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STUDIES ON ALCOHOL DIURESIS. II. THE EVALUATION OF ETHYL ALCOHOL AS AN INHIBITOR OF THE NEUROHYPOPHYSIS^{1, 2}

By CHARLES R. KLEEMAN, MILTON E. RUBINI,³ EZRA LAMDIN,⁴ AND FRANKLIN H. EPSTEIN

(From the Department of Internal Medicine, Yale University School of Medicine, New Haven, Conn.)

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In the normally hydrated semi-recumbent individual, alcohol causes a rise in urine flow that is characterized by an increase of free water clearance (C_{H_2O}) and a decreased excretion of sodium, potassium and chloride (1). The evidence to date strongly suggests that the rise of C_{H_2O} is caused by inhibition of the release of antidiuretic hormone.

The present study was undertaken to evaluate the effect of alcohol on the excretion of water and solutes in physiologic states in which alterations in the activity of antidiuretic hormone have been demonstrated or suggested. The following states were studied: 1) Minimal antidiuretic hormones (ADH) activity produced by sustained positive loads of water; 2) Increased ADH activity produced by infusions of hypertonic sodium chloride solutions, a) during water diuresis with high urine flows (10 to 15 cc. per min.), and b) in subjects with low urine flows (1 to 2 cc. per min.); 3) Increased antidiuretic activity produced by venous congestion of the limbs.

MATERIALS, METHODS AND RESULTS

Subjects were normal males, aged 25 to 32. No control of diet prior to the day of study was attempted. One to one and a half hours after a light breakfast the subjects voided and reclined in a semi-recumbent position. All studies were begun at 8:30 to 9:00 A.M.; diurnal variations in urinary flow and composition (2) were therefore presumably similar in all experiments. Alcohol was given as 120 cc. of 100 proof bourbon whisky imbibed over a 10-minute period. Techniques for collection of blood and urine and chemical methods have been described in the previous paper (1). In all studies insensible water loss was assumed to be approximately 50 cc. per hour. Changes in extracellular

space were calculated approximately from changes in the chloride space (3), assuming an initial extracellular volume of 20 per cent of body weight. Changes in plasma volume were calculated from changes in hemoglobin and hematocrit (1). Urine flow was divided into two fractions:

Osmolar clearance (C_{osm})

$$= \frac{\text{milliosmols per kilo of urine}}{\text{milliosmols per kilo of plasma}} \times \text{urine flow (cc. per min.)}$$

$$\text{Free water clearance } (C_{H_2O}) = \text{urine flow} - C_{osm}$$

Group I. Effect of alcohol during water diuresis (Table I, Figure 1C)

A positive water balance was induced in two semi-recumbent subjects by drinking one liter of water, and was maintained by infusing 4 per cent fructose solution intravenously and administering supplemental water by mouth. The accuracy of this technique was checked by weighing the subject at the beginning and the end of each experiment. Fructose solution was chosen because of its minimal effect on the total hexose in the blood. By limiting the rate of infusion to 8 cc. per minute or less, no reducing substances could be detected in the urine by qualitative test with Benedict's solution. After a maximal steady urine flow had been maintained for at least two 30-minute periods, alcohol was imbibed. Urine was collected at 15 to 30-minute intervals during the next three hours.

Under these circumstances, alcohol did *not* induce a further increase in urine flow or C_{H_2O} (Figure 1C). If large positive loads of water (1000 cc.) completely inhibit ADH release ("physiologic diabetes insipidus"), this result would be expected.⁵ The rates of excretion

⁵ The statement that maximum water diuresis is associated with complete inhibition of ADH release or so-called "physiologic diabetes insipidus" probably is true for the recumbent and semi-recumbent positions only. In unpublished experiments the authors have demonstrated that when a positive water load of 1000 cc. is maintained, the maximum urinary flow and free water clearance (C_{H_2O}) attained in the standing or 45° position were further increased by lying down. This suggests a continual "tonic" release of ADH in the upright positions in spite of the sustained water load or non-hormonal factors blocking the maximum rise in urinary flow.

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³ Major, MC, USA.

⁴ Postdoctorate Research Fellow of the U. S. Public Health Service.

