JCI The Journal of Clinical Investigation

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J Clin Invest. 1960;39(6):876-884. https://doi.org/10.1172/JCI104108.

Research Article



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THE RESPONSE TO THE ADMINISTRATION OF AN ISOTONIC SODIUM CHLORIDE-LACTATE SOLUTION IN PATIENTS WITH ESSENTIAL HYPERTENSION

BY SOLOMON PAPPER,* JOSEPH L. BELSKY AND KENNETH H. BLEIFER †

(From the Medical Service and the Research Laboratory, Veterans Administration Hospital, and the Departments of Medicine, Boston University School of Medicine and Tufts University School of Medicine, Boston, Mass.)

(Submitted for publication December 21, 1959; accepted January 8, 1960)

Many studies indicate that patients with essential hypertension have a greater natriuretic response to rapidly administered sodium chloride solutions than have normotensive individuals (1–7). However, in most instances other factors known to influence the rate of sodium excretion in the normal subject such as diet, posture, and time of day have not been rigidly controlled (8, 9). Dietary control is of particular importance in view of observations suggesting that patients with hypertension may habitually ingest more salt than do normotensive individuals (10– 11). If this is indeed the case it might well be responsible for the enhanced response of the hypertensive patient to administered salt.

The present report is concerned with a comparison of the response of hypertensive and normotensive individuals to the intravenous administration of an "isotonic-balanced" salt solution under rigidly controlled conditions and at three different levels of dietary salt ingestion. The results indicate that patients with essential hypertension excrete the infused sodium load more rapidly than do normotensive individuals at each level of salt consumption.

METHODS

Four normal Caucasian males aged 29 to 36 and 6 Caucasian patients aged 24 to 63 with essential hypertension were studied. The patients were selected on the basis of their maintaining a resting diastolic blood pressure of at least 100 mm Hg while hospitalized and consuming a diet containing approximately 10 mEq of sodium daily. Five of the 6 subjects were observed in this manner for 13 to 47 days prior to study, while Patient 8 received the low salt diet for 8 days prior to study. No patient had congestive heart failure, although no. 5 had had a myocardial infarct 6 years earlier. All 6 had electrocardiographic evidence of left ventricular strain, but were free of gross cardiomegaly on radiographic examination. Three patients had a history of mild hemiparesis 3 months to 10 years before study; Patients 5 and 8 made complete recovery while Patient 9 had mild neurological residua. Ocular fundi varied from normal to Grade 2¹ arteriolar narrowing without hemorrhage, exudate or papilledema. Renal function (including endogenous creatinine clearance, phenolsulfonephthalein (PSP) excretion, intravenous pyelogram and concentration test) was normal except in Patients 9 and 10 who had modest reduction in creatinine clearance.

Three of the 4 normal subjects and 5 hypertensive patients were provided a diet containing 10 to 15 mEq of sodium daily (low salt diet). After equilibrium was established (i.e., a minimum of 4 days of dieting and 2 consecutive days during which time the 24 hour urinary sodium excretion did not exceed 15 mEq) the following two studies were done within a period of 3 days.

1. "Blank Day." The subject had his usual breakfast including 500 ml of water. At 8 a.m. he assumed the recumbent position and remained so until 3 p.m. except to void. From 10 a.m. to 2 p.m. each subject drank 100 ml of water hourly and ingested 5 g of carbohydrate each half hour. Spontaneously voided urine was collected at one-half hour intervals. Venous blood was collected at least twice (10 a.m. and 2 p.m.).

2. "Infusion Day." The protocol was essentially the same as on Blank Day except that 2,000 ml of a solution containing 130 mEq per L of sodium, 105 mEq per L of chloride and 25 mEq per L of lactate was administered intravenously from 10 to 11:30 a.m., and hourly drinking commenced at 12 noon. Venous blood samples were collected immediately prior to and at the end of the infusion period and again at 2 p.m. An additional normal subject (no. 4) and one hypertensive patient (no. 10) were studied as described for Infusion Day without a prior Blank Day.

^{*} Present address: Medical College of Virginia, Richmond, Va.

[†]USAF (MC).

¹ The grading system employed is that recommended to the American Ophthalmological Society by the committee on Classification of Hypertensive Disease of the Retina: Wagener, H. P., Clay, G. E., and Gipner, J. F. Classification of retinal lesions in the presence of vascular hypertension. Trans. Amer. Ophthal. Soc. 1947, 45, 57.

