

Genetic Mapping of Two Blood Pressure Quantitative Trait Loci on Rat Chromosome 1

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Abstract

A genetic map for rat chromosome 1 was constructed using 66 microsatellite markers typed on either or both of two populations derived from inbred Dahl salt-sensitive (S) rats: $F_2(\text{LEW} \times \text{S})$ $n = 151$, and $F_2(\text{WKY} \times \text{S})$ $n = 159$. These populations had been raised on a high salt (8% NaCl) diet. Systolic blood pressure and heart weight were found to be genetically linked to two separate regions on rat chromosome 1 in the $F_2(\text{LEW} \times \text{S})$ population. One region was centered around the anonymous SA locus and accounted for 24 mmHg of blood pressure. The other region was 55 cM from the SA locus centered around a cluster of cytochromes P450 loci, and accounted for 30 mmHg of blood pressure. Since blood pressure and heart weight were highly correlated these same regions were also linked to heart weight. These results were cross-specific as linkage of these chromosome 1 regions to blood pressure and heart weight was not observed in several other F_2 populations derived by crossing S and other normotensive control strains. This is presumably due to different alleles and/or different genetic backgrounds in the various populations. The SA region of chromosome 1 was also found to influence body weight in $F_2(\text{LEW} \times \text{S})$ rats. Combining the present data with our previously published data on the $F_2(\text{LEW} \times \text{S})$ population showed that four separate quantitative trait loci with additive effects accounted for 106 mmHg and 38% of the total variance of blood pressure and for 506 mg and 34% of the total variance of heart wt. (*J. Clin. Invest.* 1996. 97:777-788.) Key words: Dahl rats • hypertension • body weight • SA locus • cytochrome P450

Introduction

Identification of the multiple genetic loci controlling continuously varying (quantitative) genetic traits such as blood pressure, is a major challenge. Using many genetic markers and linkage analysis it is possible to identify chromosomal regions that contain quantitative trait loci (QTL)¹ (1, 2). The identification of the actual QTL, as opposed to the chromosomal region containing the QTL, remains a more formidable obstacle

(3, 4). This paper deals with the identification of chromosomal regions influencing blood pressure using an animal model of hypertension, the Dahl salt-sensitive (S) rat. This hypertensive strain and others have been used previously to locate such regions (4).

Among the loci known to cosegregate with blood pressure is a gene of unknown function called the SA gene. This gene was originally cloned by Iwai and Inagami (5) based on its greater expression in the kidneys of spontaneously hypertensive rats (SHR) compared to Wistar-Kyoto (WKY) normotensive controls. The SA gene has been linked to markers on rat chromosome 1 (6, 7). Alleles at the SA locus were shown to cosegregate with blood pressure in F_2 populations derived from an SHR \times WKY cross (8, 9), a stroke prone-SHR \times WKY cross (6), and an F_2 population derived from an S \times Lewis cross (10). The SA locus in humans was linked to blood pressure in a Japanese population (11) but not in a European population (12).

In an attempt to improve the genetic map of rat chromosome 1, and to determine if the blood pressure effect in fact localizes maximally at or very close to the SA locus, we developed an improved genetic map for chromosome 1. We have found that rat chromosome 1 contains two blood pressure QTL, one very close or identical to the SA locus and a second QTL about 55 centiMorgans (cM) from the SA locus.

Methods

Animal procedure. Inbred Dahl salt-sensitive (SS/Jr) and inbred Dahl salt-resistant (SR/Jr) rat strains (13) were from our colony at Medical College of Ohio and will be referred to hereafter as S and R rats. Lewis rats (LEW/NCrLBR) were obtained from Charles River Laboratories (Wilmington, MA) and are referred to as LEW. Spontaneously hypertensive rats (SHR/NHsd), Wistar-Kyoto (WKY/NHsd) and Brown Norway (BN/SsNHsd) rats were obtained from Harlan Sprague-Dawley (Indianapolis, IN) and will be referred to as SHR, WKY, and BN, respectively. The Milan normotensive strain (MNS) was obtained from the Veterinary Resources Branch of the National Institutes of Health (Bethesda, MD). Albino surgery (AS) rats were obtained from C. Heatherington (National Institute for Medical Research, The Ridgeway, Mill Hill, UK).

Five large F_2 populations of male rats were used for blood pressure cosegregation analysis and have been described in previous work (14). Briefly, the F_2 populations were obtained by crossing S males with various contrasting strain (LEW, WKY, R, BN, and MNS) females, and then intercrossing the F_1 progeny to produce F_2 rats. The

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1. *Abbreviations used in this paper:* AS, Albino surgery rat; BN, brown Norway rat; DOP, degenerate oligonucleotide primer; LEW, Lewis rat; MNS, Milan normotensive strain; QTL, quantitative trait locus; R, inbred Dahl salt-resistant rat; S, inbred Dahl salt-sensitive rat; SHR, spontaneously hypertensive rat; WKY, Wistar-Kyoto rat.