TABLE I
Group I—1000 cc. positive H₂O load with and without alcohol

Subject	Time	Urine										Blood															
		pH					Excretion rates/min.					Clearances/min.					Osmolarity										
		Min.	Units	μEq.	μEq.	μOsm.	Na	K	Cl	NH ₄	TA	Osm.*	Vol.	cc.	Osm.	Free	Creat.	Na	K	Cl	Alc.	pH	Obs.	Corr.*	PV†	PV†	
K.	73	6.5	113	79	121	19	4	650	1.6	2.2	-0.6	126	146.7	4.64	105.0	0	303	303									
	32	6.1	98	50	110	40	9	945	9.8	3.2	6.6	130															
	30	6.1	88	41	111	36	7	820	10.0	2.8	7.2	121															
	32	6.2	93	41	107	38	7	865	10.3	2.9	7.4	127															
	27	6.3	114	38	107	35	9	880	10.5	3.0	7.5	132															
Exp. 1	31†	5.9	120	33	129	39	10	890	10.9	3.1	7.8	148	140.6	4.78	99.5	290	290										
	31	5.9	127	33	123	37	7	840	10.3	2.9	7.4	145	142.1	3.50	101.2	303	288										
	28	5.8	122	19	122	36	7	840	10.2	2.9	7.3	139	142.1	3.30	102.5	313	287										
	32	5.8	105	16	98	37	9	785	9.5	2.7	6.9	145	142.2	3.07	100.7	314	285										
	31	6.2	101	14	93	34	8	795	9.7	2.7	7.0	140	142.0	3.49	103.0	312	290										
H.	57	6.0	149	49	157	33	11	860	0.6	3.0	-2.4		140.5	4.00	105.5	0	288	288									
	38	6.9	162	87	180	28	4	900	2.0	3.1	-1.2																
	18	6.6	100	104	133	27	10	1,030	7.8	3.6	4.2	132															
	24	6.6	113	94	117	25	16	1,020	11.3	3.5	7.8	138															
	27	6.3	96	71	94	28	8	930	12.2	3.2	9.0	130															
Exp. 2	37	6.3	72	43	64	28	7	770	9.9	2.7	7.2	121															
	37	6.2	79	38	66	30	8	760	9.7	2.7	7.0	128	137.0	3.90	101.5	0	284	284									
	51†	6.2	79	32	60	30	16	670	8.8	2.4	6.4	125	138.0	4.12	101.5	105	280	280									
	43	5.9	87	20	64	31	20	720	9.4	2.6	6.8	136															
	40	6.1	89	21	62	31	21	710	9.1	2.5	6.6	136															
41	6.0	78	21	48	27	19	650	8.2	2.3	5.9	122	137.0	3.92	101.7	70	301	280										

* Observed osmolarity—osmolar contribution of alcohol. † PV = Plasma volume. ‡ Alcohol imbibed during first 10 minutes of this period.

of Na and Cl did not decrease after alcohol ingestion and although the urine became more acid, excretion of ammonium was not enhanced. These results contrast sharply with the findings when alcohol is administered to subjects with low or moderate urinary flows (1).

Group IIa. Effect of the simultaneous administration of alcohol and hypertonic saline to water-loaded subjects (Table IIA, Figure 1A and 1B)

Positive water balances were achieved in four experiments in a manner similar to that described for Group I. In two control tests (No. 1 and No. 2 Table IIA) 500 cc. of hypertonic saline (5 to 6 per cent) was infused without alcohol. In two further experiments (No. 3 and No. 4, Table IIA), alcohol was imbibed simultaneously with the beginning of the hypertonic infusion.

The administration of hypertonic saline without alcohol, at the height of a water diuresis, was followed by a prompt decrease in urine flow and free water clearance (Figure 1A). In contrast, the subjects receiving alcohol not only failed to show an antidiuresis, but actually increased their flow of urine above the levels reached during maximal water diuresis (Figure 1B). Free water clearance (CH₂O) increased in spite of a 4 to 5 per cent rise in the osmolarity of the serum. It is apparent that alcohol effectively blocked the antidiuretic response to hypertonic saline.

