Phylogenetic and structural analysis of the HbA ($\alpha^A/\beta^A$) and HbD ($\alpha^D/\beta^A$) hemoglobin genes in two high-altitude waterfowl from the Himalayas and the Andes: Bar-headed goose (Anser indicus) and Andean goose (Chloephaga melanoptera)

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A B S T R A C T

Two species of waterfowl living at high altitude provide a prominent example of parallel adaptation at the molecular level. The bar-headed goose (Anser indicus) breeds at high elevations in central Asia and migrates across the Himalayas, where the partial pressure of oxygen ($O_2$) is one-third of sea level. In South America, the distantly related Andean goose (Chloephaga melanoptera) is endemic to the high Andes. Both species exhibit increased blood--O$_2$ affinity, which has been attributed to the effects of single amino acid substitutions in the major hemoglobin. Here we present phylogenetic analyses of the swans and geese (Anserinae) and South American sheldgeese (Anatinae) using the three genes that encode the major (HbA) and minor (HbD) hemoglobin isoforms. We sought to determine whether two amino acid substitutions that have been the focus of extensive biochemical analysis (Ala-$\alpha^{A119}$ and Ser-$\beta^{A55}$) are uniquely derived in bar-headed goose and Andean goose, respectively, and to examine evidence of molecular adaptation at other positions in hemoglobin genes by comparing these two high-altitude taxa to their closest relatives. Bayesian analysis of the $\alpha$- and $\beta$-subunit genes produced well-resolved phylogenies, with high posterior probabilities and bootstrap values for most genera. The bar-headed goose is likely sister to all other Anser species. Andean goose, the sole highland representative of the South American sheldgeese, is either sister to the other Chloephaga species or sister to Neochen. In the bar-headed goose, four derived substitutions were observed in HbA ($\alpha^{A12}$, 18, 63, 119) and two in HbD ($\alpha^{D2}$, 47). Four derived substitutions in Andean goose include three in HbA ($\alpha^{A8}$, 77; $\beta^{A86}$) and two in HbD ($\alpha^{D9}$; $\beta^{A86}$). Considering both highland species, four substitutions (Ala-$\alpha^{A2}$, Ala-$\alpha^{A12}$, Ser-$\alpha^{A18}$, Leu-$\alpha^{A55}$) were located at adjacent positions on the $A$ helix (or $AB$ corner) of the $\alpha$-chains, three others (Thr-$\alpha^{A77}$, Ser-$\beta^{A86}$, Ser-$\alpha^{D2}$) were in close proximity to inositolpentaphosphate (IP$_5$) binding sites, and Ala-$\alpha^{A119}$ occurred at an $\alpha$-inter-subunit contact. Ser-$\beta^{A55}$, which is involved in the same $\alpha$-$\gamma$ inter-subunit contact and was previously shown to increase Hb--$O_2$ affinity, is not unique to Andean goose but is a synapomorphy of the South American sheldgeese, a clade of predominantly lowland waterfowl. Our findings illustrate the importance of understanding phylogenetic relationships and polarity of character-state changes when making inferences about adaptive evolution.

1. Introduction

Independent evolutionary lineages challenged by similar environmental extremes provide an ideal context for the study of adaptive evolution. A critical component of this approach is comparative phylogenetic analysis of sequence evolution at functionally relevant genetic loci in closely related species inhabiting contrasting environments. When combined with structural and biochemical information, phylogenetic patterns of amino acid conservation and change can be used to identify changes of likely functional significance and to distinguish parallel evolution in independent lineages from shared ancestral traits (Colosimo et al., 2005; Jost et al., 2008; Feldman et al., 2009; McCracken et al., 2009a).

