

# Neonatal oxygenation, pulmonary hypertension, and evolutionary adaptation to high altitude (2013 Grover Conference series)

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**Abstract:** Andeans and Tibetans have less altitude reduction in birth weight than do shorter-resident groups, but only Tibetans are protected from pulmonary hypertension and chronic mountain sickness (CMS). We hypothesized that differences in neonatal oxygenation were involved, with arterial O<sub>2</sub> saturation (SaO<sub>2</sub>) being highest in Tibetans, intermediate in Andeans, and lowest in Han or Europeans, and that improved oxygenation in Andeans relative to Europeans was accompanied by a greater postnatal decline in systolic pulmonary arterial pressures (P<sub>pa,sys</sub>). We studied 41 healthy (36 Andeans, 5 Europeans) and 9 sick infants at 3,600 m in Bolivia. The SaO<sub>2</sub> in healthy babies was highest at 6–24 hours of postnatal age and then declined, whereas sick babies showed the opposite pattern. Compared to that of 30 Tibetan or Han infants studied previously at 3,600 m, SaO<sub>2</sub> was higher in Tibetans than in Han or Andeans during wakefulness and active or quiet sleep. Tibetans, as well as Andeans, had higher values than Han while feeding. The SaO<sub>2</sub>'s of healthy Andeans and Europeans were similar and, like those of Tibetans, remained at 85% or above, whereas Han values dipped below 70%. Andean and European P<sub>pa,sys</sub> values were above sea-level norms and higher in sick than in healthy babies, but right heart pressure decreased across 4–6 months in all groups. We concluded that Tibetans had better neonatal oxygenation than Andeans at 3,600 m but that, counter to our hypothesis, neither was SaO<sub>2</sub> higher nor P<sub>pa</sub> lower in Andean than in European infants. Further, longitudinal studies in these 4 groups are warranted to determine whether neonatal oxygenation influences susceptibility to high-altitude pulmonary hypertension and CMS later in life.

**Keywords:** Andean, cardiopulmonary transition, Ethiopian, European, genetic adaptation, Han, hypoxia, Tibet.

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Studies at high altitude have long provided a useful model for persons interested in the pulmonary circulation.<sup>1-4</sup> While pulmonary arterial pressures (P<sub>pa</sub>'s) are generally elevated at high altitude, there is considerable population variation in both P<sub>pa</sub> and pulmonary circulation-related disorders such as chronic mountain sickness (CMS; Fig. 1A)<sup>11,36</sup> and subacute infantile mountain sickness.<sup>37</sup> Specifically, directly measured P<sub>pa</sub> and hypoxic pulmonary vasoconstrictor responses are lowest in Tibetans, intermediate in Andeans, and highest in European-derived residents of Colorado (Fig. 1B). Recent estimates using echocardiography also suggest intermediate values in Ethiopians.<sup>15</sup> While multiple factors contribute to the etiology of CMS and no well-controlled comparative epidemiologi-

cal studies exist, a reduced prevalence among Tibetans is consistent with the lower hemoglobin levels seen in Tibetans, relative to Andeans, in multiple reports.<sup>38-40</sup> Tibetans are also protected, relative to Han, from subacute infantile mountain sickness, a syndrome characterized by severe pulmonary hypertension in 4–6-month-old infants that has also been reported in European-derived populations in Colorado.<sup>37,41,42</sup> Whether subacute infantile mountain sickness is less common in Andeans than in Europeans is unknown, but lower P<sub>pa</sub>'s have been reported in Andean than in European children.<sup>43</sup>

One factor influencing P<sub>pa</sub> during adulthood is perinatal hypoxia, as shown by human and experimental animal studies in which greater hypoxia in utero or during

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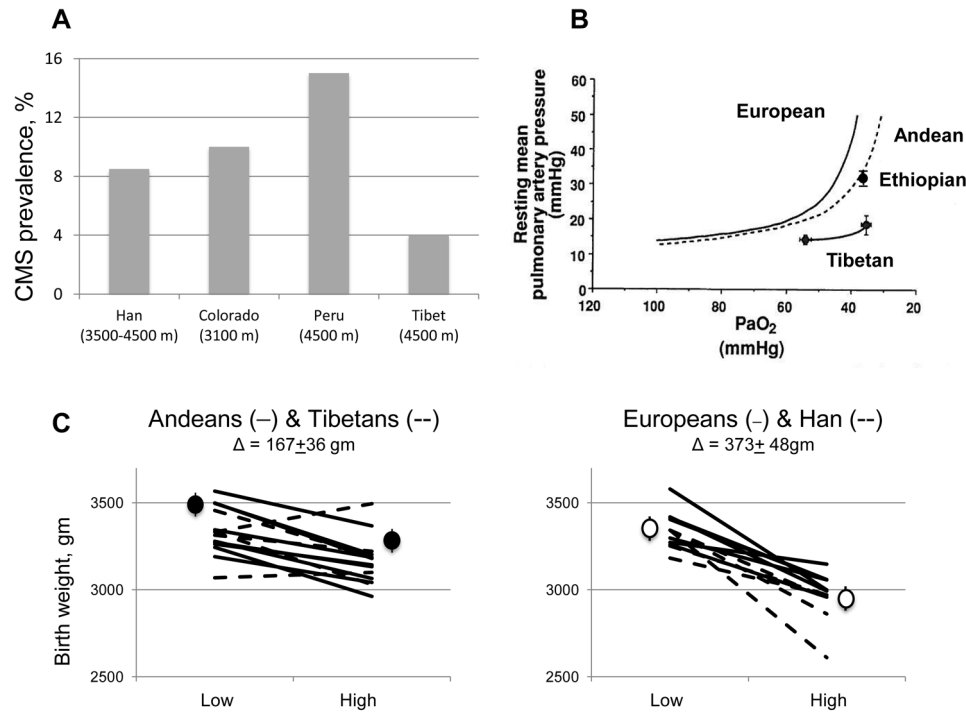


Figure 1. A, Chronic mountain sickness (CMS) is a debilitating condition occurring chiefly in men and characterized by excessive polycythemia in the absence of chronic obstructive pulmonary disease or other known causes and by symptoms of dyspnea, palpitations, sleep disturbance, cyanosis, dilatation of veins, paresthesia, headache, and/or tinnitus.<sup>5</sup> Studies in a total of 14,643 male high-altitude residents show a much lower prevalence of chronic mountain sickness (CMS) in Tibetans than in Han, Coloradans, or Andeans, especially when the altitude of residence is taken into account. Data are from 13,233 Han in Qinghai Province, China,<sup>6</sup> 110 European-derived residents of Leadville, Colorado (I. Asmus, personal communication); 213 Andeans in Peru,<sup>7,8</sup> and 1,087 Tibetans in the Tibet Autonomous Region;<sup>9</sup> plot adapted from Moore et al.<sup>10</sup> B, In healthy adult males, the PaO<sub>2</sub>-induced rise in resting pulmonary artery pressure, as measured by cardiac catheterization, is least in 5 Tibetans at 3,600 m,<sup>11</sup> intermediate in 58 Andeans at 3,700–4,500 m,<sup>12,13</sup> and greatest in 28 European-derived residents at 3,100 m in Colorado;<sup>14</sup> plot adapted from Groves et al.<sup>11</sup> Measurements obtained by echocardiography in 76 Amhara males at 3,700 m in Ethiopia<sup>15</sup> at a PaO<sub>2</sub> estimated from the reported SaO<sub>2</sub> are similar to Andean values. Circles and lines represent mean ± SEM values. C, Lines show the altitude-associated decline in birth weight in a total of 305,935 infants from 15 studies in which birth weights were collected by the same investigator at both low (sea level) and high altitude (~3,600 m) and 5 in which birth weights collected at one altitude are compared to the group value at the other altitude;<sup>16,17-33</sup> plots adapted from Moore.<sup>34</sup> Circles depict mean ± SEM values for the ancestry groups shown. The 167 ± 36-g birth-weight reduction in Andeans or Tibetans (left; solid and dashed lines, respectively) is half (45%) the 373 ± 48-g reduction seen in European or Han high-altitude residents (right; solid and dashed lines, respectively). No differences in altitude-associated birth-weight reductions are apparent between Andeans and Tibetans when ancestry is controlled using surnames or gene markers, unlike a previous summary based on literature data in which population admixture was likely present.<sup>35</sup>

the neonatal period increased the reactivity of the pulmonary vasculature to hypoxia later in life.<sup>44,45</sup> Perinatal hypoxia is greater at high than at low altitudes, as demonstrated by the known reduction in birth weight resulting from slowed fetal growth,<sup>46</sup> Doppler indices of fetal hypoxia,<sup>47</sup> and lower postnatal arterial O<sub>2</sub> saturations (SaO<sub>2</sub>'s).<sup>48</sup> Perinatal hypoxia also varies among populations; the altitude-associated reductions in fetal growth are only half (45%) as great in Tibetans or Andeans as in Han or Europeans residing at the same altitudes (Fig. 1C), and Tibetans have considerably higher SaO<sub>2</sub>'s than Han over the first 4 months of postnatal life.<sup>49</sup>

In line with the concept of developmental origins of adult disease<sup>50</sup> and observations that perinatal hypoxia influences P<sub>pa</sub> later in life, we considered that altitude-associated disorders may be interrelated. That is, babies who were growth restricted in utero or otherwise hypoxic during the perinatal period may have lower SaO<sub>2</sub>'s as a result of impaired lung development or altered respiratory control. Such lower SaO<sub>2</sub>'s could then lead to higher P<sub>pa</sub> as the result of delayed or absent regression of muscularized pulmonary arterioles that, together with the factors contributing to lower SaO<sub>2</sub>, could render persons susceptible to developing CMS later in life.