Group IIb. Effect of alcohol on the antidiuresis following hypertonic saline in subjects with low urine flows (Table IIB, Figure 2)

In two subjects, 300 cc. of 5 to 6 per cent saline were infused intravenously after a suitable control period. Approximately 30 minutes after starting the infusion alcohol was imbibed.

In neither subject did a water diuresis occur after alcohol. Prior administration of hypertonic saline, with a consequent increase in the effective osmotic pressure of extracellular fluid, presumably induced the release of increased amounts of ADH from the posterior pituitary (4). Since alcohol affects neither exogenous ADH nor the ability of the tubules to respond to this hormone (5) an excess of circulation ADH probably masked the inhibitory effect of alcohol upon the supraopticohypophysial system in these experiments.

Group III. Effect of alcohol on the antidiuresis of venous congestion (Table III, Figure 3)

The effect of alcohol on the antidiuresis produced by venous congestion of the limbs was tested in three subjects in whom sphygmomanometer cuffs were inflated about the thighs to a pressure of 70 to 80 mm. Hg. (In all, a positive water load of 500 cc. was established and maintained throughout the experiment.) In two studies (No. 1 and No. 2, Table III) a control period of venous congestion for 30 minutes, instituted after maximal urine flow had been attained, produced a prompt fall in urine flow, CH₂O, and C_{osm}, as well as in the rates of excretion of sodium, potassium, chloride, and creatinine. These