Table I. Microsatellite PCR Primers and Rat Strain Variation

Locus name	Gene name	GenBank accession number	Primers (5'-3')	Length variation of the PCR-product among 8 inbred strains
<i>ALDOA</i>	Aldolase A	X04264	CAGCCCCTGCAATAAACAGC CCTACCCTTCTCAAAGCTCTGC	LEW<BN<WKY<S=R=SHR<MNS=AS
<i>ATPIA3</i>	ATPase alpha 3 subunit	M90659	TGAGCTTCTGGTTGAAGGATCG CTCCACATATACCACCAAAGGC	BN=WKY<SHR<S=R=LEW=MNS=AS
<i>ATP4A</i>	Gastric H, K-ATPase	M63963	CTCTTCTGATGGTGGAAC ATGAGGCTTCGACTTACC	S=R=BN=LEW=AS<WKY=SHR=MNS
<i>CALM3</i>	Calmodulin III	X14265	CCACAACCCTCTCTACTTGC TATGCCGAAAGGATTTATGGCG	BN=LEW=AS<S=R=MNS<WKY=SHR
<i>CEAR</i>	Carcinoembryonic antigen-related protein (CGM4)	M32475 J05417	GCTCATCATCATCATCACC CCCCTGCTTCTTATTACC	BN=LEW=AS<WKY=SHR<S=R=MNS
<i>CYPE</i>	Cytochrome P450e	Y00410	AGGAAAAGCATATAGAACACGC TTTACTTACTAGGGCATGGGAT	S<MNS<WKY=SHR<R=BN=LEW=AS
<i>CYP2A1</i>	Cytochrome P450 IIA1	M33312	AAGAGAGAAACACTCTGTGG TTCCAGTAGGAGACTAAAGTGG	S=WKY=SHR=MNS<R=BN=LEW=AS
<i>CYP2A2</i>	Cytochrome P450 IIA2	M33313	TATAAAAAGCCCCATGCTGGAGC GGCAAATAGGAAGAAAGGTGGC	WYK=SHR=MNS<S<R=BN=LEW<AS
<i>CYP2A3</i>	Cytochrome P450 IIA3	M33190	TACACATGCACCTGCATGCA TCTTGCCAGTTGCTAAAAATGAG	S=MNS<WKY=SHR<R=BN=LEW=AS
<i>CYP2B2</i>	Cytochrome P450 2B2	S51970	TGGCTTGGAGCAGAAAAGTTGG CTTCTTGAAGGTTGTGTCCACG	S=BN<AS<R=LEW=MNS<WKY=SHR
<i>CYP2C12</i>	Cytochrome P450 15-beta gene	M33545	GAAAAGGCTACTAAAGGACTGG ATAGAGTGCCAACTATGCC	BN<WKY=SHR<S=R=MNS<LEW=AS
D1Mco1	Random cloned SSR	U19349	CTTATCAGCGAATGGATAC CCTATTTGTATTTTCATCCC	AS<LEW=BN=WKY=SHR=MNS<S=R
D1Mco2	Random cloned SSR	U19350	ATGTGGTAAGAGTTTGAATC GAAATAATAGAGTATGTGTGGG	LEW=MNS<S=R<BN<WKY=SHR=AS
D1Mco3	Random cloned SSR	U19351	GTAAAGTGGTCTGTGCC TTGCATGAAATGATGTGTG	S<R=LEW=BN=MNS=AS<WKY=SHR
D1Mco4	Random cloned SSR	U19352	CGGTGTACAAGATTTGTC GAGTATGATACCAGGACTC	S=R=WKY=MNS=AS<LEW=BN=SHR
D1Mco5	Random cloned SSR	U19357	ACGGTCACTGGGTTCTTG TCACTAGCTTCCTAGGGC	WKY=SHR<LEW<S=R=BN=AS<MNS
D1Mco6	Random cloned SSR	U29197	CTTCAGTGAGAGAAGATGCACC CCAACATTTCCCTCATCCC	WKY<LEW=BN=SHR=AS<S=R
D0Mco1*	Random cloned SSR	U19353	CTTTATGTGGGTTCTGAGG AGGTATTGAGATGTAGGATATG	S=R=LEW=BN=WKY=SHR=MNS=AS
D0Mco2 [‡]	Random cloned SSR	U19354	CAGCCTTCATTCTCACAC TGACTTCTGTGAGCTCCTAC	MNS<WKY=SHR=AS<S=R=LEW<BN
D0Mco3*	Random cloned SSR	U19355	CTCAGAGACAAAGCTAAGG AGGAAACATACAGACCCAG	S=R=LEW=WKY=NB=SHR=MNS=AS
D0Mco4 [‡]	Random cloned SSR	U19356	GGCACGTGTACATACTGATG TCTATTCAGGCCTCTGAC	LEW=BN<S=R=WKY=SHR=MNS=AS
D1Mgh1*	Random cloned SSR	Research Genetics		BN<SHR<S=R=LEW=WKY<MNS=AS
D1Mgh2	Random cloned SSR	Research Genetics		WKY=SHR<S=R=LEW=BN=MNS=AS
D1Mgh3	Random cloned SSR	Research Genetics		BN<WKY=SHR<S=R=MNS=AS<LEW
D1Mgh4	Random cloned SSR	Research Genetics		LEW=BN=MNS=AS<S=R<WKY=SHR
D1Mgh6	Random cloned SSR	Research Genetics		S=R=LEW=BN=MNS=AS<WKY=SHR
D1Mgh7	Random cloned SSR	Research Genetics		LEW=BN<MNS<WKY=SHR<S=R=AS

Table I. Continued

Locus name	Gene name	GenBank accession number	Primers (5'-3')	Length variation of the PCR-product among 8 inbred strains
D1Mgh8	Random cloned SSR		Research Genetics	LEW<S=R=MNS<BN<WKY=SHR<AS
D1Mgh9	Random cloned SSR		Research Genetics	LEW=MNS<S=R=WY=SHR=AS<BN
D1Mgh10	Random cloned SSR		Research Genetics	S=R=LEW=BN=AS<MNS<WKY=SHR
D1Mgh11	Random cloned SSR		Research Genetics	WKY=SHR<S=LEW=BN=AS<MNS<R
D1Mgh12	Random cloned SSR		Research Genetics	LEW=AS<S=R<BN=MNS<WKY=SHR
D1Mgh13	Random cloned SSR		Research Genetics	BN=MNS<AS<S=R=LEW<WKY<SHR
D1Mgh14	Random cloned SSR		Research Genetics	BN<S=SHR<R=LEW=WKY=MNS=AS
D1Mgh15*	Random cloned SSR		Research Genetics	S=R=LEW=WKY=SHR=MNS=AS<BN
D1Mgh16	Random cloned SSR		Research Genetics	LEW=WKY=SHR=MNS<S=R=AS<BN
D1Mgh17*	Random cloned SSR		Research Genetics	S=R=LEW=BN=WKY=SHR=MNS=AS
D1Mgh18‡	Random cloned SSR		Research Genetics	R<WKY<SHR<S=LEW=BN=MNS=AS
D1Mgh19	Random cloned SSR		Research Genetics	WKY=SHR<S=R=BN=AS<LEW=MNS
D1Mgh20	Random cloned SSR		Research Genetics	BN<AS<S=R=WKY=SHR<MNS<LEW
D1Mgh21	Random cloned SSR		Research Genetics	LEW<S=R=WKY<SHR=MNS<BN=AS
D1Mit2*	Random cloned SSR		Research Genetics	S=LEW=WKY=AS<SHR=MNS<BN<R
D1Mit3	Random cloned SSR		Research Genetics	SHR=MNS<S=R=LEW<WKY=AS<BN
D1Mit4	Random cloned SSR		Research Genetics	S=R=BN<LEW=MNS<WKY=SHR=AS
D1Mit5	Random cloned SSR		Research Genetics	LEW=BN<WKY=SHR<S=R=MNS=AS
D1Mit7	Random cloned SSR		Research Genetics	SHR<WKY=AS<S=R=LEW=MNS<BN
D1Mit8	Random cloned SSR		Research Genetics	WKY=SHR=MNS=AS<LEW=BN<S=R
D1Mit9	Random cloned SSR		Research Genetics	WKY=SHR<MNS<S=R<LEW=BN=AS
D1Mit10*	Random cloned SSR		Research Genetics	BN<S=R=LEW=WKY=SHR=MNS=AS
D1Mit12	Random cloned SSR		Research Genetics	AS<BN<S=R<WKY=SHR<LEW=MNS
D1Mit14	Random cloned SSR		Research Genetics	S=R=SHR<LEW=WKY<MNS<AS<BN
D1Mit15*	Random cloned SSR		Research Genetics	B<S=R=LEW=WKY=SHR=AS<MNS
D1N40	DNA segment, STMS	L08080	TGACCTAGCATGAGCAGG GCCCAACTTGCTTTTGG	WKY=SHR<S=R=BN=LEW=MNS=AS
D1N64	DNA segment, STMS	L08081	ATGTGTTTGTGGGTGGTTCCAGCAGG TCTCTCTCTCTCTCGTGTGTGT	LEW=WKY=SHR=MNS=AS<S=R<BN
D7Mit45	Random cloned SSR		Research Genetics	BN=LEW<WKY=SHR<S=R=MNS=AS

