Effect of chromosome substitution on intrinsic exercise capacity in mice [version 2; peer review: 2 approved]

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Abstract
Previous research identified a locus on Chromosome 14 as an important regulator of endurance exercise capacity in mice. The aim of this study was to investigate the effect of chromosome substitution on intrinsic exercise capacity and identify quantitative trait loci (QTL) associated with exercise capacity in mice. Mice from a chromosome substitution strain (CSS) derived from A/J and C57Bl/6J (B6), denoted as B6.A14, were used to assess the contribution of Chromosome 14 to intrinsic exercise capacity. All mice performed a graded exercise test to exhaustion to determine exercise capacity expressed as time (min) or work (kg·m). Exercise time and work were significantly greater in B6 mice than B6.A14 and A/J mice, indicating the presence of a QTL on Chromosome 14 for exercise capacity. To localize exercise-related QTL, 155 B6.A14 x B6 F₂ mice were generated for linkage analysis. Suggestive QTL for exercise time (57 cM, 1.75 LOD) and work (57 cM, 2.08 LOD) were identified in the entire B6.A14 x B6 F₂ cohort. To identify putative sex-specific QTL, male and female F₂ cohorts were analyzed separately. In males, a significant QTL for exercise time (55 cM, 2.28 LOD) and a suggestive QTL for work (55 cM, 2.19 LOD) were identified. In the female cohort, no QTL was identified for time, but a suggestive QTL for work was located at 16 cM (1.8 LOD). These data suggest that one or more QTL on Chromosome 14 regulate exercise capacity. The putative sex-specific QTL further suggest that the genetic architecture underlying exercise capacity is different in males and females. Overall, the results of this study support the use of CSS as a model for the genetic analysis of exercise capacity. Future studies should incorporate the full panel of CSS using male and female mice to dissect the genetic basis for differences in exercise capacity.

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Introduction
Cardiorespiratory fitness measured during a graded exercise test is inversely related to the relative risk of cardiovascular disease\(^1,2\). Results from human cross-sectional, twin, and prospective studies indicate that genetic factors account for 25–65% of the variation in exercise capacity\(^3,4\). Because having higher levels of exercise capacity has been shown to be beneficial for reducing the onset of cardiovascular disease, the physiological factors determining exercise capacity have been widely studied\(^5\). However, the genetic contribution to exercise capacity is not completely understood. Presently, several candidate genes contributing to improved exercise capacity have been proposed based on genome wide studies\(^6,7\), but these genes account for only a small portion of the variability in exercise capacity or training responses\(^8\).

Several studies have investigated the genetic factors contributing to exercise capacity using inbred rodent models\(^9,10\). One common approach has been to screen multiple rodent strains for exercise capacity, followed by quantitative trait loci (QTL) analyses to identify loci linked to exercise capacity. This approach has been used to identify QTL for exercise capacity in rats\(^11\) and mice\(^12,13\). Research from our laboratory previously identified significant and suggestive QTL on several chromosomes that may house candidate genes that influence variation in exercise capacity\(^12,13\). These identified regions overlap with other mouse and human QTL, suggesting that these regions and/or genes are conserved among species\(^12\). Mouse Chromosome 14 (Chr 14), for example, contained a significant QTL for intrinsic (pre-training) exercise capacity, a significant QTL for exercise capacity after training, and a suggestive QTL for the change in exercise capacity in response to exercise training\(^13\). Several linkage markers for maximal oxygen consumption in the sedentary state in humans map to these exercise-related QTL on mouse Chr 14\(^13,15\). Therefore, the present study focused on characterizing the role of Chr 14 in regulating intrinsic exercise capacity.

In the current study we employed a relatively new mouse model, chromosome substitution strains (CSS) to assess the contribution of individual chromosomes to endurance exercise capacity\(^16,17\). CSS mice are made by substituting a single chromosome from a donor inbred strain on the genetic background of a host inbred strain (recipient). Therefore phenotypic differences between the recipient or background strain mice and CSS mice support the presence of a QTL on the substituted chromosome for the phenotype being measured. Results from a previous study identified the AJ strain as having low exercise capacity in comparison to the C57BL/6J (B6) strain\(^14\). Therefore we chose to use CSS mice based on A/J and B6 inbred strains.

Utilizing this CSS model, the main purposes of the present study were to investigate the effect of chromosome substitution on intrinsic exercise capacity and to identify QTL regulating intrinsic exercise capacity in mice. We hypothesized that chromosome substitution would significantly affect exercise capacity and therefore confirm the importance of Chr 14 in the genetic regulation of intrinsic exercise capacity in mice. Furthermore, we utilized linkage analysis to map QTL on Chr 14 in progeny from a cross between the CSS and host B6 strain.