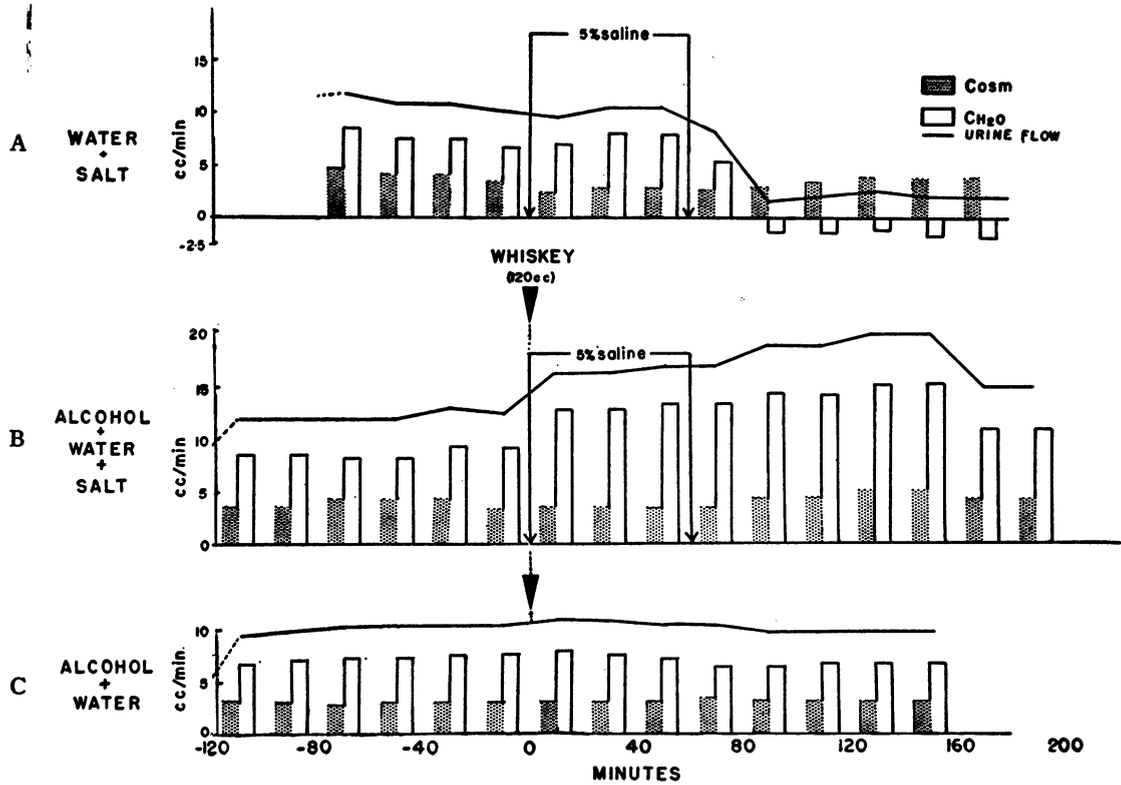


FIGURE 1

Alcohol produced no increase in an established water diuresis (Figure 1C). In contrast, when alcohol was given with an intravenous load of hypertonic saline, urine flow and CH_2O increased (Figure 1B), and the characteristic antidiuretic effect of hypertonic saline (Figure 1A) was blocked.

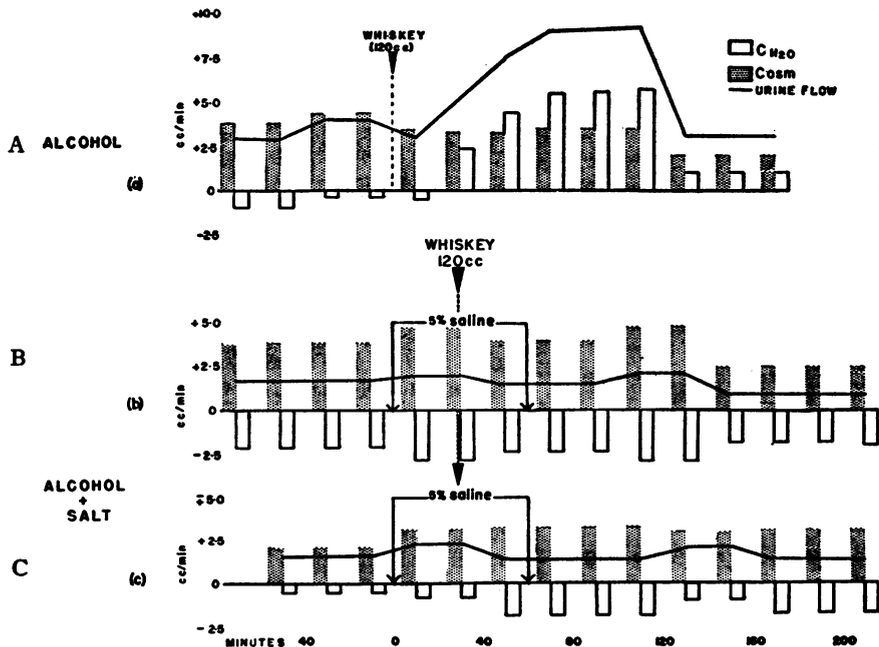


FIGURE 2

Prior administration of hypertonic saline (Figures 2B and 2C) blocked the characteristic diuresis following alcohol (Figure 2A).

TABLE IIA
Group Ila—Hypertonic salt administration with and without alcohol in subjects with maximum water diuresis