Table I. Continued

Locus name	Gene name	GenBank accession number	Primers (5'-3')	Length variation of the PCR-product among 8 inbred strains
D7Mit62 [§]	Random cloned SSR		Research Genetics	S=R=WKY=SHR=AS<LEW=BN=MNS
D7Mit66	Random cloned SSR		Research Genetics	BN<S=R=WKY=AS<MNS<LEW=SHR
D7Mit69	Random cloned SSR		Research Genetics	S=R=AS<BN=LEW=WKY=SHR=MNS
D7Mit87	Random cloned SSR		Research Genetics	S=R=AS=SHR=MNS<BN=LEW=WKY
D7Mit89 [§]	Random cloned SSR		Research Genetics	LEW=MNS=AS<S=R=BN=WKY=SHR
D7Mit99	Random Cloned SSR		Research Genetics	WKY<S=R=BN=LEW=SHR=MNS=AS
D7Mit101	Random Cloned SSR		Research Genetics	S=R=WKY=SHR=MNS<LEW=AS<BN
D7Mit109	Random Cloned SSR		Research Genetics	WKY=SHR<S=R=MNS=AS<LEW<BN
EN3C [§]			Research Genetics	SHR=MNS<BN<S=R=AS<LEW=WKY
<i>IGF2</i>	Insulin-like growth factor	X17012	GATTATACCCACACGTACATGC AAACCATGCAAAGTCTCAGGG	BN=LEW<S=R<WKY=SHR=AS<MNS
<i>KAL</i>	Kallikrein	M19647	AAATGCGAAAGTGTCTTGGC ACAAAGTTATGGGATGGCAG	WKY<BN=MNS<R=SHR<S=LEW=AS
<i>LSN</i>	Leukosianin		Research Genetics	S=R=LEW=BN=MNS=AS<WKY=SHR
<i>MTIPA</i>	Metallothionein, pseudogene A		Research Genetics	BN=LEW=WKY=AS<S=R=SHR=MNS
<i>MYL2</i>	Myosin light chain, muscle	X00975	TATAACCCAGAAAGAACTGCC ACCTGTAGTTGGAATGAGAAGC	S=R=MNS<WKY=SHR<BN=LEW=AS
<i>PBPC2*</i>	Prostatic binding protein, C2	X05034	TGTGTACAGACAAGAAGTTCC CACACTTGGCAAATTCCTTTC	S=R=BN=LEW=WKY=SHR=AS<MNS
<i>PKC</i>	Protein kinase C type I	M13707	AGAACCCTCACTGCTCACC TGAGAAAGTCCCAGAAAGTGGC	S=R=WKY=MNS<AS<BN=LEW=SHR
<i>PTH*</i>	Parathyroid hormone	K01266	TTCCTCTGTGTGCATGAG CAAACAGCAAGCCTTAGG	BN=MNS<S=R=LEW=WKY=SHR=AS
R179 [§]	Random cloned SSR		CGTTCACCGCTTTGTGTC CTCTGTGCTTGCTACTGTCC	BN<S=LEW=WKY<R<SHR<AS<MNS
R191 [§]	Random cloned SSR		TTCGGACTCAGTCTTGATTTGG GCCTTCTAGAAATACTTGGT	BN=AS<S=R=MNS<LEW<WKY=SHR
R197 [§]	Random cloned SSR		ATTGTATATTCCAGACTAGC AATTGATGTGACATTATTTTCAT	S=MNS=AS<R=BN<LEW=WKY=SHR
R401 [§]	Random cloned SSR		TGAGCCTGGGCACTATGTAG GGACAGGGACTGGAATCATC	LEW=BN=AS<S=R=MNS<WKY=SHR
R416 [§]	Random cloned SSR		GGTATTGCATGGCTATGAGG GCAGCCATAGAGTGCCAAC	LEW=BN=WKY<SHR<S=R=MNS=AS
RCA01.20 [§]	Random cloned SSR		GTAGGTGTAGAAAAGATGCTGC ATCAATGGAGGCTCTGATGGG	LEW=BN=WKY=MNS<S=R=AS<SHR
RCA07.06 [§]	Random cloned SSR		CCAATTTCTCAACATCACCCC CAATATCTATGATAGATGATGG	BN=AS<R<S=WKY<LEW=SHR=MNS
RCA09.01 [§]	Random cloned SSR		CCCCATCTATCTATCCAACGG CTCTGGGATGCTTTGTGAAGG	MNS<AS<S=R=WKY=SHR<LEW<BN
RCA17.42 [§]	Random cloned SSR		GTAGTTGTGACTGCCTTTCCTG GATGAGAACATCCTGGAGAATG	S=R=WKY=SHR=MNS=AS<LEW=BN
RCA24.16 [§]	Random cloned SSR		ATCACTCTGACCAAGAGACC TCACTTTCACATGACAGACCC	WKY=SHR<MNS<S=R=LEW=BN=AS
<i>SA(PSA1)</i>	Anonymous SA gene	U04637	CCTACACAGCAAGTTCAGAGC TACCCCATTCCTATCTCC	S=SHR=MNS=AS<R<BN<LEW=WKY
<i>SA(PSA2)</i>	Anonymous SA gene	U04638	CCTAAATTAGCTTGTAGGGAGG TACCTAGCCTTAGGGATTTCG	S=R=SHR=MNS=AS<BN<LEW=WKY

Table I. Continued

Locus name	Gene name	GenBank accession number	Primers (5'-3')	Length variation of the PCR-product among 8 inbred strains
SA(PSA3)	Anonymous SA gene	U04639	TGTTACACTCAGGCTTCTATCC ATTCTCTTCCGAAAAAGC	LEW=WKY<S=R=SHR=MNS=AS<BN

SSR, simple sequence repeat. STMS, sequence tagged microsatellite site. Primers were designed using the Primer Detective Program (Clontech, Palo Alto, CA) except *CYPE* (20), *CYP2A3* (22), and D1N64 (23). D1Mco markers were developed by screening the flow-sorted chromosome 1 library with (CA)₁₅ and (CT)₁₅ oligonucleotides. Primers purchased from Research Genetics (Huntsville, Alabama) were D7Mits which are mouse MapPair primers and D1Mits and D1Mghs which are rat MapPair primers. *D0Mco1, D0Mco3, D1Mgh1, D1Mgh15, D1Mgh17, D1Mit2, D1Mit10, D1Mit15, *PBPC2*, *PTH* and R179 could not be placed on the map in Fig. 1 because these markers were not polymorphic among S, LEW, and WKY rats. †These markers were either reported to be, or were thought to be, on rat chromosome 1, but they were not linked to any marker of the linkage group shown in Fig. 1; they are D0Mco2, D0Mco4, D1Mgh18, D7Mit62, D7Mit89, and EN3C. ‡Primers developed at the University of Oxford, England.

