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# Adaptation Mechanisms of Yak (Bos grunniens) to High-Altitude Environmental Stress

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**Simple Summary:** The yak is a multipurpose domesticated animal that serves as a protein source for local herders and a sacred carrier of culture and religion. Besides their economic significance, yaks harbor special morphological, physiological, biochemical, and genetic adaptations for tolerance to high-altitude stress. Morphologically, yaks have large hearts and lungs, compact bodies, thick outer hair covering, and nonfunctional sweat glands, which help to withstand hypoxia and cold stress. A reduced heat production, decreased respiration and sweating, reduced metabolism, and efficient nitrogen utilization are the major physiological and biochemical mechanisms for yak survival at high altitudes. Furthermore, the yak has undergone long-term natural selection and developed a unique genetic architecture that favors survival in hostile environments. The yak expresses the HIF- $1\alpha$  pathway-related genes (*ADAM17*, *ARG2*, and *MMP3*) putatively involved in hypoxia response and nutrition pathways genes (*CAMK2B*, *GENT3*, *HSD17B12*, *WHSC1*, and *GLUL*) for nutritional assimilation at high altitudes.

Abstract: Living at a high altitude involves many environmental challenges. The combined effects of hypoxia and cold stress impose severe physiological challenges on endothermic animals. The yak is integral to the livelihood of the people occupying the vast, inhospitable Qinghai—Tibetan plateau and the surrounding mountainous region. Due to long-term selection, the yak exhibits stable and unique genetic characteristics which enable physiological, biochemical, and morphological adaptations to a high altitude. Thus, the yak is a representative model for mammalian plateau-adaptability studies. Understanding coping mechanisms provides unique insights into adaptive evolution, thus informing the breeding of domestic yaks. This review provides an overview of genetic adaptations in *Bos grunniens* to high-altitude environmental stress. Combined genomics and theoretical advances have informed the genetic basis of high-altitude adaptations.

Keywords: adaptation; high altitude; hypoxia; yak



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## 1. Introduction

The Qinghai–Tibetan Plateau ecological environment is characterized by low atmospheric oxygen pressure, cold, and limited feed supplies [1,2]. Endothermic animals endure impaired oxygen supplies, which compromise cellular functions and physiologic performance under high-altitude environments. These high-altitude mountains are more sensitive and vulnerable to climate change, a huge threat to biodiversity and the ecosystem [3]. Moreover, the cold temperature increases the environmental harshness with the temperature dropping by ~6 °C per kilometer above sea level [4]. Other factors, including late winter and early spring, feed shortage, and snow cover, inevitably lead to severe malnutrition and weight loss among animals [5,6]. Species have developed special characteristic features through natural selection to adapt to extreme terrestrial environments [7,8].

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The yaks are the world's most remarkable domestic animal living freely and reproducing under the harsh plateau environment [1]. The yak was domesticated >7300 years ago from wild yak by the early nomadic people and is the only large animal that coexists with its wild ancestors in a similar environment [9]. Over 17.6 million yaks exist globally, and the majorities are found in the plateau regions of central Asia, which cover ~2.5 million km² centered around the Qinghai–Tibetan Plateau and adjacent highlands [10]. The yak supplies (milk, meat, hair, hides, and manure) and services (draft, packing, and riding) the pastoralists and agro-pastoralists occupying these areas. They are also means of financial security and cultural functions (status, dowry, religious festivals) [6]. They also offer a good framework for studying the effects of natural and artificial selection in livestock domestication and adaptation to different environments.

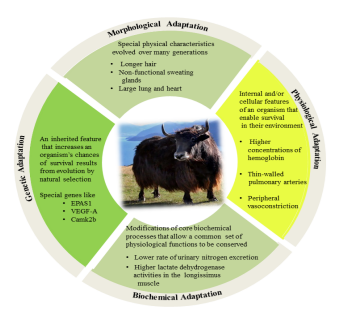
The yak is the only bovine species native to the Qinghai–Tibetan Plateau and adjacent highlands that exhibits a high adaptability to high altitudes, a low sensitivity to cold, a low oxygen pressure, and prolonged periods (approximately half a year) of food scarcity [3,6,11,12]. Natural and artificial selections from domesticated yaks resulted in breeds with distinct morphological, physiological, and adaptability traits that enhanced survival in harsh environments [13–15]. Understanding the aerobic metabolism of yaks under hypoxic environments can provide important insights into adaptive evolution [16]. Together with advanced molecular techniques and genetic research, these insights provide a basis for investigating the genetic mechanisms underlying adaptability to climate change, the current research hotspot [17,18].

