Biochemical pedomorphosis and genetic assimilation in the hypoxia adaptation of Tibetan antelope

Anthony V. Signore and Jay F. Storz*

Developmental shifts in stage-specific gene expression can provide a ready mechanism of phenotypic change by altering the rate or timing of ontogenetic events. We found that the high-altitude Tibetan antelope (*Panthelops hodgsonii*) has evolved an adaptive increase in blood-O\(_2\) affinity by truncating the ancestral ontogeny of globin gene expression such that a high-affinity juvenile hemoglobin isoform (isoHb) completely supplants the lower-affinity isoHb that is expressed in the adult red blood cells of other bovids. This juvenilization of blood properties represents a canalization of an acclimatization response to hypoxia that has been well documented in adult goats and sheep. We also found the genomic mechanism underlying this regulatory isoHb switch, revealing how a reversible acclimatization response became genetically assimilated as an irreversible adaptation to chronic hypoxia.

INTRODUCTION

When members of multigene families are developmentally regulated, shifts in the stage-specific expression of individual genes can provide a ready mechanism of phenotypic change by altering the rate or timing of ontogenetic events (heterochrony). For example, retained activity of early-expressed genes in later stages of ontogeny can result in the retention of juvenile characters into adulthood, a well-documented developmental mechanism of phenotypic evolution (1, 2). In extreme cases, deceleration of development can produce a truncation of the ancestral ontogeny, resulting in the juvenilization of the adult-expressed phenotype, a phenomenon known as pedomorphosis.

In jawed vertebrates, the subfamilies of genes that encode the \(\alpha\)- and \(\beta\)-type subunits of tetrameric hemoglobin (Hb) are developmentally regulated such that structurally and functionally distinct \(\alpha_2\beta_2\) Hb isoforms (isoHbs) are expressed during different ontogenetic stages. During mammalian development, different pre- and postnatally expressed isoHbs have evolved different oxygenation properties and perform distinct O\(_2\)-scavenging/O\(_2\)-transport tasks during different ontogenetic stages (3–6). Genetically based shifts in stage-specific isoHb expression could therefore provide a heterochronic mechanism of evolutionary change in respiratory gas transport and aerobic metabolism. Similarly, in humans, hereditary persistence of fetal Hb alleviates the severity of thalassemias and other pathologies affecting the synthesis or stability of adult Hb (7).

Prenatally expressed isoHbs of eutherian mammals typically exhibit substantially higher O\(_2\) affinities than adult-expressed isoHbs (3, 5, 6). In anthropoid primates and bovid artiodactyls, expression of a high-affinity fetal isoHb during late stages of prenatal development helps to maintain an O\(_2\) affinity difference between fetal and maternal circulations, thereby facilitating O\(_2\) transfer across the placental barrier (5, 6). Given that increased Hb-O\(_2\) affinity is generally beneficial under conditions of severe hypoxia due to the importance of safeguarding arterial O\(_2\) saturation (6, 8–11), the retention of early isoHb expression into adulthood could provide an effective mechanism of adaptation to chronic O\(_2\) deprivation. Consistent with this hypothesis, when adult goats and sheep are exposed to acute hypoxia, they up-regulate a juvenile isoHb at the expense of the normal adult isoHb (12, 13). Here, we report the discovery of a canalized version of this response in the high-altitude Tibetan antelope, *Panthelops hodgsonii* (Artiodactyla: Bovidae), a champion among mammals in aerobic exercise performance under hypoxia. This species is endemic to the Tibetan Plateau and lives at altitudes of 3600 to 5500 m above sea level. At an altitude of 5500 m, the partial pressure of O\(_2\) (P\(_{O2}\)) is roughly half the value at sea level, a level of hypoxia that severely compromises aerobic exercise performance in humans and most other mammals (14–16). However, at these altitudes, Tibetan antelope can sustain running speeds of >70 km/hour over distances of >100 km (17).