F₂ populations were weaned at 30 d of age and placed on an 8% NaCl diet (Teklad diet 82050; Teklad Premier Laboratory Diets, Madison, WI) at 37 d of age. When the highest blood pressures had reached > 200 mmHg, the blood pressures of the rats were measured several times during a 7–10-d period. Systolic blood pressure was taken by the tail cuff method on the conscious restrained rat (15) (ITC, Inc., Woodland Hills, CA) with the ambient temperature maintained at 29°C. At least three consecutive consistent readings were taken at a given session and averaged as that session's reading. Blood pressures at three separate sessions separated by 2 d were taken. These three separate sessions' pressures were averaged as the final blood pressure.

The rats were killed with an overdose of sodium pentobarbital. Pieces of liver were stored at -70°C. DNA for genotyping was extracted from frozen livers by the method of Blin and Stafford (16).

Markers from chromosome 1-sorted DNA. Rat chromosome 1 DNA was obtained by flow sorting as described by Hoebee et al. (17) and amplified by an initial PCR using a degenerate oligonucleotide-primer (DOP) (6-MW primer: 5'-CCGACTCGAGNNNNNATGTGG-3') as described by Telenius et al. (18, 19). Reamplification was carried out in a total vol of 100 µl reaction mixture containing 5 µl PCR products from the previous round of amplification, 1 × PCR buffer, 200 µM of each of the four dNTPs, 800 nM of 6-MW primer, and 1.25 U of *Taq* DNA polymerase (Perkin-Elmer Cetus, Norwalk, CT). After an initial 5-min denaturation at 95°C the reaction was subjected to 35 cycles of 1 min at 94°C, 2 min at 55°C, 3 min at 72°C, followed by an additional 10 min at 72°C for extension. 20 µl of PCR products were subjected to an 0.8% agarose gel electrophoresis to check if the DOP-PCR worked. If the DOP-PCR works, a smear of DNA between 250 bp to 1,500 bp of DNA is observed.

75 µl of DOP-PCR-amplified rat chromosome-1 DNA was digested with *Xho*I restriction enzyme in a total volume of 100 µl at 37°C for 3 h (there is an *Xho*I site in the 5'-end of the 6-MW primer). The digested DNA was concentrated in a centricon-30 microconcentrator (Amicon, Inc., Beverly, MA) by centrifugation at 1,000 g for 2 h to get about 45 µl of concentrated DNA. This DNA was cloned into the *Sa*II site of the vector pT7T3 U18 plasmid (*Sa*II and *Xho*I yield compatible ends). Clones were screened with ³²P end-labeled (CA)₁₅ and (CT)₁₅ oligonucleotides. The positive clones were purified by two additional rounds of screening and sequenced. PCR primers were designed using the Primer Detective Program (Clontech, Palo Alto, CA). Markers developed by this method are given in Table I and were: D1Mco1, D1Mco2, D1Mco3, D1Mco4, D1Mco5, D1Mco6, D0Mco1, D0Mco2, D0Mco3, D0Mco4.

Additional markers. New microsatellite based markers were also obtained from a small insert library by screening with (CA)₁₅ and (CT)₁₅ oligonucleotides. Markers that were linked to known markers on chromosome 1 in various segregating rat populations were used here. These markers are given in Table I and were: R179, R191, R197, R401, R416, RCA01.20, RCA07.06, RCA09.01, RCA17.42, RCA24.16.

Markers were also designed around microsatellite sequences in

GenBank based on information from reference [20]; see Table I for locus designation and GenBank accession number. Rat Map Pair primers (21) (Research Genetics, Huntsville, AL) are listed in Table I. Mouse chromosome 7 has homology with rat chromosome 1; and nine mouse chromosome 7 markers (D7Mit45, D7Mit62, D7Mit66, D7Mit69, D7Mit87, D7Mit89, D7Mit99, D7Mit101, D7Mit109) (Research Genetics) were found to be useful in the rat. Primers for *CYPE*, *CYP2A3*, and D1N64 were from references (20, 22, 23), respectively. Microsatellites for the SA gene were recently published (24). Rats used here had also been previously typed at the *GLUTB* and *HITH* loci (25) on rat chromosomes 5 and 17, respectively.

Genotyping. Genotyping was done by PCR amplification of DNA around microsatellites. PCR reaction vol was 25 µl which contained 1 × PCR buffer, 1–2.5 mM MgCl₂, 200 µM of each the four dNTPs, 1 µM of each primer, 0.625 U *Taq* polymerase (Perkin-Elmer Cetus), and 20–200 ng rat genomic DNA. PCR was started at 95°C for 5 min and continued for 25–35 cycles of 94°C for 1 min, 45–65°C (depending on the primer) for 2 min, 72°C for 2 min, followed by additional 72°C for 10 min. The PCR products were subjected to electrophoresis on a 4% agarose gel which contained 1 mg/ml ethidium bromide for photographing the PCR products under the ultraviolet light. In cases where the PCR allelic products were too close to separate on an agarose gel, a polyacrylamide sequencing gel was used. In this case one of the primers was end-labeled with ^γ-³²P-ATP with T4 polynucleotide kinase and used in the PCR reaction without purification. After PCR the reaction was denatured by adding an equal volume of formamide containing 0.1% bromophenol blue and 0.1% xylene cyanol and heating at 100°C for 2 min, cooling on ice, and 4 µl was loaded onto an 8% polyacrylamide sequencing gel. After electrophoresis the gel was dried and exposed 2–14 h at -80°C to X-OMAT AR film (Eastman Kodak, Rochester, NY) using a fluorescent screen, and developed.

Linkage and statistical analysis. Cosegregation of blood pressure (body weight or heart weight) with alleles at a marker locus was evaluated by comparing blood pressures among genotypes at each locus by a one-way ANOVA using SPSS programs (SPSS, Chicago, IL). When testing for dominance of an allele at a given locus the one-way ANOVA was followed by a contrast that compared the average value of blood pressure of the two homozygous groups to that of the heterozygotes. Linkage maps and QTL localization were done with the MAPMAKER Programs (26, 27) obtained from Dr. Eric Lander (Whitehead Institute, Cambridge, MA). The MAPMAKER programs also detect potential genotyping errors based on results of flanking markers. These samples were always retyped to confirm or correct the results.