Two species of waterfowl living at high altitude have served as a prominent example of parallel adaptation to hypoxia and parallel...
morphological evolution related to feeding ecology. The bar-headed goose (Anser indicus) breeds at high elevations in the steppes of central Asia and migrates twice annually across the Himalayas, where the partial pressure of oxygen \( (P_O_2) \) is one-third that of sea level (Scott and Milsom, 2007). In South America, the Andean goose (Chloephaga melanoptera) is endemic to the high Andes, where it is similarly challenged by low \( O_2 \) availability (Hiebl et al., 1987). At high elevations, the low \( P_O_2 \) of inspired air may reduce the \( O_2 \) saturation of arterial blood and limit \( O_2 \) supplies to the tissues (Powell et al., 2004). Bar-headed goose and Andean goose, like other native highland species, exhibit increased \( O_2 \)-affinity, which can be adaptive in high-altitude environments (Hall et al., 1936; Petschow et al., 1977; Monge and León-Velarde, 1991).

In each species, increased \( O_2 \)-affinity has been attributed to the effects of a single amino acid substitution in the major (HbA) hemoglobin (Jessen et al., 1991; Weber et al., 1993). In bar-headed goose, a Pro \( \rightarrow \) Ala substitution at position 119 on the \( \alpha^a \) subunit eliminates an \( \alpha \alpha \) intersubunit contact of van der Waals contact that shifts \( O_2 \) equilibrium in favor of the oxygenated (R-state) conformation of the protein and thereby increases hemoglobin–\( O_2 \) affinity (Fig. 1). Surprisingly, the same mechanism appears to increase \( O_2 \) affinity in Andean goose hemoglobin, but the effect depends on a different substitution on a different gene. In this species, a Leu \( \rightarrow \) Ser substitution at position 55 on the \( \beta^b \) subunit eliminates the same \( \alpha \beta \) intersubunit contact as in bar-headed goose (Fig. 1). The mechanistic basis of parallel evolution in these two high-altitude waterfowl species thus appears to be essentially the same, but the underlying DNA sequence and amino acid changes are independent (Fig. 2). The Andean goose intersubunit van der Waals contact that stabilizes the deoxygenated (T-state) conformation of the HbA structure. van der Waals interactions between \( \alpha^119 \) and \( \beta^55 \) are eliminated by substitution of Ala \( \rightarrow \) Ser in bar-headed goose and Andean goose. Protein crystal structures for greylag goose and bar-headed goose were illustrated using Protein Data Bank 1FAW and 1A4F, respectively. The Andean goose structure was illustrated using 1FAW, with a Leu \( \rightarrow \) Ser replacement modeled in Swiss-PdbViewer 3.7. The structures were illustrated with Python Molecular Viewer 1.5.2.

2. Materials and methods

2.1. Specimen collection, PCR, and DNA sequencing

Sixty-eight tissue samples representing 20 of the 23 species of swans and geese (\( n = 36 \)) and all six species of South American sheldgeese (\( n = 32 \)) were collected from museums or captive waterfowl collections. Among these were three bar-headed geese, including two from Mongolia collected at 1160 and 2600 m, and 20 Andean geese collected at 3743–4611 m at widespread localities in Peru, Bolivia, and Argentina (Supplementary Table 1).

DNA was isolated using DNeasy Tissue Kits (QIAGEN, Valencia, CA). Primers flanking the start and stop codons for the \( \alpha^D \), \( \alpha^A \), and \( \beta^A \) hemoglobin subunit genes were designed using duck, chicken (Gallus gallus), and other sequences in GenBank (Supplementary Table 2). PCR was performed as follows using AmpliTaq Gold PCR Master Mix (Applied Biosystems, Foster City, CA): 7 min preheat at 94 °C, followed by 45 cycles of 20 s at 94 °C, 20 s at 60–64 °C, 1–3 min at 72 °C, and a final extension of 7 min at 72 °C. The complete protein-coding region of each gene, comprising three exons and two introns, was amplified either as a single fragment or in two or three overlapping fragments. Product from an initial PCR spanning the complete \( \beta^A \) subunit was used as the template for a second nested PCR using multiple combinations of internal and primers. Gel-purified PCR products were then cycle-sequenced in both directions using BigDye Terminator Cycle Sequencing Kits on ABI 3100 automated DNA sequencers (Applied Biosystems, Foster City, CA). Sequences were reconciled using Sequencer 4.6 (Code Names, Ann Arbor, MI), and double-peaks, indicating the presence of two alleles, were coded with IUPAC degeneracy codes and treated as polymorphisms. Sequences from samples that were