Subject				Without infusion	infusion							With i	With infusion				
no.								Creati-								Clearances	5
Blood	Dietary Na	Time	Na	м	ū	Solute	Flow	clear- ance	Time	Na	К	CI	Solute	Flow	Creati- nine	Osmolar	Free water
			µEq/min	μEq/min	uEq/min	µOsm/min	ml/min	ml/min		μEq/min	µEq/min	uEq/min u0sm/min	iOsm/min	ml/min	ml/min	ml/min	ml/min
1 21	Low	8:30-10 a.m.	9.3	74	16	548	4.1 ,	106	8:30-10 a.m.		72	15	503	3.1	131	1.8	1.3
114/80		11:30–3 p.m.	1.5	74	91 4 <i>i</i>	468	1.0	122	11:30–3 p.m.	515	100 100	325	548	1.9	128	1.9	0,0
		100:21-00:11	7.7	108	11	220	7.7	771	06:2-06:1	10	06	çç	050	1.8	136	1.9	-0.1
	Medium	8:30-10 a.m. 10 -11:30 11:30-3 p.m. 9:30-10:30	406 364 186 439	96 126 45	367 339 397	1,305 1,257 704 1.402	5.5 2.7 2.7	127 124 128	8:30-10 a.m. 10 -11:30* 11:30-3 p.m. 10:30-11:30	273 416 280 460	90 129 61	294 381 399	1,061 1,366 894 1.443	4.6 3.9 4.6	150 146 144	3.7 3.1 0.2	0.9 0.8 3.4
	High	8:30-10 a.m. 10 -11:30	502 422	8 8 8	400 400	1,459 1,306	3.7 3.7	145 134	8:30-10 a.m. 10 -11:30		69 113	508 718	$1,582 \\ 2,199$	0.8 9.5	149 143	5.5	3.3 1.8
		11:30-3 p.m. 9:30-10:30	228 518	54 87	227 474	833 1,488	2.9 3.2	122 137	11:30-3 p.m. 10:30-11:30	461 883	57 119	420 742	1,278 2,329	4.0 11.5	137 142	4.5 8.2	-0.5 3.3
2	I.ow	8:30–10 a.m.		126	34	852	1.5	156	8:30-10 a.m.		67	14	553	2.5	143	2.0	0.5
29		10 -11:30		126	49	914	3.6	149	10 -11:30*‡		142	36	730	4.8	156	2.6	2.2
177/80		11:30-3 р.т. 9 -10	39	89 133	38 38	700 853	1.3 1.3	152	11:30-3 p.m. 2 -3	88 88	83 83	31 27	000 808	1.9	150 166	2.5 2.5	0.3 -0.6
	Medium	8:3	267	110	246	1,067	3.1	162	8:30-10 a.m.		98	199	945	4.7	160	3.3	1.4
		10 -11:30 11:30-3 p.m.	308 221	155 75	350 195	1,339 834	3.8 0.4 0.8	$149 \\ 139$	10 -11:30*‡ 11:30-3 p.m.	316 267	173	363 253	1,303 1.018	0.5 11.3	154 144	4.6 3.6	1.9
		9:30-10:30	402	148	358	1,377	3.5	157	10 -11		167	388	1,306	3.4	160	4.6	-1.2
	solution c :30 a.m. u alics indic	* A solution of 2,000 ml containing app 10 to 11:30 a.m. unless otherwise indicated. † Italics indicate a 60-minute period of	aining a indicate period	tpproxim d. of maxin	ately 1. 1um rat	30 mEq/ e of sodi	L of so um exc	dium, 1 ¹ retion (1	* A solution of 2,000 ml containing approximately 130 mEq/L of sodium, 105 mEq/L chloride and 25 mEq/L lactate was administered intravenously from o 11:30 a.m. unless otherwise indicated. † Italics indicate a 60-minute period of maximum rate of sodium excretion (two consecutive 30 minute periods were pooled).	ide and 2 30 minut	25 mEq/	L lactat s were p	e was ad ooled).	Iministe	red intr	avenous	sly from
1	nfusion giv	en 10:15 to 11:	55 a.m.														