Results

Fig. 1 shows the genetic maps constructed for the F₂(WKY × S) and F₂(LEW × S) populations and the composite map for

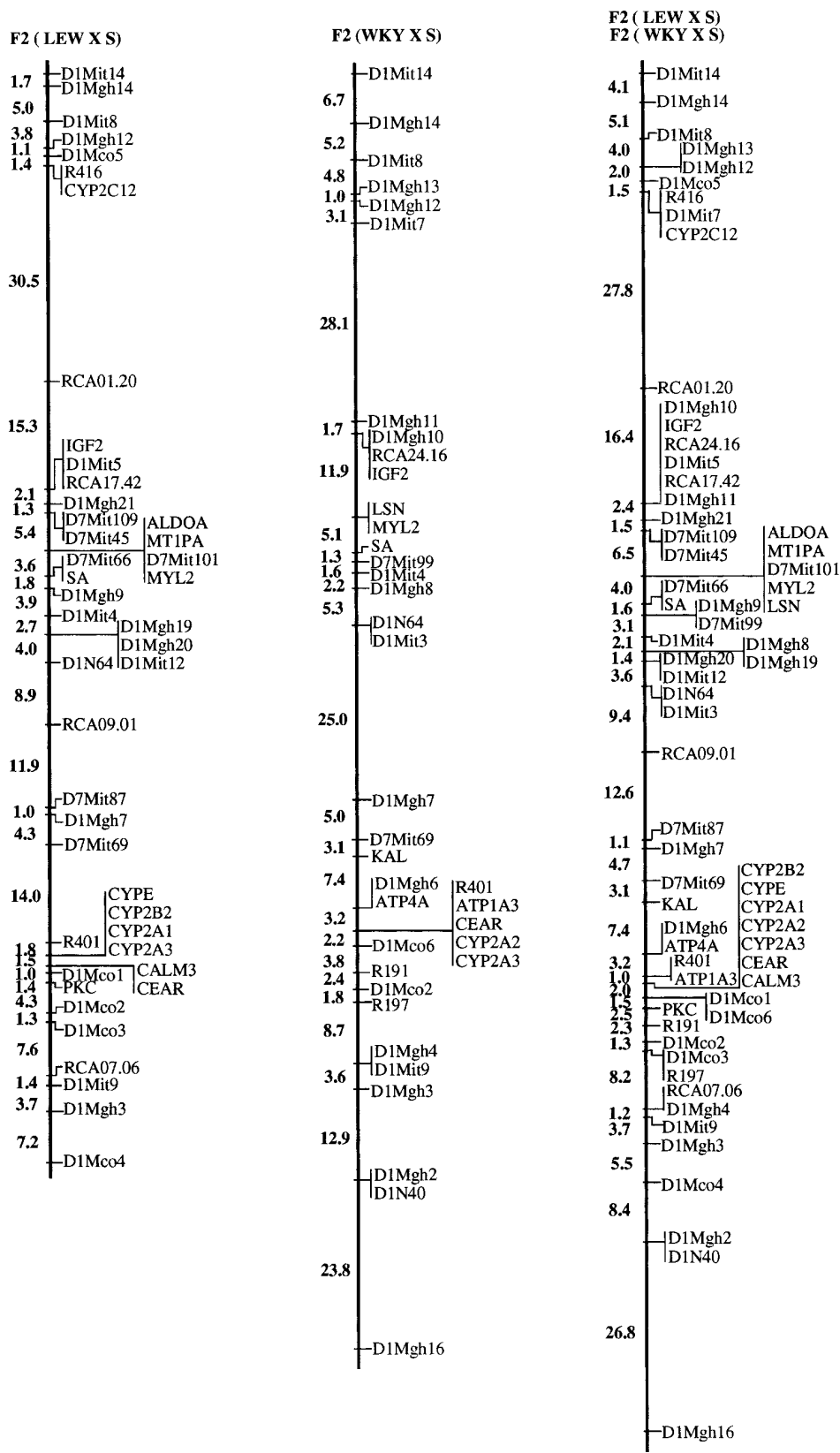


Figure 1. Linkage maps for rat chromosome 1. The left panel is the map for an F_2 (LEW \times S) population of 151 rats, center panel for an F_2 (WKY \times S) population of 159 rats, and the right panel is a map for the two populations combined. The position of the centromere is unknown. Genetic distances are in centiMorgans using the Haldane correction. The map was constructed using the MAP-MAKER programs (26, 27).

the combined populations. Table II gives the blood pressure of each of the three genotypes segregating at selected loci on chromosome 1 for the F_2 (LEW \times S) population. The analysis of this blood pressure data for the F_2 (LEW \times S) population is

summarized in Fig. 2 in conjunction with a map of selected loci for chromosome 1.

Note in Fig. 2 that the blood pressure effect (the blood pressure of rats homozygous [SS] for the S-rat allele minus the

Rat Chromosome 1 Linkage Map	Blood Pressure Effect	One-way ANOVA F	P	QTL Location
7.1	D1Mit14	-0.1	0.36	0.6986
	D1Mit8	-0.9	0.44	0.6433
6.6	CYP2C12	+6.4	0.56	0.5698
30.1				
	RCA01.20	+10.8	0.94	0.3922
15.3				
	IGF2	+11.3	0.16	0.32
9.4				
4.2	MYL2	+25.3	5.62	0.0044
4.4	SA	+24.0	5.61	0.0045
6.1	D1Mit4	+21.2	4.05	0.0194
8.6	D1N64	+13.2	2.75	0.067
	RCA09.01	+22.6	4.57	0.0119
11.6				
	D7Mit87	+19.3	2.61	0.0771
5.1	D7Mit69	+26.0	5.42	0.0053
14.6				
2.7	R401	+27.1	5.91	0.0034
5.8	D1Mco1	+30.4	7.87	0.0006
	D1Mco2	+24.1	4.94	0.0084
9.0				
	RCA07.06	+14.7	1.72	0.183
5.3	D1Mgh3	+14.7	2.22	0.1119
7.1				
	D1Mco4	+12.4	1.21	0.3019

Figure 2. Linkage map of rat chromosome 1 and associated blood pressure linkage data for the F₂(LEW × S) population of 151 rats using selected markers from Fig. 1. Rats were raised on 8% NaCl diet for 9 wk starting at 37 d of age. Genetic distances are cM with the Haldane correction; the position of the centromere is unknown. "Blood pressure effect" is the difference in blood pressure between rats homozygous for the S-rat allele, and those homozygous for the Lewis-rat allele at a given marker. Values for the F statistic (F) and probability (P) from an ANOVA are given to evaluate the effect of genotype at each locus on blood pressure. "QTL location" and the map itself were generated using the MAPMAKER/QTL programs (26, 27). The arrow indicates the position of the maximum LOD score and the vertical bar indicates ±1 LOD for linkage to blood pressure.

blood pressure of rats homozygous [LL] for the Lewis-rat allele) is positive and reaches a maximum in the region of the SA gene. The values decrease in the middle of the map and then increase markedly toward the bottom of the map. The F statistic and the associated probability value from a one-way ANOVA at each locus mirror these changes, the most significant effects occurring near the SA locus (P = 0.0045) and near the D1Mco1 locus (P = 0.0006). The SA and D1Mco1 loci are ~55 cM apart (using the Haldane mapping function), which is far enough to segregate essentially independently of each

Table II. Systolic Blood Pressure by Genotype for Selected Loci on Rat Chromosome 1 in the F₂(Lew × S) Population Raised on 8% NaCl Diet