Methods
Animals
All procedures adhered to the established National Institutes of Health guidelines for the care and use of laboratory animals and were approved by the Institutional Animal Care and Use Committee at Texas A&M University. Seven week-old inbred mice (A/J, C57BL/6J (B6), and Chr 14 substitution mice (C57BL/6J-Chr 14\(^{14}\)/NaJ, abbreviated B6.A14)) (n = 12/strain, 6 male and 6 female mice) were purchased from Jackson Laboratory (Bar Harbor, ME.). Upon arrival at Texas A&M, all mice were given one week to acclimate to their new environment before assessing exercise capacity. A separate group of male B6 mice were crossed with a separate group of female chromosome substitution B6.A14 mice to generate (B6.A14 × B6) F\(_1\) mice. The F\(_1\) mice were then intercrossed to produce 155 F\(_2\) generation mice (67 male and 88 female mice). All mice were housed in standard hanging polycarbonate cages (43 cm long × 21.5 cm wide × 15 cm high) with hardwood chip bedding and allowed food (Standardized Laboratory Rodent Diet) and water \textit{ad libitum}. Mice were housed 1–5 mice per cage depending on sex and lineage and maintained on a 12 hr light/dark schedule at an ambient temperature of 22–24°C.

Exercise performance test
At 8 weeks of age, all mice were familiarized for two days at 9.0 m/min and 10.0 m/min at 10° for 10 minutes to run on a motorized rodent treadmill (Columbus Instruments, Columbus, OH), with an electric grid (160 V, 0–2 mA) at the rear of the treadmill as described previously\(^13,14\). Each mouse then completed two graded exercise tests separated by 48 hrs. Mean values for each mouse were used for statistical analyses. For each performance test, the treadmill was started at 9.0 m/min at 0° grade for 9 minutes as a warm-up. The grade was then increased 5° every 9 minutes up to a final grade of 15° and speed was increased 2.5 m/min from a starting speed of 10 m/min every three minutes until exhaustion. Exercise continued until each mouse refused to run, defined as an inability to maintain running speed in spite of repeated contact with the electric grid\(^13,14\).

At exhaustion, each mouse was immediately removed from the treadmill and returned to its home cage. Exercise capacity was estimated for each animal using time (minutes) and work (kg·m). Work performed (kg·m) or vertical work was calculated as a product of body weight (kg) and vertical distance (meters), where vertical distance = (distance run)(\sinθ), where θ is equal to the angle of the treadmill from 0° to 15°\(^13,14\).

Genotyping
At least 24 hours after the last graded exercise test, all mice were anesthetized by intraperitoneal injection of a ketamine (80 mg/kg) - xylazine (5 mg/kg) cocktail. Mice were subsequently euthanized by exsanguination due to removal of the heart and aorta. Heart, gastrocnemius, plantaris, soleus muscle and liver tissue were excised from mice, washed in ice-cold (4°C) saline, blotted dry to remove
excess liquid, and snap frozen in liquid nitrogen. DNA was isolated from 25 mg of liver tissue with a DNeasy Blood and Tissue kit (Qiagen Science, Germantown, Maryland) according to the manufacturer’s instructions and quantified using NanoDrop spectrophotometry. Genotyping was performed using competitive allele-specific polymerase chain reaction (PCR) single nucleotide polymorphism (SNP) genotyping (KBiosciences, Hoddesdon, UK). All 155 F₂ mice were genotyped using 12 SNPs spaced at approximately 5 cM intervals.

**QTL identification**

QTL analyses were performed using R/qtl. One-dimensional scans were performed on the entire F₂ cohort with no additional covariates and with sex included as an additive and interactive covariate. Permutation tests (1,000 repetitions) were used to identify threshold values for logarithm of odds (LOD) scores for each condition (i.e., with or without covariates) and exercise phenotype. LOD scores were defined as significant if they surpassed the P < 0.05 threshold and suggestive if they surpassed the P < 0.63 threshold. If suggestive or significant QTL were identified using sex as interactive covariate, then one-dimensional scans were performed on male and female mice separately to identify potential sex-specific QTL. A two-dimensional scan also was performed on the entire F₂ cohort to identify additive or interacting QTL on Chr 14. QTL confidence intervals were determined using the 1.5 LOD support interval.

**Statistics**

All data are represented as mean ± SE. Statistical significance for phenotype comparisons was denoted by P < 0.05. Two-way analysis of variance was used to determine the effect of sex and strain on exercise capacity, which is defined as time (minutes), or work (kg·m) (JMP 9.0, SAS, Cary, NC). If significant main effects were found for strain, Dunnett’s post hoc test was used to determine significant strain differences compared with B6. If significant main effects were found for sex, t-tests were used to identify sex differences within each strain. Comparisons among parental strains and F₂ offspring and across genotypes for allelic effects were made using one way analysis of variance (strain or genotype) followed by Tukey’s post-hoc analysis. T-tests were used to identify sex differences in F₂ offspring. Linear regression was used to determine the contribution of body mass to exercise performance.

