

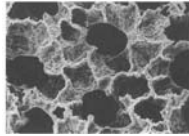
Lecture 21  
05 March 2008

Vertebrate Physiology  
ECOL 437 (MCB/VetSci 437)  
Univ. of Arizona, spring 2008

Kevin Bonine & Kevin Oh



1. Respiration (Ch 20-21)



**FIGURE 20.16** Gills. Drawing illustrates a cross-section of a fish gill. The gill consists of a central gill arch, which is supported by a cartilaginous structure called the gill raker. The gill arch is divided into two parts: the anterior part, which is the gill arch proper, and the posterior part, which is the gill arch extension. The gill arch proper is further divided into the gill arch body and the gill arch base. The gill arch extension is further divided into the gill arch tip and the gill arch base. The gill arch body is further divided into the gill arch body proper and the gill arch body extension. The gill arch body proper is further divided into the gill arch body proper anterior and the gill arch body proper posterior. The gill arch body extension is further divided into the gill arch body extension anterior and the gill arch body extension posterior. The gill arch tip is further divided into the gill arch tip proper and the gill arch tip extension. The gill arch tip proper is further divided into the gill arch tip proper anterior and the gill arch tip proper posterior. The gill arch tip extension is further divided into the gill arch tip extension anterior and the gill arch tip extension posterior. The gill arch base is further divided into the gill arch base proper and the gill arch base extension. The gill arch base proper is further divided into the gill arch base proper anterior and the gill arch base proper posterior. The gill arch base extension is further divided into the gill arch base extension anterior and the gill arch base extension posterior.

[http://eebweb.arizona.edu/eeb\\_course\\_websites.htm](http://eebweb.arizona.edu/eeb_course_websites.htm)

Housekeeping, 05 March 2008

Upcoming Readings

Wed 05 Mar: Ch20, 21 (respiration)  
LAB Wed 05 Mar: Dickinson reading on website  
Fri 07 Mar: Ch 21  
Mon 10 Mar: Ch 21, 22  
Wed 12 Mar: Ch 23 (circulation)  
LAB Wed 12 Mar: no reading  
Fri 14 Mar: EXAM TWO (through respiration)  
SPRING BREAK



Lab discussion leaders: 05 March  
1pm - Julia, Matt C.  
3pm - Dalziel, Nick

Lab discussion leaders: 26 Mar  
1pm - Vangie & Christina  
3pm - Prasun & Ajay

**PHYSIOLOGY**  
C. J. Heckman, Ph.D.  
Professor  
Department of Physiology  
Northwestern University

“Control of spinal neuron excitability: diffuse descending neuromodulation, specific local inhibition”

Friday, March 7, 2008 11:00 a.m.  
AHSC Room 5403  
Refreshments will be served

Also available on line at  
<http://www.physiology.nyu.edu>  
Hosted by Training Grant from NIH, 5R01 NS046000, cjh@northwestern.edu

Wednesday, March 5  
Life's Technological Edge: The Singularity is Near: When Humans Transcend Biology  
Ray Kurzweil, via Teleportec Teleporter  
Founder, Chairman and Chief Executive Officer, Kurzweil Technologies  
Humanity is on the edge of a vast transformation, when what it means to be human will be both enriched and challenged. Inventor and futurist Ray Kurzweil will introduce this radically optimistic singularity, an era when we break our genetic shackles to create a nonbiological intelligence trillions of times more powerful than today. In this new world, humans will transcend biological limitations to achieve entirely new levels of progress and longevity.  
This lecture co-sponsored by: UA College of Engineering and UA College of Science

The Edges of Life - 7pm at Centennial Hall

These do not count as physiology lectures.

The Journal of Physiology

Species differences in Cl<sup>-</sup> affinity and in electrogenicity of SLC26A6-mediated oxalate-Cl<sup>-</sup> exchange correlate with the distinct human and mouse susceptibilities to nephrolithiasis

Jeffrey S. Clark, David H. Vandorpe, Marina N. Chernova, John F. Hingorani, Andrew K. Stewart and Seth L. Alper

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DOI: 10.1111/jphysiol.2007.143222

This information is current as of March 5, 2008

This is the final published version of this article. It is available at  
<http://jphysiol.org/physoc/physoc/586/1291>

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Species differences in Cl<sup>-</sup> affinity and in electrogenicity of SLC26A6-mediated oxalate/Cl<sup>-</sup> exchange correlate with the distinct human and mouse susceptibilities to nephrolithiasis