Locus	Blood Pressure by Genotype		
	SS	SL	LL
D1Mit14	190.6±6.32 (38)	185.7±3.93 (71)	190.7±5.66 (40)
D1Mit8	190.3±6.03 (39)	185.3±4.08 (69)	191.1±5.29 (43)
CYP2C12	193.5±6.11 (37)	186.2±4.02 (74)	187.1±5.40 (40)
RCA01.20	193.0±5.99 (34)	189.3±4.07 (77)	182.2±5.33 (40)
IGF2	191.8±5.17 (47)	189.8±4.40 (69)	180.5±5.22 (35)
MYL2	200.7±5.30 (48)	185.5±4.33(72)	175.5±3.63 (31)
SA	200.7±5.18 (50)	184.7±4.55 (68)	176.7±3.20 (33)
D1Mit4	197.8±5.07 (55)	185.6±4.52 (66)	176.6±3.44 (30)
D1N64	197.8±5.38 (49)	183.2±4.20 (69)	184.6±5.02 (33)
RCA09.01	198.1±5.31 (53)	187.7±4.24 (64)	175.5±4.17 (33)
D7Mit87	195.0±5.76 (44)	189.5±3.83 (80)	175.7±5.24 (26)
D7Mit69	196.9±5.90 (42)	190.2±3.89 (79)	170.9±4.55 (30)
R401	202.6±6.13 (38)	187.0±3.69 (78)	175.5±5.48 (35)
D1Mco1	205.6±6.34 (38)	185.7±3.51 (78)	175.2±5.62 (34)
D1Mco2	203.8±6.88 (36)	186.9±3.46 (74)	179.7±5.61 (37)
RCA07.06	196.2±6.23 (41)	187.6±3.75 (73)	181.5±4.90 (35)
D1Mgh3	192.8±6.69 (35)	190.5±3.90 (75)	178.1±4.73 (40)
D1Mco4	195.2±6.85 (33)	188.5±4.02 (73)	182.8±4.85 (45)

Blood pressure data is mean±SE in mmHg for each genotype. The number of rats in each genotypic class is given in parenthesis. S, allele for S rats, L, allele for LEW rats. See Table I for locus designations, and Fig. 2 for statistical analysis of blood pressure data.

other. The MAPMAKER/QTL program gave LOD score peaks for linkage to blood pressure in the region of the SA and D1Mco1 loci of 2.45 and 3.38, respectively.

There was a strong correlation in the F₂(LEW × S) population between heart weight and blood pressure (Fig. 3). A linkage analysis of heart wt, body wt and heart wt/body wt ratio with selected loci along chromosome 1 is given in Table III. Heart weight showed basically the same pattern as blood pressure. The S allele increased heart weight in two regions, one around the SA locus and a second region around the D1Mco1 locus. Note, however, that the effect on heart wt of the S allele at the SA locus (+82.3 mg, P = 0.039) appears relatively weak compared to the effect of the D1Mco1 locus (+130.4 mg, P = 0.0036). This could be due to the lower blood pressure associated with the SA region compared to D1Mco1. It could also be related to the fact that the S allelic region around the SA locus had a negative effect on body wt, which would be expected to reduce absolute heart size. Correcting heart wt for differences in body wt by using the heart wt/body wt ratio in Table III yielded a very strong and positive effect of the S allelic region around the SA locus on heart wt/body wt ratio. Using body wt as a covariate in an ANOVA yielded a more significant (P = 0.001) effect of the SA region on heart wt than did the analysis of uncorrected heart wt.

It was possible to test both the chromosome 1 region marked by the SA locus and the new chromosome 1 blood pressure QTL region marked by D1Mco1 for linkage to blood pressure in populations derived from crosses of S rats with various normotensive strains. The SA locus cosegregated with

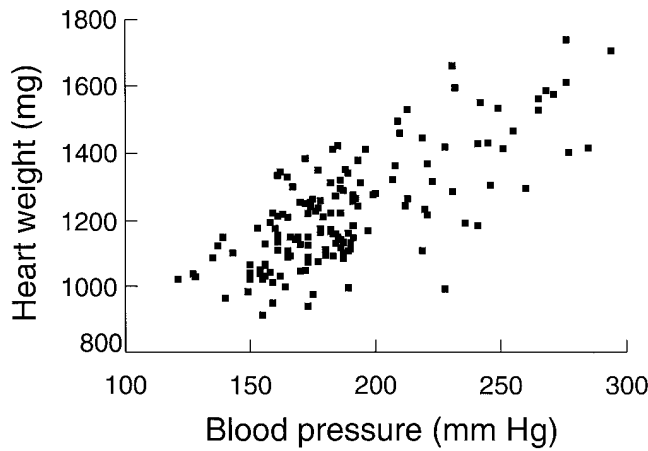


Figure 3. Relationship between heart weight and blood pressure in the F_2 (LEW \times S) population raised on 8% NaCl diet for 9 wk starting at 37 d of age. $r = 0.75$, $P < 0.001$, $n = 151$ rats.

blood pressure only in the F_2 (LEW \times S) population, but not in F_2 (WKY \times S) or in F_2 (R \times S) or in a backcross to the S, F_1 (R \times S) \times S (Table IV). Using loci near D1Mco1 (*CYP2A3* and *CYP2B2* were used), five F_2 populations were tested for linkage of this chromosomal region to blood pressure. Only the F_2 (LEW \times S) gave a clearly significant linkage to blood pressure (Table V).

The F_2 (LEW \times S) population is remarkable because, besides the two QTL on chromosome 1, we know of two other

chromosome regions with substantial effects on blood pressure. These are marked by *GLUTB* on chromosome 5, and *HITH* on chromosome 17, as reported previously (25). The effects of all four chromosomal regions (marked by *SA*, *D1Mco1*, *GLUTB*, and *HITH*) on blood pressure and heart weight are summarized in Tables VI and VII, respectively. There were no interactions between any two of these loci on either blood pressure or heart weight by two-way factorial ANOVA analyzing the loci pairwise, or in a four-way factorial analysis with all four loci simultaneously. The effects of these loci are therefore additive, or, at least they are additive on the genetic background of the F_2 (LEW \times S) population. As shown in Tables VI and VII these four loci accounted for 106 mmHg and 38% of total variance for blood pressure, and 506 mg of heart wt (the average heart wt for all rats in the population was 1,217 mg) and 34% of total variance for heart wt.

The frequency distribution for blood pressure in the F_2 (LEW \times S) population was significantly skewed to the right; the g_1 statistic for skewing was 0.98 which is statistically significant ($P < 0.005$). Correcting for the blood pressure effects of genotypes at the *SA*, *D1Mco1*, *GLUTB*, and *HITH* loci yielded a "corrected" distribution which was still skewed to the right ($g_1 = 0.41$, $0.025 < P < 0.05$) but not as badly as the uncorrected distribution. The variance of the uncorrected blood pressure distribution was $1,226 \text{ (mmHg)}^2$ and for the corrected distribution 883 (mmHg)^2 . Because the uncorrected blood pressure distribution was skewed to the right, all ANOVAs reported above were repeated on log transformed data and by a nonparametric Kruskal-Wallis one-way ANOVA. Neither procedure altered the conclusions given above.