In addition to documenting the phenotypic consequences of developmentally displacing the low-affinity adult isoHb with a higher-affinity juvenile isoHb—a form of biochemical pedomorphosis—we also found the genomic mechanism by which the up-regulation of the juvenile isoHb became canalized in Tibetan antelope. Specifically, we document how a reversible acclimatization response to acute hypoxia—as observed in modern-day sheep and goats—became genetically assimilated as an irreversible adaptation to chronic hypoxia.

RESULTS AND DISCUSSION

We characterized the genomic organization of globin genes in Tibetan antelope and other bovid artiodactyls using published genome assemblies (18). Among mammals, bovid artiodactyls are unusual in that the entire \(\beta\)-globin gene cluster has undergone multiple rounds of en bloc duplication involving the same set of pre- and postnatally expressed \(\beta\)-type globin genes (Fig. 1) (19–21). Cows (*Bos taurus*) have two duplicated gene blocks, each containing separate paralogs of the \(\beta\)-globin gene, \(\beta^A\) and \(\beta^B\), in the 5′ and 3′ blocks, respectively (Fig. 1). As with other eutherian mammals, the product of \(\beta^A\) is incorporated into an adult-expressed isoHb, Hb\(A\) (\(\alpha_2\beta^A_2\)), whereas the \(\beta^B\) gene has been recruited for prenatal expression and is incorporated into a fetal isoHb, Hb\(F\) (\(\alpha_2\beta^B_2\)) (22). Goats (*Capra hircus*) and sheep (*Ovis aries*) have an additional gene block at the 5′ end of the cluster that contains a third \(\beta\)-globin paralog, \(\beta^\prime\) (Fig. 1) (19–21). Whereas the \(\beta^A\) and \(\beta^B\) genes in goats and sheep have retained the same developmental expression profiles as their respective orthologs in cow, the \(\beta^C\) gene has been recruited for a new ontogenetic stage of expression during the first few months of neonatal life, and its product is incorporated into a juvenile isoHb, Hb\(C\) (\(\alpha_2\beta^C_2\)) (23).
The β-globin gene cluster of Tibetan antelope appears superficially similar to that of cow in terms of gene content (Fig. 1), suggesting that the Tibetan antelope inherited the same pair of βA- and βF-containing gene blocks. The alternative hypothesis is that Tibetan antelope inherited the additional en bloc duplication observed in goats and sheep but one of the triplicated gene blocks was secondarily deleted, in which case the sole remaining pair of β-globin genes would be represented by one of three possible combinations: βA + βF (a reversion to the ancestral gene complement observed in cow), βC + βF, or βC + βF (Fig. 2, A to C). Either of the latter two combinations would implicate a novel isoHb profile that is not observed in other bovid taxa. To distinguish among these three alternative scenarios, we estimated the phylogeny of bovid βC, βA, and βF genes and the pair of Tibetan antelope β-globin paralogs. Estimated phylogenies (Fig. 2D and fig. S1) demonstrate that the 5′ and 3′ β-globin genes of Tibetan antelope are orthologous to bovid βC and βF, respectively, consistent with the scenario illustrated in Fig. 2B. This result indicates that Tibetan antelope inherited the triplicated set of βC-, βA-, and βF-containing gene blocks observed in goats and sheep (Fig. 1) and that the middle gene block containing βA was secondarily deleted. This phylogenetic inference is unambiguously corroborated by patterns of conserved synteny and pairwise sequence matches (Fig. 3), as the βC- and βF-containing gene blocks of goat and sheep match the 5′ and 3′ gene blocks in Tibetan antelope. This comparative genomic analysis revealed that a ~45-kb region of the Tibetan antelope β-globin gene cluster was deleted—a gene region that contained the ortholog of the βA gene that encodes the β chain of adult Hb in bovids and all other mammals.