**Fig. 1.** Illustration of the \( \alpha^119/\beta^55 \) intersubunit contact in bar-headed goose, greylag goose, and Andean goose. Pro-\( \alpha^119 \) and Leu-\( \beta^55 \) lie adjacent to each other on different polypeptide subunits, and in the greylag goose make a van der Waals contact that stabilizes the deoxygenated (T-state) conformation of the HbA structure. van der Waals interactions between \( \alpha^119 \) and \( \beta^55 \) are eliminated by substitution of Ala-\( \alpha^119 \) in bar-headed goose and Ser-\( \beta^55 \) in Andean goose. Protein crystal structures for greylag goose and bar-headed goose were illustrated using Protein Data Bank 1FAW and 1A4F, respectively. The Andean goose structure was illustrated using 1FAW, with a Leu \( \rightarrow \) Ser-\( \beta^55 \) replacement modeled in Swiss-PdbViewer 3.7. The structures were illustrated with Python Molecular Viewer 1.5.2.
heterozygous for an indel were resolved by comparing forward and reverse strands, using the unambiguous 5’ sequence on one strand to interpret the offset peaks 3’ of the indel on the other strand (see Peters et al., 2007). Sequences were aligned by eye using Se-Al 2.0a11 (Rambaut, 2007) and are deposited in GenBank (a) HM125082–HM125128; (b) GQ271002–GQ271003, GQ271019–GQ271050, GQ271481–GQ271513; (c) GQ271748–GQ271749, GQ271768–GQ271792, GQ272207–GQ272227). The αa, αD, and βb subunit sequence alignments are provided as supplementary NEXUS files.

2.2. Phylogenetic analyses

Phylogenetic analyses were conducted separately for each gene, and using concatenated sequences from all three genes. Gene trees were constructed with MrBayes 3.1 (Ronquist and Huelsenbeck, 2003). Each MrBayes analysis included two independent Markov chain Monte Carlo runs of 2 heated chains and 1 cold chain; 10 million generations were completed with trees sampled every 1000 steps. The first 90% of sampled trees were discarded, and the final 1000 trees from each of the two independent runs were combined to compute posterior probabilities for each clade. Tracer 1.3 (Rambaut and Drummond, 2005) was used to verify convergence across independent runs. Substitution models were selected using the Bayesian Information Criterion (BIC; Schwarz, 1978) as calculated by ModelTest 3.7 (Posada and Crandall, 1998) using output from PAUP* (Swofford, 2002). Seven different data partitions (1st, 2nd, and 3rd codon positions, all three codon positions combined, intron 1, intron 2, intron 1 and 2 combined) and five mixed models were evaluated using hierarchical BIC, with gaps coded separately as binary characters. The best-fit models for the single data partitions were the HKY + I (αa), HKY + G (αD), and TrN + G (βb), respectively, and the best-fit model overall for each locus was the five-partition model with separate rate parameters for 1st, 2nd, and 3rd positions, as well as intron 1 and 2, modeled separately (Supplementary Table 3). For the concatenated data set, we used a parameter-rich model, with five data partitions for each subunit. We also performed a non-parametric bootstrap of the concatenated data set using 1000 unweighted parsimony replicates in PAUP*, with 10 random addition replicates each limited to 100
Fig. 3. Fifty-percent majority-rule consensus trees for the $\alpha^A$, $\alpha^D$, and $\beta^A$ hemoglobin subunits of the swans and geese, based on mixed model Bayesian phylogenetic analyses. Posterior probabilities are shown for each clade. Highland lineages are shown in gray boxes. The positions of amino acid replacements within each polypeptide sequence are shown in light (lowland lineage) or dark (highland lineage) circles on each branch.
trees per replicate. Start and stop codons and identical sequences were omitted from each analysis. Trees were rooted on the branch separating the swans and geese (Anserinae) from the sheldgeese (Anatinae) because the reciprocal monophony of these two groups is strongly supported by varied data and analyses (Livezey, 1986; Sibley and Ahlquist, 1990; Sorenson et al., 1999; Donné-Goussé et al., 2002; Bulgarella et al., 2010). Two monotypic genera of African “sheldgeese” (Cyanochen, Alopochen) were not included in our analysis. Cyanochen is not a true sheldgoose (Sorenson et al., 1999; Gonzalez et al., 2009; Bulgarella et al., 2010), and Alopochen is more closely related to shelducks (Tadorna) than to the South American sheldgeese (Donné-Goussé et al., 2002; Gonzalez et al., 2009; Bulgarella et al., 2010).

