffusion as electrophoresis progressed.

SDS polyacrylamide gel electrophoresis was performed according to the method of Laemmli (1970) and by using 7.5 or 4-20% gradient gels.

Lateral Flow Immunoassay

The Seratec Hemdirect lateral flow test from Serological Research was used to test immunoreactivity of hemoglobins with a monoclonal antibody to human hemoglobin. The manufacturer’s instructions were followed. In brief, 5 drops of hemoglobin solutions from different animal species were added to the well of the lateral flow immunoassay device and allowed to flow along the strip and thus rehydrate colloidal gold particles coupled to one monoclonal antibody to hemoglobin test. The strip contained a test zone (T) immediately after the sample well that contains a second monoclonal antibody to human hemoglobin. If the hemoglobin also reacted with the second monoclonal antibody to hemoglobin the colloidal gold particles adhered to the test zone on the strip and formed a colored line. This lateral flow immunoassay contains a control zone (C) to the left of the test zone. This control zone contains antibodies to mouse immunoglobulin and thus binds the colloidal gold-anti-hemoglobin particles in the presence and absence of hemoglobin. It serves to indicate that the colloidal gold particles with anti-human hemoglobin are functional. A colored line in the test strip and the control strip indicated the presence of hemoglobin in the sample that was reactive with the anti-human hemoglobin.

Preparation of Llama Hemoglobin-Sepharose CL-4B Affinity Matrix

Llama hemoglobin was covalently linked to cyanogen bromide-activated Sepharose CL-4B (CN Br-
antibodies to human hemoglobin was used to assess hemoglobin in a Lateral Flow Immunoassay. React with a Monoclonal Antibody to Human Llama and other Non-Human Hemoglobins do not from Different Species.

Results

on gels. volume of SDS gel sample buffer and 30 µL were loaded chromatography materials were mixed with an equal Hb-Sepharose CL-4B and unmodified Sepharose (sodium dodecyl sulfate) gel electrophoresis, the llama the plasmas from various species. For analysis by SDS horse plasmas chromatography were resuspended to a final volume of 100 µL of llama-Hb-Sepaharose and control Sepharose contained pelleted llama Hb-Sepharoses from the llama and horse plasmas. The final pelleted llama Hb-Sepharose was washed with 100 mL of 25 mM Tris 250 mM glycine pH 8.3 and finally stored in 15 mL of the same buffer at 4°C. Llama Hemoglobin-Sepharose CL-4B Chromatography

Llama hemoglobin covalently coupled to Sepharose CL-4B (llama Hb-Sepharose) was used in batch affinity chromatography of llama, horse and human plasmas. Unmodified Sepharose CL-4B was used for as control. A volume of 0.8 mL of human plasma was incubated with 100 µL of llama-Hb Sepharose and 4 mL of llama plasma or horse plasma were each incubated with 300 µL of llama-Hb-Sepaharose or control Sepharose at 4°C for 32 h. The llama-Hb-Sepaharose and Sepharoseincubations were washed twice by centrifugation with 1 mL of PBS for the incubations with human plasma and 2 mL of PBS for the incubations with llama and horse plasmas. The final pelleted llama Hb-Sepharose from the human plasma chromatography was resuspended to a final volume of 100 µL with PBS and the pelleted llama Hb-Sepharoses from the llama and horse plasmas chromatography were resuspended to a final volume of 300 µL with PBS. The final washed llama Hb-Sepharose and control Sepharose contained whatever proteins had bound after the incubation with the plasmas from various species. For analysis by SDS (sodium dodecyl sulfate) gel electrophoresis, the llama Hb-Sepharose CL-4B and unmodified Sepharose chromatography materials were mixed with an equal volume of SDS gel sample buffer and 30 µL were loaded on gels.

Results

Differences and Similarities Among Hemoglobins from Different Species

Llama and other Non-Human Hemoglobins do not React with a Monoclonal Antibody to Human Hemoglobin in a Lateral Flow Immunoassay

A lateral flow immunoassay that used monoclonal antibodies to human hemoglobin was used to assess similarities and differences in immunochemical reactivity among hemoglobins from llama, anteater and human (Fig.3). Human hemoglobin bound to the monoclonal anti-hemoglobin antibodies but none of the other animals’ hemoglobins bound.

Llama and Alpaca Hemoglobins Differ from Hemoglobins from Other Animal Species in Agarose Gel Electrophoretic Mobility

Agarose gel electrophoresis was done for hemoglobins from different species in order to compare them for differences that might affect their binding to non-native haptoglobins (Fig. 4). The gel was not stained for protein and the hemoglobins were detected by their visible intrinsic red color. Llama and alpaca hemoglobins displayed the slowest electrophoretic mobilities.