Deletion of the adult βA gene in the ancestor of Tibetan antelope effectively truncated the ancestral ontogeny of globin gene expression, such that juvenile HbC completely supplanted HbA in adult red blood cells. Thus, blood-O2 transport in Tibetan antelope has been juvenilized relative to the ancestral phenotype of adult bovids. To examine the effects of this pedomorphic change, we measured the oxygenation properties of purified recombinant Hb from Tibetan antelope and purified native Hbs from adult specimens of 10 other bovid species (Fig. 4 and Table 1). The adult red cells of these other taxa contain HbA alone or in combination with HbC as a minor component (fig. S2). We measured the Hb-O2 affinity of purified total Hb from each bovid species in both the absence (stripped) and presence of 100 mM Cl− (in the form of KCl). The stripped treatment provides a measure of intrinsic Hb-O2 affinity, whereas the +KCl treatment provides a measure that is relevant to in vivo conditions in bovid red cells, as Cl− ions are the principal allosteric regulators of Hb-O2 affinity (i.e., heme reactivity is modulated oxygenation-linked binding of Cl− ions at sites remote from the heme iron) (6, 22, 24). Results of our in vitro experiments revealed that Hb of Tibetan antelope has a substantially higher O2 affinity than that of all other bovid taxa (Fig. 4 and Table 1). Hbs of all taxa were similarly responsive to Cl−, as the average P50 (the PO2 yielding 50% Hb-O2 saturation) was 27.1% higher (i.e., Hb-O2 affinity was lower) in the +KCl treatment (Table 1).

![Fig. 1. Genomic organization of bovid β-globin gene clusters. Colored boxes represent individual genes. Labels denote previously annotated βC-, βA-, and βF-globin genes.](http://advances.sciencemag.org/)

![Fig. 2. Phylogenetic analyses of bovid βC-, βA-, and βF-globin genes. Alternative histories of gene deletion in Tibetan antelope yield testable phylogenetic hypotheses: (A) Deletion of βC, (B) deletion of βA, and (C) deletion of βF. (D) Estimated maximum likelihood phylogeny of bovid β-type globin genes indicates that Tibetan antelope has retained copies of βC and βF and that βA has been secondarily lost. Bootstrap support values are shown for relevant nodes.](http://advances.sciencemag.org/)
As a follow-up experiment, we isolated and purified HbC and HbA from two of the bovid species expressing both components, and we measured isoHb-specific O₂-binding properties to determine how blood-O₂ affinity would be affected by elimination of the major HbA isoHb (as would occur with the deletion of the β⁺-globin, thereby leaving HbC as the sole-expressed isoHb in adult red cells). There was very little among-species variation in the measured O₂ affinities of either juvenile HbC or adult HbA (Table 1 and fig. S2), but the O₂ affinity of HbC exceeded that of HbA by a consistent margin (average, 10.6 torr) in all species (Table 1 and fig. S2). Moreover, O₂ affinity of HbC alone was always substantially higher than that of the composite HbA + HbC mixture (with the two isoHbs present in their naturally occurring relative abundance) (Table 1 and fig. S2), reflecting the fact that the lower affinity HbA is always present as the major isoHb in adult red cells (average HbA/HbC ratio = ~80:20).

The higher Hb-O₂ affinity of Tibetan antelope relative to that of other bovid species is entirely attributable to a difference in isoHb composition: They only express the high-affinity HbC instead of jointly expressing HbA and HbC (with the lower-affinity HbA present as the major isoHb). To infer the direction of evolutionary change in isoHb-specific O₂ affinities and to reconstruct the phenotypic effect of deleting β⁺-globin (thereby leaving HbC as the sole-expressed isoHb in adult red cells), we reconstructed the ancestral bovid β⁺ and β⁻ genes as well as their single-copy, preduplication progenitor (β⁺⁻) (Fig. 5 and fig. S3). Triangulated comparison of O₂ affinities of the three recombinantly expressed ancestral isoHbs, AncHb-β⁺, AncHb-β⁻, and AncHb-β⁺⁻ (all of which had identical α-chains), revealed that the juvenile AncHb-β⁻ evolved a slight increase in O₂ affinity relative to the estimated ancestral state (represented by AncHb-β⁺⁻), whereas adult HbA evolved a slight reduction in O₂ affinity (Fig. 5). These data indicate that if HbA and HbC were present in a 80:20 ratio in the red cells of the Tibetan antelope ancestor (as in extant bovids), then the deletion of β⁺-globin and the consequent elimination of HbA from the “HbA + HbC” composite mixture would result in a 13.5% increase in Hb-O₂ affinity in the presence of 100 mM Cl⁻ (P₅₀ decreased from 18.5 to 16.0 torr). Theoretical and experimental results indicate that an increase in Hb-O₂ affinity of this magnitude—if accompanied by a corresponding enhancement of tissue O₂ diffusion capacity—would likely translate into a physiologically important enhancement of aerobic exercise performance under hypoxia (6, 8–11).
of aerobic metabolism (25–33). In other case studies of high-altitude vertebrates, evolved increases in Hb-O₂ affinity have been traced to adaptive under conditions of severe hypoxia (especially in highly vertebrates, evolved increases in Hb-O₂ affinity have been traced to 6.2020; Signore and Storz, Sci. Adv. : eabb5447 17 June 2020). Here, we document a unique case in which an evolved change in Hb-O₂ affinity has been accomplished via a heterochronic shift in globin gene expression, such that a high-affinity, juvenile isoHb supplants the lower-affinity, adult isoHb. This juvenilization of blood properties represents a novel mode of biochemical adaptation and highlights the utility of heterochrony as an adaptive mechanism, whereby “…the existing channel of ordinary ontogeny already holds the raw material in a particularly effective state for evolutionary change” (2).

