

Effect of Within-Strain Sample Size on QTL Detection and Mapping Using Recombinant Inbred Mouse Strains

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Increasing the number of mice used to calculate recombinant inbred (RI) strain means increases the accuracy of determining the phenotype associated with each genotype (strain), which in turn enhances quantitative trait locus (QTL) detection and mapping. The purpose of this paper is to examine quantitatively the effect of within-strain sample size (n) on additive QTL mapping efficiency and to make comparisons with F_2 and backcross (BC) populations, where each genotype is represented by only a single mouse. When 25 RI strains are used, the estimated equivalent number of F_2 mice yielding the same power to detect QTLs varies inversely as a function of the heritability of the trait in the RI population (h_{RI}^2). For example, testing 25 strains with $n = 10$ per strain is approximately equivalent to 160 F_2 mice when $h_{RI}^2 = 0.2$, but only 55 when $h_{RI}^2 = 0.6$. While increasing n is always beneficial, the gain in power as n increases is greatest when h_{RI}^2 is low and is much diminished at high h_{RI}^2 values. Thus, when h_{RI}^2 is high, there is little advantage of large n , even when n approaches infinity. A cost analysis suggested that RI populations are more cost-effective than conventional selectively genotyped F_2 populations at h_{RI}^2 values likely to be seen in behavioral studies. However, with DNA pooling, this advantage is greatly reduced and may be reversed depending on the values of h_{RI}^2 and n .

KEY WORDS: Recombinant inbred strain; quantitative trait locus; chromosome mapping; quantitative genetics; mouse.

INTRODUCTION

A common question in the design of experiments is to how to allocate subjects to cells in an experimental design to obtain the highest relative efficiency (RE), where RE reflects the power to detect treatment effects (smallest error term) when different designs of the same sample size are compared (Sokal and Rohlf, 1995). RE can also be expressed as the ratio of sample sizes for experimental designs yielding the same power. For example, if we

are limited to 100 mice, is it better to test 100 recombinant inbred (RI) strains with one mouse per strain or 25 strains with four mice per strain? In general, the first is the better choice (Sokal and Rohlf, 1995; Knapp and Bridges, 1990), but when every available strain is routinely tested, as is usually the case in the mouse, there is no option concerning the number of RI strains. The only choice is the selection of the within strain sample size, or n . In this paper, the RE at varying values of n is examined when the number of strains is fixed at the maximum number available, and comparisons are made with segregating F_2 or BC populations.

For QTL mapping studies in the mouse, the major disadvantage of RI strains compared to segregating populations is the limited number of gen-

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otypes available. Since each genotype is represented by a single RI strain, the largest existing RI sets, the B \times D, A \times B/B \times A, and LS \times SS sets, are limited to no more than 26 or 27 distinct genotypes (Taylor, 1995; DeFries *et al.*, 1989). Since one or two strains per set often reproduce too poorly to include in most RI studies, 25 strains is a reasonable upper limit for the existing mouse RI sets. In contrast, segregating (F_2 or BC) populations can involve any number of genotypes, since each genotype is represented by a single mouse, and large numbers can be generated.

However, RI strains have at least three advantages over segregating populations (Bailey, 1981; Plomin and McClearn, 1993). First, they have all the advantages of any inbred strain, including direct comparisons of genetic and phenotypic information across time, traits, and laboratories on the same set of readily available and stable genotypes. Second, RI strains are homozygous at all loci, which is more informative for QTL mapping than the usually intermediate-scoring heterozygotes comprising half of F_2 populations, or the total absence of one homozygote class in BC populations. Third, by testing several animals per strain, replicate measurements on the same genotype can be made to assess more accurately the phenotype associated with each genotype. This increases the power to detect QTLs compared to testing only one mouse per genotype (Knapp and Bridges, 1990; Soller and Beckman, 1990). In an F_2 or BC, in contrast, each genotype is represented by only a single mouse that cannot be replicated. The implications of these differences are examined below.