Jeffrey S. Clark, David H. Vandorpe, Marina N. Chernova, John F. Hingorani, Andrew K. Stewart and Seth L. Alper

*Molecular and Cellular Biochemistry*, 2008, Vol 302, Issue 1, pp 107-114

The mouse is refractory to lithogenic agents active in rats and humans, and we have here traditionally considered a poor experimental model for nephrolithiasis. However, recent studies have identified a defect in an oxalate nephropathy gene in the mouse. Here we tested one earlier demonstration of different oxalate sensitivities of the vertebrate renal and human SLC26A6 orthologues to investigate the correlation between species-specific differences in SLC26A6-mediated oxalate transport properties as expressed in Xenopus oocytes and in reported nephropathy susceptibility. We find that human SLC26A6 mediates maximal rates of Cl<sup>-</sup> exchange for Cl<sup>-</sup> uptake on human, but rates of oxalate/Cl<sup>-</sup> exchange roughly equivalent to those of mouse SLC26. Both transporters exhibit highly cooperative dependence of oxalate release on an extracellular Cl<sup>-</sup>, but whereas the K<sub>0.5</sub> for extracellular Cl<sup>-</sup> is only 100 μM for mouse SLC26, that for human SLC26A6 is 42 μM. This lower value approximates the reported mean normal Cl<sup>-</sup> of postprandial human serum albumin and reflects contributions from both transmembrane and C-terminal cytoplasmic domains of human SLC26A6. Human SLC26A6 versus Y398H exhibits altered Cl<sup>-</sup> dependence and reduced rates of oxalate/Cl<sup>-</sup> exchange. Moreover, mouse SLC26-mediated bidirectional electrogenic oxalate/Cl<sup>-</sup> exchange, human SLC26A6-mediated oxalate transport appears to be electrogenic. We hypothesize that the low extracellular Cl<sup>-</sup> affinity and apparent electrogenicity of oxalate release characterizing human SLC26A6 may partially explain the high human susceptibility to nephrolithiasis relative to that of mouse. SLC26A6 appears to represent a low oxalate risk molecule for nephrolithiasis.

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Key words: Cl<sup>-</sup> affinity, electrogenicity, oxalate, SLC26A6, nephrolithiasis, mouse, human

It has been noted (Holmes et al. 2001), indeed, recent studies have demonstrated minimal impact of dietary oxalate on the frequency of stone disease (Taylor & Curhan, 2007). Most urinary oxalate arises in the course of normal metabolism of the oxalate precursors glycine, glycolate, hydroxyproline, and ascorbate. Normal human serum free oxalate concentrations of ~1.5 μM (Harris et al. 2004), can rise in the setting of end-stage renal disease to predialysis values of 35 μM and higher, and to 130 μM or more in the context of familial primary hyperoxalurias (Yamauchi et al. 2001). Eighty-nine to 99% of intravenously injected oxalate is cleared by the kidney (Osswald & Hautmann, 1979; Ribays & Gershoff, 1982), but colonic oxalate secretion can be up-regulated in the presence of renal insufficiency, leading to increased

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Muldowney FP, Freaney R & Barnes E (1994). Dietary chloride and urinary calcium in stone disease. *QJM* **87**, 501–509.

Osswald H & Hautmann R (1979). Renal elimination kinetics and plasma half-life of oxalate in man. *Urologia Intis* **34**, 440–450.

Restrepo D, Cronise BL, Snyder RB & Knauf PA (1992). A novel method to differentiate between ping-pong and simultaneous exchange kinetics and its application to the anion exchanger of the HL60 cell. *J Gen Physiol* **100**, 825–846.

Ribaya JD & Gershoff SN (1982). Factors affecting endogenous oxalate synthesis and its excretion in feces and urine in rats. *J Nutr* **112**, 2161–2169.

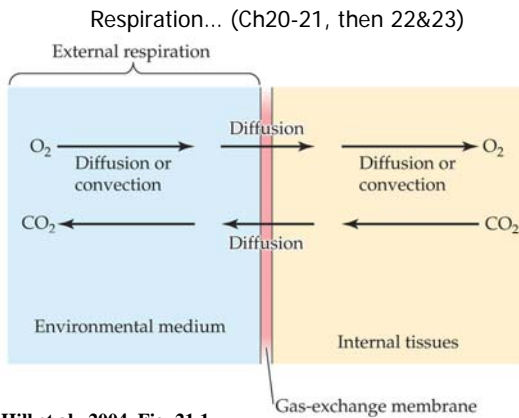
Schaechinger TJ & Oliver D (2007). Nonmammalian orthologs of prestin (SLC26A5) are electrogenic divalent/chloride anion exchangers. *Proc Natl Acad Sci U S A* **104**, 7693–7698.