Knowledge of the mechanisms underlying adaptation to various agro-ecosystems is essential for the effective management of farm animal genetic resources [7,19]. Intriguingly, multiple studies have focused on yak adaptation, allowing researchers to understand the morphological, physiological, biochemical, and genetic mechanisms of adaptation to extremely high altitudes [13,15,20–22]. Although research on the adaptation mechanisms of the yak in high altitudes has increased exponentially, review studies on the comprehensive, adaptive mechanisms remain scarce. Therefore, this review attempts to collate and synthesize current knowledge on the mechanisms of yak adaptation to high altitudes. Furthermore, it can also provide new avenues for in vitro and in vivo studies to further test hypotheses arising from previous investigations and options for designing and implementing interventions for improved yak productivity and resilience in high altitudes.

#### 2. High Altitude Adaptation Mechanisms of Yak

High altitudes negatively impact the normal bodily functions of individuals, whether they are accustomed or unaccustomed to such environments. Mishra and Ganju reviewed high-altitude environmental factors, such as cold and hypobaric hypoxia, which affect the immune system, making it more susceptible to cancer, infection, and autoimmune disease [23]. Inadequate hypoxia treatment affects reproduction and fertility traits, including reduced intrauterine growth in sheep [24] and impaired development and function of corpus luteum [25]. There should be an increased focus on breeding and managing animals for an improved resilience to applied stressors [26]. To adapt to high-altitude environments, plateau-dwelling mammals have developed some distinct characteristics. The yak, a unique breed that inhabits the alpine pastoral area of the Tibetan Plateau, is one of the rare bovine breeds adapted to high altitudes and cold climates [12,27]. The adaptive process is extremely complex, consisting of several components that exhibit stable and unique genetic characteristics for regulating the physiological, biochemical, and morphological mechanisms of adaption to a high altitude (Figure 1).

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**Figure 1.** Schematic representation of the yak adaptation to high-altitude environmental stress in comparison to a closely related species (*Bos indicus* or cattle).

## 2.1. Morphological Adaptations

Morphological adaptations are physical changes that occur over many generations of animals to enhance fitness in a given environment. Over many generations, the native high-altitude *B. grunniens* successfully adapted to the chronic hypoxia of high altitudes despite belonging to the genus *Bos*, closely related to cattle [28]. The exceptional adaptation of the yak to high altitudes is related to evolved special morphological mechanisms (Table 1). Compared with close relatives such as cattle that live at lower altitudes, yaks have relatively larger lungs and hearts [29]. Furthermore, the yak has longer, wider, and rounder pulmonary-artery endothelial cells with little smooth muscles, which allows for improved functioning in high-altitude environments compared to cattle [28,30].

Special Morphological Structures	Function	References	
Compact body, thick outer hair covering, and nonexistence of functional sweat glands	Minimize dissipation of body heat during winter	[6]	
Thin-walled pulmonary arteries with little smooth muscles	Facilitate superefficient O <sub>2</sub> flow under hypobaric hypoxia	[16]	
Larger lungs and hearts	Aid oxygen uptake	[17]	
Shorter tongue and greater lingual prominence	Improve forage digestibility through efficient grinding of food	[31]	

**Table 1.** Key morphological adaptations of yaks to high-altitude environments.

Not only does the hypoxic environment impact life at high altitudes, freezing temperatures and scarce food supply also contribute to the harsh environment. The alpine habitat of yaks is at altitudes of 3000–6000 m. As a result, there is no frost-free period throughout the year. The yaks are well-adapted to the cold, high-altitude environment; their bodies are compact with a relatively reduced skin surface area per unit of body weight  $(0.016 \text{ m}^2/\text{kg})$ . They also have thick outer hair coats and no functional sweat glands [6]. The absence of sweating in the yak enhances cold tolerance [32].