METHODS AND RESULTS

Only additive effects of QTLs are considered, since dominance effects do not occur in RI strains due to the absence of heterozygotes. It is presumed throughout that all populations are tested for the same trait and are derived from the same two inbred strains, thus allelic frequencies of $p = q = 0.5$ are assumed at all loci. We define n as the number of mice per genotype (strain), N_{str} as the number of strains, and N as the total number of animals in an experiment. Since each genotype in an F_2 or BC is represented by a single mouse, $n = 1$ in all cases, but $n > 1$ is typical in RI experiments. It is assumed for simplicity that n is the same for all RI

strains, therefore, $N_{\text{RI}} = n \times N_{\text{str}}$. When n is unequal, average n can be substituted.

Quantitative Genetic Considerations. When data from individual mice (not strain means) are used in quantitative genetic analyses, we can partition the variance in the usual way for an RI population as follows. This partitioning is similar to that in a segregating population when there is no dominance variation [Eq. (A)]: $V_P = V_A + V_E$, and the heritability (h_{RI}^2) is given by $V_A/(V_A + V_E)$, where V_P is the phenotypic (trait) variance, V_A is the additive genetic component of variance, and V_E is the environmental component of variance. The value of h_{RI}^2 can be estimated in several ways. The first is to use R^2 from a one-way ANOVA by RI strain, or $SS_{\text{strain}}/SS_{\text{total}}$. The second way is to use components of variance between and within strains calculated from the same one-way ANOVA (Hegmann and Possidente, 1981; Belknap *et al.*, 1996). The third method, and perhaps the simplest, is to use the variance of strain means to estimate V_A and divide by V_P . In this case, adjustments are often needed to correct for the fact that the variance of strain means contains a portion of V_E (Hegmann and Possidente, 1981).

V_{QTL} is the additive genetic variance due to a QTL and is calculated in an RI population as $(M_{A_1} - M_{A_2})^2/4$, where $M_{A_1} - M_{A_2}$ is the difference in phenotypic means between the two homozygote classes at a QTL or closely linked marker. In Falconer's terminology, $M_{A_1} - M_{A_2}$ is equal to twice the average effect of a single gene substitution (Falconer and Mackay, 1996). One-half of this value, or $(M_{A_1} - M_{A_2})^2/8$, gives an estimate of V_{QTL} to be expected in an F_2 population for the same QTL, and half of the F_2 estimate gives the expected BC estimate (Kearsey and Pooni, 1996). To determine h_{QTL}^2 , these V_{QTL} estimates are divided by V_P , the phenotypic variance in each population.

While V_A in an RI population can be expected to be double that in a comparable F_2 and quadruple that in a BC (Kearsey and Pooni, 1996), what can we expect for the heritability? The heritabilities in each population for the same trait will also differ approximately in proportion to V_A , that is, $h_{\text{RI}}^2 \approx 2h_{\text{F}_2}^2 \approx 4h_{\text{BC}}^2$, if V_P remains about the same in all three populations. However, V_P may not be equal, especially when h_{RI}^2 is large. The twofold greater value of V_A in RI vs. F_2 populations (i.e., $V_{A(\text{RI})} = 2V_{A(\text{F}_2)}$) can be expected to cause V_P to be larger in RI populations by an amount equal to $1/2V_{A(\text{RI})}$.

Thus, $V_{P(F_2)} = V_{P(RI)} - 1/2V_{A(RI)}$. From this, a more accurate estimate of $h_{F_2}^2$ from RI data can be obtained by taking into account the expected inequality in V_P , as follows [Eq. (B)]: $h_{F_2}^2 = 1/2V_{A(RI)}/(V_{P(RI)} - 1/2V_{A(RI)}) = 1/2h_{RI}^2/(1 - 1/2h_{RI}^2)$. [For a BC, $h_{BC}^2 = 1/4h_{RI}^2/(1 - 3/4h_{RI}^2)$.] Therefore the ratio of $h_{RI}^2/h_{F_2}^2$ will be $2(1 - 1/2h_{RI}^2)$ rather than 2. The same is also true at the QTL level; the heritability of a QTL (h_{QTL}^2), or V_{QTL}/V_P in an RI population, is expected to be somewhat less than double that in a comparable F_2 and quadruple that in a comparable BC. More accurately, when the inequality of V_P noted above is accounted for [Eq. (C)], $h_{QTL(F_2)}^2 = 1/2h_{QTL(RI)}^2/(1 - 1/2h_{RI}^2)$ for a given QTL. The above equations assume that each individual mouse is a data point for ANOVA, and that V_E is approximately the same in RI, F_2 , and BC populations. The equality of V_E assumption may be reasonable for some behavioral traits and not for others (e.g., Hyde, 1973).