But if lower birth weights predispose persons to pulmonary hypertension and CMS, what then accounts for the apparent Andean-Tibetan differences in  $P_{pa}$  and CMS prevalence, given that both groups are similarly protected from altitude-associated reductions in birth weight (Fig. 1)? We considered that differences in neonatal oxygenation could be involved. Specifically, we hypothesized (1) that  $SaO_2$  was highest in Tibetans, intermediate in Andeans, and lowest in Han or Europeans and (2) that improved oxygenation in Andeans, relative to that in Europeans, was accompanied by a greater postnatal decline in  $P_{pa}$ . While the long period over which CMS develops precluded assessing these infants' later-in-life susceptibility to CMS, consistent with such a possibility are our observations that persons with CMS weighed less at birth than healthy controls<sup>51</sup> and that young men with a preclinical form of CMS have more sleep-disordered breathing, nocturnal hypoxemia, and markers of oxidative stress consistent with impaired ventilatory control.<sup>52</sup> To test the first hypothesis, we measured  $SaO_2$  from 6–24 hours to 4–6 months of postnatal age in healthy Andeans born at 3,600 m and compared their data with those acquired previously in healthy Tibetan and Han infants at the same altitude. For the second hypothesis, we estimated  $P_{pa}$  as the systolic  $P_{pa}$  ( $P_{pa_{sys}}$ ), using quantitative echocardiography from 1 week to 4–6 months of age in healthy Andean and European infants at 3,600 m in Bolivia, and compared the time course of  $P_{pa_{sys}}$  changes to those seen in  $SaO_2$  across the same time span. Finally, to broaden the range of variation in  $SaO_2$  and  $P_{pa_{sys}}$ , we also studied a group of “sick” Bolivian babies at 3,600 m who had been diagnosed with fetal distress, newborn respiratory distress syndrome, and/or pulmonary hypertension and compared their  $SaO_2$  and  $P_{pa_{sys}}$  values to those observed in healthy neonates at the same altitude. We considered that such findings could help prompt additional longitudinal studies to determine whether the postnatal decline in  $P_{pa_{sys}}$  varies among these groups and correlates with population differences in the frequencies of pulmonary hypertension and CMS during adulthood.

## MATERIAL AND METHODS

### Ethical approval

All participants in the Bolivian studies gave their written informed consent to study procedures, which had been approved by the human subject review committees of the Colorado Multiple Institutional Review Board of the University of Colorado Denver (COMIRB) and the Colegio Médico, its Bolivian counterpart. The studies conducted in Lhasa, Tibet Autonomous Region, China, were

approved by the Tibet Institute of Medical Sciences and COMIRB.<sup>49</sup>

### Subjects

A total of 50 women and their infants were recruited in La Paz (3,600 m) and El Alto (4,100 m), Bolivia, from June 2001 to May 2003. They comprised (1) five (5) healthy European and 15 healthy Andean infants studied longitudinally from 1 week to 4–6 months of age, (2) 21 healthy Andean infants studied cross-sectionally from 6–24 hours to 1 month of age, and (3) nine (9) sick infants studied longitudinally from 1 week to 4–6 months of age. The healthy, longitudinal Andean mother-infant pairs were a subset of a larger group described previously.<sup>53</sup> Healthy babies were considered those born at term to Andean women without pregnancy complications (i.e., preeclampsia, gestational hypertension, or gestational or other forms of diabetes) and without a diagnosis of fetal distress, newborn respiratory distress, or neonatal pulmonary hypertension. The same criteria were applied to the healthy cross-sectionally studied babies, except that prenatal records were not available, and hence they were judged healthy on the basis of not requiring resuscitation at birth or not having respiratory distress, as judged by treatment with  $O_2$ . Sick babies were born to mothers of European, Andean, or mixed (*mestiza*) ancestry and had received a diagnosis of fetal distress as documented by meconium-stained amniotic fluid, newborn respiratory distress, or respiratory depression as evidenced by requiring  $O_2$  treatment at the time of birth and the first days of life or pulmonary hypertension documented by echocardiography.

The inclusion criterion for all Bolivian subjects was current residence at high altitude (La Paz, 3,600 m, or the neighboring city of El Alto, 4,100 m). The exclusion criterion was a lack of birth-weight or gestational-age data. Additional inclusion criteria for the healthy mother-infant pairs were the absence of maternal smoking, absence of known risk factors for maternal complications of pregnancy (chronic hypertension, gestational diabetes, diabetes), and absence of hypertensive complications in the current pregnancy (gestational hypertension, preeclampsia, eclampsia). All but one of the Bolivian women had lived at high altitude throughout the current pregnancy; that woman had moved there in her seventh month, but neither her nor her infant's characteristics differed from those of the other subjects, and hence their data are included here. The women were classified as Andean, European, or *mestiza* on the basis of self-identification, and classification was verified by reference to parental and grandparental surnames and, in the case of the 20 longi-

tudinally studied healthy Andeans or Europeans, a panel of 100 ancestry-informative genetic markers (AIMs), as described previously.<sup>16,54</sup> A woman was classified as Andean if she had 3 or more Andean parental surnames and/or at least 76% AIMs of indigenous American origin. European ancestry was assigned if 3 or more parental surnames were of non-Hispanic European origin, she was of European nationality, or the majority of her AIMs were of European or other low-altitude-population origin. Women not meeting these criteria were considered *mestiza*. Inclusion criteria for the 15 Tibetan and 15 Han infants studied in Lhasa, Tibet Autonomous Region, China (3,600 m), were that the births occurred consecutively in June 1991 and that the babies were healthy at the time of enrollment.<sup>49</sup>

### Study design

This was a prospective, observational cohort study in which pulse oximetry and echocardiographic measurements (with Doppler) were carried out at high altitude in Bolivia in healthy or sick babies born to women classified as Andean, European, or *mestiza*. Longitudinal studies in healthy European and Andean babies were carried out at 1 week and 1, 2–3, and 4–6 months of postnatal age, with the actual infant ages at the time of study being  $13 \pm 2$  and  $12 \pm 1$  days (1 week),  $39 \pm 4$  and  $40 \pm 2$  days (1 month),  $99 \pm 4$  and  $97 \pm 2$  days (2–3 months), and  $184 \pm 20$  and  $183 \pm 5$  days (4–6 months), respectively. Echocardiographic studies in the same infants were conducted at the same time points, with the actual infant ages at the time of study being  $9 \pm 2$  and  $15 \pm 2$  days (1 week),  $39 \pm 4$  and  $44 \pm 4$  days (1 month),  $97 \pm 2$  and  $103 \pm 4$  days (2–3 months), and  $186 \pm 16$  and  $187 \pm 5$  days (4–6 months), respectively. The actual infant ages for the longitudinal oxygenation and echocardiography studies of sick babies were  $13 \pm 2$  and  $5 \pm 4$  days (1 week),  $43 \pm 6$  and  $39 \pm 7$  days (1 month),  $97 \pm 6$  and  $101 \pm 3$  days (2–3 months), and  $174 \pm 17$  and  $187 \pm 12$  days (4–6 months), respectively. For purposes of comparing data for the Andeans to previously acquired Tibetan and Han data at earlier time points (6–24 and 24–48 hours), we conducted additional cross-sectional respiratory studies in healthy Andean babies at  $0.5 \pm 0.3$  days (6–24 hours,  $n = 4$ ),  $2.2 \pm 0.2$  days (24–48 hours,  $n = 4$ ),  $8 \pm 1$  days (1 week,  $n = 10$ ), and  $34 \pm 2$  days (1 month,  $n = 4$ ). The Tibetan and Han data were also acquired with a prospective, observational cohort design in which studies had been conducted at the same time points as in Bolivia, namely,  $0.6 \pm 0.2$  and  $0.7 \pm 0.2$  days (6–24 hours),  $1.6 \pm 0.3$  and  $1.6 \pm 0.3$  days (24–48 hours),  $7 \pm 1$  and  $7 \pm 1$  days (1 week),  $30 \pm 6$  and  $30 \pm 3$  days (1 month),  $55 \pm 4$  and  $59 \pm 53$  days (2–3 months), and  $120 \pm 7$  and  $123 \pm 4$  days (4–6 months), respectively.<sup>49</sup>

### Variables and instrumentation

Information on maternal age, self-identified ancestry, and residential and reproductive history was collected by questionnaire or from medical records. Infant birth weight, gestational age, sex, whether the babies were small for gestational age as judged by being less than the tenth centile of sea-level values for a given gestational age and sex,<sup>55</sup> Apgar scores, and postnatal complications were obtained from delivery and newborn records. Babies born before 37 weeks gestation were considered preterm.

Arterial oxygenation studies were conducted in Bolivia over a 2–3-hour period that included quiet and active sleep, as well as periods of wakefulness and feeding, and used pulse oximetry (Nellcor N-200, Hayward, CA) in the clinic, the hospital, or the subject's home, with the probe placed on the lateral aspect of the foot. The  $\text{SaO}_2$  was recorded in beat-to-beat mode and reported as the average of the stable values recorded. To simplify reporting and group comparisons, the pulse oximetry data from the cross-sectionally studied Andeans were combined with those from the longitudinally studied Andean infants. The same protocol was followed in Tibet, except that a Biox 3740 oximeter (Ohmeda, Louisville, CO) was employed.