2.3. Structural analyses of amino acid replacements

The position of each αA- and βA-subunit amino acid replacement was located on the oxy (R-state) crystal structure of the bar-headed goose or greylag goose (Anser anser) HbA hemoglobin (Zhang et al., 1996; Liang et al., 2001; Protein Data Bank 1A4F and 1FAW). Since no HbD structure is available for any species of waterfowl, we used Swiss-PdbViewer 3.7 (Guex and Peitsch, 1997) and the SWISS-MODEL protein structure homology-modeling server (Arnold et al., 2006) to construct a model of the αD subunit and the HbD tetramer using the chicken HbD oxy (R-state) hemoglobin (Knapp et al., 1999; 1HBR) as a structural template. Inter-atomic distances between neighboring amino acids were calculated using CNS 3.4 (National Institutes of Health, Bethesda, MD) and Swiss-PdbViewer 3.7, using a maximum van der Waals distance of 4.1 Å or less (Dall’Acqua et al., 1998). Changes in chemical properties such as polarity, isoelectric point, molecular volume, and normalized van der Waals volume were calculated using published values (e.g., Grantham, 1974). Amino acid replacements were reconstructed on each gene tree using unweighted parsimony in MacClade 4.6 (Maddison and Maddison, 2000). Protein crystal structures were illustrated using Python Molecular Viewer 1.5.2 (Scripps Research Institute, La Jolla, CA). The solvent accessible molecular surface was calculated using the default software settings with a probe radius of 1.5 Å. The electrostatic charge on the surface of the molecule was calculated using the Adaptive Poisson–Boltzmann Solver (APBS) v0.5.1 software (Baker et al., 2001).

3. Results

3.1. Summary of the data

Sequences of the αA, αD, and βA subunits varied from 662 to 680, 873 to 928, and 1563 to 1591 bp, respectively, encoding 141, 140, and 146 amino acids. Several small indels were observed in the two introns of each gene, but no codon insertions or deletions were observed. Amino acid sequences of the αA and αD subunits were 57.1–60.7% identical to each other within each species we sampled, whereas the αD subunits were 80.7–83.8% identical to the chicken αD subunit (Supplementary Table 4).

3.2. Phylogenetic relationships

Concatenated αA, αD, and βA sequences produced a well-resolved phylogeny of the swans and geese at the generic level, with high posterior probabilities and bootstrap values for most clades (Fig. 2). Cereopsis and Coscoroba are each other’s closest relatives and the sister group of other swans and geese. Due to uncertain placement of Cygnus melmocoryphus, the genus Cygnus was paraphyletic in most analyses, with the αA, αD, and βA gene trees yielding conflicting results (Fig. 3). A sister relationship between the goose genera Anser and Branta was well supported in the combined analysis, and the bar-headed goose is likely sister to all other Anser species (but see Gonzalez et al., 2009).