Effect of Within-Strain Sample Size, n. What happens when $n > 1$ and strain means are used in the analysis rather than individual mice? In this case, the genotypic value is the mean of measurements on n mice per strain, providing a more accurate assessment of the phenotype associated with each genotype compared to $n = 1$. For RI strains, the variance partitioning based on strain means (\bar{x}) is as follows [Eq. (D)]: $V_{P\bar{x}} = V_A + V_E/n$ (Soller and Beckmann, 1990), and the heritability (h_{RI-x}^2) is given by $V_A/(V_A + V_E/n)$, where $V_{P\bar{x}}$ is the phenotypic variance of strain means and h_{RI-x}^2 is the heritability of strain means. h_{RI-x}^2 reflects the degree to which the variance of strain means, $V_{P\bar{x}}$, is due to genetic sources of variation. As n increases, the contribution of the environmental component to $V_{P\bar{x}}$ decreases by a factor of $1/n$ (Soller and Beckmann, 1990). This has the effect of increasing the heritability based on strain means in an RI population as a function of n . As n becomes very large, h_{RI-x}^2 approaches 1.0 because the contribution of the environmental component of variance is approaching zero, causing $V_{P\bar{x}}$ to approach V_A in value. When this happens, the variance of phenotypic strain means, $V_{P\bar{x}}$, provides a good estimate of V_A . However, in many reports in the literature, it is often overlooked that this estimate is biased upward when h_{RI-x}^2 is considerably less than unity. Corrections for this bias can be made by multiplying $V_{P\bar{x}}$ by the estimate of h_{RI-x}^2 taken from Fig. 1, which eliminates this source of bias. Much the same is

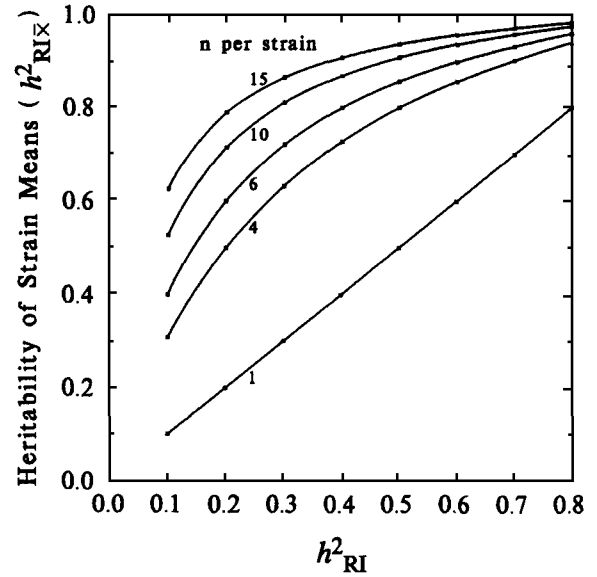


Fig. 1. The expected heritability of strain means (h_{RI-x}^2), or the proportion of the variance in strain means due to additive genetic sources, plotted as a function of the heritability of the trait (h_{RI}^2), or the proportion of the total variance due to additive genetic sources. Plots for varying values of the within strain sample size, n , are shown. h_{RI-x}^2 is based on the mean of n mice per strain, while h_{RI}^2 is based on individual mouse scores. In the special case where $n = 1$ per strain, $h_{RI-x}^2 = h_{RI}^2$. As n becomes very large (approaches infinity), h_{RI-x}^2 approaches 1.0 and the variance of strain means approaches V_A in value.

true for standard (non-RI) inbred strains when the analysis is carried out within and between strains in the same manner.

