Genomic scan for maximal oxygen uptake and its response to training in the HERITAGE Family Study

CLAUDE BOUCHARD,1 TUOMO RANKINEN,1 YVON C. CHAGNON,2 TREVA RICE,3 LOUIS PÉRUSSE,2 JACQUES GAGNON,2 INGRID BORECKI,3 PING AN,3 ARTHUR S. LEON,4 JAMES S. SKINNER,5 JACK H. WILMORE,6 MICHAEL PROVINCE,3 AND D. C. RAO3,7

Genomic scan for maximal oxygen uptake and its response to training in the HERITAGE Family Study. J. Appl. Physiol. 88: 551–559, 2000.—This study aimed to identify human genomic regions that are linked to maximal oxygen uptake (VO2max) in sedentary individuals or to the responsiveness of VO2max to a standardized endurance training program. The results of a genomic scan based on 289 polymorphic markers covering all 22 pairs of autosomes performed on the Caucasian families of the HERITAGE Family Study are presented. The mean spacing of the markers was 11 cM, and a total of 99 families and 415 pairs of siblings were available for the study. VO2max in the sedentary state was adjusted for the effects of age, sex, body mass, fat mass, and fat-free mass, whereas the VO2max response was adjusted for age and baseline level of the phenotype. Two analytic strategies were used: a single-point linkage procedure using all available pairs of siblings (SIBPAL) and a multipoint variance components approach using all the family data (SEGPATH). Results indicate that linkages at P values of 0.01 and better are observed with markers on 4q, 8q, 11p, and 14q for VO2max before training and with markers on 1p, 2p, 4q, 6p, and 11p for the change in VO2max in response to a 20-wk standardized endurance training program. These chromosomal regions harbor many genes that may qualify as candidate genes for these quantitative traits. They should be investigated in this and other cohorts.

MAXIMAL OXYGEN uptake (VO2max) varies considerably among sedentary adults. In a recent report (3), we have shown that there is significant familial aggregation for VO2max in the sedentary state even when the data are adjusted for age, sex, body mass, and body composition. These observations were derived from the HERITAGE Family Study, and they indicate that the heritability of VO2max among sedentary adults after adjustment for the above covariates could be as high as 50% of the total phenotypic variance, although this value is undoubtedly inflated by nongenetic familial factors.

Twin and other family studies have been published in the peer-reviewed literature on this topic over the past three decades (6, 12, 17–21, 23, 31). Despite the fact that these studies vary considerably in sample size, phenotype measurements, data adjustment procedures, and analytic strategies, they generally concluded that there is a significant genetic component to VO2max in the untrained state. Moreover, there is evidence that the trainability of VO2max is characterized by a significant level of familial aggregation. For instance, members of the same pairs of identical twins are significantly more alike than unrelated individuals in the VO2max increase following exposure to a standardized training program. This was confirmed by the results of three different experimental studies with identical twins (16, 26, 29). The findings of these studies are remarkably concordant: the intraclass correlations for the intrapair resemblance in the VO2max changes with training range from 0.65 to 0.77. The F ratios of the between-pair variance in VO2max gain to the within-pair variance are quite similar, with a range from 6 to 9 (4). Such results have been further supported by the findings from the HERITAGE Family Study (1). The VO2max response to the standardized training regimen of the HERITAGE Family Study exhibited familial aggregation, with some families characterized by a high-trainability pattern and others by low responsiveness. The maximal heritability of VO2max trainability among the HERITAGE families of Caucasians reached 47% (1).

Ultimately, the nature of the genetic effects on human variation in VO2max in the untrained state and in response to training will have to be resolved at the gene and molecular level. A handful of studies have consid-
ered the role of a few candidate genes on $\text{VO}_{2\text{max}}$ in sedentary adults or $\text{VO}_{2\text{max}}$ responsiveness to training (7) with mixed results. Another approach is to focus on the detection of quantitative trait loci (QTL) affecting these phenotypes. In a preliminary paper based on this strategy, we have reported the results of a first screen of chromosome 22 using seven polymorphic markers (13). No evidence for a QTL affecting $\text{VO}_{2\text{max}}$ in the sedentary state or the changes in $\text{VO}_{2\text{max}}$ with training was uncovered.

