PHYSIOLOGICAL CORRELATES OF BURROWING IN RODENTS

RICHARD C. CHAPMAN* AND ALBERT F. BENNETT

Department of Zoology, University of California, Berkeley, California 94720, U.S.A.

(Received 11 April 1974)

Abstract—1. Blood properties of valley pocket gophers, Thomomys bottae, and laboratory rats, Rattus norvegicus, were examined to determine blood buffering capabilities.

2. Hematocrit, plasma proteins and inorganic phosphate levels were not significantly different between these species.

3. Oxygen capacity of the pocket gophers, 23.1 vol.%, was greater than that of the rats, 20.8 vol.%. 

4. Bicarbonate concentration of pocket gopher blood, 28.1 mM/l., was significantly greater than that of the rats, 19.8 mM/l.

5. The non-carbonic buffering strength as determined from CO₂ titration curves of whole blood of pocket gophers, -2.67 log pCO₂/pH unit, was greater than that of the rats, -1.39 log pCO₂/pH unit.

6. The greater buffering capacity of pocket gopher blood may account for the reduced sensitivity of ventilation rate in response to elevated burrow CO₂ concentrations.

INTRODUCTION

The gas composition of rodent burrows is often very different from that of environmental air, creating situations of potential respiratory difficulty for these fossorial mammals. Field observations of pocket gopher burrows have revealed CO₂ concentrations ranging from 0.6 to 3.8 per cent and O₂ concentrations from 6.0 to 21.0 per cent (Kennerly, 1964; McNab, 1966; Darden, 1972). Maximum CO₂ concentrations of 6 per cent have been found in burrows of thirteen-lined squirrels (Spermophilus tridecemlineatus) (Studier & Proctor, 1971). Normal concentrations of CO₂ and O₂ in air are 0.04 and 20.95 per cent, respectively. The poor circulation of air within burrows accompanied by the respiration of the rodents and soil micro-organisms accounts for the accumulation of high concentrations of CO₂ and depletion of O₂ within the burrows.

Ventilation in non-burrowing mammals is sensitive to the concentration of CO₂ in the inspired air. When the concentration of CO₂ in the inspired air is increased, a lowering of body fluid pH results. The lowered pH stimulates the respiratory control centers of the central nervous system resulting in an increase in ventilation rate, such that the partial pressure of CO₂ (pCO₂) in alveolar air remains almost unchanged (Siggaard-Andersen, 1965). The increment in ventilation rate in response to elevated CO₂ concentrations in two fossorial rodents, the valley pocket gopher, Thomomys bottae (Darden, 1972), and Merriam's kangaroo rat, Dipodomys merriami (Soholt et al., 1973), has been shown to be significantly less than the response of non-fossorial mammals to the same level of CO₂ concentration. Other effects of high CO₂ concentrations are heart rate depression and a drop in body temperature. These responses develop only at very high levels of CO₂; however, the magnitude of these responses in burrowing mammals also appears to be less than in laboratory mammals (Soholt et al., 1973).

Darden (1972) suggests that the greater insensitivity of fossorial mammals to CO₂ may be due to a greater buffering capacity in the body fluids than that found in non-fossorial mammals. Another potential mechanism of increased tolerance is a reduced sensitivity of the respiratory control receptors of fossorial mammals to changes in body fluid pH. Soholt et al. (1973) also believe that the former suggestion seems more likely, since enzymatic function is dependent upon stable pH. However, pH change might also facilitate enzymatic adaptation to burrowing conditions. In any case, neither theory has been experimentally verified.

The purpose of the present investigation was to examine the blood buffering capabilities of a fossorial rodent, T. bottae, compared to the capabilities of a non-fossorial rodent, R. norvegicus. Measurements were made of the following blood properties: bicarbonate concentration, inorganic phosphate concentration, oxygen capacity, hematocrit, plasma total protein and non-albumin and albumin fractions, and titration curves for non-carbonic buffers.

* Present address: Alaska Co-operative Wildlife Research Unit, University of Alaska, College, Alaska 99701, U.S.A.

† Present address: School of Biological Sciences, University of California, Irvine, California 92666, U.S.A.
If buffering capacities are indeed greater in fossorial rodents, an examination of these parameters should reveal differences between these species, particularly in carbonic or non-carbonic buffering ability.