The same relationship between h_{RI}^2 and h_{RI-x}^2 outlined above also applies to the heritability of a QTL based on individual mice compared to that calculated from strain means. The heritability of a QTL is the proportion of the phenotypic variance due to a QTL and is given by $h_{QTL}^2 = V_{QTL}/V_P = V_{QTL}/(V_A + V_E)$ when data from individual mice are the basis for the analysis. When strain means are used, the heritability of a QTL becomes [Eq. (E)]: $h_{QTL-x}^2 = V_{QTL\bar{x}}/V_{P\bar{x}} = V_{QTL}/(V_A + V_E/n)$, which increases as n increases in a directly parallel manner to h_{RI-x}^2 . In other words, as n increases, the proportionate increase in h_{RI-x}^2 and h_{QTL-x}^2 will be the same; thus the present analysis applies to both.

Using Eqs. (A) and (D) above, the relationship between h_{RI-x}^2 and h_{RI}^2 is shown in Fig. 1 as a function of n . Six values of n are plotted: 1, 4, 6, 10, 15, and ∞ . The ratio of h_{RI-x}^2/h_{RI}^2 represents the gain in heritability in an RI set when strain means are

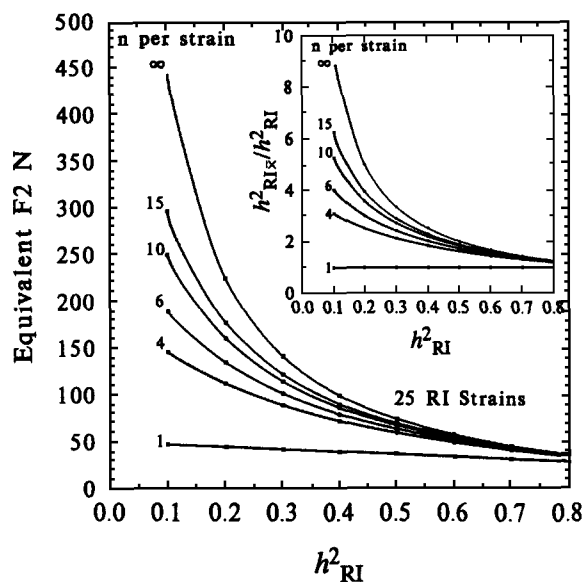


Fig. 2. Inset: The ratio of $h_{RI_x}^2/h_{RI}^2$ as a function of h_{RI}^2 for varying values of n , the within-strain sample size. This ratio represents the gain in heritability when strain means based on n mice are used compared to when individual mice are used in partitioning the phenotypic variance. This same ratio also applies to the heritability of a QTL. **Main body:** The estimated F_2 sample size approximately equivalent in power to 25 RI strains is plotted as a function of the heritability of the trait in the RI population (h_{RI}^2), and n , the within-strain sample size. The plotted values were calculated from Eq. (F) in the text. It is assumed that all phenotyped mice are also genotyped for both RI and F_2 mice. When selective genotyping is used in the F_2 , the plotted N_{F_2} values should be increased by 2.3-, 1.5-, 1.3-, 1.15-, or 1.08-fold, respectively, when the selection fraction for selective genotyping in both tails is 10, 20, 30, 40, or 50% of the total population (Darvasi and Soller, 1992).

used compared to when individual mice are used in the analysis, as in a segregating population. This ratio is plotted in Fig. 2 (inset) as a function of h_{RI}^2 .

The same $h_{RI_x}^2/h_{RI}^2$ ratios in Fig. 2 (inset) also apply to the heritability of a QTL, or h_{QTL}^2 . This is because the ratio $h_{RI_x}^2/h_{RI}^2$ is equal to the ratio $h_{QTL_x}^2/h_{QTL}^2$, i.e., the effect of increasing n is proportionately the same for both the heritability of all QTLs in the aggregate and the heritability of an individual QTL.

Equivalent F_2 Sample Size. From the $h_{RI_x}^2/h_{RI}^2$ ratio, it is possible to estimate the approximate equivalent F_2 sample size, N_{F_2} , when 25 RI strains are tested with varying n . This estimated N_{F_2} will have the same power to detect additive effects of a QTL as the RI population. [Methods for cal-