Echocardiography was performed with an ATL HDI 3000 equipped with a 5–7.5-MHz sectorial transducer, continuous and pulsed-wave Doppler, and color Doppler flow (Royal Philips Electronics, Amsterdam). All studies were carried out in the Clínica del Sur (Southern Clinic) in La Paz, Bolivia. The measurements were made in a warm environment in the quiet, awake state without the use of sedation and with Doppler measurements across the pulmonary valve being made early in the study to avoid elevation of estimated pressures because of agitation. The initial study included detailed documentation of anatomy, namely, great-vessel relationships, pulmonary venous return, interatrial and interventricular communications, and the presence of a patent ductus arteriosus. All studies included a review of cardiac anatomy; morphometric measurements (ventricular dimensions, septal and free-wall thicknesses, valve dimensions); ventricular shortening and ejection fractions; Doppler flow measurements across the aortic, mitral, tricuspid, and pulmonary valves; and presence of patent foramen ovale (PFO). Right ventricular systolic intervals (pre-ejection period [PEP], acceleration time [AT], and ejection time [ET]) were measured, as well as the R-R interval for calculating heart rate (HR). Measurements were made on a minimum of 3 complete waveforms; the maximal and minimal values (reflecting respiratory variation) were then averaged for the reported result. In healthy infants, systolic time intervals were incorporated into the regression equation of Li<sup>56</sup> to

estimate  $P_{pa_{sys}}$ , as previously published in studies of infants at high altitude.<sup>57</sup> When tricuspid regurgitation was present and waveforms were complete, right ventricular pressures were estimated by quantification of the tricuspid regurgitation jet with the modified Bernoulli equation.<sup>58</sup> When a ductus arteriosus was present, the ductal shunt size was estimated, the direction of flow was documented, and the aortopulmonary pressure difference was derived from maximal velocity detectable in the ductal-flow jet.<sup>59</sup>

### Statistics

Data are reported as the mean  $\pm$  standard error of the mean in the text, tables, and figures or as the mean and 95% confidence intervals for proportions, except when indicated otherwise. Comparisons between groups at a single time were carried out with one-way analysis of variance (ANOVA), with Sidak multiple comparisons to identify the source of the differences observed. The effects of time, ancestry, and their interaction on  $SaO_2$  or echocardiographic characteristics were assessed by means of 2-way ANOVA with Tukey's multiple comparisons. Differences in proportions between groups were tested with  $\chi^2$ . All statistics were conducted in GraphPad software Prism 6.0 (GraphPad, La Jolla, CA). Comparisons are reported as significant when the 2-tailed  $P < 0.05$  and as trends when  $0.05 < P < 0.10$ .

## RESULTS

### Maternal and infant characteristics

By study design, the mothers of the healthy Andean and Tibetan babies had more often been born at high altitude and had lived there longer than the healthy Han or European women (Table 1). Values were intermediate in the mothers of sick babies, with fewer being born at high altitude than in the healthy Andean group but more than was the case for healthy Europeans. The mothers of the sick babies had lived at high altitude for as long as the healthy Andeans and longer than the healthy Europeans. Maternal age was similar in all groups, as was parity, with the exception of higher parity in the mothers of the longitudinally studied Andean babies. By study design, none of the healthy mothers experienced pregnancy complications, but most of the sick babies were born to women with complications of pregnancy or labor (Table 2). Neither parity nor maternal age was related to infant birth weight,  $SaO_2$ , or  $P_{pa_{sys}}$  in any group.

Birth weights and gestational ages were similar in all the healthy groups, although absolute birth weights were somewhat lower in the Han (Table 1). Sick babies were more often born prematurely. Approximately equal sex ratios were seen in Tibetans and Han, but there were more

male than female babies in the healthy European and sick-baby groups, and more females in the healthy, longitudinal, and cross-sectional Andean groups. Apgar scores were similar among the various groups, with the exception of the sick babies, whose values were somewhat lower. Approximately half the sick babies required supplemental  $O_2$  (Table 2).

### Pulse oximetry studies

The  $SaO_2$  showed consistent variations according to infant age and activity, and values were below sea-level norms at all study times (Figs. 2–4). Healthy Tibetans, Han, and Andeans had their highest  $SaO_2$ 's 6–24 hours after birth (Fig. 2). Across all study times and healthy groups,  $SaO_2$  was highest in the awake state ( $88.7\% \pm 1.8\%$  [standard deviation]), intermediate while feeding and during active sleep ( $86.4\% \pm 2.8\%$  and  $86.4\% \pm 3.5\%$ ), and lowest during quiet sleep ( $85.9\% \pm 3.5\%$ ).

When healthy Tibetans, Han, and Andeans were compared across all study times,  $SaO_2$  was higher in the Tibetan than the Han or Andean babies during wakefulness, active sleep, and quiet sleep (Fig. 2). The  $SaO_2$  while feeding was higher in the Tibetans or Andeans than in the Han. Lower  $SaO_2$  values in the Han babies were chiefly the result of greater age-related declines, whereas the Tibetan-Andean differences were due to small but consistent differences at each study time.

There were no group differences in  $SaO_2$  between the healthy Andean, healthy European, and sick babies from 1 week to 4–6 months of age (Fig. 3). There was, however, significant interaction between the effects of group and time, which stemmed largely from the lower and more variable values seen in the sick babies. The European and sick babies had a greater age-related rise in  $SaO_2$  during active sleep and more variable values during quiet sleep than the Andean babies.

### Echocardiography studies

All babies had structurally normal hearts on initial exam. Several minor anatomic variants without physiologic significance were identified, such as supernumerary chordae tendinae, Chiari network, persistent left superior vena cava draining to coronary sinus, and mild branch pulmonary stenosis. One small apical ventricular septal defect became apparent on the final study, after estimated right-sided pressures had declined from 48 to 29 mmHg, presumably allowing visualization of a left-to-right shunt across the defect. None of the healthy infants had a measurable tricuspid regurgitation jet. Thus, right heart pressures were estimated with systolic time intervals of the pulmonary valve.



Table 1. Characteristics of Han, Tibetan, Andean, and European healthy and sick mother-infant pairs studied at high altitude (3,600 m)

	Healthy					Sick Bolivian babies	P, all groups
	Han longitudinal	Tibetan longitudinal	European longitudinal	Andean longitudinal	Andean cross-sectional		
<i>n</i>	15	15	5	15	21	9	
Mothers							
Born at HA, <i>n</i>	0	15	0	14	17	3	...
Time at HA, yr	2 ± 1 <sup>A</sup>	25 ± 1 <sup>B</sup>	9 ± 4 <sup>A</sup>	22 ± 2 <sup>B</sup>	24 ± 2 <sup>B</sup>	21 ± 5 <sup>B</sup>	<0.0001
Age range, yr	20–28	20–30	31–40	18–38	19–40	23–35	...
Parity, <i>n</i>	1.3 ± 0.1 <sup>B</sup>	1.3 ± 0.1 <sup>B</sup>	2.2 ± 0.6 <sup>A</sup>	3.5 ± 0.5 <sup>C</sup>	2.3 ± 0.3 <sup>A</sup>	1.8 ± 0.4 <sup>B</sup>	<0.0001
Infants							
Birth weight, gm	2,773 ± 92	3,067 ± 107	3,020 ± 245	3,114 ± 102	3,074 ± 109	2,834 ± 249	NS
Gestational age, wk	38.9 ± 0.4 <sup>A</sup>	38.9 ± 0.4 <sup>A</sup>	38.9 ± 0.4 <sup>A</sup>	39.6 ± 0.4 <sup>A</sup>	38.7 ± 0.3 <sup>A</sup>	36.9 ± 1.2 <sup>B</sup>	<0.03
Sex, F/M	8/7	9/6	1/4	10/5	12/9	3/6	...
Apgar score, 1 min	8	8	8	8	7	6	...
Apgar score, 5 min	10	9	10	9	8	8	...

Note: Data are counts or mean ± SEM. HA: high altitude (>2,500 m); NS: not significant. Different letters signify differences between groups at  $P < 0.05$ .

Table 2. Clinical characteristics of sick babies

ID	Ancestry (yr at HA)	BW, g	Sex	GA, wk	Antecedent factors	Presenting signs	Status at discharge	Clinical course	P <sub>pa</sub> , mmHg				PFO present, age
									1 wk	1 mo	3 mo	6 mo	
20	Eur (1)	2,310	M	33–34	Placenta previa, hemorrhage	RDS, surfactant deficiency	Medical evac on O <sub>2</sub> ; return on RA at 3 mo	Deterioration at 36 hr clinically consistent with PPHN	NA	NA	38	24	Never
55	Andean (23)	2,000	F	30	PPROM, chorioamnionitis	RDS, pneumonia	RA	Malnutrition, hy- pertonia, enlarg- ing OFC, SHAPH; died at 5 mo	NA	NA	68	NA	Never
64	Andean (23)	2,430	F	37	Meconium- stained amn fluid	Polycythemia, jaundice, PPHN	On O <sub>2</sub>	NA	53	NA	NA	NA	1 wk
115	Andean (30)	1,730	M	35	Grade 3+ meconium	RDS	RA	O <sub>2</sub> for 6 hr, incubator for 40 hr	NA	NA	NA	NA	NA
201	Mestiza (NA)	3,960	M	38	Meconium- stained amn fluid	Retained fetal lung fluid, PPHN	RA	O <sub>2</sub> for 3 d	61	42	42	34	3 d
202	Mestiza (NA)	3,430	M	40	None	Polycythemia, hyperviscosity; PPHN, jaundice	Home O <sub>2</sub> ; SaO <sub>2</sub> of 55% at 6 d, poor feeding; rehospitalized	Continuous home O <sub>2</sub> at 2–6 wk; descent to 2,500 m; return to 3,000 m	49	31	52	38	All times
203	Mestiza (29)	3,050	M	40	Prolonged labor; perinatal depression	Hypoxic, bradycardic at 6 hr; PPHN; PFO, PDA, perimembranous VSD < 2 mm (closed by 6 wk)	Home O <sub>2</sub> , SaO <sub>2</sub> of 75% at 2 wk on RA	NA	48	45	29	38	1 wk, 1 mo, 3 mo
204	Eur (NA)	3,400	F	NA	NA	NA	RA	Healthy at 8-d exam	NA	49	NA	NA	1 mo
205	Mestiza (NA)	3,200	M	NA	NA	Jaundice	RA	URI at 1 mo, otherwise normal	47	54	NA	NA	1 wk, 1 mo

Note: amn: amniotic; anc: mother's ancestry; BW: birth weight; evac: evacuation; Eur: European; GA: gestational age; HA: high altitude (>2,500 m); NA: not available; OFC: occipital frontal circumference; PDA: patent ductus arteriosus; PFO: patent foramen ovale; P<sub>pa</sub>: pulmonary artery pressure; P<sub>pa<sub>sys</sub></sub>: systolic P<sub>pa</sub>; PPHN: persistent pulmonary hypertension of the newborn; PPRM: preterm rupture of the membranes; RA: room air; RDS: respiratory distress syndrome; SHAPH: symptomatic high-altitude pulmonary hypertension; SaO<sub>2</sub>: arterial O<sub>2</sub> saturation; URI: upper respiratory infection; VSD: ventricular septal defect.