**MATERIALS AND METHODS**

**Experimental animals**

A total of thirteen adult valley pocket gophers, Thomomys bottae, and approximately fifteen adult laboratory rats, Rattus norvegicus, were used. They were given Berkeley Diet Mouse Breeder Food supplemented every other day with waste vegetable material (potato and carrot peelings, lettuce, celery, etc.). Water was provided ad lib.; however, most of the water needed by the pocket gophers was obtained from the vegetable matter. Temperature was maintained at approximately 20°C with a natural photoperiod. The experiments were conducted from October 1973 to March 1974.

Approximately fifteen adult laboratory rats, Rattus norvegicus (mean weight, 396 g), were also used. They were given Berkeley Diet Mouse Breeder Food and water ad lib. Temperature was maintained at approximately 20°C. The photoperiod was 12L12D.

**Blood samples**

Prior to obtaining blood samples, all experimental animals were anesthetized with Nembutal (sodium pentobarbital) in concentrations of 50 mg/kg body weight. All blood samples were obtained by ventricular puncture with either No. 24, 26 or 27 sized needles. No. 24 sized needles proved the most satisfactory. Coagulation was prevented by the addition of crystalline sodium heparin to the blood samples.

Blood samples of approximately 0.25 ml were obtained from nine pocket gophers and nine rats. The samples were placed in heparinized capillary tubes, sealed and stored on ice. For each curve, 0.25 ml of blood was obtained from each of four pocket gophers. For each curve the blood samples were pooled, mixed thoroughly and stored on ice between determinations. The equilibration system previously described was utilized. The temperature of the system was regulated at 38°C. Gas mixtures of varying proportions of CO₂, O₂, and N₂, saturated with water vapor, were prepared in a 9 l capacity spirometer (pCO₂ = 35 torr, pO₂ = 145 torr, pN₂ = 50 torr) and equilibrated with the gas mixture as described previously for oxygen capacity measurements. The pH, in an atmosphere of 9 1. capacity spirometer (pCO₂ = 35 torr, pO₂ = 145 torr, pN₂ = 50 torr), acidulating the sample with a gas mixture of 95% air and 5% carbon dioxide saturated with water vapor at 38°C (pCO₂ = 145 torr, pO₂ = 35 torr, pN₂ = 50 torr) was slowly metered into the flask for about 4 min. The flask was tilted and rotated at about 20 rev/min, so that the blood sample formed a thin layer on the periphery of the sample. After the equilibration period, the sample was removed and placed in a 25 ml volumetric flask. A gas mixture of 91% air and 9% carbon dioxide saturated with water vapor at 38°C (pO₂ = 145 torr, pCO₂ = 35 torr, pN₂ = 50 torr) was slowly metered into the flask for about 10 min. At the end of the equilibration period, the sample was removed and analyzed for oxygen content according to the method of Roughton & Scholander (1943). The oxygen content was expressed as vol% corrected to STPD conditions.

**Plasma proteins**

Red and white blood cell fractions were taken from the plasma fractions of the capillary tubes used in the hema tocrit determinations. Plasma samples of 0.1 ml from four pocket gophers and four rats were used in this experiment. Total proteins were determined with the biuret procedure. The non-albumin proteins were precipitated with ether and the albumin-containing solvent was again analyzed with the biuret procedure. Protein content was expressed as g/100 ml plasma.

**Phosphate buffers**

The phosphate buffers were assayed by determination of the total amount of inorganic phosphate by the spectrophotometric method outlined by Hawk et al. (1947). The method consisted of equilibrating a 0.10 ml blood sample with a gas mixture of 95% air and 5% carbon dioxide saturated with water vapor at 38°C (pCO₂ = 145 torr, pO₂ = 35 torr, pN₂ = 50 torr), acidulating the sample and measuring the liberated carbon dioxide manometrically. Bicarbonate concentrations were expressed in mM/l. Blood samples were obtained from seven pocket gophers and eight rats.

**Bicarbonate concentrations**

The bicarbonate-carbonic acid buffer systems were assayed by the determination of bicarbonate concentration under standard conditions. This was done according to the method outlined by Umbreit et al. (1964). The method consisted of equilibrating a 0.10 ml blood sample with a gas mixture of 95% air and 5% carbon dioxide saturated with water vapor at 38°C (pO₂ = 145 torr, pCO₂ = 35 torr, pN₂ = 50 torr), acidulating the sample and measuring the liberated carbon dioxide manometrically. Bicarbonate concentrations were expressed in mM/l. Blood samples were obtained from seven pocket gophers and eight rats.