culating power for F_2 populations are given by Darvasi and Soller (1992, 1994), and for RI populations by Belknap *et al.* (1996).] Since N_{F_2} required is proportional to the heritability of a QTL (Lander and Botstein, 1989; Soller and Beckmann, 1990; Belknap *et al.*, 1996), this estimate was calculated by multiplying the appropriate $h_{RI_x}^2/h_{RI}^2$ ratio (which equals the $h_{QTL_x}^2/h_{QTL}^2$ ratio) from Fig. 2 (inset) times 25 (N_{str}) times $2(1 - 1/2h_{RI}^2)$, the ratio of $h_{RI}^2/h_{F_2}^2$ when $n = 1$ for both. Thus [Eq. (F)], $N_{F_2} = 2(1 - 1/2h_{RI}^2)(h_{RI_x}^2/h_{RI}^2)N_{str}$. These N_{F_2} estimates are shown in Fig. 2 as a function of h_{RI}^2 . The plotted values presume that all phenotyped mice are also genotyped for both RI and F_2 populations. If selective genotyping is used in an F_2 population, where only the extreme tails of the trait distribution are genotyped to reduce cost (Lander and Botstein, 1989; Darvasi and Soller, 1992), the equivalent N_{F_2} will need to be larger than shown in Fig. 2 to offset the loss in power due to the restricted sample (Lander and Botstein, 1989). For example, with a selection fraction (both tails) of 10, 20, 30, 40, or 50% of the total population, the plotted N_{F_2} values shown in Fig. 2 should be increased by a factor of 2.3-, 1.5-, 1.3-, 1.15-, or 1.08-fold, respectively (Darvasi and Soller, 1992).

Relative Efficiency Ratio (RE). An important question is the relative efficiency (RE) of F_2 vs. RI populations, which can be expressed as the ratio of N_{F_2}/N_{RI} when the power to detect additive QTLs is the same in both populations. Thus, F_2 populations are more efficient (require fewer mice) than RI populations when RE is <1.0 and less efficient when RE >1.0 . This ratio can be calculated by determining the ratio of the estimated equivalent F_2 N for an RI population (Fig. 2) divided by the actual N used in the RI population. These results are shown in Fig. 3 in the case of no selective genotyping. When selective genotyping is used, the plotted RE values should be multiplied by a factor that varies with the selection fraction, as noted in the previous section.

For example, in the absence of selective genotyping, the estimated equivalent F_2 N when $h_{RI}^2 = 0.4$ and $n = 6$ for 25 RI strains is 80 (from Fig. 2), while the actual RI N is $25 \times 6 = 150$. The RE for the RI experiment is thus $80/150 = 0.53$, or 53%. Thus, an RI population under these conditions is only about half as efficient (requires twice as many mice) as an F_2 of the same power. If selective genotyping is used in the F_2 where (for ex-

ample) 1/8 of the population is genotyped (1/16 at each tail), the plotted N_{F_2} values should be increased by 2.0-fold to offset loss of power (Darvasi and Soller, 1992), yielding $N_{F_2} = 160$. In this case, $RE = 160/150 = 1.07$, thus RI has a slight advantage (requires 7% fewer mice). As can be seen in Fig. 3, RE is determined by h_{RI}^2 , n , and the F_2 selection fraction. RE (or N_{F_2}/N_{RI} when power is equal) is increased when either n , h_{RI}^2 , or the selection fraction is reduced in magnitude, and vice versa. The same considerations for h_{RI}^2 also apply to the heritability of a QTL (not shown).

The newest genotyping cost-saving measure for segregating populations is DNA pooling. The extreme ends of the trait distribution are genotyped, as in conventional selective genotyping, except that DNA pooled from all individuals in an extreme tail is genotyped rather than individual mouse samples. Darvasi and Soller (1994) discuss the cost benefit and genetic implications of this approach in the general case. Since the PCR reaction is only semiquantitative, a loss in accuracy occurs in estimating allele frequencies in each tail, resulting in an increase in both Type I (false positive) and Type II (false negative) errors (reduced power) that can vary widely from marker to marker. A further power loss occurs because interval mapping, as implemented by standard programs such as Map-Maker QTL, cannot be used with pooled data. (If a high density of markers is used, this power loss is minimal, but the additional genotyping required diminishes the cost savings.) To offset these shortcomings, (1) DNA pooling can be used as a low-cost preliminary screen of the genome, followed by individual genotyping for only those markers (say, 15%) showing evidence of QTL linkage (e.g., Taylor and Phillips, 1996), or (2) a larger F_2 population can be grown and phenotyped to offset the loss in power. (In a cost analysis example given below, it is assumed that a 20% larger population is needed.) With either option, the overall cost remains roughly the same in our example (calculations not shown).