Within the South American sheldgeese, the concatenated tree (Fig. 2) and all three Hb gene trees (Fig. 4) yielded strong support for a clade of lowland Chloephaga excluding Andean goose. The relationships among Neochen, Andean goose, and the other Chloephaga varied among genes (Fig. 4); the combined analysis suggests a sister relationship between Neochen and Andean goose, although this was not strongly supported (Fig. 2).

3.3. Amino acid replacements in highland lineages

Six derived amino acid replacements were found in the bar-headed goose, but not other species of the genus Anser (Fig. 5). These included four replacements on the αA subunit (Gly → Ala-αA-12, Gly → Ser-αA-18, Ala → Val-αA-63, Pro → Ala-αA-119) and two on the αD subunit (Thr → Ser-αD-2, Leu → Val-αD-47). One individual collected at 2600 m in Mongolia (UWBM 57875) was heterozygous for Gly/Ala-αA-12; the other individual was homozygous for Gly. No derived replacements were observed on the βA subunit of bar-headed goose. Glu-βA-121 is a synapomorphy for Anser, and Thr-βA-4 and Asp-βA-125 are the presumably ancestral states found also in Cygnus and Branta (but not other Anser). The only other amino acid replacement occurring within the genus Anser was a Thr → Ala-βA-12 substitution in one of two greylag geese collected in Mongolia (UAM TB 1513); this specimen was homozygous for Ala-βA-12.

Four derived amino acid replacements were observed in Andean goose but not Neochen or other Chloephaga species
These included two replacements on the αA subunit (Thr → Ala-α8, Ala → Thr-α77), one replacement on the αD subunit (Ile → Leu-α13, Ala → Val-α95), and one replacement on the βA subunit (Ala → Ser-β98). All 20 Andean geese that were sampled were homozygous for all of these residues. Ser-β55, which has been the focus of previous analyses of Andean goose hemoglobin, is a synapomorphy for Neochen and Chlorophaga and is thus present in five additional sheldgeese species that are presently found only in lowland environments. Two nonsynonymous substitutions were found in Neochen (Val → Ile-α13, Ala → Val-α95), but none were found in any of the five lowland Chlorophaga species.
4. Discussion

4.1. Taxonomic conclusions

Our findings based on the α4, αD, and β4 hemoglobin subunits are generally in agreement with previous analyses of waterfowl based on morphology (Livezey, 1996a,b, 1997). Monophyly and the close relationship of Anser and Branta were strongly supported, whereas our data were equivocal regarding the monophyly of Cygnus due to an uncertain placement of C. melanocoryphus. Consistent with a recent analysis based on mtDNA (Gonzalez et al., 2009), the hemoglobin data indicate that Cytmus atratus is sister to Cygnus olor, rather than to C. melanocoryphus (see Livezey, 1996a). Likewise, both hemoglobin and mtDNA indicate that the South American C. melanocoryphus is a divergent lineage that is likely the sister taxon of the remaining Cygnus species (Gonzalez et al., 2009). Constraining Cytmus monophyly in our analysis of the concatenated data produces this result and requires just one additional parsimony step, an insignificant increase in tree length (P > 0.52; Kishino–Hasegawa (1989) test).

As in other molecular analyses, we also found strong support for a sister relationship between Cereopsis and Coscoroba (Laskowski and Fitch, 1989; Donne-Goussé et al., 2002; Gonzalez et al., 2009), a result confirmed by shared CR1 retrotransposons (St. John et al., 2005). Similarly, the hemoglobin data suggest that Cereopsis and Coscoroba form the sister group of all other swans and geese. In contrast, Livezey (1996a) placed Cereopsis alone as the sister group to other Anserinae and placed Coscoroba sister to Cygnus based on six synapomorphic characters in the synsacrum, sternum, coracoid, and pelvis. Similarities between Coscoroba and Cygnus have thus likely arisen through morphological convergence.