Subject	Time	pH	Urine										Blood									
			Excretion rates/min.					Clearances/min.					Na	K	Cl	CO ₂	Alc.	pH	Osmolarity		PV [†] / PV _i	Cl space
			μEq.	μEq.	μEq.	μEq.	cc.	cc.	cc.	cc.	cc.	cc.							mEq./L.	mEq./L.		
KI.	117	7.0	188	78	169	17	2.3	1,640	16.5	7.3	9.2	126	144.7	4.08	108.5	290	290	0	14.1			
Exp. 1	19	6.3	414	147	242	42	16.5	1,640	16.5	7.3	9.2	141										
	15	6.4	457	146	272	34	16.6	1,580	18.3	7.7	10.6	118										
	17	5.6	415	104	160	45	15.80	1,990	13.2	6.8	12.3	123										
	20	5.6	261	78	45	54	19.00	1,930	13.2	4.7	8.5	119										
	38	5.5	215	82	40	57	14.1	855	10.6	4.1	7.5	121										
	30	5.9	160	61	62	46	8.55	885	9.1	3.5	6.6	125										
	20	5.8	132	48	68	57	4.2	810	11.0	2.9	7.0	121										
	36	5.9	277	31	194	32	8.00	8.1	2.7	5.4	110	142.0	4.30	111.4	289	289	-5	15.4				
	20	295	38	269	35	1.6	875	1.6	3.0	1.4	123	146.0	4.30	114.8	289	289	0	16.2				
	24	308	70	371	39	1.05	2.7	3.8	1.4	1.27	127											
25	354	106	372	38	1.085	2.1	3.8	1.1	1.27	127												
21	328	120	385	38	1.085	2.1	3.8	1.1	1.27	127												
31	324	123	371	37	1,110	2.1	3.9	1.8	1.23	123												
109																						
Re.	57	6.6	257	163	328	52	10	1,335	6.8	4.6	2.2	156	143.8	4.15	112.7	288	288	-2	16.2			
Exp. 2	34	6.8	271	201	361	43	10	1,325	12.9	5.0	7.9	152	143.0	4.59	105.0	290	290	0				
	40	6.2	198	127	147	47	10	1,420	17.1	3.9	13.2	140										
	47	6.5	138	82	124	46	9	1,145	16.8	3.6	13.2	144	139.3	4.38	101.0	273	273	+3				
	68	6.1	197	90	182	39	9	1,320	15.2	3.3	11.9	140										
	46	6.0	288	86	304	42	7	1,220	11.1	3.4	7.7	138	147.2	4.26	110.4	332	332	+8				
	37	6.4	304	122	304	49	8	1,335	11.0	4.2	6.8	142										
	31	6.2	347	122	416	40	8	1,340	9.4	4.6	4.8	139	143.5	4.61	107.6	297	297					
	31	6.4	346	125	390	33	8	1,535	11.0	5.2	5.8	145										
	109																					
	111	7.5	435	205	477	24	11	1,685	7.1	6.0	1.1	161	133.0	4.20	101.5	283	283					
Exp. 3	56	7.2	358	163	379	22	1	1,470	6.9	5.2	1.7	155										
	35	6.3	165	89	151	25	4	970	12.0	3.6	8.4	142										
	40	6.3	195	55	84	36	9	1,100	12.0	3.9	8.1	132										
	30	6.3	196	31	64	42	6	1,050	13.0	3.8	9.2	140										
	24	6.6	160	20	68	37	13	885	12.3	3.2	9.1	140	133.0	3.68	100.0	276	276	+11	14.3			
	41	6.6	295	36	194	35	6	990	16.2	3.5	12.7	150	133.9	3.18	108.0	302	302	+21	14.6			
	33	6.6	231	11	165	39	15	970	16.7	3.3	13.4	166	144.8	3.43	113.0	311	311	+32	14.8			
	36	6.5	304	23	188	35	8	1,250	18.7	4.3	14.4	135										
	37	6.7	332	180	266	6	9	1,420	20.0	4.9	15.1	135	135.3	4.20	107.5	296	296	+24	15.4			
	43	6.2	279	15	138	35	9	1,195	14.9	4.1	10.8	138										
Exp. 4	65	7.1	174	90	167	22	0	960	3.7	3.1	0.6	117	139.1	4.30	98.5	307	307		15.6			
	27	6.6	171	59	126	24	0	1,055	11.1	3.5	7.6	121										
	23	5.5	166	71	75	43	27	1,245	11.5	4.1	7.4	128										
	17	5.3	237	47	61	54	19	1,530	13.8	5.1	8.7	111										
	19	5.3	275	28	85	51	18	1,660	14.5	5.0	9.5	108										
	20	5.2	234	26	73	50	8	1,285	13.3	4.3	9.0	115	139.8	4.32	95.0	302	302	+4	16.0			
	19	5.5	203	21	93	43	11	1,130	13.7	3.9	9.8	111										
	25	5.4	280	28	148	31	7	1,130	14.3	3.7	10.6	109										
	31	5.5	239	21	177	26	6	1,125	14.8	3.6	11.2	123	145.5	4.10	103.0	326	312		16.1			
	25	6.0	309	20	257	23	6	1,335	16.7	4.2	12.5	110	145.3	4.31	103.5	340	322		16.9			
18	5.8	513	21	402	16	25	1,625	16.6	5.0	11.6	122											
17	5.8	616	25	493	37	37	1,740	17.2	5.4	11.8	117	143.3	4.23	105.2	338	318	+10	16.5				
21	5.5	452	20	399	35	22	1,405	15.8	4.4	11.1	117	143.3	4.23	105.2	338	318		16.5				
20	5.5	526	26	401	36	22	1,555	15.5	4.9	10.9	118	143.3	4.08	105.0	332	316		16.3				
21	5.5	535	15	368	45	28	1,570	16.2	5.0	11.2	116											
30		317	11	263	36	18	1,065	6.7	3.4		3.3											