There has been debate in the literature regarding the relative importance of regulatory versus coding changes in genetic adaptation (34) and phenotypic evolution in general (35–36). In the case of Tibetan antelope, the evolved increase in Hb-O₂ affinity was caused by an unusual combination of regulatory and structural changes. Specifically, a marked regulatory switch in protein isoform expression (via truncation of the ancestral ontology of globin gene regulation) was caused by a large-scale chromosomal deletion, highlighting the unexpected diversity of genetic mechanisms and substrates of phenotypic evolution.

### MATERIALS AND METHODS

#### Collection of blood samples

Frozen erythrocytes from 10 bovid species were provided by the San Diego Zoo Institute for Conservation Research (Uniform Biological Material Transfer Agreement BR2017063). This sample included six species in the subfamily Caprinae (Capra aegagrus cretica, Capra caucasica caucasica, Ovis nivicola, Capra nubiana, Ovis orientalis musimon, and Ovis canadensis nelsoni), two species in the subfamily Alcelaphinae (Damaliscus pygargus phillipsi and Connochaetes gnou), and one species of each from Hippotraginae (Addax nasomaculatus) and Bovinae (Bos gaurus).

#### Sequencing of bovid globin genes

RNA was extracted from ~100 μl of flash-frozen erythrocytes using an RNeasy Universal Plus Mini kit (QIAGEN). Complementary DNA (cDNA) was synthesized from freshly prepared RNA using SuperScript IV reverse transcriptase (Invitrogen). Gene specific primers were used to amplify the α- and β-type globin transcripts. Polymerase chain reaction (PCR) reactions were conducted using