The hemoglobin data suggest that bar-headed goose is the sister group of all other Anser species included in this study. In contrast, Gonzalez et al. (2009) found bar-headed goose to be embedded within Anser based on sequences from two mitochondrial genes. In our study, intra-generic relationships within Anser and Branta were poorly resolved, owing to short branch lengths, with different gene trees supporting different topologies. In the case of the three hemoglobin genes sequenced here, it is likely that the small number of variable sites and incomplete lineage sorting both contribute to lack of resolution within genera. Additional data from the mitochondrial genome or many more nuclear loci likely will be required to resolve these relationships.

Like Livezey (1997), we also found strong support for a clade of lowland Chloephaga excluding Andean goose. The concatenated data suggest that Neochen is sister to Andean goose, and that these two species from northern South America form a clade, sister to the four lowland Chloephaga species endemic to southern South America. Additional data will also be required to resolve the relationships of these taxa.

4.2. Amino acid replacements in highland lineages

Structural changes in the HbA isoform arising from substitutions at α18, α63, α119, and β125 have been examined extensively using the protein crystal structures of bar-headed goose and greylag goose (Zhang et al., 1996; Wang et al., 2000; Liang et al., 2001a,b; Liu et al., 2001). Additionally, the effects of Ala-α119 also have been measured experimentally using site-directed mutagenesis to incorporate this substitution into human hemoglobin (Jessen et al., 1991; Weber et al., 1993). Our findings provide insight into the evolutionary pathways by which these amino acid replacements occurred within the Anserinae. Ser-α18, Val-α63, and Ala-α119 are each derived residues, present only in bar-headed goose and absent in greylag goose and other Anser species. In contrast, Asp-β125 is not derived, but is the ancestral residue present in most Anserinae and other basal waterfowl lineages (McCracken et al., 2009a). Based on phylogenetic analysis of the concatenated hemoglobin sequences, Glu-β125 is a synapomorphy for a clade comprising all Anser species except bar-headed goose. We also observed a Gly → Ala-α12 replacement in a single bar-headed goose specimen from Mongolia. This residue occurs at an exterior, solvent-accessible position on the A helix of the α subunit (Fig. 6), a region of the HbA isoform that has experienced parallel changes in three other high-altitude waterfowl species, including the Andean goose (as discussed below; see also McCracken et al., 2009a,b). Thus, it would be worthwhile to measure the frequency of this polymorphism in the bar-headed goose population and also determine if it affects hemoglobin-O2 affinity. Additionally, we do not yet know whether this polymorphism represents two allelic copies of a single α4-subunit gene, or whether bar-headed goose possesses duplicate α4-subunit genes distinguished by a single amino acid change. An example of this kind of similarity between gene copies is maintained by recurrent gene conversion in the

Fig. 6. HbA structure illustrating three amino acid substitutions (Thr → Ala-α18, Gly → Ala-α12, and Gly → Ser-α18 shown in yellow) at external, solvent-accessible positions on the A helix and AB corner of the α subunit (shown in blue). Substitutions at two adjacent sites that occurred in other highland waterfowl species (McCackren et al., 2009a) are shown in faded gray. The top panel shows the internal three-dimensional ‘ribbon’ structure of the A helix and AB corner, whereas the bottom panel shows the external, solvent-accessible surface of the protein with a probe radius of 1.5 Å. The protein crystal structure was illustrated using Protein Data Bank 1FAW and Python Molecular Viewer 1.5.2. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
duplicated α and β globin genes of deer mice (Peromyscus maniculatus; Storz et al., 2007, 2009).

