The Journal of Clinical Investigation

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J Clin Invest. 1930;8(3):317-323. https://doi.org/10.1172/JCI100266.





STUDIES IN CONGESTIVE HEART FAILURE

III. THE BUFFERING POWER OF THE BLOOD AND TISSUES

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(Received for publication July 12, 1929)

The results reported in a previous paper of this series (Harrison and Pilcher, 1930) led us to accept tentatively the hypothesis of Eppinger, Kisch and Schwarz (1927) that the buffering power of the tissues is diminished in congestive heart failure.

Since the evidence from our previous work was indirect, it seemed wise to study the matter further by more direct methods. Accordingly, the pH of the blood, determined by Cullen's (1922) method, using the micro-modification of Hawkins (1923) and the bicolor standards of Hastings and Sendroy (1924), and the carbon dioxide content of the blood, determined by analysis in the Van Slyke-Neill (1924) constant-volume apparatus were studied in patients with cardiac failure and in normal subjects, in four series of observations.

- a. The pH and carbon dioxide content of the blood at rest was found to be usually within normal limits in both the "decompensated" patients and the control subjects, as shown in the left-hand parts of tables 2, 3 and 4. However, that a state of "cardiac acidosis" may exist is shown in table 1.
- b. The effect of the oral administration of 15 to 17.35 grams of ammonium chloride over a period of 12 hours was found to be approximately the same on the arm venous blood (drawn without stasis) of normal and "decompensated" individuals. The data are shown in table 2. These observations were suggested by the work of J. B. S. Haldane (1924), who showed that ammonium chloride given in large amounts by mouth caused a diminution in the alkaline reserve of the blood.

c. The effect of breathing a mixture of 5 per cent carbon dioxide and 95 per cent oxygen was found to be slightly greater on the pH of the blood of three of the four "decompensated" patients than on that of any of the normal subjects. The carbon dioxide content was usually increased somewhat in both controls and patients (table 3). One patient (Ada D.) who had nephrosis with massive edema, but no evidence of heart disease, showed the greatest diminution in pH found.

TABLE 1
Spontaneous acidosis in patients with heart failure

Patient	Blood from	CO ₂ content of blood	Clinical data
		volumes per cent	
	Artery	33.0	Marked dyspnea; massive edema (weight 33 pounds greater than when edema-free)
A. R. {	Artery	39.3	5½ hours later; 3 hours following administration of 20 grams sodium bicarbonate. Dyspnea di- minished
	Artery	31.1	August 4, 1928. Luetic aortic insufficiency; severe pulmonary edema; Cheyne-Stokes respiration; cyanosis of fingers; massive edema
н. с.	Artery	61.4	August 5, 1928. Greatly improved. Edema diminished; respirations rapid but regular. (Diuresis following "Salyrgan")
	Artery	65.2	August 7, 1928. Has broncho-pneumonia and severe anoxemia (47.5 per cent arterial oxygen saturation). Edema stationary
R. R.	Arm vein	30.8	Aortic insufficiency; terminal heart failure; blood drawn a few minutes before death; pH 7.08

d. The effect of exercise was determined in three patients with no circulatory disorder, in one patient with heart disease but neither dyspnea nor edema, and in five patients with varying degrees of "decompensation." Blood was drawn from the femoral vein of the subject before and within 30 to 60 seconds after a standardized exercise of the leg. Observations were made on both legs (with an interval between) of one control subject and of two patients with congestive failure.

TABLE 2

-0.07 Luetic aortic regurgitation; massive edema; slight -0.04 Hypertensive heart disease; no dyspnea; slight ease; marked Cheyne-Stokes respiration; no -0.11 Hypertensive heart disease; paroxysmal dyspnea; -12.6 7.11 |-0.15 Hypertensive heart disease; luetic aortitis; slight -0.04 Hypertensive heart disease; paroxysmal dyspnea; Hypertensive heart disease; chronic pulmonary disslight edema of lower half of tibia. (Duration, Hypertensive heart disease; auricular fibrillation; no edema. Liver enlarged. Vomited 3 times No dyspnea or edema. No vomiting Clinical data slight edema; slight dyspnea The effect of ammonium chloride on the carbon dioxide content of the arm venous blood edema and dyspnea No edema or dyspnea Normal controls dyspnea 6 days) edema -0.11 -0.08 -0.03 -0.12-0.03Change Arm venous blood after NH,Cl -15.4 7.23 -14.7 7.27 7.23 -16.4 7.33 7.36 7.20 -6.4 | 7.22-9.2 | 7.40-6.5|7.25-14.5 7.31 펁 -6.7 -12.7 -16.8-16.2Change 38.0 CO₂ 41.7 44.0 40.1 33.2 33.9 38.8 volumes per cent 35.8 42.3 38.4 36.8 17.35 17.35 17.35 17.35 15.00 15.00 15.00 15.00 19.00 15.00 15.00 15.00 NH'C grams in 12 hours 7.40 7.33 7.36 7.30 7.36 7.34 7.30 7.37 7.32 Arm venous blood before NH.Cl 7.33 7.39 Ηd 56.6 CO₂ 48.7 50.0 54.6 volumes per cent 53.4 50.4 47.2 47.5 47.6 55.2 50.1 43.5 February 18, 1929 February 10, 1929 February 17, 1929 February 4, 1929 February 5, 1929 March 12, 1929 March 10, 1929 March 12, 1929 March 12, 1929 March 12, 1929 March 12, 1929 March 4, 1929 Date M. A. M. T. Subject G. C. T. H. D. K. C. P. A. A. A. D. T. P. F. C.