**Results**

*Inbred strains.* Exercise capacity, defined as mean run time during two graded exercise treadmill tests, for inbred and CSS mice is shown in Figure 1. Exercise times in A/J and B6.A14 mice were significantly less (P < 0.0001) than that in B6 mice. A significant effect of sex also was identified in all strains (A/J, B6.A14, and B6). For each strain, female mice ran significantly longer than male mice from the same strain (Figure 1A). When exercise capacity was expressed as work, A/J and B6.A14 strains were significantly different from B6 (P < 0.0001), with mice from both strains performing less work than B6 mice (Figure 1B). In contrast to exercise time, there was no significant main effect for sex (P = 0.1) on exercise capacity defined as work. Significant differences among the strains were primarily limited to differences in exercise phenotypes. Body mass was significantly less in B6.A14 mice compared with B6 (P < 0.0008) (Table 1). There were no significant differences in absolute tissue mass among A/J, B6, and B6.A14 strains (Table 1).

Within each strain, body mass was significantly lower in females compared to males. Accordingly, tissue masses were lower in female mice compared to male mice from the same strain (Table 1). For each strain there was a significant negative correlation between body mass and exercise time (B6, r = -0.71, P = 0.0096; A/J, r = -0.77, P = 0.0035; B6.A14, r = -0.86, P = 0.0003).

The sex-specific distributions for exercise time and work in F₂ mice are shown in Figure 2. Both time and work varied significantly between male and female F₂ mice (Table 2). On average, female mice ran approximately 2.5 min longer than male mice. These differences in run time between male and female mice were offset by a significantly higher body mass in male mice (P < 0.0001) resulting in comparable levels of work in male and female mice. Body mass was approximately 5 g higher in male mice compared with females (P < 0.0001), which likely accounts for the similar levels of work performed. Similar to body mass, tissue masses were significantly smaller in female F₂ mice compared to their male counterparts.
Table 1. Physical characteristics of female and male inbred and B6.A14 CSS mice.

|                  | A/J     | B6.A14  | C57BL/6J |
|------------------|---------|---------|----------|
|                  | female  | male    | female   | male    | female  | male    |
| Body mass, g     | 19.3 ± 0.2† | 22.7 ± 0.8 | 17.1 ± 0.0† | 20.7 ± 0.4* | 18.5 ± 0.6† | 22.8 ± 0.7 |
| Heart mass, mg   | 94.0 ± 3.5† | 105.8 ± 3.6 | 93.7 ± 2.0† | 116.0 ± 3.5 | 104.3 ± 2.8† | 122.0 ± 5.9 |
| HM:BM, mg/g      | 4.94 ± 0.17 | 4.63 ± 0.10 | 5.44 ± 0.12 | 5.55 ± 0.08* | 5.68 ± 0.21† | 5.03 ± 0.13 |
| Gastrocnemius mass, mg | 120.0 ± 4.2 | 124.0 ± 3.1 | 99.8 ± 4.6† | 130.0 ± 2.8 | 113.3 ± 3.4† | 151.8 ± 6.0 |
| GM:BM, mg/g      | 6.30 ± 0.21 | 5.44 ± 0.20 | 5.79 ± 0.27 | 6.23 ± 0.18 | 6.18 ± 0.29 | 6.27 ± 0.15 |
| Soleus mass, mg  | 7.7 ± 0.3 | 8.5 ± 0.5 | 8.0 ± 0.4 | 8.3 ± 0.3 | 7.5 ± 0.3† | 9.5 ± 0.2 |
| SM:BM, mg/g      | 0.40 ± 0.02 | 0.37 ± 0.02 | 0.46 ± 0.02† | 0.40 ± 0.01 | 0.41 ± 0.03 | 0.40 ± 0.01 |
| Plantaris mass, mg | 21.0 ± 1.1 | 22.7 ± 1.1 | 17.8 ± 0.3† | 21.0 ± 1.0 | 17.8 ± 0.4† | 20.3 ± 0.4 |
| PM:BM, mg/g      | 1.10 ± 0.05† | 0.99 ± 0.05* | 1.03 ± 0.02 | 1.01 ± 0.05* | 0.97 ± 0.04† | 0.84 ± 0.02 |

Values are mean ± SE. n = 6 per group, except n = 5 for B6 males. HM:BM, heart mass-to-body mass ratio; GM:BM, gastrocnemius mass-to-body mass ratio; SM:BM, soleus mass-to-body mass ratio; PM:BM, plantaris mass-to-body mass ratio.

*P < 0.05 significant main effect for strain compared with C57BL/6J.
†P < 0.05 significantly different from male mice of same strain.

(B6.A14 × B6) F₂ mice.