Here, we report the results of a genomic scan, based on 289 markers covering all autosomes, performed on the Caucasian families of the HERITAGE Family Study with a view to identify the genomic regions encoding genes contributing to human heterogeneity in $\text{VO}_{2\text{max}}$ in the untrained state and its trainability.

**MATERIALS AND METHODS**

**Sample**

The HERITAGE Family Study was designed to investigate the role of the genotype in cardiovascular, metabolic, and hormonal responses to aerobic exercise training and the contribution of regular exercise to changes in selected cardiovascular disease and diabetes risk factors. Five centers located at Indiana University, Pennington Biomedical Research Center, University of Minnesota, Texas A&M University, and Washington University are presently involved in the HERITAGE Family Study consortium. The study design, sample, and protocol have been described earlier (5).

A total of 481 individuals from 99 two-generation families of Caucasian descent (236 men, 245 women) were available for this study. The following criteria were applied to screen subjects for participation. First, individuals were required to be between the ages of 17 and 65 yr (17–40 yr of age for offspring and ≤65 yr of age for parents). Second, all participants were required to be sedentary at baseline. Third, individuals with a body mass index >40 kg/m² were excluded unless they were able to meet the demands of the exercise tests and exercise training program. Fourth, resting blood pressure levels could not exceed 159 mmHg for systolic and 99 mmHg for diastolic. Antihypertensive drug therapy was also a cause for exclusion. Participants were required to be in good physical health and able to complete the 20-wk exercise program. Further details about the study, including inclusion and exclusion criteria, can be found in Bouchard et al. (5).

Informed written consent was obtained from all subjects. The study was approved by the Institutional Review Board of each participating institution.

**Exercise Training Program**

The training program was conducted on cycle ergometers (Universal Aerobicycle, Cedar Rapids, IA) interfaced with a Mednet computer system (Universal Gym Mednet, Cedar Rapids, IA) to control the power output of the ergometers so that constant training heart rates could be maintained. Subjects started training at a heart rate associated with 55% of their initial $\text{VO}_{2\text{max}}$, after which constant training heart rates could be maintained. Subjects started training at a heart rate associated with 55% of the initial $\text{VO}_{2\text{max}}$ for 30 min/day and gradually progressed to 75% of the heart rate of the initial $\text{VO}_{2\text{max}}$ for 50 min/day at the end of 14 wk. They maintained this intensity and duration throughout the remaining 6 wk. Frequency was maintained at three sessions per week throughout the 20-wk training program. The power output of the cycle ergometer was adjusted automatically to the heart rate response of the subject at all times during all training sessions. All training sessions were supervised on site. A detailed description of the training program can be found elsewhere (5, 30).

**Maximal Oxygen Uptake Measurements**

Each individual was examined with a battery of measurements before and after the 20-wk standardized exercise program. Two maximal exercise tests designed to lead to $\text{VO}_{2\text{max}}$ on a cycle ergometer were performed on 2 separate days at baseline and again on 2 separate days after training on a SensorMedics 800S (Yorba Linda, CA) cycle ergometer connected to a SensorMedics 2900 metabolic measurement cart. The tests were conducted at about the same time of day, with at least 48 h between the two tests. An electrocardiogram was used to monitor heart rate. Gas exchange variables [O₂ consumption (VO₂), CO₂ production (VCO₂), minute ventilation, and respiratory exchange ratio (RER)] were recorded as a rolling average of three 20-s intervals. The criteria for $\text{VO}_{2\text{max}}$ were RER >1.1, plateau in VO₂ (change of <100 ml/min in the last three 20-s intervals), and a heart rate within 10 beats/min of the maximal heart rate predicted for that age. All subjects achieved a $\text{VO}_{2\text{max}}$ by at least one of these criteria in at least one of the two tests, both pre- and posttraining. In the first test, subjects exercised at a power output of 50 W for 3 min, followed by increases of 25 W each subsequent minute until volitional exhaustion. For older, smaller, or less fit individuals, who were generally the older mothers among the family members, the test was started at 40 W, with increases of 10–20 W each every 2 min thereafter. In the second test, subjects exercised for ~10 min at an absolute (50 W) and then for ~10 min at a relative power output equivalent to 60% $\text{VO}_{2\text{max}}$. They then exercised for 3 min at a relative power output that was 80% of their $\text{VO}_{2\text{max}}$ after which resistance was increased to the highest power output attained in the first maximal test. If subjects were able to pedal after 2 min, power output was increased every 2 min thereafter until they reached volitional fatigue.