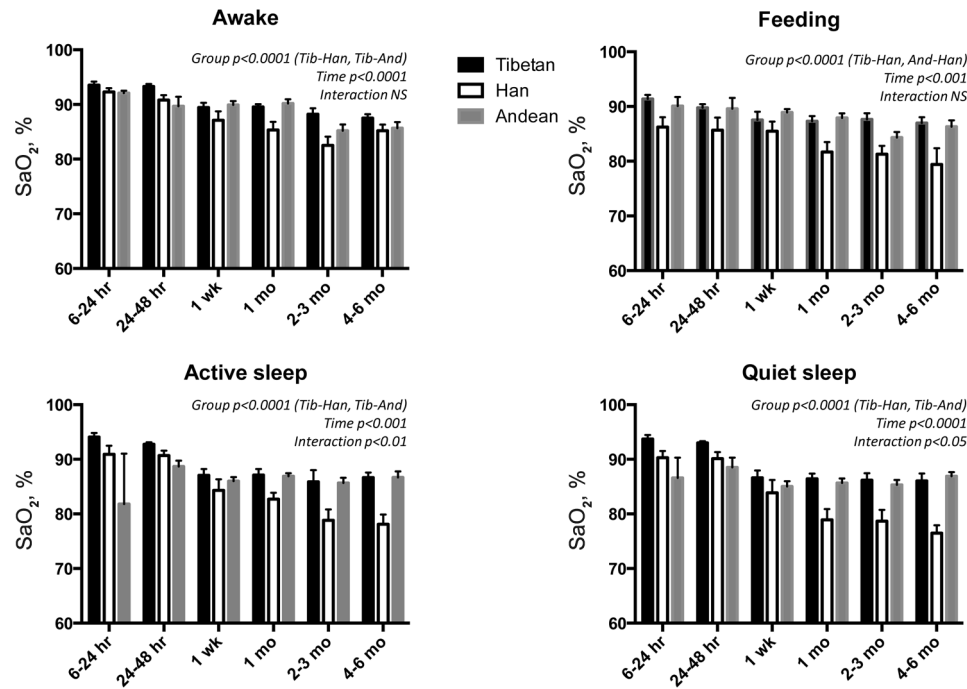


Figure 2. Mean  $\pm$  SEM arterial O<sub>2</sub> saturation (SaO<sub>2</sub>) in healthy Tibetan (Tib;  $n = 15$ ), Han ( $n = 15$ ), and Andean (And;  $n = 36$ ) babies at 3,600 m declined with advancing age and during quiet sleep relative to that in the other activity states. Across time, values were higher in Tibetan than in Han babies in all states<sup>49</sup> and higher than those in the Andeans in the awake and active- and quiet-sleep states. Tibetan and Andean values during feeding were similar and higher than those observed in Han infants. NS: not significant.

The healthy European and Andean babies had similar PEP, AT, ET, AT/ET, PEP/ET, HR, and  $P_{pa_{sys}}$  (Table 3). There was a gradual rise in AT, ET, and AT/ET and a decline in PEP/ET consistent with a decrease in estimated right heart pressure over the period of study. The  $P_{pa_{sys}}$  at 4–6 months remained above the sea-level range.<sup>60</sup> Most infants had a persistent PFO, with the prevalence of PFO declining such that at 4–6 months, approximately half the healthy infants had PFO. No healthy infants had a persistently patent ductus arteriosus.

Compared with healthy babies, the sick babies had lower PEP, lower AT, a trend toward lower AT/ET, higher HR, and higher  $P_{pa_{sys}}$  (Table 3). All but one survived. The  $P_{pa_{sys}}$  declined to values seen in healthy Andean or European babies at 4–6 months.

### Changes in $P_{pa}$ with respect to SaO<sub>2</sub> and clinical course

The  $P_{pa_{sys}}$  gradually declined across the 4–6-month study period in the healthy European and Andean babies despite the age-related decline in SaO<sub>2</sub> (Fig. 4). This pattern contrasted with that observed in the sick babies, in whom SaO<sub>2</sub> was lowest and  $P_{pa_{sys}}$  highest at 1 week, with  $P_{pa_{sys}}$  then declining at the higher SaO<sub>2</sub>'s seen at advancing ages. The  $P_{pa_{sys}}$ , PEP/ET, and SaO<sub>2</sub> values were similar

in the infants with PFO and those without PFO (data not shown). All the sick babies had PFO at the first exam, with closure being achieved in all cases by 4–6 months of age (Table 3).

### DISCUSSION

At 3,600 m, Tibetans had higher levels of neonatal oxygenation than Andean or Han babies while awake or asleep, and Andean values were higher than Han values during feeding, but neither SaO<sub>2</sub> nor  $P_{pa_{sys}}$  differed in European versus Andean babies. Newborns with fetal distress, neonatal respiratory distress, or pulmonary hypertension had lower SaO<sub>2</sub> at initial exam and higher  $P_{pa_{sys}}$  at 3,600 m in Bolivia, supporting the ability of our study techniques to detect elevated  $P_{pa_{sys}}$ . The  $P_{pa_{sys}}$  and the frequency of PFO declined in the healthy or sick European or Andean babies with advancing age, but  $P_{pa_{sys}}$  remained above the normal sea-level range, and half the infants at 4–6 months still had PFO. Thus, the data were consistent with our hypothesis that Tibetans maintained better neonatal oxygenation than Andeans, which may contribute to their protection from high-altitude pulmonary hypertension and CMS later in life. But counter to our expectation that Andean values would differ from Europeans', neither was SaO<sub>2</sub> higher nor  $P_{pa_{sys}}$  lower in the Andean than in the



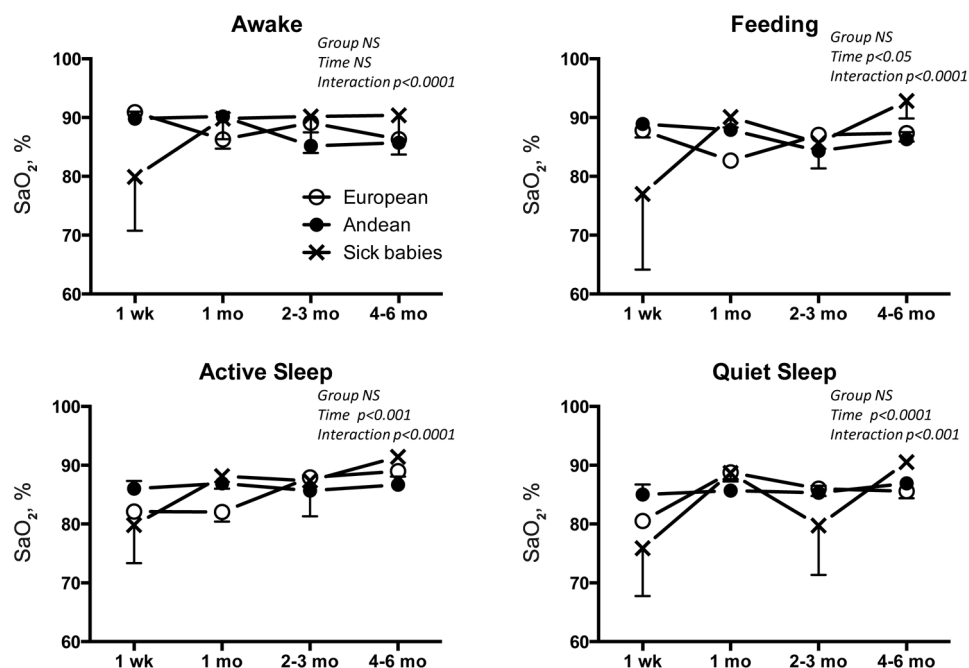


Figure 3. Mean  $\pm$  SEM arterial O<sub>2</sub> saturations (SaO<sub>2</sub>) are shown for 15 healthy Andeans, 5 healthy Europeans, and 9 babies diagnosed with fetal distress, newborn respiratory distress, or pulmonary hypertension (“sick babies”) of Andean, European, or *mestiza* descent who were studied longitudinally at 3,600 m. While values did not vary by group, the effects of group interacted with those of time in each activity as a result of greater desaturation in the sick babies, especially at the earliest time point, and the greater age-related rise and more variable values during quiet sleep in European than in Andean babies. NS: not significant.