Table 2. Exercise capacity and physical characteristics of female and male (B6.A14 × B6) F₂ mice.

|                  | Female (n = 88) | Male (n = 67) |
|------------------|-----------------|---------------|
| Time, min        | 31.9 ± 0.1*     | 29.3 ± 0.1    |
| Work, kg·m       | 1.70 ± 0.02     | 1.68 ± 0.03   |
| Body mass, g     | 19.9 ± 0.2*     | 25.0 ± 0.2    |
| Heart mass, mg   | 110.8 ± 1.3*    | 129.3 ± 1.6   |
| HM:BM, mg/g      | 5.43 ± 0.04*    | 5.13 ± 0.05   |
| Gastrocnemius mass, mg | 127.7 ± 1.4*  | 155.8 ± 1.8   |
| GM:BM, mg/g      | 6.28 ± 0.07     | 6.19 ± 0.06   |
| Soleus mass, mg  | 9.1 ± 0.1*      | 9.5 ± 0.1     |
| SM:BM, mg/g      | 0.45 ± 0.01*    | 0.38 ± 0.01   |
| Plantaris mass, mg | 19.2 ± 0.3*    | 22.1 ± 0.3    |
| PM:BM, mg/g      | 0.94 ± 0.01*    | 0.87 ± 0.01   |

Values are mean ± SE. HM:BM, heart mass-to-body mass ratio; GM:BM, gastrocnemius mass-to-body mass ratio; SM:BM, soleus mass-to-body mass ratio; PM:BM, plantaris mass-to-body mass ratio. * P < 0.05 significantly different from male mice.

Relative to the progenitor strains, eight week old F₂ mice ran an average of 30.8 ± 0.1 min, which was significantly longer (P < 0.0001) than B6.A14 (28.5 ± 0.2 min) and not different from B6 (31.0 ± 0.1 min) mice. F₂ mice also performed significantly more work (1.69 ± 0.02 kg·m, P < 0.0001) than B6 (1.39 ± 0.04 kg·m) and B6.A14 (0.95 ± 0.02 kg·m) strains. Body mass also was compared across strains and generations. F₂ mice had average body mass of 22.1 ± 0.2 g, which was significantly greater than (P < 0.0001) the progenitor B6 (20.7 ± 0.8 g), and B6.A14 (18.9 ± 0.6 g) strains, respectively. Similar to the inbred strains, there was a significant negative correlation between exercise time and body

Figure 2. Frequency distribution for (A) time and (B) work in male and female (B6.A14 × B6) F₂ mice. All F₂ mice (n = 155) performed a graded exercise test to exhaustion to assess exercise capacity. Mice assorted into 1min/0.25 kg·m buckets. n = 67 for males and n = 88 for females.
mass \( r = -0.74, P = 0.02 \). However, this relation was not main-
tained when the F2 population was divided by sex. In male F2 mice
the correlation between exercise time and body mass was -0.47
\( P = 0.001 \), but there was no significant relation between these
variables in female F2 mice \( r = 0.14, P = 0.18 \).

QTL analysis. Significant differences in exercise capacity between
B6.A14 and B6 strains indicate the presence of a QTL on Chr 14.
To fine-map the QTL, F2 mice were generated from B6.A14 and
B6 strains. Single chromosome-wide scans for time and work are
shown in Figure 3. Suggestive QTL for time (LOD = 1.75, P = 0.131) and work (LOD = 2.08, P = 0.063) with no covariates were
identified on Chr 14. When sex was included as an interacting
covariate, a significant QTL was identified at 56 cM for time (LOD
= 3.8, \( P = 0.048 \)) (Figure 3A). A suggestive QTL for work (LOD
= 3.69, \( P = 0.07 \)) was identified at the same location (Figure 3B).
Because significant and suggestive QTL were identified using sex as
an interacting covariate, chromosome-wide scans were performed
on male and female mice separately. In male mice, a significant
QTL for time (LOD = 2.28, 1.5 LOD = 49.0 – 58.9 cM, \( P = 0.049 \))
and a suggestive QTL for work (LOD = 2.19, 1.5 LOD = 38.0 –
58.9 cM, \( P = 0.056 \)) were identified at 55 cM. In female mice,
no QTL were identified for time (Figure 4A). However, a
suggestive QTL for work was identified at 16 cM (LOD = 1.8,
\( P = 0.106 \)).

Figure 3. QTL analyses on mouse Chromosome 14 for intrinsic
capacity expressed as time (min, A) and work (kg·m, B) in 155
(B6.A14 × B6) F2 mice. Three analyses were performed for each
phenotype: 1) with no covariates, 2) with sex as an additive covariate,
and 3) with sex as an interactive covariate. For time, significant
\( P = 0.05 \) logarithm of odds (LOD) thresholds are 2.22 with no
covariates, 2.12 with sex as an additive covariate, and 3.78 with
sex as an interactive covariate. For work, significant \( P = 0.05 \) LOD
thresholds are 2.22 with no covariates, 2.27 with sex as an additive
covariate, and 3.95 with sex as an interactive covariate. LOD
thresholds were determined using 1000 permutations. Chromosome-
wide scans and permutation analyses were performed using R/qtl.