* Observed osmolarity—osmolar contribution of alcohol.
 † PV = Plasma volume.
 ‡ Alcohol imbibed during first 10 minutes of this period.
 § Period of infusion of hypertonic salt.

II. ETHANOL INHIBITION OF THE NEUROHYPOPHYSIS

TABLE III
Group III—Venous congestion of the lower extremities with and without alcohol

Subject	Time	pH	Urine						Blood						PV _f † PV _i	% Δ				
			Excretion rates/min.			Clearances/min.			pH	Alc.	CO ₂	Cl	K	Na			mEq./L. mEq./L. mEq./L.	mOsm./L.	mOsm./L.	Corr.*
			μEq. μEq. μEq.	μEq. μEq. μEq.	cc. cc. cc.	Osm. Free Creat.														
μEq.	μEq.	μEq.	μEq.	μEq.	μEq.	cc.	cc.	cc.	cc.	cc.	cc.	mEq./L.	mEq./L.	mEq./L.	mEq./L.	mEq./L.	mEq./L.	mEq./L.		
N. Exp. 1	76	6.8	241	148	226	20	20	984	1.4	3.5	-2.1	166	140.0	4.10	105.3	280	280	280	0	
	77	7.0	39	168	32	23	850	12.4	3.2	6.2	164	140.6	4.21	104.0	275	275	272	+5		
	26	6.8	196	62	170	23	955	16.0	3.5	12.5	160	138.0	4.05	103.3	272	272	272			
	21	6.9	210	57	173	31	1,000	20.0	3.0	16.3	159									
	36	6.9	167	42	153	23	820	8.0	3.0	5.0	159									
	46	6.7	145	30	88	24	675	9.6	2.5	7.1	130									
	22	6.7	211	45	141	31	1,010	20.7	3.7	17.0	184									
	17	6.7	219	41	146	29	1,000	19.4	3.7	15.7	179									
	20	6.8	228	39	152	28	1,060	20.8	3.9	16.9	173									
	25†	6.9	236	37	160	27	915	19.0	3.4	15.6	145									
	25	7.0	248	31	174	20	932	19.4	3.5	15.9	147									
	28	6.3	161	22	111	51	720	15.0	2.7	12.3	146									
36	6.4	168	24	120	61	845	17.6	3.2	14.4	133										
26	6.5	115	18	114	65	810	16.9	3.1	13.8	130										
20	6.6	107	19	88	18	730	15.2	2.8	12.4	134										
20	6.6	122	20	91	22	825	17.2	3.1	14.1	137										
W. Exp. 2	65	6.8	245	219	314	31	1,230	6.6	4.4	2.2	160	139.4	3.70	103.7	280	280	280	0		
	47	6.8	268	123	298	26	1,190	13.2	5.3	7.9	168									
	26	6.5	342	152	392	33	1,480	16.8	5.4	11.4	166									
	35	6.5	350	171	399	22	1,650	17.5	6.1	11.4	162									
	32	6.5	221	108	274	33	1,060	10.9	3.9	7.0	151									
	42	6.3	178	75	204	35	870	9.6	3.2	6.4	131									
	64	6.5	189	63	204	30	816	8.9	3.0	5.9	148									
	23†	6.7	273	89	288	31	1,015	11.3	4.1	7.2	155									
	21	6.6	274	63	271	33	940	11.0	4.0	7.0	151									
	32	6.9	222	44	222	37	750	8.9	2.9	6.0	122									
	31	6.2	222	32	225	40	770	8.9	3.0	5.9	145									
	32	6.0	191	19	184	46	700	8.1	2.7	5.4	140									
38	6.1	194	25	186	33	845	9.4	3.2	6.2	158										
31	6.2	171	16	156	34	825	9.8	3.1	6.7	166										
S. Exp. 3	65	6.6	175	79	179	20	905	1.2	3.1	-1.9	145	145.6	3.96	105.8	26.9	292	292	0		
	34	6.9	267	146	278	26	1,300	8.1	4.5	3.6	124									
	21	6.9	287	164	290	25	1,430	14.8	5.0	9.8	124									
	19	6.8	332	174	333	22	1,575	17.9	5.5	12.4	129									
	20	6.7	345	185	354	25	1,615	19.7	5.6	14.1	133									
	18	6.7	356	181	365	24	1,615	20.3	5.6	14.7	134									
	24	6.6	289	154	290	30	1,395	18.1	4.9	13.2	122									
	17	6.5	256	141	262	31	1,305	17.6	4.6	12.0	131									
	18	6.3	199	113	204	33	1,115	15.9	3.9	12.0	134									
	22	6.3	185	106	185	35	1,065	15.4	3.7	11.7	141									
	18	6.3	173	88	162	33	1,035	15.0	3.5	11.5	145									
	21	6.2	168	78	139	38	1,028	14.2	2.9	11.3	143									
21	5.4	162	50	97	28	815	10.5	2.6	7.9	127										
20	5.5	152	44	94	27	745	10.5	2.8	7.7	134										
20	5.6	110	28	66	28	800	10.8	2.6	8.2	139										
20	5.6	106	21	65	27	740	10.9	2.5	8.4	131										
17	5.6	88	24	45	28	720	10.6	2.5	8.1	139										
21	5.6	94	21	48	30	740	11.2	2.6	8.6	143										
20	5.8	92	20	48	28	720	11.3	2.5	8.8	134										
20	5.9	112	20	54	26	750	12.5	2.6	9.9	139										
20	6.0	122	21	62	24	755	13.0	2.6	10.4	136										
20	6.9	133	22	75	28	755	14.0	2.6	11.4	141										