Hemoglobins from Different Animal Species Interact with Human Plasma in a Way Consistent with Haptoglobin Binding

An experiment was done to test if different animal hemoglobins could bind with human haptoglobin by using agarose gel electrophoresis (Fig. 5). A second fainter hemoglobin band was present in all samples in the presence but not in the absence of plasma (Fig. 5). The second band containing llama hemoglobin migrated more slowly than those of the other species second hemoglobins (dashed arrow, Fig. 5). Llama hemoglobin alone (arrowhead, Fig. 5) migrates towards the anode, opposite to the movement of other species hemoglobins in agarose gel electrophoresis.

Different Animal Species Haptoglobins form Complexes with Different Mobilities with two Different Human Plasmas as Detected by Electrophoresis

Hemoglobins from llama, dog, human and pig were mixed with a second sample of human plasma (plasma B) as well as plasma A and analyzed by agarose gel electrophoresis in order to further demonstrate that hemoglobins from different animal species could bind human haptoglobin. The results again showed faster moving hemoglobin bands in the presence of the plasmas but not in the negative controls which lacked plasma (Fig. 6). In the case of plasma B, two faster moving hemoglobin bands were seen in the presence of plasma (arrow and dash, Fig. 6) as opposed to the one faster band seen with hemoglobins mixed with plasma A. One of the hemoglobin bands seen with plasma B migrated similarly to the faster band seen with hemoglobins in the presence of plasma A, but the second plasma B fast hemoglobin bands migrated less. It was evident that one of the presumptive llama hemoglobin-haptoglobin bands migrated more slowly than those of the other species (arrowhead, Fig. 6).
**Fig. 3:** Llama and anteater hemoglobins do not react with a monoclonal antibody to human hemoglobin in a lateral flow immunoassay:

A lateral flow immunochromatographic assay based on monoclonal antibodies to human hemoglobin was used to test hemoglobins from human, anteater, and llama. The presence of two lines in the assay device indicated a positive reaction for human hemoglobin and the presence of only one line indicated a lack of reaction for anteater and llama hemoglobins.

**Fig. 4:** Llama and alpaca hemoglobin differ from hemoglobins of other animals in mobility in agarose gel electrophoresis: Gel A. samples loaded from left are hemoglobins from: (1) Dog (2) Cat (3) Human (4) Anteater (5) Pig (6) Bovine (7) Llama. Gel B. samples from left are hemoglobins from: (7) Llama (8) Alpaca (6) Bovine. The cathode is at the top. Faint bands near the top of the gel in samples 1-6 are the bromophenol blue marker dye. Left arrow indicates positions of non-llama hemoglobins, left arrowhead indicates llama hemoglobin. Right arrow indicates position of bovine hemoglobin and right arrowhead indicates positions of llama and alpaca hemoglobins.

**Fig. 5:** Hemoglobins from different interact with human plasma in a way consistent with haptoglobin binding: Agarose gel electrophoresis was done with hemoglobin samples from different species in the presence (+) and absence (-) of single donor human plasma. From left the samples are hemoglobins from: (1) Llama (2) Human (3) Pig (4) Dog. Sample Br is the marker dye bromophenol blue alone. The solid arrow indicates the position of the second hemoglobin band in the presence of plasma in samples 2-4. The hatched arrow indicates the position of the second hemoglobin band in the presence of plasma for the llama sample. The arrow head indicates the position of the native (first) band of llama hemoglobin versus that of other species hemoglobins.
Different Animal Hemoglobins Interact with Horse Plasma in a Way Consistent with Haptoglobin Binding

Llama, human, pig and dog hemoglobins all formed a second faster migrating band in the presence of horse plasma (Fig. 7). The faster moving hemoglobin in the presence of horse plasma moved at a similar position to the slower of the fast hemoglobin bands seen in the presence of human plasma B (dash, Fig. 7). The second faster moving hemoglobin band seen in the mixtures of different species hemoglobins with human plasma B had no counterpart in the hemoglobins mixed with horse plasma. It was also observed that the slower moving hemoglobin band in the llama sample (arrowhead, Fig. 7) migrated more slowly than the corresponding band formed with human, pig and dog hemoglobins. In the absence of plasma, no faster moving hemoglobin bands were seen.