Shcheynikov N, Wang Y, Park M, Ko SB, Dorwart M, Naruse S, Thomas PJ & Muallem S (2006). Coupling modes and stoichiometry of  $\text{Cl}^-/\text{HCO}_3^-$  exchange by slc26a3 and slc26a6. *J Gen Physiol* **127**, 511–524.

7

# Vertebrate Respiration

8



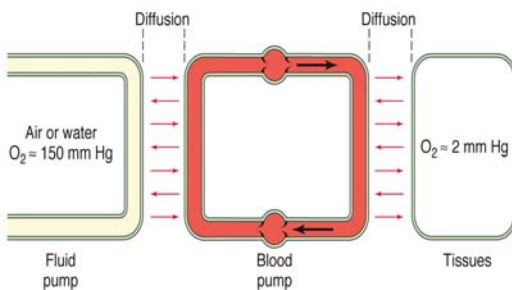
Hill et al., 2004, Fig. 21.1

ANIMAL PHYSIOLOGY, Figure 21.1 © 2004 Sinauer Associates, Inc.

## Gas transfer

1. **Breathing** (supply air or water to respiratory surface)
2. **Diffusion of  $\text{O}_2$  &  $\text{CO}_2$  across resp. epithelium** (humans = 50-100 $\text{m}^2$  SA)
3. **Bulk transport** of gases by blood
4. **Diffusion across capillary walls (blood  $\rightarrow$  mitochondria)**

10



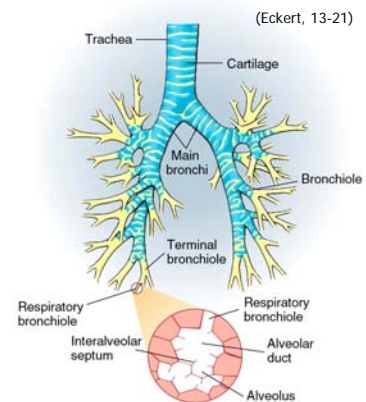
11

## Lung Anatomy

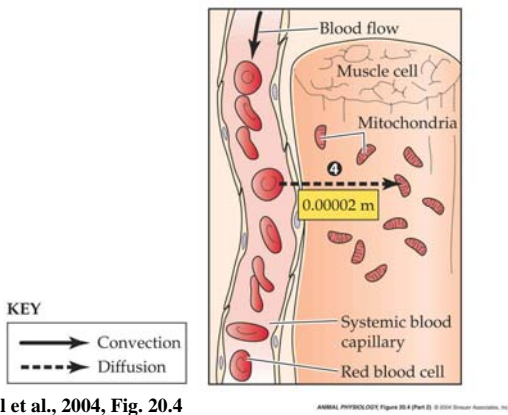
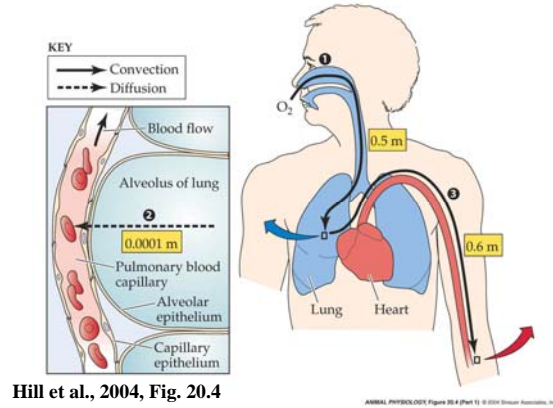
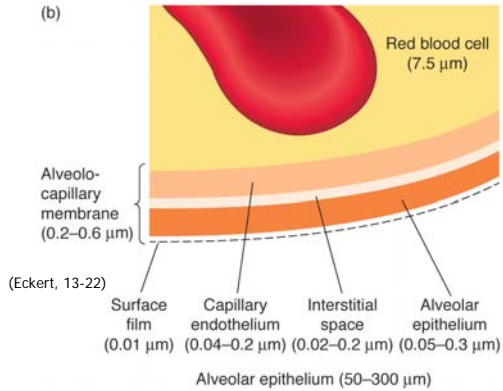
- Nonrespiratory
- Trachea ->
  - Bronchi ->
  - Bronchioles ->