TABLE I

Fluid and electrolyte excretion in subjects with normal blood pressure

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1	1		= -								
	es	Free water	ml/min — 4 0	-2.5	-3.7	-0.1	-0.4	-2.6	-2.8	-2.4 -0.3 -3.9	0.2 1.1 1.0
	Clearances	Osmolar	ml/min 7 A	5.8	4.3 7.7	2.5	2.5	4.1 5.4	5.6	5.4 3.7 6.9	2.4 2.1 2.2
		Creati- nine	ml/min 168	146	151 <i>159</i>	170 155	153 158	175 162	167 160	192 182 175 186	194 144 130 <i>132</i>
		Flow	ml/min 3 A	3.3	5.4 4.0	1.5 3.6	2.2 2.2	4.6 2.8	3.4 2.8	3.0 3.4 3.0 3.0	2.6 3.2 3.2
fusion		Solute	µОsm/min 2 176	1,653	1,216 2,221	712 697	715 722	1,154 1,514	1,005 1,562	1,561 1,052 1,122 1,973	660 504 575 598
With infusion		C	μEq/min 658	461	265 <i>695</i>	30 28	45 47	251 291	416	463 539 304 623	36 58 58
		ĸ	uEq/min uEq/min 644 1.34	142	100 146	98 162	901 80	102 194	115 184	105 129 82 123	118 134 58
		Na	μEq/min 644	446	329 673	47	108 <i>136</i>	247 409	243 446	458 593 329 710	16 33 118 147
		Time	8 · 30–10 a m	10 -11:30*	11:30–3 p.m. $g -10$	8:30-10 a.m. 10 -11:30*	11:30–3 p.m. 1:30–2:30	8:30-10 a.m. 10 -11:30*	11:30-3 p.m. 10 -11	8:30-10 a.m. 10 -11:30* 11:30-3 p.m. 10 -11	9 -10 a.m. 10 -11:30* 11:30-3 p.m. 2 -3 p.m.
	Creati-	clear- ance	ml/min 161	148	146 156	146 139	138 140	167	161	173 155 158 162	
		Flow	ml/min 2 9	3.6	5.0 3.5	2.1 1.6	1.4 1.4	4.4 3.0	3.1 3.4	5.1 3.5 3.0	
		Solute	μOsm/min ml/min 1 742 2 9	1,993	1,005 2,223	718 695	539 702	$938 \\ 1,340 \\ 20$	888 1,403	1,323 1,329 1,034 1,464	
nfusion		C	μEq/min 532	640	239 738	20 23	12 25	156 336 236	337	323 283 271 299	
Without infusion		К	Eq/min 101	149	76 128	81 103	51 102	70 141	84 145	42 93 67	
		Na	µЕд/тіп µ. 512	629	222 731	22	30 30	167 373	209 384	335 377 269 453	
		Time	8 · 30–10 a m	10 -11:30	11:30–3 p.m. <i>9:30–10:30</i>	8:30-10 a.m. 10 -11:30	11::30-3 p.m. 9:30-10:30	8:30-10 a.m. 10 -11:30	11:30-3 p.m. 10 -11	8:30-10 a.m. 10 -11:30 11:30-3 p.m. 9:30-10:30	
	•	Dietary Na	Hioh			Low		Medium		High	Low
Cubicot	no.	Blood				3 29	120/78				4 36 112/68

TABLE I (Continued)

Three normal subjects and three hypertensive patients were similarly studied after equilibrium was established while taking the same 10 mEq sodium diet with approximately 35 mEq of sodium chloride (non-enteric coated tablets) added to each meal and again at bedtime for a total of 150 mEq sodium intake daily (medium salt diet). On both Blank Day and Infusion Day each subject ate his usual breakfast and 35 mEq of additional salt in tablet form. At the end of the Blank Day experimental period, the subjects were given sufficient food and sodium to maintain caloric intake and the 150 mEq daily quantity of sodium.

Three normal subjects and three hypertensive patients were similarly studied while they were taking approximately 300 mEq of sodium daily (high salt diet).

Blood pressure was determined at one-half hour intervals in the hypertensive patients during both experimental days at each dietary level, and less often in the normal subjects.

Serum and urine were analyzed for sodium, potassium, chloride, creatinine and total solute content by methods employed in this laboratory and previously described (12). Serum protein, blood hemoglobin concentration and hematocrit were also determined.

RESULTS

1. On Blank Day there was no significant difference between the normotensive and hypertensive subjects, taking the low sodium diet, in the quantities of sodium and chloride excreted from 10 a.m. to 3 p.m. In five of six instances, while provided with medium and high salt intakes, the hypertensive patients excreted more sodium and chloride than did the normal individuals from 10 a.m. to 3 p.m. (Tables I and II, Figure 1).

2. On Infusion Day the preinfusion rates of sodium excretion were no higher in the hypertensive than in the normal group at each dietary level of sodium ingestion. In fact, the hypertensive patients had slightly lower rates of sodium excretion prior to infusion while taking the medium salt diet (Tables I and II, Figure 1).

3. In each instance at all levels of salt intake, the hypertensive patient had a far greater natriuresis after intravenous salt loading than had the normal. The maximal rates of sodium excretion after salt loading occurred more promptly in the hypertensive patient at the low level of salt intake (Tables I and II, Figures 1 and 2).