We also observed two derived substitutions on the αD subunit of bar-headed goose (Thr → Ser-αD2 and Leu → Val-αD47) that were previously identified by protein sequencing (Hiebl et al., 1986). Thr → Ser-αD2 occurs at the N-terminus of the αD subunit in a region of the protein tetramer that may function in inositolpentaphosphate (IP5) binding (Fig. 7), based on corresponding IP5 binding sites identified in the South Polar skua (Catharacta macroura; Tamburrini et al., 2000; Riccio et al., 2001). IP5 is the principal allosteric effector in avian red blood cells. It influences the equilibrium between deoxygenated (T-state) and oxygenated (R-state) hemoglobin by binding to positively charged sites between in the central cavity between the N- and C-termini of the α- and β-chain subunits, thereby stabilizing the deoxygenated conformation of the protein with salt-bridges. Changes that suppress phosphate binding are thus expected to reduce hemoglobin–O2 affinity (Storz et al., 2009).

The total number of derived amino acid substitutions in the HbA and HbD isoforms of bar-headed goose was thus four and two, respectively. No replacements were observed on the βA subunit, which is a component of both HbA and HbD.

Andean goose exhibited a total of four derived amino acid substitutions. Two substitutions occurred on the αA subunit (Thr → Ala-αA8 and Ala → Thr-αA77), one occurred on the αB subunit (Ile → Leu-αB9), and one occurred on the βB subunit (Ala → Ser-βB66). Thr-αA77 and Ser-βB66 each occur in close proximity to IP5 binding sites and in regions of positive charge (Fig. 7; Wang et al., 2000). Substitution of threonine and serine each result in non-polar to polar changes that correspond to the addition of a hydroxyl group and increased molecular volume. It is thus possible that the hydroxyl group substitutions influence IP5 binding indirectly; IP5 is a negatively charged ligand. The addition of threonine or serine might also enhance hemoglobin solubility, and this could be beneficial in desiccating, high-altitude environments. The likely functional significance of Thr-αA77 is also supported by the fact that the same substitution is independently derived in four other waterfowl species inhabiting high-altitude regions (McCracken et al., 2009a,b).

Intriguingly, the two other derived replacements we observed in Andean goose, Ala-αA8 and Leu-αA9, occur at nearby positions on the A helix of the αA and αB subunits (αA9 corresponds to αB10 in HbD and HbA, respectively, due to one amino acid difference in the length of the αD- and αA-chain subunits). Highland populations of two other Andean waterfowl species also have derived amino acid substitutions in this structural domain of the HbA tetramer; crested duck (Lophophetra spectabilioides; Thr-αA8 and Ala-αA78) and cinnamon teal (Anas cyanoptera; Ser-αA9; McCracken et al., 2009a). One noteworthy difference, however, is that while αA8, 9, 12, and 18 each occupy external, solvent accessible positions in the HbA structure (Fig. 6), αA9 occurs at an internal position within the chicken HbD structure.

Finally, the finding that Ser-βB55 is not derived in Andean goose, but is a synapomorphy for the South American sheldgeese (Neochen + Chloephaga) raises questions about its physiological function in species native to lowland environments. Perutz (1983, 1989) hypothesized that Leu → Ser-βB55 destabilizes the deoxygenated (T-state) conformation of the protein by eliminating a van der Waals contact at an important αI intersubunit contact (Fig. 1), thereby shifting the allosteric equilibrium in favor of the oxygenated (R-state) structure. When Ser-βB55 was experimentally introduced into recombinant human hemoglobin, this pre-