Figure 4. QTL analyses for the effect of sex on intrinsic exercise
capacity in male and female (B6.A14 × B6) F2 mice expressed as
time (A) and work (B). Single chromosome-wide scans for time
(in min) and work (in kg·m) were performed separately on male
and female F2 mice. Dashed lines in the upper graph represent the
suggestive (0.82, \( P = 0.63 \)) and significant (2.21, \( P = 0.05 \)) logarithm
of odds (LOD) thresholds for time in males. Dashed lines in the lower
graph represent the suggestive (0.82, \( P = 0.63 \)) and significant (2.25,
\( P = 0.05 \)) LOD thresholds for work in males. In females, suggestive
and significant LOD thresholds for time were 0.84 (\( P = 0.63 \)) and
2.15 (\( P = 0.05 \)), respectively; and for work 0.81 (\( P = 0.63 \)) and 2.08
(\( P = 0.05 \)), respectively. LOD thresholds were determined using 1000
permutations. Chromosome-wide scans and permutation analyses
were performed using R/qtl.
The two-QTL analyses for time showed limited evidence for additive QTL at 0 cM and 58 cM (LOD = 2.74, P = 0.19) on Chr 14. No significant additive or interacting QTL were identified for work. QTL scans also were performed for all physical characteristics and no significant QTL were identified.

The allelic effects for suggestive and significant QTL are shown in Table 3. In the entire F2 cohort, heterozygous mice had the highest average exercise time and work. For both phenotypes there was no significant difference between homozygous A and B groups. A similar pattern was observed for time and work in the male F2 cohort. In this group, mice with parental genotypes had significantly lower exercise time than mice carrying the heterozygous genotype (Table 3). In the female F2 cohort, work was significantly higher in homozygous A mice compared with homozygous B mice. Heterozygous female F2 mice had an intermediate phenotype.

### Table 3. Allelic effects for significant and suggestive QTL for exercise time and work in sex-specific and entire F2 cohorts.

| Cohort | Trait      | Position, cM | Marker         | Genotype | p-value |
|--------|------------|--------------|----------------|----------|---------|
|        |            |              |                |          |         |
|        | Time, min  | 58           | rs3685710      | A        | 30.4 ± 0.3 | 31.1 ± 0.2 | 30.4 ± 0.3 | 0.986 | 0.078 | 0.067 |
|        | Work, kgm  | 57           | rs3685710      | B        | 1.68 ± 0.03 | 1.74 ± 0.02 | 1.63 ± 0.03 | 0.532 | 0.244 | 0.019* |
|        | Time, min  | 55           | rs3715673      | A vs. B  | 28.9 ± 0.2 | 29.7 ± 0.3 | 28.8 ± 0.2 | 0.945 | 0.048* | 0.042* |
|        | Work, kgm  | 55           | rs3715673      | A vs. H  | 1.62 ± 0.05 | 1.76 ± 0.04 | 1.59 ± 0.08 | 0.933 | 0.107 | 0.070 |
|        | Work, kgm  | 16           | rs3696080      | B vs. H  | 1.78 ± 0.05 | 1.72 ± 0.02 | 1.65 ± 0.03 | 0.035* | 0.530 | 0.107 |

Position, location of peak marker in cM; Marker, SNP marker closest the LOD peak; Genotype, genotype at peak marker; A, homozygous for A/J allele; B, homozygous for B6 allele; H, heterozygous; p-value, p-value from means comparison using Tukey post-hoc analysis with specific allelic comparisons indicated (e.g., A vs. B), significant p-values are indicated by*.
the substituted chromosome shifted the phenotype toward the donor strain, the effect of chromosome substitution on exercise capacity was less than expected. Based on previous CSS surveys, chromosome substitution can produce phenotypic effects of 75% or more of the difference between parental strains. Nevertheless, the significant difference between B6 and B6.A14 for exercise time and work suggest the presence of one or more QTL on Chr 14 for exercise capacity.