4. By the morning after salt loading the hypertensive patients had excreted more sodium than had the normal subjects at each dietary level. This difference is attributed to the prompt re-

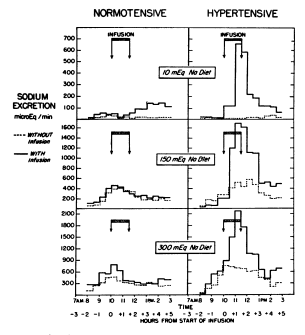


FIG. 1. SODIUM EXCRETION WITH AND WITHOUT IN-FUSION IN ONE NORMOTENSIVE (SUBJECT 3) AND ONE HYPERTENSIVE (SUBJECT 6) WHILE PROVIDED WITH LOW, MEDIUM AND HIGH SALT DIETS. The ordinate scale for the 150 mEq and 300 mEq sodium diets is double and triple, respectively, the scale for the 10 mEq sodium diet.

sponse (10 a.m. to 3 p.m.) rather than to any continued difference in sodium excretion throughout the remainder of the day (Table III).

5. Endogenous creatinine clearance generally increased in both normal and hypertensive subjects when dietary salt was increased from low to medium salt intake levels. The change in clearance was less conspicuous between the medium and high salt intakes. It is also apparent that the 8:30 to 10 a.m. endogenous creatinine clearance often varied significantly in the two studies carried out in a single individual on different days. The differences in natriuretic response observed were not consistently or uniformly correlated with preinfusion differences in endogenous creatinine clearance or with change in clearance following infusion (Tables I and II).

6. In nine experiments in patients with hypertension, the infusion of sodium chloride-lactate solution was not associated with a rise in blood pressure. In the remaining two experiments a rise in diastolic blood pressure of 10 to 12 mm Hg was observed following infusion.