**Titration curves for all non-carbonic buffers**

A titration curve for all non-carbonic buffers in whole blood was constructed by varying the pCO₂ of the atmosphere in which an in vitro blood sample was exposed and measuring the resulting pH. Two separate curves were constructed for both the pocket gophers and the rats. Approximately 0.6 ml of blood was obtained from each of four rats for the first curve; approximately 1.0 ml from each of four rats for the second curve. For both pocket gopher curves approximately 1.0 ml of blood was obtained from each of four pocket gophers. For each curve the blood samples were pooled, mixed thoroughly and stored on ice between determinations. The pH of each solution previously described was utilized. The temperature of the system was regulated at 38°C. Gas mixtures of varying proportions of CO₂, O₂, and N₂ saturated with water vapor, were prepared in a 9 l capacity spiro meter (pCO₂ = 50 torr, pO₂ = 142 torr, pN₂ = 35 torr). A 0.4 ml sample of blood was placed in a 10 ml volumetric flask and equilibrated with the gas mixture as described previously for oxygen capacity measurements. The pH, in an atmosphere of 9 1. capacity spirometer (pCO₂ = 35 torr, pO₂ = 145 torr, pN₂ = 50 torr) was slowly metered into the flask for about 4 min. The flask was tilted and rotated at about 20 rev/min, so that the blood sample formed a thin layer on the periphery of the sample. After the equilibration period, the sample was removed and placed in a 25 ml volumetric flask and measured the total amount of oxygen bound to the hemoglobin of the blood sample, and the total amount of inorganic phosphate by the spectrophotometric method outlined by Hawk et al. (1947).
Statistics
All linear regressions are the best computed least-squares fit to the data. Mean values are reported with standard errors; 95 per cent confidence limits were used to estimate significance. The difference between mean values was tested with a Student's t-test.

RESULTS
Hematocrit and oxygen capacity
The average hematocrit for the pocket gophers, 46.1% (±1.3), is not significantly greater (P > 0.1) than that for the rats, 43.7% (±1.3).

The average oxygen capacity of pocket gopher blood, 23.1 vol% (±0.33), is slightly greater (0.05 > P > 0.02) than that of rat blood, 20.8 vol% (±0.55).

Plasma proteins
The average total protein content of blood plasma for the pocket gophers, 6.3 g/100 ml (±0.13), is identical to that for the rats, 6.3 g/100 ml (±0.20). Values of non-albumin protein are slightly higher in the pocket gophers, 0.99 g/100 ml (±0.072), than in the rats, 0.70 g/100 ml (±0.074) (0.05 > P > 0.02).

Phosphate buffers
The average concentration of inorganic phosphate in the blood of the rats, 4.9 mg % (±0.33), is not significantly different (P > 0.01) from that of the pocket gophers, 4.2 mg % (±0.15).

Bicarbonate concentration
The average bicarbonate concentration in the blood for the pocket gophers, 28.1 mM/1 (±0.89), is significantly greater (P < 0.001) than that for the rats, 19.8 mM/1 (±1.0).

Titration curves for all non-carbonic buffers
Figures 1 and 2 show the regression lines of the titration curves for all non-carbonic buffers for the pocket gophers and rats, respectively. Siggaard-Andersen (1965) has found linear approximation represents a satisfactory description of this relationship of pH as a function of log pCO2. The average non-carbonic buffering strength of pocket gopher blood is -2.67 log pCO2/pH unit, and of rat blood, -1.39 log pCO2/pH unit. The non-carbonic buffering systems of the pocket gophers are, therefore, more effective in retarding pH change than those of the rats.

DISCUSSION
Hematocrit and oxygen capacity
The hematocrits measured are very similar to those reported for laboratory rats (46%) and other non-ungulate mammals (40-53%) (Altman & Ditter, 1971). The oxygen capacity values of the rats in this study are higher than those previously reported for laboratory rats (46%) and other non-ungulate mammals (40-53%) (Altman & Ditter, 1971).

Fig. 1. The pH of whole blood of the pocket gophers (T. bottae) at 38°C as a function of pCO2. Experimental points for regression line 1 (pH = 7.90 - 0.363 log pCO2); O, regression line 2 (pH = 8.10 - 0.387 log pCO2).