Furthermore, the thick fleece covering the entire yak body enhances heat conservation. The thick fleece comprises an outer coat of long hair and an undercoat consisting of a dense layer of fine down fibers that appear in the colder season to contain body heat and repel moisture [6]. Undoubtedly, feeding mechanisms are an important factor in determining

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the success and survival of vertebrate species within their environment [33]. The alpine habitats at high altitudes are marked by a severe climate and very short growing seasons with limited grazing resources and often treacherous terrain. Together, these factors lead to severe malnutrition and weight loss among animals [6,34]. Yaks have developed shorter tongues with greater lingual prominence, larger and more numerous conical papillae, and thicker keratinized epithelium than domestic cattle. These attributes enable the yak to consume a wider variety of pasture plant species [31]. Furthermore, the yak rumen is unusually large relative to omasum. This large rumen allows it to consume large quantities of low-quality food at a time and to ferment it longer in order to extract more nutrients during times of nutritional scarcity [6].

## 2.2. Physiological Adaptations

Physiology can be viewed as mechanisms and processes that allow organisms to deal with internal challenges (for example, exercise, growth, and reproduction) and external stress (e.g., variations in temperature, oxygen, water availability, salinity, pressure, radiation, and heavy metals, etc.). The yak inhabits the entire Qinghai–Tibetan Plateau, and physiological adaptations have contributed to their success in surviving hostile environmental conditions [35]. Chronic hypoxia is the primary stressor of high-altitude conditions and limits the efficient functioning of respiratory and cardiovascular systems in mammals and birds [36]. Interestingly, the prolonged exposure of yaks to high altitudes increases their physiological response to chronic hypoxia because yaks have a larger pulmonary alveolar area per unit area, thinner alveolar septum, thinner blood–air barrier [37], larger hearts and lungs, and higher concentrations of erythrocytes and hemoglobin than other cattle species [29]. Thin-walled pulmonary arteries with little smooth muscle and absent right ventricular hypertrophy are additional hypoxic adaptations observed in yaks [16]. Indeed, these characteristics and changes in the cardiovascular system compensate for the hypobaric high-altitude environment [5].

Furthermore, this adaptation is probably due to natural selection, which enhances the hypoxic pulmonary vasoconstrictor to respond with no hypoxemic stimulus for increased red blood cell production and hemoglobin concentrations [16]. Previous research demonstrates that chronic exposure to hypoxic conditions raises the blood/erythrocyte volume for high-altitude native animals [38]. As a result, hyperventilation, hemoconcentration, and stimulated erythropoiesis are physiological responses that warrant oxygen delivery to tissues [39,40].

Adaptation often occurs at the expense of performance, and survivability is often better in "low" performance animals because of their low input needs (especially feed) and moderate internal heat production [41]. In their natural habitat, yaks must maintain normal energy production under hypoxic conditions [5,11] and optimize nutritional assimilation as a consequence of the cold stress [5] and limited feed [6]. Under cold stress, heat loss is prevented by peripheral vasoconstriction and heat production through shivering and uncoupled mitochondrial activity [42]. Furthermore, multiple cold-adapted species can temporarily slow their metabolisms in response to harsh environmental conditions, leading to torpor or, in extreme cases, hibernation [43]. The significant reduction in heat production of yaks during winter might result from their adaptation to low oxygen concentrations in the air, the cold environment, and the long-term under nutrition prevalent in the sixmonths-long cold season of the Tibetan plateau [5].

#### 2.3. Biochemical Adaptations

The cold, hypoxic conditions of high-altitude habitats impose severe metabolic demands on endothermic vertebrates. Understanding how high-altitude endotherms cope with the combined effects of hypoxia and cold can provide important insights into the process of adaptive evolution. Biochemical adaptions provide fascinating insights into how organisms "work" and how they evolve to sustain physiological function under a vast array of environmental conditions. Their high blood hemoglobin concentrations enable

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yaks to adapt to tolerating low atmospheric partial pressures of oxygen characteristic of the >2000 m Qinghai–Tibetan plateau [5,6]. The lower energy metabolism of yaks might result from adaptation to a low O<sub>2</sub> concentration in the air, a cold environment, and the long-term undernutrition during the annual six-months-long cold season of the Qinghai–Tibetan plateau. The yak rumen microorganisms ferment approximately 70–80% of the feed intake and produce volatile fatty acids, providing 60–75% of the required metabolic energy. This phenomenon may be a coevolutionary coping strategy for low feed resources in cold environments [44]. Compared to indigenous cattle, yaks have a lower rate of urinary nitrogen excretion (a possible adaptation to a poor feed supply) and more efficient nitrogen utilization, which is at least in part due to a greater microbial protein production in the rumen [45]. This adaptability is believed to assist in the rapid recovery of body weight over the summer grazing period [46,47]. In addition, the low maintenance protein requirements [48] and low surface area of the yak body [49] result in a low metabolic rate. Altogether, these attributes are beneficial for the yak's survival in the harsh environmental conditions of the Tibet plateau.