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$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	Subject				Withou	Without infusion							With i	With infusion				
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	no. Age								Creati- nine						•		Clearance	(0)
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Blood	Dietary Na	Time	Na	K	IJ	Solute	Flow	clear- ance	Time	Na	К	C	Solute	Flow		Osmolar	Free water
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$				μEq/min	1 4		µOsm/min	ml/min	ml/min		μEq/min	μEq/min		uOsm/min	ml/min	ml/min	ml/min	ml/min
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	ю		8:30-10 a.m.	1.5		7.3	359	1.0	94	8:30-10 a.m.	6	31	14	298	$\frac{2.0}{2.0}$	87	1.0	1.0
$ \begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	03		10 -11:30 11:30-3 n.m.	35		28 28	409 409	1.0	60	10 -11:50 ⁺	345 470	11/ 88	288 400	1,180	۰ ۲. 4 4	4 7 1	4.1 4.3	5.5 2.7
			12:30-1:30	45		34	452	1.7	90	11:30-12:30	683	130	599	1,409	9.2	93	4.9	4.3
High 8:30-10 a.m. 73 65 650 1756 113 113 -173 500 920 253 113 113 -122 173 500 920 253 113 113 -123 173 500 920 253 113 113 -122 173 500 920 253 113 113 -133 90 920 253 114 2.6 -133 90 920 253 114 2.6 113 93 153 113 93 133		Medium	8:30-10 a.m.	114 472	55 00	136	606 1 465	4.0	115	8:30-10 a.m.	104 6 2 /	43 73	127	565 1 508	2.2	109	2.0	0.2
High 8:30-10 a.m. 170 45 214 666 1.0 11:30 493 124 11:30 493 125 114 216 125 117 205 125 114 216 125 117 205 125 117 205 135 5.2 111 11.30 203 125 137 20 137 20 231 756 137 205 117			11:30-3 p.m.	289	64 8	255 458	1 632		97 113	11:30-3 p.m.	778 1 1 2 3	198	620 020	1,795	6.5 15.0	100	0.3 6.3	0.5
High $8:30-10 \text{ a.m.}$ 170452146661.0124 $8:30-10 \text{ a.m.}$ 189422317561.41142.610:-11:30-10:30499621,3194.511711:30-330110712.78012.51179.810:-0:30-11:30525664851,3355.211111: -12 2,0371121,6984,35820.81165.510:-0:30-11:3065694391.2991011:30*2947124881.81081.710:-11:3065694391.2991011:30*2947112313111::0-3 pm.15641.310011::0-3 pm.201711658632.91183011::0-3 pm.15641.310011::0-3 pm.201711658632.91183011::0-3 pm.201711659431.310011::0-3 pm.201711658632.91183011::0-3 pm.201711659431.31318::30-10 a.m.105441190.71191182.311::0-3 pm.2021031121310011::0-3201712051192.32.31112.311::0-3 pm.			11_ 01	200	2	074	1006	1.1	CTT	77 11	001,1	22	740	040,70	0.01	011	v.0	1.0
Ii: 30-3 pm. 399 42 360 1/33 5.2 111 11 -12 2/37 112 1/698 4,358 2/08 116 15.3 3.5 111 11 -12 2/37 112 1/698 4,358 2/08 116 15.3 3.5 111 11 -12 2/037 112 1/698 4,358 2/08 116 15.3 4.1 9.1 6.1 10.3 8.5 11 -12 2/037 112 1/698 7.7 123 4.1 9.1 6.1 13.5 6.4 13 100 11:30-3 201 11 15 6.4 13 100 11:30-3 201 11 15 6.4 13 100 11:30-3 201 11.6 5.2 11.8 13 30 13 13 33 13 13 8:30-10 a.m. 203 14 13 20 14 23 20 14 23 20 14		High	8:30-10 a.m. 10 -11:30	170 499	45 62	214 472	666 1.319	1.0 5.5	124	8:30-10 a.m. 10 -11:30*	1.218 1.218	42 79	231 1.071	756 2.780	1.4	114 117	2.6 9.8	-1.2
Low 8:30-10 a.m. 8 45 11 503 2.1 94 8:30-10 a.m. 7,30 7,00 1,00 1,10 1,10 1,10 1,10 1,10 1,10 1,10 1,10 1,130 6 56 9 10 -11:30* 2.94 100 2.37 1,186 7.7 123 4,11 12 1,86 7.7 123 4,11 133 6.9 11 -11:30* 2.94 100 2.37 1,186 7.7 123 4,11 12 4,83 3.0 133 3.0 11 -12 6.21 140 520 1,967 11.9 133 6.9 11:30-3 p.m. 201 45 1,303 2.4 123 10 11:30* 10.0 12.3 140 5.20 134 13 5.3 131 6.9 Medium 8:30-10 a.m. 201 4,41 123 10.0 12.3 1,40 5.20 1,41 2.3			11:30–3 p.m.	399	42	360	1,035	4.5	107	11:30–3 p.m.	1,037	12	943	2,411	9.1	103	8.5 2.5	0.6