European infants at 3,600 m. To determine whether post-natal declines in  $P_{pa}$  differ in Andeans versus Tibetans or between long- and short-resident groups, future studies are required in which  $P_{pa}$  and SaO<sub>2</sub> are assessed longitudinally with well-controlled methodologies in larger numbers of infants from all 4 populations (Andean, European, Han, and Tibetan).

Strengths of our study were the ability to measure SaO<sub>2</sub> in each of the 4 recognized activity states and to study it, as well as  $P_{pa_{sys}}$ , longitudinally in healthy Andean and European babies of well-defined maternal ancestry. Although the Tibet and Bolivia studies were performed approximately 10 years apart, the two sites are located at the same altitude (3,600 m), the same protocols were followed, and the same investigator (SN) conducted both studies. A single operator conducted all echocardiography exams in Bolivia (MPA-M), making it unlikely that differences in study techniques influenced the results obtained. However, our study also suffered from various weaknesses. We were limited by the small size of our European sample, because of the small numbers of pregnant Europeans living in La Paz during the period of study. Having to recruit subjects who delivered at multiple sites also prevented us from conducting measurements in the longitudinal groups at the earliest (6–24- and 24–48-hour) time points. We were able to overcome this for the Andeans by

performing additional cross-sectional studies, but this necessitated forgoing the additional sensitivity provided by a repeated-measures design. Different oximeters were employed in the two studies, but the error in their precision is less than 3% for SaO<sub>2</sub>'s greater than 83%,<sup>61</sup> which is generally the range present in our study. Group differences were chiefly due to different patterns of change with time and therefore were not likely affected by mean differences between instruments. While we measured SaO<sub>2</sub> in each activity,  $P_{pa_{sys}}$  could be assessed only while the infants were awake, which is when SaO<sub>2</sub> was highest, and therefore our values may have underestimated  $P_{pa_{sys}}$  at other times. In addition, while the babies born to the larger group of European women participating in the maternal O<sub>2</sub> transport studies weighed less than the Andeans,<sup>16</sup> birth weights did not differ in the subsets presented here, thus limiting our ability to assess the effects of fetal growth on neonatal oxygenation. Ancestry was based on maternal self-identification, parental surnames, and AIMs in the longitudinal group; however, this meant that paternal ancestry was not controlled. We were also not able to measure SaO<sub>2</sub> and  $P_{pa_{sys}}$  at all study times or conditions because of clinical considerations or inability to achieve the desired activity state at the time of study. Finally, for ethical considerations, we used indirect echocardiographic measures rather than cardiac catheterization to document

Table 3. Echocardiography studies in healthy and sick European and Andean babies at 3,600 m in La Paz, Bolivia

Measure, age	European healthy	Andean healthy	Sick babies	Group <i>P</i>	Time <i>P</i>	Interaction <i>P</i>
PEP, m				0.06 (NS) <sup>a</sup>	0.08 (NS)	<0.05
1 wk	46 ± 4 (2)	52 ± 2 (15)	35 (1)			
1 mo	42 ± 3 (4)	46 ± 2 (14)	42 ± 2 (6)			
2–3 mo	42 ± 3 (4)	43 ± 1 (15)	47 ± 2 (5)			
4–6 mo	38 ± 3 (3)	38 ± 1 (14)	41 ± 5 (4)			
AT, ms				<0.01 <sup>a</sup>	<0.0001	NS
1 wk	53 ± 2 (3)	60 ± 3 (15)	40 (1)			
1 mo	59 ± 3 (5)	60 ± 2 (14)	54 ± 4 (6)			
2–3 mo	68 ± 4 (5)	67 ± 3 (15)	58 ± 6 (5)			
4–6 mo	71 ± 3 (4)	73 ± 2 (15)	69 ± 3 (4)			
ET, ms				NS	<0.0001	NS
1 wk	196 ± 16 (3)	228 ± 6 (15)	182 (1)			
1 mo	197 ± 7 (5)	220 ± 7 (14)	214 ± 8 (6)			
2–3 mo	232 ± 9 (5)	241 ± 7 (15)	230 ± 3 (5)			
4–6 mo	236 ± 15 (4)	251 ± 6 (15)	256 ± 20 (4)			
AT/ET				0.08 (NS)	NS	NS
1 wk	0.27 ± 0.01 (3)	0.27 ± 0.02 (14)	0.22 (1)			
1 mo	0.28 ± 0.03 (5)	0.28 ± 0.02 (14)	0.25 ± 0.02 (6)			
2–3 mo	0.29 ± 0.02 (5)	0.28 ± 0.01 (15)	0.25 ± 0.02 (5)			
4–6 mo	0.30 ± 0.03 (4)	0.29 ± 0.01 (15)	0.27 ± 0.01 (4)			
PEP/ET				NS	<0.001	NS
1 wk	0.24 ± 0.03 (3)	0.23 ± 0.01 (15)	0.19 (1)			
1 mo	0.22 ± 0.02 (5)	0.21 ± 0.01 (14)	0.19 ± 0.00 (6)			
2–3 mo	0.18 ± 0.01 (5)	0.18 ± 0.01 (15)	0.21 ± 0.02 (5)			
4–6 mo	0.16 ± 0.02 (4)	0.15 ± 0.01 (15)	0.16 ± 0.01 (4)			
HR, bpm				<0.05 <sup>a</sup>	<0.05	NS
1 wk	149 ± 12 (3)	147 ± 4 (15)	176 (1)			
1 mo	161 ± 10 (5)	146 ± 3 (14)	156 ± 6 (6)			
2–3 mo	140 ± 11 (5)	137 ± 5 (15)	154 ± 14 (5)			
4–6 mo	146 ± 6 (3)	131 ± 3 (15)	144 ± 11 (4)			
P <sub>pa<sub>sys</sub></sub> , mmHg				<0.05 <sup>a</sup>	<0.0001	NS
1 wk	47 ± 5 (3)	48 ± 2 (15)	51 ± 3 (4)			
1 mo	39 ± 2 (5)	41 ± 1 (14)	42 ± 4 (6)			
2–3 mo	33 ± 2 (5)	35 ± 2 (15)	46 ± 7 (5)			
4–6 mo	29 ± 3 (4)	28 ± 1 (15)	32 ± 3 (4)			
PFO, %						
1 wk	33 [0, 90] (3)	88 [62, 98] (15)	100 [40, 100] (4)	NS		
1 mo	80 [28, 99] (5)	71 [42, 92] (14)	83 [36, 99] (6)	NS		
2–3 mo	60 [15, 95] (5)	73 [45, 92] (15)	60 [15, 95] (5)	NS		
4–6 mo	50 [7, 93] (4)	47 [21, 73] (15)	0 [0, 60] (4)	NS		

Note: Data are mean ± SEM or mean [95% confidence interval], with sample sizes in parentheses. AT: acceleration time, ET: ejection time, HR: heart rate, NS: not significant; PEP: pre-ejection period, PFO: patent foramen ovale, P<sub>pa<sub>sys</sub></sub>: estimated systolic pulmonary artery pressure.

<sup>a</sup> Pairwise differences are present between the sick babies and the healthy European or Andean groups.

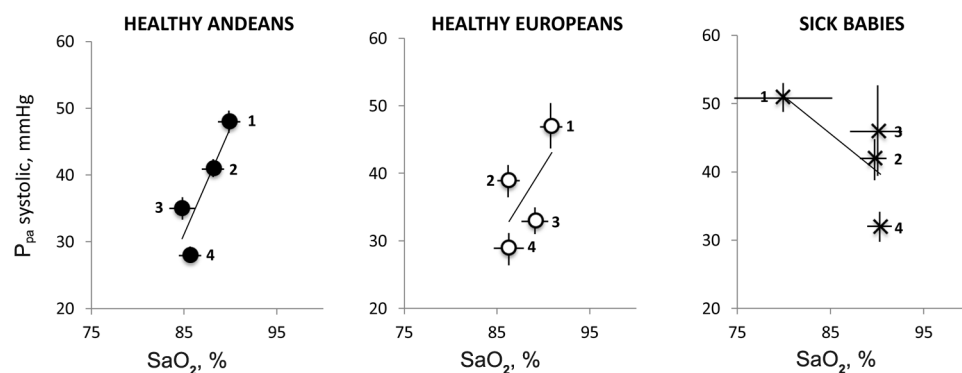


Figure 4. Mean  $\pm$  SEM values for estimated pulmonary artery systolic pressure ( $P_{pa_{sys}}$ ) in relation to arterial  $O_2$  saturation ( $SaO_2$ ) are compared at 1 week (1), 1 month (2), 2–3 months (3), and 4–6 months (4) of postnatal age in babies at 3,600 m in La Paz, Bolivia.  $P_{pa_{sys}}$  declined as  $SaO_2$  rose in healthy Andean ( $n = 15$ ) or healthy European ( $n = 5$ ) babies but showed a different relationship in sick babies ( $n = 9$ ), with the highest  $P_{pa_{sys}}$  being seen at the lowest  $SaO_2$  at 1 week of age.

the  $P_{pa}$ , which is in keeping with all prior high-altitude infant studies but the Gamboa report.<sup>48,57,62,63</sup>