To localize the QTL on Chr 14, linkage analysis was performed in F$_1$ mice from B6 and B6.A14 strains. Suggestive QTL were identified for both time and work at 58 cM (Figure 3). The 1.5 LOD interval for each of these QTL spanned nearly the entire chromosome, so these QTL overlapped with previously reported QTL for pre-training and post-training work. However, the peak markers for pre-training work (4 cM) and post-training work (26 cM) QTL localize to positions distant from the QTL identified in the current study and likely represent different QTL. Further analysis using sex as an interactive covariate provided evidence for sex-specific QTL; therefore, male and female F$_1$ cohorts were analyzed separately. Significant and suggestive QTL for time and work, respectively, were identified in the male cohort only and were similar to those identified in the entire cohort. Conversely, there was less evidence for exercise-related QTL on Chr 14 in female mice. This is somewhat surprising given the differences between B6 and B6.A14 female mice were comparable to those in male mice (Figure 1). However, the peak marker for the suggestive QTL for work in the female F$_1$ cohort is in close proximity to a syntenic human region linked to maximal oxygen consumption in the sedentary state in the HERITAGE Family Study. We previously reported a significant effect of sex on exercise capacity after 4 weeks of exercise training and the responses to training in mice. Sex-specific QTL also have been reported for voluntary wheel running and exercise-related traits such as muscle mass. Furthermore, Wang et al. identified several QTL related to fat mass in the mouse which were influenced by sex. Global gene expression analysis of liver tissue in the same population of mice revealed that a large percentage of expression QTL also were influenced by sex. These data suggest that sex can affect the genetic regulation of gene expression as well as clinical phenotypes. Therefore sex-specific effects should be considered when investigating the genetic regulation of phenotypes, especially those such as exercise capacity that are known to differ between males and females.

One potential explanation for the limited evidence for exercise QTL in the F$_1$ cohort is that the number of animals was insufficient for detecting multiple QTL with small effects. However, the number of mice included in the entire F$_1$ cohort or each sex-specific cohort is comparable to most intercross populations utilizing a CSS and B6 strains and should have been sufficient to detect at least 1 QTL. Similar to the current study, Burrage et al. were also unable to localize QTL in CSS × B6 intercross populations for several traits showing significant differences between parental CSS and B6 mice. They concluded that multiple QTL with opposing effects might be present on individual chromosomes and that congenic strains might be more advantageous for QTL detection and mapping than larger intercross populations. Alternatively, a close inspection of the allelic effects for each exercise QTL suggests that alleles derived from the A/J stain contribute to increasing exercise capacity (Table 3).

This was most evident in the female F$_1$ cohort. The suggestive QTL for work identified in this population mapped to a position (16 cM) that was different from that observed in the entire F$_1$ and male-only cohorts. In females, mice homozygous for the parental A allele performed significantly greater work that mice homozygous for the parental B allele. Heterozygous mice were intermediate and not significantly different from either parental genotyping suggesting an additive inheritance pattern with the A allele conferring increasing exercise capacity. In the full F$_1$ and male-only cohorts, there was no significant difference between mice homozygous for the parental genotypes and heterozygous mice had the highest exercise capacity. Thus, at some locations A and B alleles can interact to elicit a phenotype greater that either parental genotype.

Collectively, these data support the use of CSS as a model for the genetic analysis of exercise capacity. They also provide evidence that genetic factors on Chr 14 contribute to the variation in exercise capacity. Based on the complexity of the exercise phenotype, a survey of the complete C57BL/6J-Chr$^{AJ}$/NaJ CSS panel will likely identify multiple chromosomes of interest and potential QTL for exercise capacity. Furthermore, the sex-dependent differences in exercise capacity and the putative sex-specific QTL imply that the genetic architecture underlying exercise capacity might be different between males and females. Thus, any such survey should be conducted in male and female mice to elucidate the potential genotype by sex interaction underlying differences in exercise capacity between males and females. Once strong candidate genes are identified, the link between exercise capacity and cardiorespiratory fitness, and the mechanistic basis for diseases associated with low cardiorespiratory fitness can be explored.

Data availability
Figshare: Effect of chromosome 14 substitution on intrinsic exercise capacity in mice: R/qtl linkage analysis and phenotype data, http://dx.doi.org/10.6084/m9.figshare.893581.

Author contributions
SC designed and conducted the study, analyzed the data, and wrote the manuscript. SC critically revised the manuscript and agreed to its publication. MM designed the study, analyzed the data, and wrote the manuscript. MM critically revised the manuscript and agreed to its publication.

Competing interests
No competing interests were disclosed.

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Version 2

Reviewer Report 23 June 2014

https://doi.org/10.5256/f1000research.4599.r3115

Reuben Howden
Department of Kinesiology, University of North Carolina at Charlotte, Charlotte, NC, USA

I appreciated and enjoyed the interesting conversation with these authors on an important and engaging topic. I have no further comment regarding this manuscript.

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 1

Reviewer Report 11 April 2014

https://doi.org/10.5256/f1000research.3335.r4461

Reuben Howden
Department of Kinesiology, University of North Carolina at Charlotte, Charlotte, NC, USA

The authors give the impression that a higher, genetically controlled, exercise capacity should equal lower disease risk based on previous work. However, I wonder if a genetic predisposition to a higher exercise capacity is more or less important that habitual exercise itself. It would be interesting to investigate disease risk in inbred mice using training programs of differing volume/intensity so the effect of environmental factors (e.g. habitual exercise) can be assessed in
a controlled environment. Habitual exercise and disease risk reductions may be achievable without developing an exceptional exercise capacity. Prof. Edward Howley’s recent comments about recommending vigorous exercise, needed for high exercise capacity, from the end of his lecture\(^1\) may be pertinent here.