Fig. 2. The pH of whole blood of the laboratory rats (R. norvegicus) at 38°C as a function of pCO2. Experimental points for regression line 1 (pH = 9.02 - 0.862 log pCO2); O, regression line 2 (pH = 8.59 - 0.622 log pCO2).
The bicarbonate–carbonic acid system, as assayed by the bicarbonate content of the blood, appears much better developed in the pocket gophers than in the rats. Many of the values reported for other mammals are not significantly different from that of the pocket gophers—man, 27.0 mM/l.; cat, 29.4 mM/l.; cattle, 32.0 mM/l.; guinea pig, 22.0 mM/l.; hamster, 37.3 mM/l.; hibernating hamster, 42.4 mM/l.; horse, 38.1 mM/l.; rabbit, 22.8, 18.0 mM/l.; rat, 20.9, 24.0 mM/l.; sheep, 26.3 mM/l.; ground squirrel, 30.5 mM/l.; hibernating ground squirrel, 38.6 mM/l. (Altman & Dittmer, 1971). The pocket gopher bicarbonate content is, however, higher than all other rodents examined except the hamster and hibernating ground squirrel. The greater levels of bicarbonate in the latter animals may be directly related to hibernation.

The greater buffering ability of pocket gopher blood is also manifested in the titration curve for non-carboxic buffers. The buffering value for the pocket gophers, –2.67 log pCO2/pH unit, is greater than that of man, –1.57 log pCO2/pH unit, and also that reported for man, –1.57 log pCO2/pH unit (Siggaard-Andersen, 1965). The intercept (vertical position) of the titration curve is influenced by diet or the addition of non-volatile acids (e.g. lactic acid), but the slope of the curve is thought to reflect the concentration and composition of the non-carbonic blood buffers: hemoglobin, plasma proteins, phosphates and non-protein thiol groups. Since inorganic phosphate levels are low in both pocket gopher and rat blood, it might be concluded that the protein buffers are responsible for its greater pH stability. Quantitatively, there was no difference in the plasma proteins of the two animals investigated; however, qualitative differences (i.e. differences in amino acid composition) have not been studied. Hemoglobin concentrations are perhaps higher in pocket gopher blood than in most other mammals (see above). Although the blood buffering capabilities appear to be better in pocket gophers, no whole-body buffer curves, measuring the response of blood pH to in vivo carbon dioxide exposure, have been constructed for any fossorial mammal. The determination of such a response is clearly indicated.

Physiological correlates of burrowing

The problem facing fossorial mammals encountering high CO2 concentrations is one of chronic respiratory acidosis. The extent of change in pH of the body fluids in a mammal challenged by high levels of CO2 in the inspired air is determined by the effectiveness of three compensatory mechanisms: (1) the magnitude of the ventilatory response evoked, (2) the capacity of the blood and tissue buffer systems and (3) the efficiency of the renal adjustment (Darden, 1972).

The magnitude of the ventilatory response evoked by high levels of CO2 is reduced in fossorial mammals (Darden, 1972; Sobol, et al., 1973). The reduced breathing may prevent excessive respiratory water loss (however, the humidity in most burrow systems is high) or excessive dust inhalation while digging (Scholander et al., 1943; Irving et al., 1956). Also, it appears that fossorial mammals may experience a wider range of CO2 concentrations at a reduced mechanical energy cost of breathing (Milic-Emili & Petil, 1960; Darden, 1972). However, reduced ventilation does not facilitate loss of CO2 through the lungs. Therefore, adaptations in the renal and buffer systems of the body may be necessary. (Darden 1972) indicated that the rate of renal adjustment during chronic hypercapnia, as indicated by excretion of NH4+, Cl−, Na+ and K+, is rapid. The present investigation has shown relatively high bicarbonate concentrations and a greater strength of buffer systems in pocket gophers. This tends to confirm the hypothesis that blood buffering is greater in pocket gophers than in non-fossorial mammals. Indeed, it appears that the blood buffers and the renal system are compensating for the depressed response of the respiratory system to elevated CO2 concentrations.

Acknowledgements—This paper is part of a Senior Honors thesis submitted to the Department of Zoology, University of California, Berkeley, by R. C. C. We wish to thank Dr. Paul Licht for the use of equipment and Dr. James L. Patton for the use of pocket gopher traps and the Museum of Vertebrate Zoology animal room.

REFERENCES

Physiological correlates of burrowing in rodents


Key Word Index—Bicarbonate; blood; buffering; burrowing; carbon dioxide; fossorial; hematocrit; oxygen capacity; pH; phosphate; plasma protein; Rattus; rodent; Thomomys.