Low 8:30-10 a.m. 8 45 11 503 2.1 94 8:30-10 a.m. 9 47 12 488 1.8 108 1.7 10 -11:30 6 56 9 439 1.2 90 10 -11:30* 294 100 237 1,186 7.7 123 4.1 11:30-3 p.m. 15 64 19 464 1.3 100 11:30-3 p.m. 201 71 165 863 2.9 118 3.0 11:30-2:30 2.9 77 25 581 1.9 108 11 -12 621 140 520 1,967 11.9 138 5.0 1967 11.9 138 6.9 11 1:30-3 p.m. 339 67 349 1,303 2.4 123 10 -11:30* 1,026 92 959 2,685 9.3 141 2.3 11:30-3 p.m. 339 67 349 1,153 2.9 112 11:30-3 p.m. 779 81 756 2,063 5.7 123 7.2 11:30-3 p.m. 339 67 349 1,153 2.9 112 11:30-3 p.m. 779 81 756 2,063 5.7 123 7.2 11 -12 468 88 472 1,560 3.7 122 11 -12 1,650 145 1,545 3,935 15.1 133 13.7 54 1130 -11:30 2,050 3.7 122 11 -12 1,650 145 1,545 3,935 15.1 133 13.7 11:30-3 p.m. 477 69 481 1,582 2.6 137 5.4 1130 -11:30 3.2 1,383 2.1 123 11:30-3 p.m. 477 69 481 1,582 2.6 137 5.4 1130 -11:30 3.5 1,383 2.1 123 11:30-3 p.m. 477 69 481 1,582 2.6 137 5.4 1130 -11:30 3.5 1,383 2.1 123 11:30-3 p.m. 477 69 481 1,582 2.6 137 5.4 1130 -11:30 3.5 1,383 2.1 123 11:30-3 p.m. 887 68 882 2,476 6.2 117 8.7 12.5 11.30-3 p.m. 779 10 -11:30 723 90 702 1,989 2.5 132 10 -11:30* 1,506 156 2,787 4,441 175 128 15.5 100 -11.50 16.5 117 8.7 10 -11:30 723 90 702 1,989 2.5 132 10 -11:30* 1,506 156 2,787 4,441 175 128 15.5 100 -11:30 -11:			00:11-00:01	C7C	00	604	CCC'1	7.0	111	71- 11	100,2	211	1,098	4,338	50.8	011	5.01	<i>c.c</i>
High 8:30-10 a.m. 317 65 136 581 10 11:30-3 p.m. 201 71 165 783 29 118 30 30 138 6.9 138 139 137 131 <td></td> <td>Low</td> <td>8:30-10 a.m. 10 -11:30</td> <td>% \c</td> <td>45</td> <td>110</td> <td>503 430</td> <td>2.1</td> <td>94 8</td> <td>8:30–10 a.m. 10 –11:30*</td> <td>9 204</td> <td>47 100</td> <td>12 237</td> <td>488 1 186</td> <td>1.8</td> <td>108 173</td> <td>1.7</td> <td>0.1 2.6</td>		Low	8:30-10 a.m. 10 -11:30	% \c	45	110	503 430	2.1	94 8	8:30–10 a.m. 10 –11:30*	9 204	47 100	12 237	488 1 186	1.8	108 173	1.7	0.1 2.6
$ \begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	174/112		11:30-3 p.m.	15	84	19	464	1.3	100	11:30–3 p.m.	201	11	165	863	2.9	118	3.0	-0.1
um 8:30-10 a.m. 201 45 226 938 1.3 131 8:30-10 a.m. 105 44 110 672 0.8 141 2.3 10 $-11:30$ 387 61 384 1,303 2.4 123 10 $-11:30^*$ 100 672 0.8 141 2.3 11 $-0.1:30$ 387 61 384 1,303 2.4 123 10 $-11:30^*$ 92 959 2,685 9.3 141 0.3 11: -12 468 88 472 1,560 3.7 122 11 -12 1,650 145 1,545 3,935 15.1 133 13.7 8:30-10 a.m. 317 69 320 1,176 1.6 143 8:30-10 a.m. 474 69 481 1,582 2.6 137 5.4 10 -11:30 723 90 702 1,989 2.5 130 10.56 2,787 3,491 175 125 11 2.5 10 -11:30 723			1:30-2:30	29	17	25	581	1.9	108	11 –12	621	140	520	1,967	11.9	138	6.9	5.0
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$		Medium	8:30-10 a.m.	201 387	45 61	226 384	938 1 303	1.3 1.3	131	8:30-10 a.m.	105	44 0	110	672	0.8	141	2.3	-1.5
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$			11:30–3 p.m.	339	67	349	1,153	2.9	112	11:30–3 p.m.	793	81	756	2,063	5.7	141	7.2	-1.5
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$			11 -12	468	88	472	1,560	3.7	122	11 -12	1,650	145	1,545	3,935	15.1	133	13.7	1.4
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		High	8:30-10 a.m.	317	69	320	1,176	1.6	143	8:30-10 a.m.	474	69	481	1,582	2.6	137	5.4	-2.8
737 92 717 2,013 2.6 135 11 –12 1,950 156 2,787 4,441 17.5 128 15.5			11:30-3 p.m.	475	<u>8</u> 0	475	1,383	2.1	121	11:30–3 p.m.	887	08 89	2,030 852	2,476	6.2	1117	8.7	-2.5
			II- 0I	737	26	717	2,013	2.0	135	11 -12	1,950	156	2,787	4,441	17.5	128	15.5	2.0