If both values were within 5% of each other, the average $\text{VO}_{2\text{max}}$ from these two tests was taken as the $\text{VO}_{2\text{max}}$ for that subject and used in this analysis. If they differed by more than 5%, the higher $\text{VO}_{2\text{max}}$ value was used. The reproducibility of $\text{VO}_{2\text{max}}$ in these subjects was examined and was characterized by an intraclass correlation coefficient of 0.97 for repeated tests with a coefficient of variation of 5% and no difference among clinical centers (3, 30). The $\text{VO}_{2\text{max}}$ response was defined as the difference (ml O₂/min) between posttraining $\text{VO}_{2\text{max}}$ and baseline $\text{VO}_{2\text{max}}$ (i.e., $\text{VO}_{2\text{max}}$ response = posttraining $\text{VO}_{2\text{max}}$ minus baseline $\text{VO}_{2\text{max}}$).

**Molecular Studies**

Genomic DNA was prepared from permanent lymphoblastoid cells by the proteinase K and phenol/chloroform technique. DNA was dialyzed four times against 10 mM Tris-1 mM EDTA (pH 8.0) buffer for 6 h at 4°C, and ethanol was precipitated.

Choice of markers. Microsatellite markers (di-, tri-, and tetranucleotide repeat) were selected from different sources but mainly from the Marshfield panel version 8a (http://www.marshmed.org/genetics) and the Location Database (LDB) map (January 1999 version) from Southampton, UK (http://cedar.genetics.soton.ac.uk). The LDB summary map was also used to look for candidate genes 10 cM on each side of the markers yielding the strongest results. The mean spacing for the 289 markers was 11 cM with a range from 0.2 to 51 cM.

PCR conditions. PCR conditions and genotyping have been described in detail elsewhere (9). Briefly, PCR reactions were conducted using 250 ng of genomic DNA, 0.1 pmol of the forward primer coupled to the infrared tag IRD800 or IRD700.
(LICOR), and 0.4 pmol of the reverse untagged primer (Research Genetics), 125 µM dNTPs, and 0.3 units Taq polymerase (Perkin-Elmer or Pharmacia) in PCR buffer (100 mM Tris-HCl, pH 8.3, 15 mM MgCl₂, 0.5 M KCl, and 0.01% gelatin) for a final volume of 10 µl. PCR cycles followed a two-step procedure: 1 cycle at 93°C for 5 min, 10 cycles at 93°C for 20 s and 57°C for 60 s, and 24 cycles at 93°C for 20 s and 52°C for 60 s. Annealing temperature was defined according to the melting temperature of the primers used and ranged between 52 and 67°C, with a difference of 5°C being maintained between the two annealing temperatures.

Marker analysis. Automatic infrared DNA sequencers from LICOR were used to detect the PCR products. One microliter of each of the 52 samples, including up to five different markers, spaced with standards at each of the four samples, was applied to the gel. Standards were produced by PCR from a PUC19 plasmid using a tagged M13 forward primer (LICOR) and corresponding untagged reverse reverse primer for each standard. The 18-cm long gel was run for 1.5–2 h. At the end of the run, an electronic image of the gel was produced and used for genotyping.