* Observed osmolarity—osmolar contribution of alcohol.

† PV = Plasma volume.

‡ Alcohol imbibed during first 10 minutes of this period.

§ Period venous congestion.

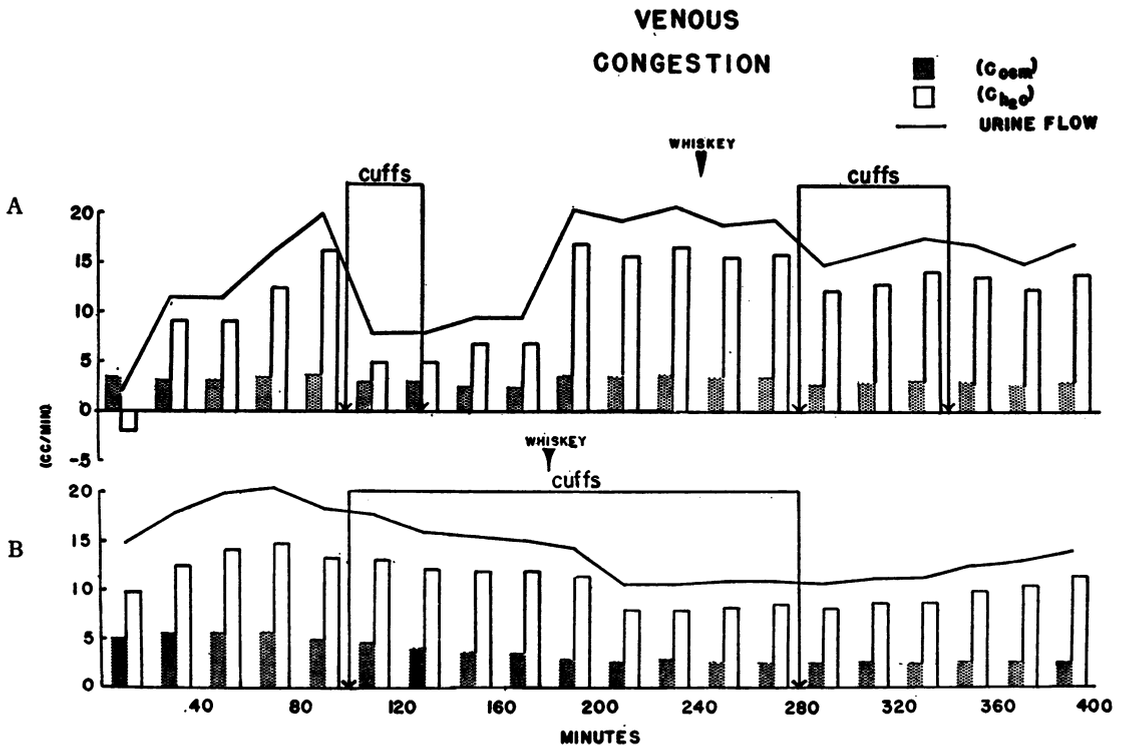


FIGURE 3

Alcohol minimized the antidiuretic effect of venous congestion when given *prior* to the latter (Figure 3A). In contrast, alcohol had little effect when given *during* the period of venous congestion (Figure 3B).

secondary to variations in the activity of the neurohypophysis.⁶

Alcohol had little effect on urine flow and free water clearance in those experiments in which it was given *after* an infusion of hypertonic saline or venous congestion had been initiated. Once the neurohypophysis has been stimulated the resultant excess of circulating antidiuretic hormones might, until it is inactivated or destroyed, mask the temporary inhibition of the posterior pituitary by alcohol. These experiments suggest that alcohol diuresis may be blocked by a prior rise in circulating endogenous ADH as well as by the administration of exogenous Pitressin® (5, 6).

In the experiments of Group I, when the release of ADH was presumably completely inhibited by a positive water load in the semi-recumbent position, administration of alcohol caused no further rise in urine flow and free water clearance.

⁶In a study published since completion of this paper Newman (12) demonstrated that alcohol could effectively block the antidiuresis of quiet standing.

The increase in CH₂O which occurred in the water-loaded subjects of Group IIb, to whom alcohol was given simultaneously with an infusion of hypertonic saline, was therefore unexpected. The situation in these experiments is probably comparable to the rapid administration of large solute loads to patients with diabetes insipidus, in whom an increased volume of isosmotic fluid is suddenly delivered to a distal tubular segment in which water reabsorption is blocked but where further reabsorption of