Table III. Analysis of the Effect of Rat Chromosome 1 on Body Weight (BW), Heart Weight (HW), and Heart Weight/Body Weight Ratio for the Male F_2 (LEW \times S) Population Raised on 8% NaCl Diet

Locus	Effect	HW		Effect	BW		HW/BW Ratio		
		F	P		F	P	Effect	F	P
	mg			g			mg/g		
D1Mit14	+33.4	1.44	0.24	-7.8	1.12	0.33	+0.176	1.56	0.21
D1Mit8	+41.9	1.41	0.25	-6.8	0.72	0.49	+0.183	2.45	0.090
<i>CYP2C12</i>	+64	2.50	0.086	-13.4	2.34	0.095	+0.336	5.91	0.0034
RCA01.20	+57.7	1.08	0.34	-13.3	2.68	0.072	+0.291	3.22	0.043
<i>IGF2</i>	+36.8	0.48	0.62	-17.2	4.44	0.013	+0.274	3.21	0.043
<i>MYL2</i>	+71.4	2.66	0.073	-18.7	4.95	0.0083	+0.408	8.23	0.0004
<i>SA</i>	+82.3	3.36	0.037	-18.0	4.80	0.0096	+0.427	9.38	0.0001
D1Mit4	+82.7	3.78	0.025	-14.3	2.76	0.066	+0.392	8.40	0.0004
D1N64	+38.5	2.48	0.087	-17.6	4.78	0.0097	+0.326	7.62	0.0007
RCA09.01	+64.9	1.93	0.15	-16.9	5.50	0.0050	+0.377	8.35	0.0004
D7Mit87	+50.6	0.90	0.41	-12.1	3.11	0.047	+0.296	2.84	0.062
D7Mit69	+78.3	3.25	0.041	-6.9	2.43	0.092	+0.329	3.98	0.021
R401	+106.6	3.76	0.025	-5.0	0.38	0.68	+0.365	5.05	0.0076
D1Mco1	+130.4	5.84	0.0036	-4.1	0.23	0.79	+0.418	6.72	0.0016
D1Mco2	+111.8	4.26	0.016	-4.9	0.42	0.66	+0.368	5.38	0.0056
RCA07.06	+65.2	1.44	0.24	-10.5	2.33	0.10	+0.291	3.39	0.036
D1Mgh3	+76.0	3.13	0.047	-11.1	2.02	0.14	+0.340	5.36	0.0057
D1Mco4	+51.1	1.03	0.36	-9.7	1.25	0.29	+0.247	2.42	0.092

"Effect" is the difference in the phenotypic trait mean of homozygous *SS* rats minus the phenotypic trait mean of homozygous *LL* rats at each locus where *S* is the allele from S rats and *L* is the allele from Lewis rats. F is the F statistic, and P the probability, from a one-way analysis of variance of phenotype by genotype at each locus.

Table IV. Cosegregation of Systolic Blood Pressure with Genotypes at the SA Locus in Various Segregating Populations Raised on 8% NaCl

Population	Blood pressure by SA genotype			Blood pressure effect	One-way ANOVA probability
	SS	SL	LL		
F ₂ (LEW × S)*	200.7±5.18 (50)	184.7±4.55 (68)	176.7±3.20 (33)	+24.0	0.0045
F ₂ (WKY × S) [‡]	177.0±4.32 (40)	169.3±2.58 (72)	179.9±3.89 (44)	-2.9	0.057
F ₂ (R × S)	167.5±3.85 (25)	161.8±2.67 (56)	166.0±4.13 (30)	+1.5	0.44
F ₁ (R × S) × S	163.0±2.95 (35)	166.3±3.28 (35)			0.46

Blood pressure data are mean±SE in mmHg for each genotype. S, S allele; L, LEW allele; W, WKY allele; number of rats in each genotypic class are in parenthesis; "Blood Pressure Effect" is the difference in blood pressure of homozygous SS minus blood pressure of homozygotes for the contrasting allele. *These data differ slightly from previous published data on this population (10) because the present data includes more rats; also genotyping errors based on restriction fragment length polymorphisms were corrected. [‡]These data were published previously (10), but are included here for comparative purposes.

Discussion

Identification of the chromosomal regions containing blood pressure QTL by linkage analysis is a fundamental first step in understanding genetic hypertension, but the statistical nature of this localization is limiting. The best that can be expected under reasonably favorable conditions (a QTL with a moderate to large effect, a dense genetic map, and a very large population) is to localize the QTL to a region ~ 10 cM in size (28). Thus, it is obvious that if two QTL are on the same chromosome and close together, problems arise in differentiating them.

Using the MAPMAKER/QTL program and simulated data, when two QTL are far apart they were readily recognizable (27). If the two QTL were closer, then simulated data predict that a "ghost" QTL between the actual QTL will be observed (29). Rat chromosome 1 in the F₂(LEW × S)

population provides a reasonably favorable situation in that the two putative QTL are far enough apart to be recognized as distinct. The estimated distance between them is 55 cM (Haldane corrected), which is presumably far enough apart for them to be differentiated statistically. Interference of the two putative QTL in our data, however, may account for the paradoxical increase in blood pressure effect at marker RCA09.01 in Fig. 2, or more likely it is just statistical variation. For comparative purposes the reader is referred to our previous work on rat chromosome 2 (30), which illustrates a reasonable genetic map showing only one QTL, and to work on rat chromosome 10 where there is evidence for two QTL too close to be adequately resolved (31).

In general it is desirable to have the linkage of a chromosomal region to the phenotypic trait supported by a LOD score of ≥ 3 or a one-way ANOVA of $P \leq 0.001$ (2, 27). This is because in searching for QTL so many markers are used that

Table V. Cosegregation of Systolic Blood Pressure with Genotypes at the CYP2A3 (or CYP2B2) Locus in Various F₂ Rat Populations Raised on 8% NaCl

Population	Blood pressure by CYP2A3 genotype			Blood pressure effect	One-way ANOVA probability
	SS	SL	LL		
F ₂ (LEW × S)	203.7±6.00 (41)	186.1±3.55 (77)	174.2±5.72 (33)	+29.5	0.0009
F ₂ (WKY × S)	178.1±5.95 (30)	176.2±2.38 (93)	164.7±3.20 (36)	+13.4	0.034
F ₂ (R × S)	163.8±3.35 (39)	163.4±3.06 (47)	166.4±3.90 (25)	-2.6	0.83
F ₂ (BN × S)	168.9±3.58 (28)	176.4±3.38 (43)	176.0±4.21 (23)	-7.1	0.30
F ₂ (S × MNS)*	175.9±3.49 (50)	171.7±2.19 (74)	173.2±3.06 (43)	+2.7	0.56

Blood pressure data are mean±SE in mmHg for each genotype. S, S allele; L, LEW allele; W, WKY allele; R, R allele; B, BN allele; M, MNS allele; number of rats in each genotypic class are in parenthesis; "Blood pressure effect" is the difference in blood pressure of homozygous SS minus blood pressure of homozygotes for the contrasting allele. *The genotype of the F₂(S × MNS) population was done with CYP2B2 primers as there was no polymorphism between S and MNS using CYP2A3 primers.