Genotyping. Automatic genotyping was performed using the computer software SAGA (Rick M ndoe, Roger Bumgarner, and Russ Welti, University of Washington, Seattle, WA; LICOR). In this process, sample and standard lanes were automatically found, the different markers were located on the gel, bands for each sample were identified, and genotyping was done simultaneously for a given marker on all the subjects of the study. After a manual edition of the typing, results were exported directly to a local dBase IV database (GENEMARK) in which a procedure had been developed to check for misinheritance (incompatibilities among the genotypes) within nuclear families and extended pedigrees. Subjects with incompatibilities were excluded from the database by GENEMARK and retyped completely for the incompatible markers, i.e., from the PCR reaction to the genotyping. Less than 5% of the subjects had to be retyped (mainly because of insufficient DNA amplification).

Data Adjustment

Two VO₂max phenotypes are considered. VO₂max at baseline was analyzed after adjustment for the effects of age (up to a cubic polynomial), sex, body mass, fat mass, and fat-free mass using stepwise multiple regression procedures, separately in each of the four sex-by-generation groups, and retaining only those terms that were significant at the 5% level. Fat mass and fat-free mass were estimated from an underwater weighing assessment of body density as recently described (33). The resulting squared residuals were similarly regressed on another polynomial in age (to adjust for heterocedasticity). The phenotype used in the genetic analysis was thus adjusted for relevant covariate effects in both the mean and variance. The phenotype was standardized to a mean of zero and a standard deviation of one. The percentage of variance accounted for by these covariates was quite large, ranging from 42.2% to 57.6%.

The response of VO₂max to training was adjusted for age and baseline VO₂max using a stepwise multiple regression procedure. Briefly, the response variable was regressed on up to a cubic polynomial in age plus baseline value within four sex-by-generation groups (fathers, mothers, sons, and daughters). Only terms significant at the 5% level were retained. The resulting squared residuals were similarly adjusted for effects on the variance; the final adjusted phenotype was standardized to a mean of zero and a standard deviation of one. Significant terms (percentage of variance) in each of the sex-by-generation groups for the VO₂max response phenotype were seen only in sons (9.2%) and daughters (5.0%). SAS (version 6.08) for PC was used for the regression analysis.

Statistical Analysis

First, single-point linkage analysis using all available sibling pairs was performed on these two phenotypes (SIBPAL; SAGE version 3.0). Second, analyses were carried out using the multipoint variance components analytic strategy implemented in SEGPATH (25). This is an extension of a path model for analysis of family resemblance (24) in which correlations among family members are modeled as functions of the allele sharing at a marker locus and a residual component including genetic as well as shared environmental effects. In addition to the trait locus heritability ($h^2$) and the residual heritability ($h^r^2$), the spouse correlation and excess sibling resemblance beyond that predicted under the genetic model were estimated by maximum likelihood. Linkage was tested by the likelihood ratio test of the null hypothesis $h^2 = 0$. Multipoint estimation of identical by descent was used in the SEGPATH analysis, whereas a single-point approach was implemented in SIBPAL. The number of families available for the SEGPATH analysis was 99. The SIBPAL linkage analyses were undertaken with 415 pairs of siblings at baseline and 327 for the response to training. Suggestive evidence of linkage was defined as $P < 0.01$, whereas weaker but potentially useful evidence of linkage was identified as $0.01 < P < 0.05$.

RESULTS

General trends in the linkage results are depicted in Figs. 1 and 2 for the 22 autosomes. In Fig. 1, the evidence for linkage (P values) from SEGPATH and SIBPAL is illustrated for the baseline VO₂max (adjusted for age, body weight, fat mass, and fat-free mass within each sex). Suggestive evidence of linkage can be seen for chromosomes 4, 8, 11, and 14 ($P < 0.01$). The training response (adjusted for age and baseline level by sex) results are shown in Fig. 2. Again, suggestive evidence of linkage is observed for chromosomes 1, 2, 4, 6, and 11.

The details of all suggestive linkages for the pretraining VO₂max are summarized in Table 1. The region with the highest significance level with SEGPATH is located on the long arm of chromosome 4 (4q12) at map position 61.658 cM. Three additional regions for baseline VO₂max were identified with SIBPAL. These regions are located on chromosome 14q21.3 at map position 49.255 cM, chromosome 8q24.12 (128.157 cM), and chromosome 11p15.1 (21.174 cM). Moreover, 11 regions on seven chromosomes (chromosomes 1, 3, 4, 7, 8, 11, 13) with SEGPATH and 10 markers on six chromosomes (chromosomes 1, 2, 3, 4, 7, 22) with SIBPAL showed weaker but possibly useful evidence of linkage ($0.01 < P < 0.05$) with VO₂max in the sedentary state (Table 2).