- Respiratory
- Terminal bronchioles ->
  - Respiratory bronchioles ->
  - Alveoli

-Cilia and Mucus



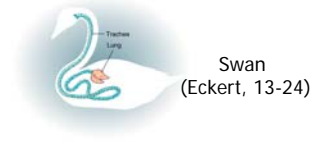
-Gas Diffusion Barriers:



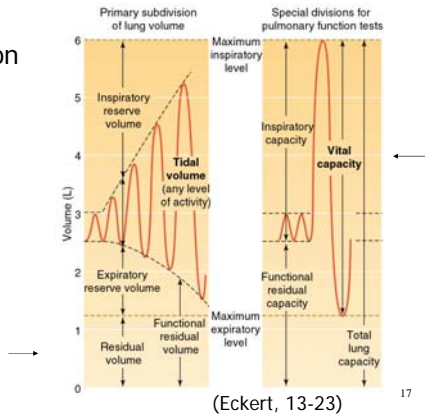
Lung Ventilation

-Small mammals with greater per gram  $O_2$  needs and therefore greater per gram respiratory surface area?

-Dead Space (anatomic and physiological)



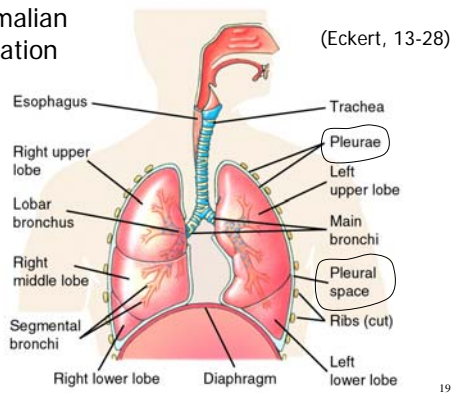
Lung Ventilation



Mammalian Ventilation

- lungs are elastic bags
- suspended in pleural cavity within thoracic cage (ribs and diaphragm define, fluid lines)
- low volume pleural "space" between lung and thoracic wall
- negative pressure to inflate lungs (increase volume)
- pneumothorax

## Mammalian Ventilation

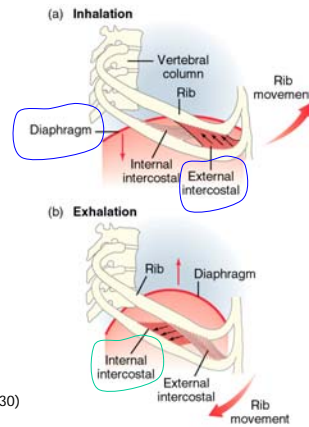


(Eckert, 13-28)

19

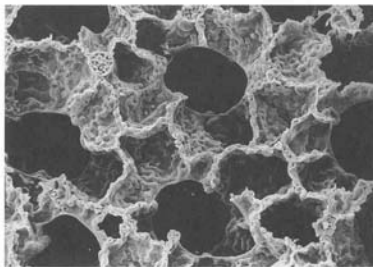
## Mammalian Ventilation

-expiration usually passive



(Eckert, 13-30)

20



**MAMMALIAN LUNG** Scanning electron micrograph of the lung structure of a wildebeest (*Connochaetes taurinus*). The capillaries, filled with red blood cells, are visible as a bulging network in the alveolar walls. The distance across the photograph corresponds to about 0.5 mm in the lung. [Courtesy of Ewald Weibel, University of Berne, Switzerland]

## Mammalian Lung



Alveoli and Capillaries

RBC (not to scale)

Knut Schmidt\_Nielsen 1997

## Bird Lung Ventilation

Unidirectional!!!

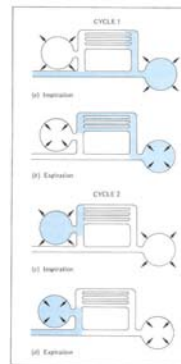
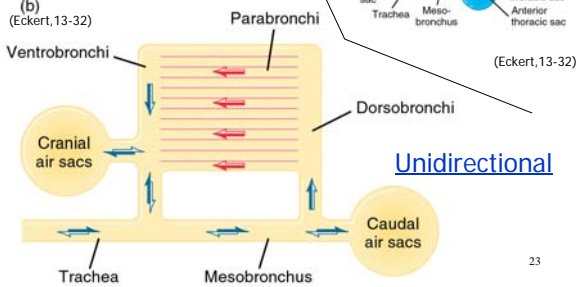