Table VI. Systolic Blood Pressure by Genotype for Selected Loci on Rat Chromosomes 1, 5, and 17 in the F₂(LEW × S) Population Raised on 8% NaCl Diet

Locus	Chromosome	Blood pressure by genotype			Blood pressure effect	One-way ANOVA probability	DOM	Percentage of total variance
		SS	SL	LL				
SA	1*	200.7±5.18	184.7±4.55	176.7±3.20	+24.0	0.0045	0.48	7.3
<i>n</i>		50	68	33				
D1Mco1	1*	205.6±6.34	185.7±3.51	175.2±5.62	+30.4	0.0006	0.40	9.6
<i>n</i>		38	78	34				
GLUTB [§]	5	202.1±5.67	192.0±4.44	168.8±2.91	+33.3	< 0.0001	0.22	13.0
<i>n</i>		38	72	41				
HITH [§]	17	187.3±4.61	195.9±4.26	168.9±4.73	+18.4	0.0015	0.002	8.4
<i>n</i>		43	79	29		(0.0008) [‡]		
				Sums:	106.1 mmHg			38.3%

Blood pressure data is mean±SE in mmHg for each genotype. *S*, allele for S rats, *L*, allele for Lewis rats; “Blood pressure effect” is the difference in blood pressure of homozygous *SS* minus blood pressure of homozygous *LL* at each locus; DOM, probability associated with the test for dominance; *n*, number of rats in each genotypic class. See Table I for locus designations. **SA* and *D1Mco1* loci are 55 cM apart, see Figs. 1 or 2. [‡]Probability from one-way ANOVA assuming dominance of *S* allele. [§]Data were published previously (25) but are included here for the purpose of summarizing additive genetic effects.

many false positives will result if significance levels too high, i.e., $P < 0.05$, are accepted. The QTL region marked by the *SA* locus is supported by a LOD of only 2.4 and $P = 0.0045$ by an ANOVA. Thus the evidence by itself is only suggestive of a QTL in this region. But since the present evidence serves to corroborate previously published data from three independent F₂ populations derived from SHR or stroke prone-SHR and WKY indicating linkage of the *SA* locus to blood pressure (6, 8, 9), confidence that the result is correct is enhanced. It is interesting that the peak LOD score is essentially at the *SA* gene. This does not, of course, establish the *SA* gene as the blood pressure QTL in this region. The β and γ subunits of the

renal epithelial sodium channel are closely linked to the *SA* locus on human chromosome 16 (32). Thus it is likely that these sodium channel subunits are closely linked to the *SA* locus on rat chromosome 1. Certain mutations in either the β or γ subunits cause activation of the sodium channel and result in a Mendelian form of inherited hypertension in humans (Liddles syndrome, pseudoaldosteronism) (32, 33). Clearly these genes are also major candidates for the blood pressure QTL in the *SA* region of rat chromosome 1.

The new putative QTL on chromosome 1 that is 55 cM from the *SA* region and marked by *D1Mco1* has a somewhat better statistical support (LOD score of 3.4 and $P = 0.0006$ by

Table VII. Heart Weight by Genotype for Selected Loci on Rat Chromosomes 1, 5, and 17 in the F₂(LEW × S) Population Raised on 8% NaCl Diet

Locus	Chromosome	Heart weight by genotype			Heart weight effect	One-way ANOVA probability	DOM	Percentage of total variance
		SS	SL	LL				
SA	1*	1,266±24.5	1,197±22.4	1,184±18.4	+82	0.037	0.30	4.4
<i>n</i>		50	68	33		(0.001) [‡]		
D1Mco1	1*	1,288±34.2	1,211±16.4	1,157±24.8	+131	0.0036	0.67	7.3
<i>n</i>		38	78	34				
GLUTB	5	1,302±30.8	1,217±19.5	1,137±16.8	+165	< 0.0001	0.91	13.0
<i>n</i>		38	72	41				
HITH	17	1,240±25.0	1,242±20.3	1,112±17.1	+128	0.0008	0.009	9.0
<i>n</i>		43	79	29		(0.0002) [§]		
				Sums:	506 mg			33.7%

Heart weight data is mean±SE in mg for each genotype. *S*, allele for S rats; *L*, allele for LEW rats; “Heart weight effect” is the difference in heart weight of homozygous *SS* minus heart weight of homozygous *LL* at each locus; DOM, probability associated with the test for dominance; *n* = number of rats in each genotypic class. See Table I for locus designations. **SA* and *D1Mco1* loci are 55 cM apart, see Figs. 1 or 2. [‡]Probability for one-way ANOVA using body weight as a covariate. This corrects for an effect of the *SA*-locus chromosomal region on body weight. [§]Probability from one-way ANOVA assuming dominance of *S* allele.

an ANOVA) than the SA region. Candidate genes in this region include a cluster of cytochromes P450. Some cytochromes P450 are involved in the production of biologically active compounds from arachidonic acid. These metabolites have important effects on vascular smooth muscle and ion transport in kidney tubule cells (34–36). Although none of the cytochromes P450 on our present map are known to have this function, the possibility certainly exists that unknown cytochrome(s) P450 with this function exist as part of this gene cluster. The Na,K-ATPase $\alpha 3$ isoform (*ATPIA3*) is also a candidate locus very close to the cytochrome P450 gene cluster (Fig. 1). The cluster of kallikrein-like proteinases (marked by the *KAL* locus) is also at the edge of this QTL region (compare Figs. 1 and 2).

The effect of the blood pressure QTL marked by the cytochromes P450 cluster on chromosome 1 was highly population dependent, being seen only in the F₂ population derived from S and LEW strains, but not in F₂ populations derived from S and other strains (Table V). This was true also of the SA locus on chromosome 1 (Table IV). Cross specificity has invariably been the case with other QTL we have described on rat chromosomes 2 (30), 3 (25, 37) 5 (25), 10 (31, 38), 13 (14), and 17 (25). We attribute this to the fact that different alleles at each QTL can obviously be segregating in each cross depending on the strain crossed with S and to effects of genetic background also dependent on the strain crossed with S. The theoretical basis of this phenomenon is discussed in detail in a recent review (4).

In the F₂(LEW \times S) population there was a strong correlation between heart weight and blood pressure. Since left ventricular hypertrophy is well known to occur with chronically increased blood pressure, the heart weight data are taken to corroborate and validate the blood pressure measurements.

The F₂(LEW \times S) population is particularly favorable since we know of four QTL marked by the SA, D1Mco1, *GLUTB*, and *HITH* loci segregating in this population. These loci collectively accounted for 106 mmHg of blood pressure and 506 mg of heart wt, which was 38 and 34% of the total phenotypic variances, respectively. We do not have a measure of genetic variance for the F₂(LEW \times S) population, so we cannot state the percentage of the genetic variance accounted for by the four known QTL. It is, however, obvious that a substantial part of the genetic variance is accounted for by these four loci.

Screening segregating populations for QTL is clearly only a first step in establishing the existence of a QTL in a chromosomal region. Because of the imprecision with which a QTL can be detected and localized it will be necessary to create congenic strains by substituting, for example, putative QTL chromosomal regions lowering blood pressure into the background of the S rats (4) and to demonstrate that such a strain actually has reduced blood pressure compared to S. In the case of chromosome 1, the two QTL proposed here should be tested by the construction of two such congenic strains, each containing nonoverlapping Lewis chromosome 1 regions, where each congenic strain is predicted to have lower blood pressure than the S strain.

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