Despite variations among species, most studies attribute metabolic adaptation under high altitudes to a decreased muscle oxidative capacity. In this regard, lactate dehydrogenase (LDH) is the crucial enzyme in anaerobic glycolysis, catalyzing the conversion between pyruvate and lactate, a critical role in energy metabolism [50]. Interestingly, unlike cattle, the yak exhibits higher LDH activities in the longissimus muscle, facilitating carbohydrate utilization under limited oxygen supplies. Thus, this provides a unique adaptive feature of yaks in high altitudes [51].

## 2.4. Genetic Background of High-Altitude Adaptations

Genetic adaptations to novel environments and climatic changes are a fundamental process for species' survival. The genetic mechanism for adaption to high altitudes appears to be more complicated than any other phenotype understood thus far [52]. Species have developed special characteristic features through long-term selection to cope with the specific stressors of extreme terrestrial environments [7,8]. Genetic variation in a population provides flexibility to adapt to changing environments and is crucial for the survival and speciation of that population over time [53]. In the past, it was customary to focus on structural sequence variation and consider each gene as a separate unit of evolution in population genetic theory and empirical practice. However, adaptive phenotypes are more likely a function of polygenic mutations. The detection of adaptive genetic signals with conventional selection methods is more complicated than detecting other phenotypes [54]. It is widely accepted that adaptive evolutionary mechanisms can evolve via changes at one or a few loci (selective sweep) with major effects or via simultaneous allele frequency shifts at many loci with small effects [55]. The major developments in sequencing and genotyping technology over the past decade have facilitated the identification and selection of population-specific genome signatures for livestock adaptation. In the yak populations, several genes putatively associated with adaptation to life at high altitudes were identified. These genes are primarily related to physiological pathways in response to hypoxia and temperature acclimatization and modifications of the cardiovascular system and energy metabolism [13,20,56]. Endothelial PAS domain-containing protein 1 (EPAS1) is the top candidate gene encoding the hypoxia-inducible transcription factor (HIF- $2\alpha$ ). This gene (EPAS1) is believed to regulate erythropoietin production, which changes with the available oxygen in the cellular environment under high-altitude conditions [13,21,57]. There are also distinct selection signatures within yaks, which suggest unique adaptation mechanisms (Table 2).

For example, the vascular endothelial growth factor-A (*VEGF-A*) gene is a key regulator of angiogenesis and an endothelial cell mitogen that regulates blood vessel size as an adaptation to high-altitude functioning [35]. Furthermore, yaks must not only maintain a normal energy production under hypoxic conditions [5,11] but must also optimize nutritional assimilation as a consequence of the limited forage resources available in their

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high-altitude environments [6]. Qiu et al. [13] reported five key genes showing positive selection in the yak nutritional and metabolic pathways. Among the five described genes, *Camk2b* gene regulates the secretion of gastric acid in the rumen, contributing to the assimilation of volatile fatty acids produced by ruminal fermentation [58,59]. In addition, *Gcnt3*, *Hsd17b12*, *Whsc1*, and *Glul* are functional in the polysaccharide, fatty acid, and amino-acid metabolism, respectively [60,61]. The positively selected changes in *Glul* may be important for the enhanced nitrogen utilization in yaks [11]. Other genes putatively involved in the yak adaptation to high-altitude environments are summarized in Table 2.

Table 2. Genes	candidate genes	underlying s	selection s	ignatures of	vak ada	ptation to high	altitudes.