	The effect of br	eathing	5 per ce.	nt carbo	n dioxi	de on th	e pH an	ed carbo	n dioxid	The effect of breathing 5 per cent carbon dioxide on the pH and carbon dioxide content of the arm venous blood
		Ä	Breathing air	ir		Breathin	Breathing 5 per cent CO2	ent CO2		
Subject	Date	pH of arm venous blood	CO2 content of arm venous blood	Total venti- lation	pH of arm venous blood	Change	COrcontent of arm venous blood	Change	Total venti- lation	Clinical data
			volumes per cent	cc. per minute			volumes per cent		cc. per minute	
G. C.	March 26, 1929 March 29, 1929	7.33	46.0	5,800 7.32 4,440 7.33	7.32	-0.01 47.4 -0.05 44.3	47.4	+1.4	+1.411,280) Normal controls
C. P.	March 17, 1929	7.32	46.7		7.29	-0.03	48.4	+1.7	+1.711,160	
A. A.	March 19, 1929	7.30	53.1	6,260	7.23	6,260 7.23 -0.07 52.6	52.6	-0.5	-0.5 19,840	Hypertensive heart disease; no edema or dyspnea for 6 weeks previously
A. D.	March 20, 1929	7.34	46.6		7.27	6,940 7.27 -0.07 48.0	48.0	+1.4	+1.414,200	Hypertensive heart disease; no edema or dyspnea for 4 weeks previously
T. P.	March 27, 1929	7.26	47.3	6,840 7.26	7.26		50.4	+3.1	17,680	+3.1 17,680 Luetic aortic regurgitation; no dyspnea; barely perceptible edema
T. B.	March 24, 1929	7.44	47.3	3,650	7.39	3,650 7.39 -0.05	48.8	+1.5	10,540	+1.5 10,540 Hypertensive heart failure; edema of feet and ankles, back of legs and thighs. Disoriented mentally
Ada D.	March 24, 1929	7.39	44.0	5,360 7.31	7.31	-0.08 46.1	46.1	+2.1	11,760	+2.1 11,760 Nephrosis; general anasarca; massive edema of 5 weeks duration. No evidence of heart disease

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The effect of exercise on the pH and on the carbon dioxide content of the femoral venous blood	Clinical data		Gastric ulcer. No circulatory disease	Lung abscess (surgically drained). No circulatory disease	Mild diabetes mellitus; moderate arteriosclero-	sis. No other circulatory symptoms	Mild diabetes mellitus; hypertension cardiac	enlargement; no dyspnea or edema	Hypertensive heart disease; moderate edema of	feet and lower halves of legs; no dyspnea at	rest	Luetic aortic regurgitation; massive edema		Hypertensive heart disease; very slight edema	(first "break" in compensation); duration, 4	days	Hypertensive heart disease; very slight noc-	turnal dyspnea until 3 days previously; slight	edema; 5th "break" in compensation	Hypertensive heart disease. General ana-	sarca; moderate dyspnea; Cheyne-Stokes	respiration
TABLE 4 ve carbon	Change	'	-0.05	-0.04	-0.03	;	40.04	4 0.0 4	-0.07			-0.18	-0.15	-0.05			-0.13			-0.18	<u>+0.0</u>	
and on th	ral venous	After exercise	7.30	7.36	7.40		7.24	7.22	7.16			7.24	7.25	7.31			7.25			7.32	7.34	
on the pH	pH of femoral venous blood	Before exercise	7.35	7.40	7.43		7.28	7.26	7.23			7.42	7.40	7.33			7.38			7.40	7.38	
of exercise	Leg						≃,	٦				2	1			•				æ	1	
The effect o	Date		February 21, 1929	February 28, 1929	March 2, 1929		January 31, 1929		February 2, 1929		\	March 11 1020	Maich 11, 1727	March 2, 1929			February 24, 1929				March 10, 1929	
	Patient	Patient		C. I.	J. W.		I. H.		A. A.			Ę	:	M. G.			A. D.				D. K.	

In five of the seven observations on "decompensated" patients, the diminution in pH was greater than any diminution found in a control subject. The average diminution in the controls was 0.04 pH and in the decompensated patients, 0.11 pH. With one exception (table 4, D. K., left leg), the degree of change in pH was proportional to the extent or duration of the edema, or both.

The changes in carbon dioxide content of the blood varied greatly in both control and "decompensated" subjects and these data are not included in table 4. Such variation is to be expected, since the carbon dioxide content of the blood during exercise varies with three different factors: the oxidation occurring in the muscles, the lactic acid "overflowing" into the blood stream and the minute ventilation.

SUMMARY AND CONCLUSION

The pH and carbon dioxide content of the blood of normal subjects and of patients with congestive heart failure has been studied before and after (a) administration of large doses of ammonium chloride, (b) breathing 5 per cent carbon dioxide and (c) a standardized exercise.

The findings in patients with heart failure at rest were usually within normal limits, but a state of acidosis was found in three patients with very severe symptoms.

The changes occurring after administration of ammonium chloride and after breathing carbon dioxide were usually within normal limits in patients with congestive failure.

The changes in pH after exercise were usually greater in "decompensated" patients than in control subjects and the degree of change was, with one exception, proportional to the extent and duration of the edema. This is believed to be compatible with the presence of diminished buffering power of the tissues in heart failure.

One compensated cardiac patient showed changes similar to those found in control subjects and one patient with "non-cardiac" edema showed changes similar to those in the "decompensated" patients. This suggests that edema itself may be related to changes in the tissues of patients with heart failure.

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