TABLE II

Fluid and electrolyte excretion in subjects with hypertension

*† See footnotes to Table I.

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I		1	ŀ														
	8	Free water	ml/min	1.5	0	!	-1.5	-0.3) 1	-2.7	-0.1	7 6	5.1	0.6 4.6	5	0.8	3.9 3.9
	Clearance	Osmolar	ml/min	2.2	9.5 0	2	2.4	6.2		4.3 6.6	7.1	76	7.1	5.1	101	2.3 6 8	5.0
		Creati- nine	ml/min	131	113	100	116 106	104		111	107	86	85	808	00	99	82 87 87 87 87 87 87 87 87 87 87 87 87 87
		Flow			4.5	1.01	0.0	5.9		1.0	6.4 7.8	10.2	12.2	5.7	0.11	3.1	5.8
fusion		Solute	uOsm/min	626	1,258	0,0,1	665 1 537	1,725		1,211	1,984	2117	2.184	1,487	101,1	649 1 024	1,407 2,266
With in		ū	LEq/min	43	352	010	84 463	622 788		350 705	907 907	79	697	427	1,016	105	446 953
		. м	LEq/min p	80	84 163	201	56 93	22 011		167 167	121 241	105	198	117 205		94 184	98 201
		Na	LEq/min	20	443	CT/(T	68 470	655		523 1 118	1,274	05	847	546	111	70 20	485 1,003
		Time		9 –10 a.m.	11:30–2:30 p.m.	77_ 77	8:30-10 a.m. 10 -11.30*	11:30–3 p.m.		8:30-10 a.m. 10 -11 ·30*	11:30-3 p.m. 11 -12	8.30-10 ° m	$10 -11:30^{*}$	11:30-2:30 p.m.	77_ 11	8:30-10 a.m.	11:30–3 p.m. 11 –12
	Creati-	clear- ance	ml/min				101 04	16		111	103 99						
		Flow	ml/min	3.3	50	ŗ	1.5	2.7		1.2	2.9	0		3.2	;		
		Solute	uOsm/min	473	423	C11	896 803	915		971 1 004	1,263	165	414	406 305	~~~		
infusion		ū	uEq/min	53	38	F	221 737	246 274		272 301	470 503	13	12	28 33	S		
Without		К			38	5	57 71	50		57	61 61	ç	101	C1 m	2		
		Na	uEq/min	==	362	2	186 105	213		234	465 515	10	10	24 31	10		
		Time		8 -10:30 a.m.	11:30–3 p.m.	····· · · · · · · ·	8:30-10 a.m.	11:30–3 p.m.		8:30-10 a.m. 10 -11:30	11:30–3 p.m. 2 –3	8.30 10 a m	0.30-10 $a.11.$	11:30–2:30 p.m.	00.3-00.1		
		Dietary Na		Low			Medium						row Low			Low	
Chint	no.	Age Blood pressure		74	172/120			182/104		-		c		230/154		10	214/136
	With infusion Without infusion	Without infusion Creati-	Without infusion Without infusion Dietary Creati- nine Na Creati- clear Dietary Creati- clear Na K Classing Creati- clear	Without infusion With infusion Creati- nine Na Time Clearances Clearances Distary Ledmin Ledmin Ledmin Ledmin Month milmin Use Na K Clearances Clearances Distary Ledmin Ledmin Ledmin Ledmin Month milmin	With infusion With infusion Distary With infusion Creation in the image of t	$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	With infusion With infusion Distary With infusion Distary With infusion Distary With infusion Creati- nine Creati- nine Creati- nine Creati- clear- clear- Creati- clear- Creati- clear- Creati- clear- Creati- clear- Creati- clear- Creati- clear- Creati- clear- Creati- clear- Creati- clear- Creati- clearin Creati- 0 smolar Low 8 -10:30 a.m. 11 Creati- clearin Creati- clearin Creati- clearin Creati- clearin Creati- clearin Low 8 1007 11:30-2:30 p.m. 4.5 Creati- clearin I:30-3 11:30-3:30 p.m. 4.5 13.3 Creati- clearin I:30-3 11:30-3 11:30-3	$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	With infusion Diferation Nation in the properties of the pro	$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	Without infusion Without infusion Vithout infusion Diteration Time Na With infusion Diteration Creation Creation Creation Creation Diteration Na K Cleanances Distant Creation Creation Creation Creation Distant Ma K Creation Creation	$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$

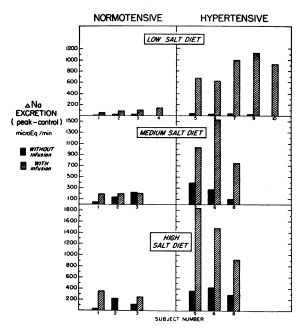


FIG. 2. THE DIFFERENCE IN THE RATE OF SODIUM EX-CRETION BETWEEN THE 60 MINUTE PERIOD OF MAXIMUM NATRIURESIS AFTER 10 A.M. AND THE MEAN SODIUM EX-CRETION BETWEEN 8:30 AND 10 A.M. (While on a high salt diet, Subject 2 had a maximal natriuresis prior to the infusion period.)

7. Potassium excretion varied considerably but there was no consistent difference between the hypertensive and normotensive subjects (Tables I and II).

8. On Infusion Days urine flow and free water clearance (as well as osmolar clearance) were greater in the hypertensive patients than in the normal subjects.

 TABLE III

 Sodium excretion* during and after infusion

	10 a.m.	[-3 p.m.		I -7 a.m.		+11) -7 a.m.
			Normo- tensive			
I. Low salt	18	131	24	41	42	172
diet	26 13	72 185	48	22 60	74	94 245
	28	158		68		226
II. Medium salt		211		168		379
diet	83	260	186	138	269	398
	87	179	183	151	270	330
III. High salt	171	327	238	634	409	961
diet	109	322	336	222	445	544
	122	367	265		387	

* Total number of milliequivalents excreted in each time interval.

9. Serum concentration of sodium and chloride did not change significantly (-1 to + 3 mEq per L) following the infusion of the sodium chloride-lactate solution. A small decrease in hematocrit (1 to 4 points) and total protein concentration (0.2 to 1.1 g per 100 ml) was observed following infusion.