solute does occur. In this case an increase in the calculated value of free water clearance (CH₂O) might be produced, not by diminished reabsorption of water in the distal tubule (Smith, 13), but by an increased distal reabsorption of solute. An increase in CH₂O during mannitol or solute diuresis in subjects with diabetes insipidus can in fact be demonstrated by recalculating the data of Brodsky and Rapoport (14). Similar increases in CH₂O, C_{osm}, and urine flow were shown by Welt, Young, Thorup, and Burnett (15) to follow the adminis-

tration of a carbonic anhydrase inhibitor to water-loaded subjects who were in a state of "physiological diabetes insipidus." Although tubular secretion of water (14) could explain such changes, there seems little reason to invoke such a concept.

A relative or absolute increase in antidiuretic hormone has been implicated in the abnormal water metabolism of such clinical states as hyponatremia, cirrhosis of the liver, congestive heart failure, adrenal insufficiency, and panhypopituitarism. The results of the present and previous studies (1, 5) suggest that the effects of alcohol in states of abnormal water metabolism might be of value in interpreting their pathophysiology. Such investigations are now in progress.

SUMMARY

1. Alcohol had no effect upon urine flow or solute excretion when given at the height of a water diuresis.

2. Alcohol blocked the antidiuretic response to hypertonic saline when both were simultaneously administered to water-loaded subjects.

3. Alcohol minimized the antidiuretic effect of venous congestion of the legs in water-loaded subjects, when imbibed before the legs were congested.

4. The characteristic diuretic response to alcohol was blocked by prior infusion of hypertonic saline or cuff congestion of the limbs.

5. When administered prior to the stimulus, alcohol will effectively block stimulation of the release of ADH.

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