Figure 1.26 The movement of a single inflated volume of gas through the avian respiratory system. It takes two full respiratory cycles to move the gas through its complete path. (Brett and Schmidt-Nielsen 1972)

Knut Schmidt\_Nielsen 1997

## Bird Ventilation

-lung volume changes very little, air sacs instead



(Eckert, 13-32)

(Eckert, 13-32)

Unidirectional

23

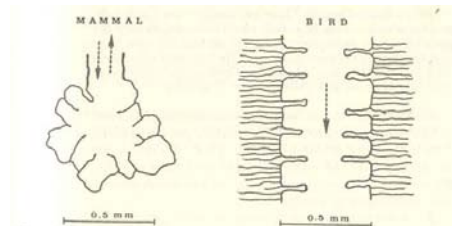


Figure 1.8. The smallest units of the lung of mammals end in blind sacs, alveoli. In birds the finest branches of the bronchi (the parabranchi) are tubes which are open at both ends and permit through-flow of air.

Knut Schmidt\_Nielsen 1972

Mammal Lung  
Alveoli

Bird Lung  
Parabranchi

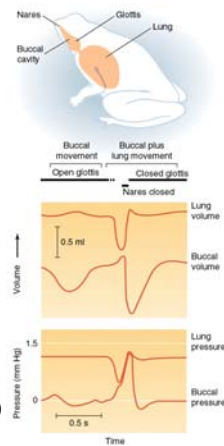
24

## Frog Ventilation

-Positive pressure ventilation

1. Into mouth (buccal cavity)
2. Close nares, open glottis and force air into lungs by raising buccal floor

(Eckert, 13-33)



## Pulmonary Surfactants

- Reduce liquid surface tension in alveoli
- Allows for compliance and low-cost expansion of lung
- Lipoproteins
- keep alveoli from getting stuck closed  
Atelectasis = collapsed lung
- premature babies may need artificial surfactant

26

## Panting Dogs?

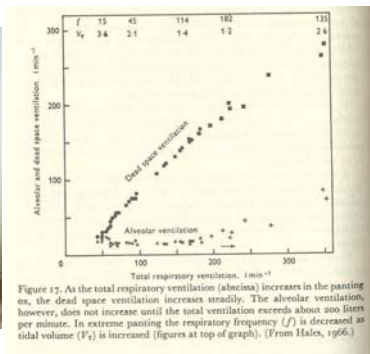
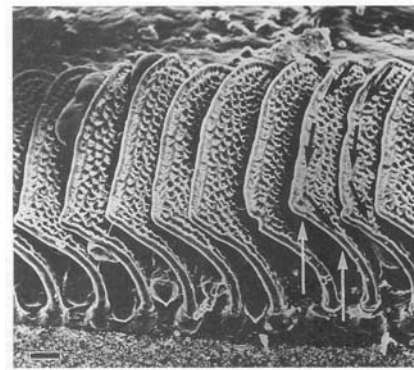


Figure 17. As the total respiratory ventilation (abscissa) increases in the panting dog, the dead space ventilation increases steadily. The alveolar ventilation, however, does not increase until the total ventilation exceeds about 200 liters per minute. In extreme panting the respiratory frequency ( $f$ ) is decreased as tidal volume ( $V_t$ ) is increased (figures at top of graph). (From Hales, 1966.)

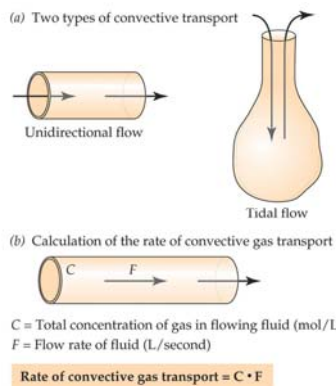
Knut Schmidt\_Nielsen 1972



FISH GILL Scanning electron micrograph of gill filaments of a sturgeon (*Acipenser transmontanus*). White arrows show the direction of water flow and black arrows the direction of blood flow. The bar in the lower left corner represents 0.05 mm. (Burggren et al. 1979; courtesy of Warren W. Burggren, University of Massachusetts)

Fish Gill

Knut Schmidt\_Nielsen 1997



Hill et al., 2004, Fig. 20.3

ANIMAL PHYSIOLOGY, Figure 20.3 © 2004 Sinauer Associates, Inc.