Fig. 7. HbA and HbD structures illustrating three amino acid substitutions (Ala → Ser-αA8, Ala → Thr-αA77, and Thr → Ser-αD2 shown in green) that occurred in close proximity to inositolpentaphosphate (IP5) binding sites (shown in yellow). The HbA IP5 binding sites include βA1, 2, 84, 104, 135, 143, 144, and 146 (Wang et al., 2000; top left), and α1, 95, 99, 134, 137, 138, and 141 (Tamburrini et al., 2000; Riccio et al., 2001; top middle); the same homologous α2 sites are shown for the HbD isoform of chicken (top-right). The electrostatic charge on the surface of the molecule is illustrated for each structure at bottom. Blue surfaces correspond to positively charged regions, and red surfaces correspond to negatively charged regions. In the HbA isoform, the IP5 binding sites shown in yellow (top row) occur within regions of dense positive charge (bottom row). Protein crystal structures for HbA and HbD were illustrated using Protein Data Bank 1FAW and 1HBR, respectively, and Python Molecular Viewer 1.5.2. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
dicted effect was observed (Jessen et al., 1991; Weber et al., 1993). This raises the intriguing hypothesis that Ser-β55 may have resulted in elevated hemoglobin–O₂ affinity in species that presently are restricted to lowland environments, such that the South American sheldgeese exhibit higher average blood–O₂ affinity than other lowland endemic waterfowl species. This hypothesis remains to be tested experimentally, but if so, it might be used to infer the direction of colonization of the Andes by the sheldgeese. If the ancestral allele for this clade confers elevated hemoglobin–O₂ affinity, the South American sheldgeese might have originated in the highlands, but more recently colonized lowland regions.

5. Conclusion

Our study of the swans and geese and the South American sheldgeese represents one of the first comprehensive molecular phylogenetic analyses of these taxa and provides a historical basis for understanding patterns of amino acid substitution in the HbA and HbD genes of bar-headed goose and Andean goose, two species of waterfowl that have featured prominently in studies of high-altitude adaptation. Bar-headed goose and Andean goose have adapted to similar hypoxic environments in the Himalayas and Andes, and thus experience the same hypobaric selection pressures. While these two species share no identical amino acid substitutions in either the Hba or Hbd subunit genes, similar types of substitutions have occurred within the same structural regions of the protein, including the same structural domains of the α and β subunits. The mechanistic basis of hypoxia resistance in these two waterfowl species has yet to be fully determined, but the patterns identified here provide an example of parallel evolution of sites within the same hemoglobin protein domains. In addition to the previously described amino acid substitutions that influence an important αβ intersubunit contact, additional substitutions that suppress the phosphate sensitivity of hemoglobin–O₂ affinity may also likely be important. Further experimental studies are required to determine how the observed substitutions influence hemoglobin–O₂ binding properties.

It is also important to identify the molecular mechanisms responsible for differences between the Hba and Hbd isoforms in their intrinsic O₂ affinity, allosteric regulation, and oxygen-linked polymerization (Cirotto and Geraci, 1975; Baumann et al., 1984; Riggs, 1998; Knapp et al., 1999; Rana et al., 2008). Hbd has been shown to have higher O₂ affinity than Hba, and the ratio of these two isoforms in avian red blood cells may play an important role in modulating blood–O₂ affinity (Hiebl et al., 1988; Weber et al., 1988). For example, it is possible that highland species maintain elevated blood–O₂ affinity by upregulating expression of the β gene, thereby increasing the relative abundance of the Hbd isoform (Hoffmann and Storz, 2007; Bulgarella et al., 2008).

In conclusion, our findings illustrate the importance of understanding phylogenetic relationships and polarity of character-state changes when making inferences about adaptive evolution. A comparative phylogenetic approach reveals the evolutionary origins of substitutions of putative functional significance. For example, we found that a substitution showing evidence of physiological adaptation was not uniquely derived in a high-altitude lineage. Similarly, recurrent patterns of substitution in relation to protein structure across independent lineages can lead to testable hypotheses about function that can be used to guide future experimental and comparative studies.

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Appendix A. Supplementary data


References


Hiebl, I., Schneeggans, D., Braunitzer, G., 1988. The primary structures of the α and β chains of the bar-headed goose (Anser indicus); the greylag goose (Anser anser) and the Canada goose (Branta canadensis). Biol. Chem. Hoppe-Seyler 367, 591–599.