Relative Cost per Mouse Ratio (RC). In addition to the relative efficiency, which reflects the number of F_2 vs. RI mice needed to obtain the same power, the relative cost per mouse, expressed as a ratio (F_2/RI), is also important in answering questions about the overall cost efficiency of F_2 vs. RI populations. We refer to this ratio as RC, in parallel with RE. Table I shows the costs per mouse in F_2 and RI populations based on data from our labo-

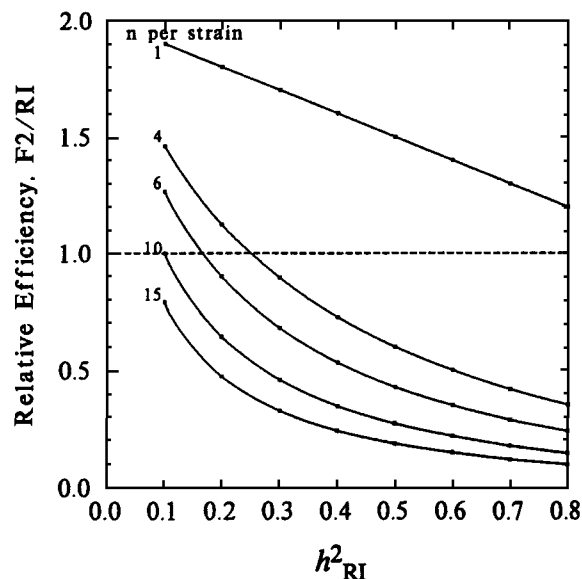


Fig. 3. The relative efficiency (RE) of F_2 vs. RI populations as a function of the heritability in the RI population (h_{RI}^2) and the within-RI strain sample size, n . RE was calculated as the ratio of the estimated F_2 N from Fig. 2 to the actual N used in the RI population, or $n \times N_{str}$. Thus, $RE = N_{F_2}/N_{RI}$ when both populations have the same power to detect additive QTLs. In the special case where $n = 1$ per strain, RE is just under 2 due to h_{RI}^2 being almost twice $h_{F_2}^2$ for the same trait. Plotted values presume that all phenotyped mice are also genotyped for both RI and F_2 populations. When selective genotyping is used in the F_2 , the RE values shown should be multiplied by a factor that depends on the selection fraction, as given in the legend to Fig. 2.

ratory, both with and without F_2 selective genotyping, and with or without DNA pooling. The optimal selection fraction to minimize overall cost has been estimated by Darvasi and Soller (1992) in the general case. For the data shown in Table I, where the ratio of genotyping cost to all other costs in the absence of selective genotyping was just over 13 (\$163/\$12.30), 1/8 is about the optimal fraction, or the extreme 1/16 at each end of the trait distribution. The “bottom line” from Table I is as follows: the total cost for an RI mouse was \$12.30 (breeding, raising, and phenotyping); for an F_2 mouse without selective genotyping, \$175.30; for an F_2 mouse with optimal selective genotyping, \$31.80; and for an F_2 mouse with DNA pooling coupled with selective genotyping, \$13.40. The relative cost ratio per mouse (F_2/RI , or RC) is therefore 14.25 without selective genotyping, 2.59 with conventional selective genotyping, and 1.09 with DNA pooling and selective genotyping combined.

Table I. Costs per Mouse for RI and F₂ Populations Based on Cost Data from Our Laboratory for a “Typical” Behavioral Trait^a