Candidate Genes	Functions	References
Camk2b, Gcnt3, Hsd17b12, Whsc1, and Glul	High level of nutrition utilization in high altitudes	[13]
HIF1A, MMP3, ADAM17, ARG2	High-altitude adaptation	[13]
DEXI, DCC, and MRP4	Adaptation to high-altitude environments	[14]
PDE4D, RPS6KA6, ITPR1, and GNAO1	Environmental information processing and environmental adaptability	[20]
EPAS1	Key transcription factor that activates the expression of oxygen-regulated genes	[21]
ABCG8, COL4A1, LOC102287650, PDCD1, and NUP210	Adaptation to high-altitude environments	[22]
VEGF-A	Regulation of blood vessel size	[35]
MMP3	Regulator of the cellular response to hypoxia	[58]
HIF-1α	Transcription of genes involved in oxygen homeostasis	[59]
AQP4 ATP8 and ATP6	Resistance to cerebral edema Mitochondrial ATPase assembly	[62] [63]
DCC, GSTCD, MRPS28, and MOGAT2	Adaptation to high-altitude environments	[64]
MT-ND1 and MT-ND2	Electron transport chain of oxidative phosphorylation	[65]
GRIK4, IFNLR1, LOC102275985, GRHL3, and LOC102275713	Physiological regulation under a hypoxic environment	[66]

#### 3. Transcriptomic Changes in Yaks Living in High-altitude Environments

Gene expression profiles indicate the activation of specific molecular pathways/networks of genes that regulate external stimuli and provide insight into the role of regulatory variation in adaptive evolution [67-69]. Recent studies have identified different genes and pathways that widely participate in various biological processes, including adaptations to hypoxia. During periods of reduced O2 availability, changes in gene expression are mediated by a specific family of transcription factors called the "master regulator" of O<sub>2</sub> homeostasis. These genes are collectively known as hypoxia-inducible factors (HIF) [70,71]. Hypoxia-inducible factors are oxygen-dependent transcriptional activators, crucial in tumor angiogenesis and mammalian development through the transcriptional regulation of oxygen homeostasis genes in response to hypoxia [72]. In mammalian cells,  $HIF\alpha$  isoforms (HIF-1 $\alpha$  and HIF-2 $\alpha$ ) are the most extensively studied and understood central mediators of cellular adaptation to hypoxia. HIF1α functions as a key regulator of O<sub>2</sub> homeostasis that coordinates oxygen sensing and intracellular responses to hypoxia by regulating the expression of hundreds of genes. Some of the  $HIF1\alpha$ -regulated genes belong to biological pathways for energy metabolism, angiogenesis, erythropoiesis, iron homeostasis, and apoptosis [73–75].

Because of their adaptation to high-altitude environments coupled with exceptional physical endurance, yaks ought to be a model animal for understanding the molecular basis of high-altitude adaptation. As a result, the cDNA of  $HIF-1\alpha$ , a subunit of  $HIF-1\beta$ 

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in the yak, was sequenced by Dolt et al. [57]. The authors observed the variant-specific expression of  $HIF-1\alpha$  in the blood and liver. In contrast, expression was absent in the lung, heart, and kidney of yaks. This tissue-specific expression might be the consequence of the alternative splicing of  $HIF-1\alpha$ , as observed in plateau pika, another high-altitude-adapted animal [76].

Furthermore, a comparison of tissue-specific expression between yaks and cattle revealed that HIF- $1\alpha$  was ubiquitously expressed, whereas HIF- $2\alpha$  expression in the yak was limited to endothelial cells in tissues (kidney, heart, lung, spleen, and liver) and blood [72]. The expression of both HIF- $1\alpha$  and HIF- $2\alpha$  was higher in the yak tissues than in cattle. Other comparative transcriptome studies between cattle and yaks revealed that, among several organs, the gene expression patterns of the heart showed the greatest differentiation between the two species [77] and that differentially expressed genes in lung and gluteus tissue were involved in red blood cell development and inhibition of blood coagulation [78].

#### 4. Conclusions

High-altitude habitats are characterized by extremely harsh climates consisting of low temperature and low oxygen pressure. The native high-altitude yak has evolved multiple unique adaptations, including morphological, physiological, biochemical, and genetic changes due to long-term selection. To gain a more holistic understanding of high-altitude adaptations, these types of studies need expansion, and efforts should be made to integrate work on DNA sequence polymorphism with analyses of transcriptional variation. Indeed, cold and hypoxia act synergistically on an organism's performance at high altitudes, yet the vast majority of studies have focused solely on adaptation to hypoxia. Hence, joint investigations of these co-occurring environmental stressors should be the priority for future research. Furthermore, the current climate change scenario characterized by rising temperature undoubtedly alters the natural habitats of yaks by creating new environmental conditions to which these animals were never before exposed. Therefore, further investigations to determine how these shifts in climate contribute to changes in the yak production and the livelihoods of highlanders are indispensable.

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