Several human and experimental animal studies demonstrate that postnatal normalization of  $P_{pa}$  is affected by residence at high altitude. At 4,540 m, the decline in  $P_{pa}$  is prolonged or even fails to occur in some infants.<sup>63</sup> Lambs gestated at 3,600 or 3,800 m also have elevated  $P_{pa}$ , which persists even after return to sea level.<sup>64,65</sup> Greater hypoxia in utero or during the neonatal period also increases the reactivity of the pulmonary vasculature to hypoxia later in life in human and experimental animal studies.<sup>44,45,64</sup> At 3,600 m in Bolivia, right ventricular wall thickness, an indicator of the shift from right to left ventricular predominance with declining  $P_{pa}$ , showed essentially no decrease during the 12 months after birth, whereas values at low altitude (400 m) decreased during the first month of life to a dimension that remained unchanged through the rest of infancy.<sup>62</sup> Despite supplemental  $O_2$  during the first 6 hours of postnatal life in Leadville, Colorado (3,100 m), echocardiographic  $P_{pa}$  indices were normal to moderately elevated in healthy term infants and did not fully normalize until 2–4 months of age.<sup>48</sup> Even healthy controls born at 2,240 m in Mexico City manifested a moderately elevated PEP/AT ratio at 15 and 30 days of age ( $0.90 \pm 0.2$  and  $0.86 \pm 0.24$ , respectively) and an elevated  $P_{pa_{sys}}$  ( $48 \pm 11$  and  $47 \pm 13$  mmHg, respectively).<sup>57</sup> There is also a greater frequency of symptomatic pulmonary hypertension (subacute infantile mountain sickness), persistent patent ductus arteriosus, and PFO later in childhood at high than at low altitude.<sup>37,41,66-72</sup>

While several reports indicate that population ancestry influences resting  $P_{pa}$  and the pulmonary vasoconstrictor response to hypoxia during adulthood (Fig. 1B),

little is known concerning population variation in  $P_{pa}$  during infancy or childhood. To our knowledge, the  $P_{pa}$  response to hypoxia has not been measured in Han, but the greater right-axis deviation observed in adult male Han relative to Tibetan residents at 3,600 m<sup>73</sup> is consistent with the likelihood of greater pulmonary hypertension. Supporting our catheterization data, recent echocardiographic studies show lower  $P_{pa}$  values in Sherpa highlanders than in acclimatized lowland controls.<sup>74</sup> Higher values in Amhara highlanders than in sea-level controls have also been observed in Ethiopia,<sup>15</sup> suggesting that the Tibetans' low  $P_{pa}$  at high altitude may be unique. The factors responsible for the Tibetans' relative protection from pulmonary hypertension at high altitude are unknown. Greater nitric oxide (NO) production and/or NO transfer has been suggested on the basis of higher levels of exhaled NO in Tibetans than in lowlanders in the United States.<sup>75</sup> This may be a body-wide response, since greater lower-limb vasodilator response to ischemia<sup>76</sup> and greater forearm blood flow in association with higher circulating NO products<sup>77</sup> have been observed in Sherpa or Tibetans than in sea-level residents of the United States or acclimatized lowlanders. Population differences in ventilatory sensitivity to hypoxia and hence arterial oxygenation are also likely involved, since Tibetans resemble acclimatized lowlanders by having a brisk ventilatory response to hypoxia, higher levels of alveolar ventilation,<sup>78</sup> and higher  $SaO_2$  in most, but not all, studies (see Gilbert-Kawai et al.<sup>79</sup> for a recent review).

We suggest that not only pulmonary hypertension but also CMS has antecedents during perinatal life. Babies born to pregnancies complicated by preeclampsia and/or who weighed less at birth are known to be at increased

susceptibility to a range of cardiovascular and pulmonary diseases later in life.<sup>50</sup> It has been shown that  $P_{pa}$ 's following ascent to high altitude are greater in young adult men who experienced neonatal hypoxia<sup>45</sup> or were born to mothers with mild to moderate preeclampsia,<sup>80</sup> relative to controls, and that those men were therefore likely subjected to greater hypoxia in utero. We have shown that adult men with CMS had lower birth weights,<sup>51</sup> and Julian et al.<sup>52</sup> recently reported that young men with a preclinical form of CMS had impaired breathing during sleep, compared with age-matched controls. Collectively, these studies are consistent with the concept that reduced fetal growth at high altitude impairs the maturation of pulmonary vascular and respiratory control systems, which in turn could predispose persons to pulmonary hypertension and CMS later in life.

Since  $P_{pa}$  estimates are not available from healthy Tibetan or Han babies at high altitude, we considered that it would be useful to compare their neonatal oxygenation with that of Andean and European infants in whom both  $SaO_2$  and  $P_{pa_{sys}}$  measurements could be obtained. Similar to previous reports, we found that  $SaO_2$  in Andeans and Europeans had a postnatal course different from that observed at low altitude. Whereas  $SaO_2$  remains relatively stable at sea level and gradually increases from 92%–93% at 24–48 hours to 93%–94% at 1–3 months at 1,600 m,<sup>81</sup> it was highest in the Andean and European babies at 3,600 m during the first week of postnatal life and then declined across the next several months. This age-related decline is similar to what has been reported at 3,100 m in Colorado<sup>48</sup> and has been attributed to the onset of periodic breathing, which is more prominent at high than at low altitude and produces greater cycles of desaturation.<sup>70,82–84</sup>

As reported previously, Tibetans have higher  $SaO_2$  than Han infants from the time of birth,<sup>49</sup> indicating a better neonatal cardiopulmonary transition. Consistent with our hypothesis, we found that Tibetans had higher  $SaO_2$  than Andeans in the awake and active- and quiet-sleep states (Fig. 2). The 2%–5% point differences in  $SaO_2$  between Tibetans and Han during wakefulness and the 2% to 8%–9% point differences during the other activity states were greater than the 1%–3% point differences between the Tibetans and Andeans, suggesting a cardiopulmonary transition in Andeans that is intermediate between those of Tibetans and Han high-altitude residents.

However, counter to our hypothesis, neonatal oxygenation did not differ between the healthy Andean and European groups (Fig. 3). Nor did our echocardiographic studies show any differences between healthy Andean and European babies in right heart function or  $P_{pa_{sys}}$  at any time

(Table 3). The absence of 6–24- and 24–48-hour  $SaO_2$  data in the Europeans may have limited our ability to detect ancestry-group differences at the earliest time points, when  $P_{pa_{sys}}$  is the highest. While echocardiographic estimates are less sensitive than direct measurements, we were able to detect elevated  $P_{pa_{sys}}$  in the small number of babies diagnosed with fetal distress, neonatal respiratory distress, or symptomatic pulmonary hypertension, indicating that our methods were sufficiently sensitive to detect pulmonary hypertension. Elevated  $P_{pa}$  values, as measured by the systolic right ventricular-to-right atrial pressure gradient, have been reported in larger numbers of European ( $n = 77$ ) versus Andean (Aymara,  $n = 200$ ) 2–17-year-olds living at 3,600 m in Bolivia. While there was considerable overlap between groups, Andean values averaged 33% lower than those in the Europeans, and both groups had values higher than those of 29 low-altitude (400 m) children in Bolivia.<sup>43</sup> The small number of European infants available for our studies and the difficulty in acquiring data in all activity states likely limited our power to detect small differences between the Andean and European groups during the first 6 months of life. Andean admixture in the European babies may also have played a role, given that while all the mothers were of low-altitude, European ancestry, 3 of the 5 fathers were of mixed low-altitude (European, Guarani, Mexican) and Andean descent. Alternatively,  $P_{pa}$  differences between Andeans and Europeans may develop after the 4–6-month period evaluated here.

In summary, the implications of our study findings are threefold. First, population differences in neonatal oxygenation exist such that Tibetans, Andeans, and healthy Europeans did not show the progressive decrease in  $SaO_2$  during sleep that characterized Han in our earlier study.<sup>49</sup> Specifically, mean saturations remained at or above 85% in these groups, whereas the Han values dropped into the 70% range and even lower in some cases. In comparisons of high-altitude residents, neonatal oxygenation was highest in Tibetans, intermediate in Andeans and Europeans, and lowest in Han. Values were similar in healthy Andean and European babies, but additional studies in larger numbers of European infants at high altitude are required to confirm (or refute) this observation. Particularly critical will be longitudinal studies of  $P_{pa}$  and  $SaO_2$  using well-controlled methodologies to determine whether the postnatal decline in  $P_{pa}$  differs between Andeans and Tibetans or between long- and short-resident groups. Second, while some babies have clinical disease, many show considerable resilience and succeed in maintaining or improving their  $SaO_2$  and lowering their  $P_{pa_{sys}}$  through infancy

despite having saturations consistently lower than those of babies at sea level. Third, perinatal life and infancy are not only times when the effects of altitude-related hypoxia are particularly acute but also periods during which developmental factors can have consequences for disease susceptibility later in life. Thus, while lower birth weights, pulmonary hypertension, and CMS have been viewed as distinct entities in prior studies at high altitude, they may be underlain by a common sequence of developmental responses to high-altitude hypoxia that begins during perinatal life and infancy. Population comparisons of physiological responses to the challenges of hypoxia during these critical periods can provide insight into the mechanisms responsible for adaptation and can yield strategies for the treatment or prevention of these hypoxia diseases or disorders that will benefit persons at any altitude and especially the 140 million residents at high altitude worldwide.