Assessment of exercise capacity by treadmill running - this may be an age old question, but it remains an important one; how do the authors know they successfully assessed exercise capacity in these strains and that differences in treadmill running time were not influenced by individual strain motivation to continue exercise? While this is not a study design flaw, it could change the question asked about factors influencing sedentary behavior and chronic disease risk in human populations.

As the authors point out, a limitation of their study is the small F\(_2\) cohort. While the reported cohort size may be concordant with previous work, that does not mean there were sufficient meiotic events to fine map their previously identified QTL on Chr14.

It is difficult to determine from this article if the Chr14 QTL was reduced in the current study. Perhaps the authors could provide more information about that.

Did the use of CSS increase the authors confidence about specific candidate genes identified previously?\(^2\) What potential candidate genes are located within the present (refined?) Chr14 QTL?

I agree that congenic strains might be useful for identifying which regions within the reported Chr14 QTL, although because of the likely complexity of factors influencing exercise capacity, demonstrating a phenotypic influence of more specific genomic regions or individual candidate genes may be challenging.

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**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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**Author Response 23 Apr 2014**

Michael Massett, Texas Tech University, USA

“The authors give the impression that a higher, genetically controlled, exercise capacity should equal lower disease risk based on previous work. However, I wonder if a genetic predisposition to a higher exercise capacity is more or less important that habitual exercise itself. It would be interesting to investigate disease risk in inbred mice using training programs of differing
volume/intensity so the effect of environmental factors (e.g. habitual exercise) can be assessed in a controlled environment. Habitual exercise and disease risk reductions may be achievable without developing an exceptional exercise capacity. Prof. Edward Howley’s recent comments about recommending vigorous exercise, needed for high exercise capacity, from the end of his lecture may be pertinent here.”

The Reviewer raises several good points in this comment. First, our goal in the Introduction was to remind the readers of the well-documented inverse relationship between cardiorespiratory fitness and morbidity and mortality in humans. This relationship appears to plateau at 10 METs, suggesting that having a fitness level above 10 METs does not confer significant additional protection (Kokkinos et al., 2008). Therefore, we agree with the Reviewer that developing high levels of exercise capacity and/or fitness are not required to achieve beneficial effects.

Regarding the question of fitness versus physical activity, the Reviewer is correct in pointing out that increasing physical activity can reduce the risk of all-cause mortality and CVD and that some benefit may be gained by increasing activity without improving fitness (maximal oxygen uptake). However, Williams (2001) did point out that cardiorespiratory fitness and physical activity should be considered independently as predictors of disease risk and that the relative risk of heart disease in the most fit group is nearly half that of the most active group. These data imply that both fitness and activity are important factors in reducing the risk of cardiovascular disease. Determining the genetic basis for each phenotype could yield important information regarding the beneficial effects of exercise.

“Assessment of exercise capacity by treadmill running - this may be an age old question, but it remains an important one; how do the authors know they successfully assessed exercise capacity in these strains and that differences in treadmill running time were not influenced by individual strain motivation to continue exercise? While this is not a study design flaw, it could change the question asked about factors influencing sedentary behavior and chronic disease risk in human populations.”

In the current study, exercise performance was assessed during a graded exercise test, which was stopped when mice exhibited pre-defined behaviors that we, and others, have defined as exhaustion. As pointed out by Booth, Laye and Spangenburg (2010), in ideal settings, exhaustion would be defined by a number of physiological markers. The large number of mice required for genetic experiments exceeded our capacity to assess maximal oxygen consumption and/or blood lactate levels at the time of testing. However, Desai et al. (1997) showed that an exercise test similar to the one utilized in the current study elicited heart rates near 750 beats per minute and a respiratory exchange ratio at or above 1.0 in mice from several inbred strains. We cannot assess an animal’s motivation to run on the treadmill so we do not know how much this might have contributed to our results. But, as demonstrated by Lerman et al. (2002) the willingness to run voluntarily is likely determined by different genetic factors than those underlying the variation in intrinsic exercise capacity in the untrained state. This latter phenotype is the focus of the current study.

“As the authors point out, a limitation of their study is the small F cohort. While the reported cohort size may be concordant with previous work, that does not mean there were sufficient
meiotic events to fine map their previously identified QTL on Chr14.”

The number of mice utilized in this study was sufficient to detect at least one QTL explaining 5% of the variance in exercise capacity with a power of 50% (Belknap, 2003). Increasing the number of mice would likely have allowed us to detect more QTL with smaller effects. However, as the reviewer points out, an experimental model with a greater number of recombination events, such as an advanced intercross line based on A and B6 strains, the hybrid mouse diversity panel, or mice from the collaborative cross, would probably have yielded a greater resolution for our QTL interval.

“It is difficult to determine from this article if the Chr14 QTL was reduced in the current study. Perhaps the authors could provide more information about that.”