Different Animal Species Hemoglobins Bind to two Alleles of Human Haptoglobin (Non-Stained Gel: Early Stage of Electrophoresis)

Hemoglobins from llama, pig, dog, goat, horse and human were mixed with purified genetic variants of human haptoglobin, Hp1:1 or Hp2:2 and the mixtures were analyzed by agarose gel electrophoresis. The results for an early stage of electrophoresis are shown for the unstained gel in which the visualization of protein bands is by means of the intrinsic red color of hemoglobin (Fig. 8). The hemoglobins from llama, pig, dog, goat, horse and human formed a second faster migrating hemoglobin band with both haptoglobins 1:1 and 2:2. The same electrophoresis gel was allowed to run for a longer period of time so that proteins migrated further into the gel and was stained for protein with Coomassie Blue (Fig. 9). Two protein bands were seen in the hemoglobin samples that were mixed with haptoglobins. However the second fast moving band (arrow A) in samples mixed with haptoglobin 1:1 migrated further than those mixed with haptoglobin 2:2 (arrow A1). It was also the case that llama hemoglobin formed more slowly migrating complexes with both haptoglobin variants compared to the hemoglobins from other species. The position of free haptoglobin alone in the last gel lane is indicated by the thick arrowhead in Fig. 9. It was also evident that where hemoglobins were run in electrophoresis in the absence of haptoglobin, only one slow moving (human hemoglobin) or backwards moving (llama) hemoglobin was present.

Llama Hemoglobin Binds Horse and Human Haptoglobin in Affinity Chromatography

Affinity chromatography by using llama hemoglobin that was covalently linked to Sepharose CL-4B was done to test for the binding of haptoglobin in plasma of llama, human and horse. As a negative control, Sepharose 4B was used in chromatography of samples of the same plasmas. The samples that bound to llama hemoglobin-Sepharose and to plain Sepharose were analyzed by SDS gel electrophoresis under reducing conditions (Fig. 10).
Fig. 7: Hemoglobins from different species interact with horse plasma in a way consistent with haptoglobin binding: Hemoglobin samples from (1) Llama (2) human (3) Pig (4) Dog were subjected to agarose gel electrophoresis in the presence of horse plasma (first 1-4) or the presence of human plasma donor B (second 1-4) or in the absence of plasma (last 1-4). The left dash marks the position of the second hemoglobin band seen in the presence of horse plasma. The left arrow marks the position of the furthest moving of the two bands seen in hemoglobins incubated with human plasma B. The left arrowhead marks the position of the second (slower moving) hemoglobin band seen in the presence of human plasma B. The fast moving hemoglobin band marked with a dash in samples mixed with horse plasma migrates in between the two faster moving hemoglobin bands seen in the presence of human plasma B. The right solid arrowhead marks the position of free llama hemoglobin and the right white arrow marks the positions of other animal hemoglobins.

Fig. 8: Hemoglobins from different species bind two alleles of human haptoglobin: Non stained agarose gel in the early stage of electrophoresis of hemoglobins from (1) Llama (2) Human (3) Pig (4) dog (5) Goat (6) Horse in the presence of human haptoglobin 1:1 (first set of samples 1-6) or human haptoglobin 2:2 (second set of samples 1-6). Samples 1’ and 2’ are llama and human hemoglobin in the absence of haptoglobin and Br is the bromophenol blue marker alone. Left arrow indicates faster moving hemoglobin band. Right arrowhead and dashed arrow indicates positions of free llama hemoglobin and free hemoglobins from other species respectively.

A protein of approximately 40 kDa was isolated from both human and horse plasma but not llama plasma by llama-hemoglobin Sepharose. This 40kDa protein was not present in the fraction bound by the negative control matrix Sepharose 4B alone but a 50kDa protein was present in the fraction retained from human plasma (Fig. 10).
Fig. 8: Hemoglobins from different species bind two alleles of human haptoglobin: Non stained agarose gel in the early stage of electrophoresis of hemoglobins from (1) Llama (2) Human (3) Pig (4) dog (5) Goat (6) Horse in the presence of human haptoglobin 1:1 (first set of samples 1-6) or human haptoglobin 2:2 (second set of samples 1-6). Samples 1’ and 2’ are llama and human hemoglobin in the absence of haptoglobin and Br is the bromophenol blue marker alone. Left arrow indicates faster moving hemoglobin band. Right arrowhead and dashed arrow indicates positions of free llama hemoglobin and free hemoglobins from other species respectively.

Fig. 9: Hemoglobins from different species bind two alleles of human haptoglobin Comassie blue-stained gel. The late stage of electrophoresis of hemoglobins from (1) Llama (2) Human (3) Pig (4) dog (5) Goat (6) Horse in the presence of human haptoglobin 1:1 (first set of samples 1-6) or human haptoglobin 2:2 (second set of samples 1-6). Samples 1’ and 2’ are llama and human hemoglobin in the absence of haptoglobin. H is haptoglobin 1:1 alone.