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## REFERENCES

- De Reuck AVS, O'Connor M, eds. Problems of pulmonary circulation: in honour of Prof. G. Liljestrand. Ciba Foundation Study Group, no. 8. Boston: Little, Brown, 1961.
- Grover RF. Pulmonary hypertension: the price of high living. In: Wagner WW Jr., Weir EK, eds. The Pulmonary Circulation and Gas Exchange. Armonk, NY: Futura, 1994: 317–342.
- Murray F, Insel PA, Yuan JX. Role of O<sub>2</sub>-sensitive K<sup>+</sup> and Ca<sup>2+</sup> channels in the regulation of the pulmonary circulation: potential role of caveolae and implications for high altitude pulmonary edema. *Respir Physiol Neurobiol* 2006; 151(2–3):192–208.
- Peñaloza D, Arias-Stella J. The heart and pulmonary circulation at high altitudes: healthy highlanders and chronic mountain sickness. *Circulation* 2007;115(9):1132–1146.
- León-Velarde F, Maggiorini M, Reeves JT, Aldashev A, Asmus I, Bernardi L, Ge RL, et al. Consensus statement on chronic and subacute high altitude diseases. *High Alt Med Biol* 2005;6(2):147–157.
- Wu T, Die W, Huo K. An epidemiological study on high altitude disease in Qinghai-Xizang (Tibet) Plateau [in Chinese]. *Chin J Epidemiol* 1987;8:65–69.
- León-Velarde F, Ramos MA, Hernández JA, De Idiáquez D, Muñoz LS, Gaffo A, Córdova S, Durand D, Monge C. The role of menopause in the development of chronic mountain sickness. *Am J Physiol Regul Integr Comp Physiol* 1997;272(1):R90–R94.
- Monge C, Arregui A, León-Velarde F. Pathophysiology and epidemiology of chronic mountain sickness. *Int J Sports Med* 1992;13(S1):S79–S81.
- Xie C, Pei S. Some physiological data of sojourners and native highlanders at three different altitudes in Xizhang. In: Liu D, ed. Geological and Ecological Studies of Qinghai-Xizang Plateau. New York: Gordon & Breach, 1981:1449–1552.
- Moore LG, Asmus I, Curran L. Chronic mountain sickness: gender and geographic variation. In: Ohno H, Kobayashi T, Masuyama S, Nakashima M, eds. Progress in Mountain Medicine and High Altitude Physiology. Matsumoto, Japan: Press Committee of the 3rd World Congress on Mountain Medicine and High Altitude Physiology, 1998:114–119.
- Groves BM, Droma T, Sutton JR, McCullough RG, McCullough RE, Zhuang J, Rapmund G, Sun S, Janes C, Moore LG. Minimal hypoxic pulmonary hypertension in normal Tibetans at 3,658 m. *J Appl Physiol* 1993;74(1):312–318.
- Cruz-Jibaja J, Banchero N, Sime F, Peñaloza D, Gamboa R, Marticorena E. Correlation between pulmonary artery pressure and level of altitude. *Dis Chest* 1964;46(4):446–451.
- Banchero N, Sime F, Peñaloza D, Cruz J, Gamboa R, Marticorena E. Pulmonary pressure, cardiac output, and arterial oxygen saturation during exercise at high altitude and at sea level. *Circulation* 1966;33(2):249–262.
- Reeves JT, Grover RF. High-altitude pulmonary hypertension and pulmonary edema. In: Yu PN, Goodwin JF, eds. Progress in Cardiology IV. Philadelphia: Febiger, 1975:99–118.
- Hoit BD, Dalton ND, Gebremedhin A, Janocha A, Zimmerman PA, Zimmerman AM, Strohl KP, Erzurum SC, Beall CM. Elevated pulmonary artery pressure among Amhara highlanders in Ethiopia. *Am J Hum Biol* 2011;23(2): 168–176.
- Julian CG, Wilson MJ, Lopez M, Yamashiro H, Tellez W, Rodriguez A, Bigham A, et al. Augmented uterine artery blood flow and oxygen delivery protect Andeans from altitude-associated reductions in fetal growth. *Am J Physiol Regul Integr Comp Physiol* 2009;296(5):R1564–R1575.
- Giussani DA, Phillips PS, Anstee S, Barker DJ. Effects of altitude versus economic status on birth weight and body shape at birth. *Pediatr Res* 2001;49(4):490–494.
- Haas JD, Frongillo EA Jr., Stepick CD, Beard JL, Hurtado L. Altitude, ethnic and sex difference in birth weight and length in Bolivia. *Human Biol* 1980;52(3):459–477.
- Mortola JP, Frappell PB, Aguero L, Armstrong K. Birth weight and altitude: a study in Peruvian communities. *J Pediatr* 2000;136(3):324–329.
- Julian CG, Vargas E, Armaza JF, Wilson MJ, Niermeyer S, Moore LG. High-altitude ancestry protects against hypoxia-



- associated reductions in fetal growth. *Arch Dis Child Fetal Neonatal Ed* 2007;92(5):F372–F377.
21. Gonzales GF, Steenland K, Tapia V. Maternal hemoglobin level and fetal outcome at low and high altitudes. *Am J Physiol Regul Integr Comp Physiol* 2009;297(5):R1477–R1485.
  22. Gonzales GF, Tapia V, Carrillo CE. Stillbirth rates in Peruvian populations at high altitude. *Int J Gynecol Obstet* 2008;100(3):221–227.
  23. Hartinger S, Tapia V, Carrillo C, Bejarano L, Gonzales GF. Birth weight at high altitudes in Peru. *Int J Gynecol Obstet* 2006;93(3):275–281.
  24. Zamudio S, Postigo L, Illsley NP, Rodriguez C, Heredia G, Brimacombe M, Echalar L, et al. Maternal oxygen delivery is not related to altitude- and ancestry-associated differences in human fetal growth. *J Physiol* 2007;582(2):883–895.
  25. Zamudio S, Droma T, Norkyel KY, Acharya G, Zamudio JA, Niermeyer SN, Moore LG. Protection from intrauterine growth retardation in Tibetans at high altitude. *Am J Phys Anthropol* 1993;91(2):215–224.
  26. Tripathy V, Gupta R. Birth weight among Tibetans at different altitudes in India: are Tibetans better protected from IUGR? *Am J Hum Biol* 2005;17(4):442–450.
  27. Smith C. The effect of maternal nutritional variables on birthweight outcomes of infants born to Sherpa women at low and high altitudes in Nepal. *Am J Hum Biol* 1997;9(6):751–763.
  28. Yangzom Y, Qian L, Shan M, La Y, Meiduo D, Hu X, Da Q, Sun B, Zetterström R. Outcome of hospital deliveries of women living at high altitude: a study from Lhasa in Tibet. *Acta Paediatr* 2008;97(3):317–321.
  29. Zhao X, Gao WX, Suo L, Chen J. Placental mitochondrial respiratory function of native Tibetans at high altitude [in Chinese]. *Zhonghua yi xue za zhi* 2007;87(13):894–897.
  30. Yip R. Altitude and birth weight. *J Pediatr* 1987;111(6):869–876.
  31. Yip R, Li Z, Chong WH. Race and birth weight: the Chinese example. *Pediatrics* 1991;87(5):688–693.
  32. Wen SW, Kramer MS, Usher RH. Comparison of birth weight distributions between Chinese and Caucasian infants. *Am J Epidemiol* 1995;141(12):1177–1187.
  33. Moore LG, Young D, McCullough RE, Droma T, Zamudio S. Tibetan protection from intrauterine growth restriction (IUGR) and reproductive loss at high altitude. *Am J Hum Biol* 2001;13(5):635–644.
  34. Moore LG. Uterine blood flow as a determinant of fetoplacental development. In: Burton GJ, Barker DJP, Moffett A, Thornburg KL, eds. *The Placenta and Human Developmental Programming*. Cambridge: Cambridge University Press, 2011:126–146.
  35. Niermeyer S, Zamudio S, Moore LG. The people. In: Hornbein T, Schoene RB, eds. *Adaptations to Hypoxia*. New York: Dekker, 2001:43–100.
  36. Qadar Pasha MA, Newman JH. High-altitude disorders: pulmonary hypertension: pulmonary vascular disease: the global perspective. *Chest* 2010;137(6\_suppl):13S–9S.
  37. Sui GJ, Liu YH, Cheng XS, Anand IS, Harris E, Harris P, Heath D. Subacute infantile mountain sickness. *J Pathol* 1988;155(2):161–170.
  38. Winslow RM, Chapman KW, Gibson CC, Samaja M, Monge CC, Goldwasser E, Sherpa M, Blume FD, Santolaya R. Different hematologic responses to hypoxia in Sherpas and Quechua Indians. *J Appl Physiol* 1989;66(4):1561–1569.
  39. Beall CM, Brittenham GM, Macuaga F, Barragan M. Variation in hemoglobin concentration among samples of high-altitude natives in the Andes and the Himalayas. *Am J Hum Biol* 1990;2(6):639–651.
  40. Wu T, Wang X, Wei C, Cheng H, Wang X, Li Y, Ge-Dong, et al. Hemoglobin levels in Qinghai-Tibet: different effects of gender for Tibetans vs. Han. *J Appl Physiol* 2005;98(2):598–604.
  41. Khoury GH, Hawes CR. Primary pulmonary hypertension in children living at high altitude. *J Pediatr* 1963;62(2):177–185.
  42. Anand IS, Wu T. Syndromes of subacute mountain sickness. *High Alt Med Biol* 2004;5(2):156–170.
  43. Stuber T, Sartori C, Salmón CS, Hutter D, Thalmann S, Turini P, Jayet PY, et al. Respiratory nitric oxide and pulmonary artery pressure in children of Aymara and European ancestry at high altitude. *Chest* 2008;134(5):996–1000.
  44. Hampl V, Herget J. Perinatal hypoxia increases hypoxic pulmonary vasoconstriction in adult rats recovering from chronic exposure to hypoxia. *Am Rev Respir Dis* 1990;142(3):619–624.
  45. Sartori C, Allemann Y, Trueb L, Delabays A, Nicod P, Scherrer U. Augmented vasoreactivity in adult life associated with perinatal vascular insult. *Lancet* 1999;353(9171):2205–2207.
  46. Jensen GM, Moore LG. The effect of high altitude and other risk factors on birthweight: independent or interactive effects? *Am J Public Health* 1997;87(6):1003–1007.
  47. Browne VA, Julian CG, Toledo-Jaldin L, Cioffi-Ragan D, Vargas E, Moore LG. Uterine artery blood flow, fetal growth and fetal hypoxia. *Philos Trans Royal Soc B* 2015;370(1663):20140068. doi:10.1098/rstb.2014.0068.
  48. Niermeyer S, Shaffer EM, Thilo E, Corbin C, Moore LG. Arterial oxygenation and pulmonary arterial pressure in healthy neonates and infants at high altitude. *J Pediatr* 1993;123(5):767–772.
  49. Niermeyer S, Yang P, Shanmina, Drolkar, Zhuang J, Moore LG. Arterial oxygen saturation in Tibetan and Han infants born in Lhasa, Tibet. *N Engl J Med* 1995;333(19):1248–1252.
  50. Barker DJ. Fetal origins of cardiovascular disease. *Ann Med* 1999;31(suppl 1):3–6.
  51. Moore LG, Niermeyer S, Vargas E. Does chronic mountain sickness (CMS) have perinatal origins? *Respir Physiol Neurobiol* 2007;158(2–3):180–189.
  52. Julian CG, Vargas E, Gonzales M, Dávila RD, Ladenburger A, Reardon L, Schoo C, Powers RW, Lee-Chiong T, Moore LG. Sleep-disordered breathing and oxidative stress in pre-clinical chronic mountain sickness (excessive erythrocytosis). *Respir Physiol Neurobiol* 2013;186(2):188–196.
  53. Wilson MJ, Lopez M, Vargas M, Julian C, Tellez W, Rodriguez A, Bigham A, et al. Greater uterine artery blood flow during pregnancy in multigenerational (Andean) than shorter-term (European) high-altitude residents. *Am J Physiol Regul Integr Comp Physiol* 2007;293(3):R1313–R1324.
  54. Shriver MD, Kennedy GC, Parra EJ, Lawson HA, Sonpar V, Huang J, Akey JM, Jones KW. The genomic distribution of population substructure in four populations using 8,525 au-