DISCUSSION

The present study confirms other observations that patients with essential hypertension have a greater natriuretic response to administered salt solution than have normal individuals. In addition it establishes the fact that this difference in sodium excretion is short-lived and is not due to differences in dietary ingestion of salt prior to the test. Indeed, the data demonstrate that the exaggerated natriuresis of the hypertensive individual is apparent at all levels of salt intake ranging from 10 to 300 mEq daily. Furthermore, the study suggests that the difference in natriuretic response is probably not due to an alteration in diurnal rhythm. Thus, on Blank Days on the low salt diet, the normal and the hypertensive subjects excreted comparable quantities of sodium. The present data do not provide final proof on this point since the change in sodium excretion (peak minus control) on Blank Days was greater in the patients with hypertension than in the normal subjects. This difference seems to be related primarily to a generally lower control (8:30 to 10 a.m.) rate of sodium excretion in the hypertensive subjects on Blank Days, compared with the normal individuals, rather than to a consistently higher peak excretory rate for sodium. That this may have been fortuitous is perhaps suggested by the fact that the preinfusion values on Infusion Days while taking the low salt diet do not bear out this difference in control values. Clearly, more studies are required to resolve the role of diurnal rhythm with complete certainty. That the hypertensive subject excreted more sodium than did the normal individual on Blank Days at medium and high dietary levels is probably due to the fact that added salt was taken on these days at breakfast time and in effect constituted a small but effective "salt load." In addition, the present study makes it quite clear that the exaggerated natriuretic response is not due to a difference in baseline rates of sodium excretion prior to infusion. Finally, the infusate was such that serum sodium concentration was not changed significantly during any experiment, thus precluding the possibility that the exaggerated natriuresis may be related to a peculiarly distorted response to hypertonic salt solutions in patients with hypertension.

While the present study documents the existence of abnormal sodium excretion in hypertension under controlled conditions, the data do not provide an understanding of the mechanisms involved. In these studies, as in others, in which there is no consistent relationship between endogenous creatinine clearance (or inulin clearance) and sodium excretion, it is virtually impossible to establish or exclude the importance of small but significant changes in glomerular filtration rate in determining differences in so-From the present studies it dium excretion. would appear that the exaggerated natriuretic response is not clearly attributable to increased glomerular filtration rate either in the basal period or in response to salt administration. Consequently, it seems reasonable to focus attention on the renal tubular handling of sodium in patients with hypertension. The possibility exists that the renal tubular cell itself is abnormal or that a normal tubular cell is responding normally to abnormal influences or to stimuli that are abnormally mediated. While a specific tubular "defect" cannot be excluded, there is little evidence in support of this concept (7). Among the variety of known and unknown extrarenal factors that might influence renal tubular handling of sodium in the hypertensive patient are hormonal factors (e.g., adrenocortical and adrenomedullary), neurogenic factors and intrarenal circulatory phenomena (8). There is little direct evidence to support the causal role of any of these at the present time. The amount of sodium in the body, perhaps as expressed in terms of "effective" extracellular fluid volume, seems to be an important determinant of sodium excretion in the normal individual (8). How the kidney is made aware of changes in this factor is not at all clear. There are data which suggest increased total body sodium as well as increased extracellular fluid volume in patients with essential hypertension, although other data are not in accord with these findings (13-17). Nonetheless, no causal relationship between alterations in extracellular volume or body sodium content and the observed exaggerated natriuretic response to administered sodium seems warranted by the data at this time. The present study does not clarify the relationship of the disturbances in sodium excretion to other aspects of the condition called "essential hypertension," including the elevation of blood pressure.

SUMMARY AND CONCLUSIONS

1. The natriuretic response to the infusion of an isotonic solution of sodium chloride-lactate was studied in four normal subjects and in six patients with essential hypertension, under conditions rigidly controlled in respect to the amount of sodium ingested, posture, and time of day.

2. At each of three levels of daily sodium ingestion (10, 150 and 300 mEq) the patients with hypertension had a far greater natriuretic response to administered sodium than had the normal individuals.

3. Without infusion, at the low salt dietary level, there was no difference in the quantity of sodium excreted between normal subjects and patients with hypertension, suggesting that variations in basic diurnal rhythm probably do not account for the enhanced rate of sodium excretion.

4. The exaggerated natriuresis is not attributable to differences in preinfusion rates of sodium excretion or to greater increase in serum sodium concentration. In addition, the difference in natriuresis following infusion between the hypertensive and normotensive subjects is not associated with clearly consistent differences in endogenous creatinine clearance or further augmentation in blood pressure.

5. The "abnormal" response to salt administration in patients with essential hypertension remains unexplained.

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