Multipoint analyses (SEGPATH) revealed two regions and single-point analyses (SIBPAL) revealed three regions with suggestive evidence of linkage with the VO₂max training response. The multipoint signals are located on chromosomes 4 (4q26, 127.793 cM) and 6 (6p21.33, 28.788 cM), whereas the single-point linkages are seen with markers on chromosomes 11 (11p14.1, 31.272 cM), 1 (1p11.2, 125.032 cM), and 2 (2p16.1, 62.602 cM). In addition, weaker but potentially useful linkages ($0.01 < P < 0.05$) were found with SEGPATH
for regions located on chromosomes 3, 6, 9, and 13 plus for 15 markers on chromosomes 1, 3, 5, 6, 7, 11, 13, 15, 19, and 21 with SIBPAL (Table 3).

DISCUSSION

From the quantitative genetic evidence gathered in twin and family studies, it is justified to search for QTLs and, eventually, genes and mutations pertaining to VO2max in the sedentary state and its responsiveness to training. The genetic dissection of these two phenotypes and their determinants will require a wide array of designs and technologies. At this time, only one linkage study has been reported in humans, but it has dealt with a limited number of markers on one chromosome only (13). Other studies have focused on candidate genes encoded in the nuclear DNA and on a few mitochondrial DNA sequences.

In the same population, we have earlier reported a maximal heritability of ~50% for the baseline VO2max data (3). On the other hand, the maximal heritability estimate of the VO2max response to training reached 47% for the same sample of families with 2.5 times more variance between families than within families. Thus the familial factors underlying VO2max in sedentary families are quantitatively similar to those underlying its response to exercise training. However, although they are quantitatively about the same, the familial and genetic factors underlying the two phenotypes appear to be different, as indicated by the noncomparability of the linkage results for baseline VO2max and VO2max response. From the earlier observations on identical twins and nuclear families plus the recent data on the HERITAGE Family Study cohort, we previously concluded that VO2max in the sedentary state and its trainability are highly familial with a significant genetic component (1, 3). It should therefore be possible to identify the genes and mutations responsible for the variability in both phenotypes. However, it must be recognized at the outset that it is unlikely that a single gene or very few loci will be sufficient to define the genetic component of VO2max and trainability.

A few reports on the topic of genetic markers and performance have appeared over the past 30 years or so. The study of elite performance by means of genetic markers on one chromosome only (13). Other studies have focused on candidate genes encoded in the nuclear DNA and on a few mitochondrial DNA sequences.

Since then, a handful of papers have dealt with the potential contribution of a few candidate genes. For instance, three studies have reported an association between the insertion allele of the angiotensin-converting enzyme (ACE) insertion/deletion (I/D) polymorphism and indicators of performance (14, 15, 22). The ACE locus is encoded on 17q23. In the present study, no evidence of linkage was observed on chromosome 17 for either baseline VO2max or its responsiveness to training. Therefore, it is not yet clear whether the ACE I/D polymorphism is truly of importance for physical performance in general or endurance performance in particular.

Skeletal muscle-specific creatine kinase (CKMM) is a legitimate candidate gene (19q13.22) to investigate in relation to endurance performance. Because CKMM activity level is two times greater in type II (fast-twitch) than in type I fibers (34), a low CKMM activity level is typical of the skeletal muscle of endurance athletes. An early study indicated that a CKMM protein charge variant was weakly associated with the ability to perform a 90-min endurance test (2). In addition, research on transgenic mice indicates that a low CKMM activity is associated with improved skeletal muscle resistance to fatigue (32). More recently, a sibling pair linkage study has shown a weak genetic linkage between the CKMM locus and changes in VO2max (age, sex, and pretraining VO2max adjusted) in the HERITAGE Family Study (28). This linkage was suggested again in the present study with a P value of 0.026 (Table 3). Moreover, we have reported a significant association between the CKMM genotype and the VO2max response to 20 wk of endurance training in both parents and adult offspring of the HERITAGE Family Study (27). One-third of all homozygotes for the less-frequent allele (CKMM Nco I polymorphism in the 3' untranslated region) were observed in the low-responder group (lowest decile of response), whereas this genotype was not seen in any high responders (upper decile of response). The CKMM genotype accounted for ~10% of the variance of VO2max response.