- tosomal SNPs. *Hum Genomics* 2004;1:274–286. doi:10.1186/1479-7364-1-4-274.
55. Williams RL, Creasy RK, Cunningham GC, Hawes WE, Norris FD, Tashiro M. Fetal growth and perinatal viability in California. *Obstet Gynecol* 1982;58(5):624–632.
  56. Li W, Du J, Ma Y, Li Y, Li X. Pulmonary artery pressure evaluated by pulsed Doppler echocardiography in children with a left-to-right intracardiac shunt. *Pediatr Cardiol* 1991; 12(1):17–19.
  57. Victoria-Oliva G, Mojarro-Ríos J, Alva-Espinosa C, Villasís-Keever MA, Labarthe-Cabrera J, Arellano-Penagos M, Velasco-Jiménez S, Alarcón-Santos S, Muro R, Díaz-Arauzo A. Ecocardiografía Doppler en recién nacidos con riesgo de hipertensión pulmonar. *Rev Mex Cardiol* 1996;7(1):25–31.
  58. Skinner JR, Stuart AG, O'Sullivan J, Heads A, Boys RJ, Hunter S. Right heart pressure determination by Doppler in infants with tricuspid regurgitation. *Arch Dis Child* 1993; 69(2):216–220.
  59. Skinner JR, Boys RJ, Heads A, Hey EN, Hunter S. Estimation of pulmonary arterial pressure in the newborn: study of the repeatability of four Doppler echocardiographic techniques. *Pediatr Cardiol* 1996;17(6):360–369.
  60. Emmanouilides GC, Moss AJ, Duffie ER, Adams FH. Pulmonary artery pressure changes in human newborn infants from birth to 3 days of age. *J Pediatr* 1964;65(3):327–333.
  61. Hannhart B, Haberer JP, Saunier C, Laxenaire MC. Accuracy and precision of fourteen pulse oximeters. *Eur Respir J* 1991;4(1):115–119.
  62. Aparicio OO, Romero Gutierrez F, Harris P, Anand I. Echocardiography shows persistent thickness of the wall of the right ventricle in infants at high altitude. *Cardioscience* 1991; 2(1):63–69.
  63. Gamboa R, Marticorena E. Presión arterial pulmonary en el recién nacido en las grandes alturas. *Arch Inst Biol Andina* 1971;4(2):55–66.
  64. Herrera EA, Riquelme RA, Ebensperger G, Reyes RV, Ulloa CE, Cabello G, Krause BJ, Parer JT, Giussani DA, Llanos AJ. Long-term exposure to high-altitude chronic hypoxia during gestation induces neonatal pulmonary hypertension at sea level. *Am J Physiol Regul Integr Comp Physiol* 2010;299(6): R1676–R1684.
  65. Papamatheakis DG, Chundu M, Blood AB, Wilson SM. Prenatal programming of pulmonary hypertension induced by chronic hypoxia or ductal ligation in sheep. *Pulm Circ* 2013; 3(4):757–780.
  66. Alzamora-Castro V, Battilana G, Abugattas R, Sialer S. Patent ductus arteriosus and high altitude. *Am J Cardiol* 1960;5 (6):761–763.
  67. Hurtado Gomez L, Calderon G. Hipoxia de altura en la insuficiencia cardiaca del lactante. *Bol Soc Boliv Pediatr* 1965; 9(1):11–23.
  68. Lin CP, Wu TY. Clinical analysis of 286 cases of pediatric high altitude heart diseases [in Chinese]. *Zhonghua yi xue za zhi* 1974;6:353–356.
  69. Mo LF, Jiang MX, Li JB, Jiao HG. Pediatric high altitude heart disease. In: People's Hospital of the Tibetan Autonomous Region, ed. *High Altitude Medicine*. Lhasa: Tibetan People's Publisher, 1983:251–259.
  70. Niermeyer S, Shaffer EM, Moore LG. Impaired cardiopulmonary transition at high altitude. *Pediatr Res* 1998;43(S4):292.
  71. Miao CY, Zuberbuhler JS, Zuberbuhler JR. Prevalence of congenital cardiac anomalies at high altitude. *J Am Coll Cardiol* 1988;12(1):224–228.
  72. Wu TY, Miao CY, Lin CP, Ma RY, Yao RL, Lian TY. Altitude zillness in children on the Tibetan Plateau. In: Ohno H, Kobayashi T, Masuyama S, Nakashima M, eds. *Progress in Mountain Medicine and High Altitude Physiology*. Matsumoto, Japan: Press Committee of the Third World Congress on Mountain Medicine and High Altitude Physiology, 1998: 195–200.
  73. Halperin BD, Sun S, Zhuang J, Droma T, Moore LG. ECG observations in Tibetan and Han residents of Lhasa. *J Electrocardiol* 1998;31(3):237–243.
  74. Faoro V, Huez S, Vanderpool RR, Groepenhoff H, de Bisschop C, Martinot J-B, Lamotte M, Pavelescu A, Guénard H, Naeije R. Pulmonary circulation and gas exchange at exercise in Sherpas at high altitude. *J Appl Physiol* 2014;116 (7):919–926.
  75. Beall CM, Laskowski D, Strohl KP, Soria R, Villena M, Vargas E, Alarcon AM, Gonzales C, Erzurum SC. Pulmonary nitric oxide in mountain dwellers. *Nature* 2001;414(6862): 411–412.
  76. Schneider A, Greene RE, Keyl C, Bandinelli G, Passino C, Spadacini G, Bonfichi M, et al. Peripheral arterial vascular function at altitude: sea-level natives versus Himalayan high-altitude natives. *J Hypertens* 2001;19(2):213–222.
  77. Erzurum SC, Ghosh S, Janocha AJ, Xu W, Bauer S, Bryan NS, Tejero J, et al. Higher blood flow and circulating NO products offset high-altitude hypoxia among Tibetans. *Proc Natl Acad Sci USA* 2007;104(45):17593–17598.
  78. Zhuang J, Droma T, Sun S, McCullough RE, McCullough RG, Cymerman A, Huang SY, Reeves JT, Moore LG. Hypoxic ventilatory responsiveness in Tibetan compared with Han residents of 3,658 m. *J Appl Physiol* 1993;74(1):303–311.
  79. Gilbert-Kawai ET, Milledge JS, Grocott MPW, Martin DS. King of the mountains: Tibetan and Sherpa physiological adaptations for life at high altitude. *Physiology* 2014;29(6): 388–402.
  80. Jayet PY, Rimoldi SF, Stuber T, Salmòn CS, Hutter D, Rexhaj E, Thalmann S, et al. Pulmonary and systemic vascular dysfunction in young offspring of mothers with pre-eclampsia. *Circulation* 2010;122(5):488–494.
  81. Thilo EH, Park-Moore B, Berman ER, Carson BS. Oxygen saturation by pulse oximetry in healthy infants at an altitude of 1610 m (5280 ft): what is normal? *Am J Dis Child* 1991; 145(10):1137–1140.
  82. O'Brien LM, Stebbens VA, Poets C, Heycock E, Southall D. Oxygen saturation during the first 24 hours of life. *Arch Dis Child Fetal Neonatal Ed* 2000;83(1):F35–F38.
  83. Deming J, Washburn AH. Respiration in infancy. I. A method of studying rates, volume and character of respiration with preliminary report of results. *Am J Dis Child* 1935; 49(1):108–124.
  84. Lubchenco LO, Ashby BL, Markarian M. Periodic breathing in newborn infants in Denver and Leadville, Colorado. *Soc Pediatr Res Program Abstr* 1964:50.