| Table 1. Oxygenation properties of purified bovid Hbs. P50 values are reported as means ± SEM. Cl⁻ effect calculated as logP50(+KCl) − logP50(stripped). |
|-----------------|-----------------|-----------------|-----------------|
| **Hb solution** | **Species**      | **P50, stripped (torr)** | **P50, +KCl (torr)** | **Cl⁻ effect** |
| --- | --- | --- | --- | --- |
| Total Hb | European mouflon, O. orientalis musimon | 12.08 ± 0.09 | 21.12 ± 0.20 | 0.24 |
| | Snow sheep, O. nivicola | 15.72 ± 0.12 | 29.91 ± 2.28 | 0.28 |
| | Desert bighorn sheep, O. canadensis nelsoni | 12.00 ± 0.42 | 22.98 ± 1.50 | 0.28 |
| | Cretin wild goat, C. aegagrus cretica | 14.43 ± 0.33 | 28.14 ± 0.10 | 0.29 |
| | Nubian ibex, C. nubiana | 11.20 ± 1.01 | 27.59 ± 0.57 | 0.39 |
| | West caucasian tur, C. caucasica | 0.996 ± 0.15 | 23.91 ± 0.21 | 0.33 |
| | White-tailed gnu, C. gnou | 14.27 ± 0.64 | 24.97 ± 0.78 | 0.24 |
| | Blesbok, D. pygargus phillipsi | 13.96 ± 1.09 | 20.97 ± 0.71 | 0.30 |
| | Addax, A. nasomaculatus | 13.07 ± 0.22 | 25.89 ± 0.62 | 0.30 |
| | Indian gaur, B. gaurus | 12.32 ± 0.14 | 27.50 ± 0.24 | 0.35 |
| Pure HbA | Nubian ibex, C. nubiana | 0.996 ± 0.19 | 23.79 ± 1.12 | 0.33 |
| | West caucasian tur, C. caucasica | 0.996 ± 0.42 | 23.79 ± 1.12 | 0.33 |
| | Goat Hb, C. hircus | 13.51 ± 0.21 | 26.33 ± 0.53 | 0.29 |
| | Nubian ibex, C. nubiana | 9.21 ± 0.22 | 19.41 ± 0.71 | 0.32 |
| Pure HbC | West caucasian tur, C. caucasica | 8.62 ± 0.12 | 18.03 ± 0.57 | 0.32 |
| | Goat Hb, C. hircus | 8.82 ± 0.33 | 18.29 ± 0.21 | 0.32 |
| | Tibetan antelope rHb, P. hodgsonii | 8.53 ± 1.10 | 18.59 ± 1.07 | 0.34 |
mean and structurally distinct chains). βα the subunit composition of the three ancestral isoHbs (which have identical
94°C for 30 s, 53° to 65°C for 30 s, 72°C for 45 s for 30 cycles, fol-
Gradient thermocycler. Following a 5-min denaturation period at
of 10× Reaction Buffer (Invitrogen), 0.75 μl of 50 mM MgCl₂, 1.25 μl of each primer (10 pmol/μl), 1 μl of Taq polymerase (Invitrogen), and
were subsequently purified with a GeneJET Plasmid Miniprep kit
were run on a 1.5% agarose gel, and bands of the correct size were
purified PCR products were ligated subsequently excised and purified using Zymoclean Gel DNA recovery
were centrifuged at 20,000 g for 10 min to remove cell debris. Buffer
were used to estimate the O₂ at half saturation (P₅₀) and the coop-
erativity coefficient (n₅₀) from the χ-intercept and slope of these plots, respectively. O₂ equilibrium curves for each Hb solution were mea-
sured in triplicate, and P₅₀ is reported as means ± SEM.

Protein purification
Blood samples (~200 μl) were added to a 5× volume of ice-cold water and incubated on ice for 30 min to lyse the red blood cells. Samples were centrifuged at 20,000 g for 10 min to remove cell debris. Buffer was added to the supernatants to a final concentration of 0.01 M Hepes/0.2 M NaCl (pH 7.4) and passed through a PD-10 desalting column (GE Healthcare) equilibrated with 25 ml of 0.01 M Hepes/0.5 mM EDTA (pH 7.2) to remove intracellular cofactors. Desalted lysates were loaded onto a HiTrap SP cation exchange column (GE Healthcare), and isoHbs were eluted using a linear pH gradient [0.01 M Hepes/0.5 mM EDTA (pH 7.2 to 7.7)]. For each species, a subsample of each isoHb was pooled to create a “Total Hb” solution. Each Hb solution was then desalted using a PD-10 column (GE Healthcare) equilibrated with 0.01 M Hepes/0.5 mM EDTA (pH 7.4), and eluates were concentrated using the Amicon Ultra-4 Centrifugal Filter Units (Millipore).

Measuring O₂-binding properties of purified Hbs
O₂ equilibrium curves for purified Hb solutions [0.1 mM Hb in 0.1 M Hepes/0.05 mM EDTA buffer (pH 7.4)] were measured at 37°C using the Blood Oxygen Binding System (Loligo Systems). O₂ equilib-
rium curves were measured in the absence (stripped) and presence of chloride ions (0.1 M KCl). Each Hb solution was sequentially equilibrated with three to five different O₂ tensions (P0₂) at saturation levels between 30 to 70%, while the absorbance was continually monitored at 430 nm (deoxygen peak) and 421 nm (oxy/deoxy isosbestic point) (43–45). Hill plots (log[fractional saturation/[1 – fractional saturation]]) versus logPO₂ constructed from these measurements were used to estimate the PO₂ at half saturation (P₅₀) and the cooperativity coefficient (n₅₀) from the x-intercept and slope of these plots, respectively. O₂ equilibrium curves for each Hb solution were mea-
sured in triplicate, and P₅₀ is reported as means ± SEM.