The chromosomal regions showing suggestive linkages with VO2max in the sedentary state or in response to endurance training in the present paper encode several potential candidate genes. The marker D4S3248 on chromosome 4, which was the strongest signal with the baseline VO2max in multipoint analyses, is located close (0.2 cM) to the β-sarcoglycan gene. β-Sarcoglycan is part of the dystrophin-glycoprotein complex, which acts as a structural link between the cytoskeleton of muscle and the extracellular matrix. It is thought to confer stability to the sarcolemma and prevent muscle cells from contraction-induced damage. A gene of another dystrophin-associated protein, syntrophin β-1, located 5.9 cM from the D8S592 marker, showed suggestive single-point linkage evidence with the baseline VO2max. In addition, the genes encoding...
Training response results (p values)

Ch 1

Ch 2

Ch 3

Ch 4

Ch 5

Ch 6

Ch 7

Ch 8

Ch 9

Ch 10

Ch 11

Ch 12

Ch 13

Ch 14

Ch 15

Ch 16

Ch 17

Ch 18

Ch 19

Ch 20

Ch 21

Ch 22

Delta \( \dot{V}_O_2 \) max, age, baseline from SEGPATH

Delta \( \dot{V}_O_2 \) max, age, baseline from SIBPAL
Table 1. Summary of the suggestive linkages (P < 0.01) with VO$_2$max in the sedentary state and in response to endurance training in the HERITAGE Family Study

<table>
<thead>
<tr>
<th>P Value</th>
<th>Chromosome</th>
<th>Marker</th>
<th>Map Position, cM</th>
</tr>
</thead>
<tbody>
<tr>
<td>SEGPATH</td>
<td>3q12</td>
<td>FABP2</td>
<td>127.793</td>
</tr>
<tr>
<td>SIBPAL</td>
<td>6p21.33</td>
<td>D6S2439</td>
<td>28.788</td>
</tr>
</tbody>
</table>

VO$_2$max, maximal oxygen uptake; cM, centimorgans.

γ-sarcoglycan (13q12.11), dystrophin-associated glycoprotein 1 (3p21.31), and lamin A/C (1q21.2) are located within 2.3 to 6.2 cM from markers, showing weak but potentially useful linkages with VO$_2$max in the sedentary state. The marker (D14S587) showing the strongest evidence of linkage with the baseline VO$_2$max in SIBPAL is located in the vicinity of the liver glycogen phosphorylase (0.6 cM) and the GTP cyclohydrolase I (1.8 cM) genes. GTP cyclohydrolase I is a rate-limiting enzyme in the synthesis of tetrahydrobiopterin, an essential cofactor of nitric oxide synthase.

A restriction fragment length polymorphism marker on chromosome 11p15.1 within the sulfonylurea receptor (SUR) gene exhibited one of the most significant linkages with VO$_2$max in SIBPAL showing the strongest evidence of linkage with the baseline VO$_2$max in SIBPAL. This marker is located in the vicinity of the liver glycogen phosphorylase (0.6 cM) and the GTP cyclohydrolase I (1.8 cM) genes. GTP cyclohydrolase I is a rate-limiting enzyme in the synthesis of tetrahydrobiopterin, an essential cofactor of nitric oxide synthase.