Fig. 10: Llama hemoglobin binds horse and human haptoglobin in affinity chromatography: An SDS gel shows samples retained by llama-hemoglobin Sepharose (set A) and Sepharose 4B only (set B) from plasmas from (1) Human (2) llama (3) Horse. The arrow indicates a protein band present only in the fractions bound to llama-hemoglobin-Sepharose.
Discussion

This study supports the hypothesis that the ability of hemoglobins from different species to bind to haptoglobins of other species is conserved. The findings contradict the idea that evolutionary changes in hemoglobins and haptoglobins prevent cross species in vitro interactions of these proteins. This study implies the occurrence of conserved protein-protein interactions sites on hemoglobin and haptoglobin.

The conserved binding hypothesis was examined by (1) showing that previously unreported differences exist among hemoglobins (2) testing the abilities of hemoglobins to bind haptoglobins of different species through electrophoresis of mixtures and affinity chromatography.

It was reported for the first time that llama and alpaca hemoglobins differed in electrophoretic mobility from hemoglobins from human, pig, dog, bovine, goat, horse, cat and anteater. The llama and alpaca hemoglobins had cathodal mobility and thus more positive charge than the other hemoglobins. The other animal hemoglobins also differed among each other in precise electrophoretic mobility indicating further subtle charge differences. This finding was also in accordance with the report that alpaca hemoglobin has only a single amino acid difference from that of llama (Piccinini et al., 1990).

A lateral flow immunoassay for human hemoglobin recognized hemoglobin from human but not hemoglobins from the other species tested including the previously unreported cases of llama and anteater. However, this assay recognizes certain non-human hemoglobins such as those of higher primates and ferrets. These results add to the documented differences and similarities in the hemoglobins from different animals (Perutz, 1983; Storz 2007; Ikehara et al., 1996; Lin et al., 1989). The immunochemical results indicate that a particular antigenic determinant on human hemoglobin was absent from hemoglobins from llama and the other animal species tested.

This study provided evidence that llama, goat, horse, dog, pig and bovine hemoglobins bind to human and horse haptoglobins because faster migrating hemoglobin bands were observed in the presence of different human plasmas, horse plasma and purified human haptoglobin variants. The formation of a second faster migrating band of hemoglobin after mixing of various animal hemoglobins with human and horse plasma is consistent with these hemoglobins binding to haptoglobin in these plasmas. Haptoglobin is considerably more acidic than hemoglobin so a complex of these proteins would migrate faster than hemoglobin but slower than haptoglobin.

The second hemoglobin band (haptoglobin-hemoglobin complex), formed by hemoglobin from different species with horse plasma, migrated more slowly than the complex of hemoglobins with human haptoglobin (Hp 1:1) (Fig. 6 dash). This differentiates the horse from human haptoglobin.

It was also found that hemoglobins from different animal species bind to two purified human haptoglobin genetic variants. Non-human species do not express the polymorphic haptoglobin variants Hp 1:1 and Hp 2:2 that exist in humans (Liau et al., 2003; Wobeto et al., 2008) Binding of animal hemoglobins to these genetic variants of human haptoglobin such as by hemoglobins from goat, horse, pig, dog, bovine and llama further supports the hypothesis of conservation of binding sites on different species hemoglobins and haptoglobins. Purified haptoglobin variants Hp 1:1 and Hp 2:2 incubated with hemoglobins from different species including human, formed only one faster mobility hemoglobin band for each haptoglobin variant (Fig. 9). The hemoglobin complexes with Hp 1:1 had faster mobility than those with Hp2:2. In contrast, in the case of hemoglobins incubated with plasma B, two new hemoglobin bands were observed. The different pattern is consistent with plasma B containing a third variant of haptoglobin – Hp2:1.

Conclusion

It is therefore concluded that hemoglobin from the different animal species studied can bind to all three genetic variants of human haptoglobin.

Affinity chromatography of plasmas from different species on llama-hemoglobin-Sepharose (Fig. 10) also isolated a protein similar in molecular mass to haptoglobin from human and horse plasmas but not from llama plasma. Not all animal plasmas contain significant amounts of haptoglobin in the healthy state as for bovine and goat plasmas and this is thus indicated true for llama plasma. Human hemoglobin-Sepharose affinity chromatography has been previously used to isolate human haptoglobin (Liau et al., 2003) but llama hemoglobin has not previously been used in affinity chromatography.

In summary it is proposed that hemoglobin-haptoglobin binding sites were optimized early in evolution and afterwards retained for different species.

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Author’s Contributions

Oluseyi A. Vanderpuye: Designed and performed experiments, wrote the paper, analysed results.

Moges Woldemeskel: Conceived basis for research wrote sections of paper, checked and revised paper, evaluated experimental results.

Ethics

This study was done in compliance with applicable university regulations and rules for the use of human and animal commercial blood products, human and animal protection. All vendors comply with Federal and State laws.

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