Construction of Hb expression vector
Globin sequences for domestic goat, Tibetan antelope, and the re-
constructed ancestral globins were synthesized by GeneArt Gene Synthesis (Thermo Fisher Scientific) after optimizing the nucleotide sequences in accordance with E. coli codon preferences. The synthesized globin gene cassette was cloned into a custom pGM vector system along with the methionine aminopeptidase gene, as described previously (46).

Expression and purification of recombinant Hbs
Recombinant Hb expression was carried out in the E. coli JM109 (DE3) strain as described previously (46–48). Bacterial cell lysates

Sequence analyses
Genomic sequences containing the complete α- and β-globin gene clusters for the domestic goat (C. hircus), sheep (O. aries), cow (B. taurus), and Tibetan antelope (P. hodgsonii) were obtained from GenBank (table S1). Sequence identity between bovid chromosomal regions containing the β-globin gene clusters was calculated using Blastn, and patterns of sequence matching were visualized using Easyfig 2.1 (37). Coding sequences of α- and β-globin genes were

extracted from genomic and cDNA sequences available on GenBank (table S1) and were combined with the newly generated cDNA sequences mentioned above (fig. S3). Sequences were aligned using MUSCLE (38) and were then used to estimate phylogenetic trees. The best fitting codon substitution model and initial tree search were estimated using IQ-TREE with the options -st CODON, -m TESTNEW, -allnni, and -bnni (39, 40). Initial trees were then subject-
to 1000 ultrafast bootstrap replicates (41). Bootstrap consensus trees (fig. S1, A and B) were used to estimate ancestral globin se-
quencies using IQ-TREE with the option -asr (figs. S1C and S3). As boid β'-globins are truncated by 9 base pairs (relative to β*), the ancestral reconstruction of indels in the β-globin gene tree was performed by FastML (42).

1 ml of cDNA template in 0.2-ml tubes containing 25 μl of reaction mixture [0.5 μl of each deoxynucleotide triphosphate (2.5 mM), 2.5 μl of 10× Reaction Buffer (Invitrogen), 0.75 μl of 50 mM MgCl₂, 1.25 μl of each primer (10 pmol/μl), 1 μl of Taq polymerase (Invitrogen), and
and presence of 0.1 M KCl at 37°C (pH 7.4) (0.1 mM Hb4). Schematic diagrams show the subunit composition of the three ancestral isoHbs (which have identical α chains and structurally distinct β chains).

Fig. 5. O₂ affinities of reconstructed ancestral bovid isoHbs. (A) Reconstructed ancestral β-globin genes (βC, βA, and βB) of boids. (B) O₂ tensions at half saturation (mean P₅₀ ± SEM, n = 3) for recombinant ancestral isoHbs in the absence (stripped) and presence of 0.1 M KCl at 37°C (pH 7.4) (0.1 mM Hb4). Schematic diagrams show the subunit composition of the three ancestral isoHbs (which have identical α chains and structurally distinct β chains).
were then loaded onto a HiTrap Q HP anion exchange column (GE Healthcare), then equilibrated with 20 mM tris/0.5 mM EDTA (pH 8.3), and eluted with a linear gradient of 0 to 0.25 M NaCl. Hb-containing fractions were then loaded on to a HiTrap SP HP cation exchange column (GE Healthcare) and eluted with a linear pH gradient (pH 6.8 to 8.4). Eluted Hb fractions were concentrated using the Amicon Ultra-4 Centrifugal Filter Units (Millipore), and oxygenation properties were measured as described above.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/6/25/eabb5447/DC1

View/request a protocol for this paper from Bio-protocol.

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