Potential candidate genes located close to markers with suggestive linkages to VO$_2$max training response include the voltage-gated potassium channel gene (KCNA4) on chromosome 11p14.1 (0.1 cM from the marker), the pancreatic colipase (CLPS, 0.8 cM) gene and the hemochromatosis locus (HFE, 1.3 cM) on chromosome 6p21.33, the fatty-acid binding protein 2 (FABP2, 0.0 cM) gene and the long QT syndrome 4 locus (LQT4, 4.0 cM) on chromosome 4q26, the calmodulin 2 (CALM2, 0.5 cM) and the calcineurin B (PPP3R1, 3.4 cM) genes on chromosome 2p16.1, and the 3-β-hydroxy-

Table 2. Summary of the potentially useful linkages (0.01 < P < 0.05) with VO$_2$max in the sedentary state in the HERITAGE Family Study

<table>
<thead>
<tr>
<th>P Value</th>
<th>Chromosome</th>
<th>Marker</th>
<th>Map Position, cM</th>
</tr>
</thead>
<tbody>
<tr>
<td>SEGPATH</td>
<td>4p14</td>
<td>D4S2397</td>
<td>32.246</td>
</tr>
<tr>
<td>0.014</td>
<td>7p11.2</td>
<td>IGFBP1</td>
<td>60.720</td>
</tr>
<tr>
<td>0.047</td>
<td>7p21.3</td>
<td>D7S513</td>
<td>16.740</td>
</tr>
<tr>
<td>0.043</td>
<td>8q24.12</td>
<td>D8S592</td>
<td>128.157</td>
</tr>
<tr>
<td>0.038</td>
<td>11q14.1</td>
<td>UCP2</td>
<td>91.588</td>
</tr>
<tr>
<td>0.033</td>
<td>13q12.11</td>
<td>D13S787</td>
<td>19.370</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>P Value</th>
<th>Chromosome</th>
<th>Marker</th>
<th>Map Position, cM</th>
</tr>
</thead>
<tbody>
<tr>
<td>SEGPATH</td>
<td>1p11.2</td>
<td>D1S534</td>
<td>125.032</td>
</tr>
<tr>
<td>0.025</td>
<td>1q21.2</td>
<td>S100A1</td>
<td>154.041</td>
</tr>
<tr>
<td>0.033</td>
<td>1q24.1</td>
<td>ATP1B1</td>
<td>181.137</td>
</tr>
<tr>
<td>0.021</td>
<td>3p21.3</td>
<td>D3S1766</td>
<td>66.879</td>
</tr>
<tr>
<td>0.048</td>
<td>3p21.31</td>
<td>D3S1447</td>
<td>58.763</td>
</tr>
<tr>
<td>0.028</td>
<td>4p14</td>
<td>D4S2397</td>
<td>32.246</td>
</tr>
<tr>
<td>0.014</td>
<td>7p11.2</td>
<td>IGFBP1</td>
<td>60.720</td>
</tr>
<tr>
<td>0.047</td>
<td>7p21.3</td>
<td>D7S513</td>
<td>16.740</td>
</tr>
<tr>
<td>0.043</td>
<td>8q24.12</td>
<td>D8S592</td>
<td>128.157</td>
</tr>
<tr>
<td>0.038</td>
<td>11q14.1</td>
<td>UCP2</td>
<td>91.588</td>
</tr>
<tr>
<td>0.033</td>
<td>13q12.11</td>
<td>D13S787</td>
<td>19.370</td>
</tr>
<tr>
<td>SIBPAL</td>
<td>0.016</td>
<td>1p11.2</td>
<td>D1S534</td>
</tr>
<tr>
<td>0.039</td>
<td>1q24.1</td>
<td>ATP1A2</td>
<td>181.720</td>
</tr>
<tr>
<td>0.035</td>
<td>1q24.2</td>
<td>D1S1677</td>
<td>182.512</td>
</tr>
<tr>
<td>0.039</td>
<td>2q37.3</td>
<td>D2S2968</td>
<td>253.130</td>
</tr>
<tr>
<td>0.027</td>
<td>3p21.1</td>
<td>D3S1768</td>
<td>39.702</td>
</tr>
<tr>
<td>0.028</td>
<td>4p14</td>
<td>D4S2397</td>
<td>61.658</td>
</tr>
<tr>
<td>0.026</td>
<td>7p14.3</td>
<td>D7S2541</td>
<td>35.660</td>
</tr>
<tr>
<td>0.049</td>
<td>7p21.3</td>
<td>D7S513</td>
<td>16.740</td>
</tr>
<tr>
<td>0.019</td>
<td>7q36.1</td>
<td>D7S2195</td>
<td>158.000</td>
</tr>
<tr>
<td>0.012</td>
<td>22q13.2</td>
<td>D22S274</td>
<td>45.091</td>
</tr>
</tbody>
</table>