The QTL for pre-training work on Chromosome 14 identified in our previous study was located at 4 cM with a 95% CI of 0-38 cM. The suggestive QTL for time and work identified in the current study were located near 58 cM and the 95% CI were very large. Although the QTL in the current study did overlap with our previously identified QTL, it is very likely that the QTL identified in the current study are different from our previous report. A brief statement regarding this issue is located in the fourth paragraph of the Discussion section.

“Did the use of CSS increase the authors confidence about specific candidate genes identified previously? What potential candidate genes are located within the present (refined?) Chr14 QTL?”

Because there was no overlap between the current and previous QTL, the use of the CSS model did not increase our confidence regarding specific candidate genes in the Chromosome 14 QTL region. However, given the complexity of the exercise capacity phenotype, performing a genome scan using a complete set of CSS might be a logical step toward identifying chromosomes of interest and potential candidate genes. These chromosomes could then be targeted for more detailed analyses.

“I agree that congenic strains might be useful for identifying which regions within the reported Chr14 QTL, although because of the likely complexity of factors influencing exercise capacity, demonstrating a phenotypic influence of more specific genomic regions or individual candidate genes may be challenging”

We agree with the Reviewer that identifying candidate genes for exercise and exercise-related traits is challenging. However, the availability of resources for genetic and genomic studies continues to increase, which should facilitate discovery of genes and gene networks that regulate variation in exercise capacity, fitness, and potentially diseases associated with low levels of fitness.

**Competing Interests:** None.
Michael Turner
Department of Kinesiology, University of North Carolina at Charlotte, Charlotte, NC, USA

This manuscript is very well written regarding a strong, well thought out research design. The research is cutting edge and moves the knowledge regarding possible genetic factors influencing intrinsic exercise capacity in active mice. Chromosome 14 is an important participant in this phenotype and the authors do an excellent job utilizing chromosome substitution to continue pursuing their research interests.

Minor comments:
- The authors point out in the first paragraph of the Results that body mass was less in the B6.A14 mice compared to the B6 mice. Could the authors perform a simple regression to assess the influence of differing body mass on intrinsic exercise capacity as a potential confounding factor to their findings?
- The third sentence in the second paragraph of the Results appears to conflict with the previous statement regarding differences in work with sex in the F2 generation. If I'm reading these two sentences incorrectly I would ask a clearer statement be made to help the reader understand the sex-related difference in work with these F2 mice.
- In the last few sentences in the fourth paragraph of the Discussion the authors present an interesting discussion regarding the different QTLs on chromosome 14 that may be playing a role in the male vs. female F2 mice and intrinsic exercise capacity. More discussion would add to the major findings and future directions for readers. This is a difficult research issue and a bit more discussion would direct the readers towards an appreciation of this position. Possibly a better transition to the argument provided in the following paragraph would assist in the interpretation of the authors' findings.

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Michael Massett, Texas Tech University, USA

The authors point out in the first paragraph of the Results that body mass was less in the B6.A14 mice compared to the B6 mice. Could the authors perform a simple regression to assess the influence of differing body mass on intrinsic exercise capacity as a potential confounding factor to their findings?

We thank the Reviewer for this suggestion. We performed a linear regression to determine
the relation between body mass and exercise capacity expressed as time. We analyzed each of the inbred strains separately as well as the F₂ population. For each of these groups, exercise time was significantly negatively correlated with body mass. In general, body mass explained 50% to 74% of the variance in exercise time in the inbred strains. In the F₂ population, the variance in exercise time explained by body mass was approximately 54%; however, this percentage decreased when the population was divided by sex. In males this percentage was 22% and in females, less than 2%. We will include the correlation data in the revised manuscript to identify both the direction and magnitude of this relation.

The third sentence in the second paragraph of the Results appears to conflict with the previous statement regarding differences in work with sex in the F generation. If I'm reading these two sentences incorrectly I would ask a clearer statement be made to help the reader understand the sex-related difference in work with these F₂ mice.

We thank the Reviewer for pointing out this confusing description. Simply stated, work is determined as the product of exercise capacity and body mass. Female mice ran longer/farther than male mice, but weighed less; therefore work was approximately equal between male and female F₂ mice. We will edit this description in the revised manuscript.

In the last few sentences in the fourth paragraph of the Discussion the authors present an interesting discussion regarding the different QTLs on chromosome 14 that may be playing a role in the male vs. female F mice and intrinsic exercise capacity. More discussion would add to the major findings and future directions for readers. This is a difficult research issue and a bit more discussion would direct the readers towards an appreciation of this position. Possibly a better transition to the argument provided in the following paragraph would assist in the interpretation of the authors' findings.

We are grateful for the Reviewer's appreciation of the complexity of the phenotype and the number of genetic and environmental factors that influence this phenotype. We will expand our discussion of some of these genetic factors in the revised manuscript.

**Competing Interests**: None
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