Potential candidate genes located close to markers with suggestive linkages to VO$_2$max training response include the voltage-gated potassium channel gene (KCNA4) on chromosome 11p14.1 (0.1 cM from the marker), the pancreatic colipase (CLPS, 0.8 cM) gene and the hemochromatosis locus (HFE, 1.3 cM) on chromosome 6p21.33, the fatty-acid binding protein 2 (FABP2, 0.0 cM) gene and the long QT syndrome 4 locus (LQT4, 4.0 cM) on chromosome 4q26, the calmodulin 2 (CALM2, 0.5 cM) and the calcineurin B (PPP3R1, 3.4 cM) genes on chromosome 2p16.1, and the 3-β-hydroxy-

Fig. 2. Overview of the linkage results for the 22 autosomes based on SEGPATH and SIBPAL for the VO$_2$max response to a standardized training program. Map distances in cM are shown on the abscissa, and P values for the strength of the linkage relationships are shown on the ordinates.
yesteroid dehydrogenase (HSD3B1, 0.1 cM) and the cardiac muscle calsequestrin (CASQ2, 5.5 cM) genes on chromosome 1p11.2. These genes are involved in cardiac contractility (KCNA4, LQT4), long-chain fatty acid absorption (CLPS, FABP2), calcium homeostasis and signaling in skeletal and cardiac muscle (CALM2, PPP3R1, CASQ2), and steroid hormone synthesis (HSD3B1). Although there is no direct evidence to support the notion that they are involved in human trainability, one could hypothesize that they contribute to interindividual variation in training response.

There was not a high degree of concordance between the single point and multipoint linkage results in the present study. The differences in the findings are likely explained by the relatively small sample sizes available for such studies, the marker spacing, which is relatively large in some chromosomal areas, and perhaps misspecification of the location of markers on the genetic map. The latter two factors could have had influences on the multipoint linkage results, particularly if errors of location exist in the current map, but would not have affected the single-point findings. Moreover, it should be recognized that, even though a panel of 289 markers is reasonably large for a first genomic screen, a more dense and accurate map will likely yield more and perhaps stronger signals in the future.

In summary, multipoint and single point linkage studies performed on the Caucasian families of the HERITAGE Family Study have revealed some indications of linkages with $V_{O2max}$ in the sedentary state and with its responsiveness to training. None of the linkages is very strong. However, several linkages were found at the 0.01 level and better with markers on 4q, 8q, 11p, and 14q for $V_{O2max}$ before training and with markers on 1p, 2p, 4q, 6p, and 11p for the gain in $V_{O2max}$ in response to 20 wk of a standardized endurance training program. These chromosomal regions encode several candidate genes that should be investigated further.

Thanks are expressed to all the coprincipal investigators, investigators, coinvestigators, local project coordinators, research assistants, laboratory technicians, and secretaries who have contributed to the study and to Diane Drolet, MSc, for assistance with the preparation of this manuscript.

The HERITAGE Family Study is supported by the National Heart, Lung, and Blood Institute through the following grants: HL-45670 (C. Bouchard, principal investigator), HL-47323 (A. S. Leon, principal investigator), HL-47317 (D. C. Rao, principal investigator), HL-47327 (J. S. Skinner, principal investigator), and HL-47322 (J. H. Wilmore, principal investigator). A. S. Leon is partially supported by the Henry L. Taylor Professorship in Exercise Science and Health Enancement.

Address for reprint requests and other correspondence: C. Bouchard, Pennington Biomedical Research Center, Louisiana State Univ., 6400 Perkins Rd., Baton Rouge, LA 70808-4124 (E-mail: bouchac@pbr.edu).

Received 31 August 1999; accepted in final form 29 October 1999.

REFERENCES