	RI	F ₂
Cost per mouse (US\$) with no selective genotyping		
1. Animal care for 6 weeks past weaning (\$0.14 × 42 days)	\$5.90	\$5.90
2. Cost of breeder pair (\$0.14 × 84 days × 2 plus purchase price per pair) × 1/30	\$2.40	\$1.40
3. Phenotyping (behavioral testing):	\$4.00	\$4.00
4. DNA extraction and purification (\$4)	None	\$4.00
5. Genotyping costs for 80 markers (\$2 × 80):	None	160.00
Total cost per mouse		
F ₂ /RI relative cost ratio (RC) = 14.25	\$12.30	\$175.30
With conventional selective genotyping in F ₂ (1/8)		
1–3. Same as above.		\$11.30
4. DNA extraction and purification (\$4 × 1/8)		\$0.50
5. Genotyping costs for 80 markers (\$2 × 80 × 1/8)		\$20.00
Total cost per mouse		
F ₂ vs. RI relative cost ratio (RC) = 2.59		\$31.80
With selective genotyping (1/8) and DNA pooling in F ₂ (N=400)		
1–4. Same as conventional selective genotyping above.		\$11.80
5. Genotyping costs for 80 markers (\$4* × 80 × 2 tails ÷ 400 mice)		\$1.60
Total cost per mouse		
F ₂ vs. RI relative cost ratio (RC) = 1.09		\$13.40

^a Each mouse was maintained for 6 weeks past weaning, or to 9 weeks of age, at \$0.14 per day animal care costs. The cost to breed each mouse includes the purchase of a mating pair from JAX at a cost of \$48 per RI breeder pair or \$18 per F₁ pair, and animal care for 84 days. Each mating pair produced an average of 30 offspring, thus the cost to breed each mouse will be 1/30 of the total mating pair cost. The costs of phenotyping a mouse are \$4, based on the ability of a \$24K/year technician to phenotype 4 mice per hr, plus \$1 per mouse for behavioral equipment and supplies. For a full genome search in an F₂, 80 markers at 20-cM intervals are needed at a cost of \$2 per marker for genotyping (labor, supplies, equipment, PCR, electrophoresis and photographic archiving costs). (This is just over half the cost of most outside contractors.) DNA extraction, purification and storage adds \$4 per mouse. When conventional selective genotyping is used, it is assumed that 1/8 are genotyped, the approximate optimal fraction to reduce cost in this example (Darvasi and Soller, 1992). Whether the relative F₂ vs. RI cost ratios per mouse (RC) shown below are applicable to other experiments will depend on the relative cost of animal production and phenotyping (categories 1, 2, and 3 together) vs. genotyping (4 and 5 below) and the selection fraction. Note that the costs per mouse do not take into account the additional mice needed to maintain the same power when selective genotyping is used; this is considered as part of relative efficiency, or RE (see text).

^b With DNA pooling, the added cost of densitometry equipment is incurred, and the per unit cost advantage of performing many genotypings per marker is largely lost. Thus, per unit cost with DNA pooling is estimated to double to \$4. It is presumed that the genotyping cost is distributed over a mapping population of N = 400 F₂ mice to estimate per mouse costs. However, the overall cost per mouse is little effected by other values of N or per-unit genotyping cost ranging from one-half to double the assumed values.

Relative Total Cost Ratio (RTC). The total cost (TC) of an experiment is given by the number of mice multiplied by the cost per mouse. The relative total cost ratio (RTC) of F₂/RI is thus RE × RC when the power to detect QTLs is the same in both populations. For example, when $h_{RI}^2 = 0.4$ ($h_{F_2}^2 = 0.25$) and $n = 10$, typical values for many behavioral traits, F₂ populations without selective genotyping will be RE (0.35 from Fig. 3) × 14.25 (from Table I), or five times as costly compared to RI populations of the same power.

When selective genotyping is used, RC declines to 2.59 in our example (Table I). In this case, RTC = RE (from Fig. 3 × 2) × RC (2.59). The

factor of 2 for RE is to offset the loss in power when selective genotyping of this magnitude (1/8) is practiced. Multiplying by 2 gives the RE value expected if no selective genotyping was used. (This is necessary because the RE values shown in Fig. 3 presume no selective genotyping.) For example, when $h_{RI}^2 = 0.4$ and $n = 10$, RTC = RE (0.35 from Fig. 3 × 2) × 2.59 (from Table I) = 1.8. Therefore, F₂ populations will be 1.8 times as costly compared to RI populations of the same power. For $n = 6$, RTC rises to almost threefold. Figure 4 shows RTC values as a function of h_{RI}^2 for $n = 6$ and 10, based on RE values taken from Fig. 3 and RC from Table I. Note that RTC will increase (F₂

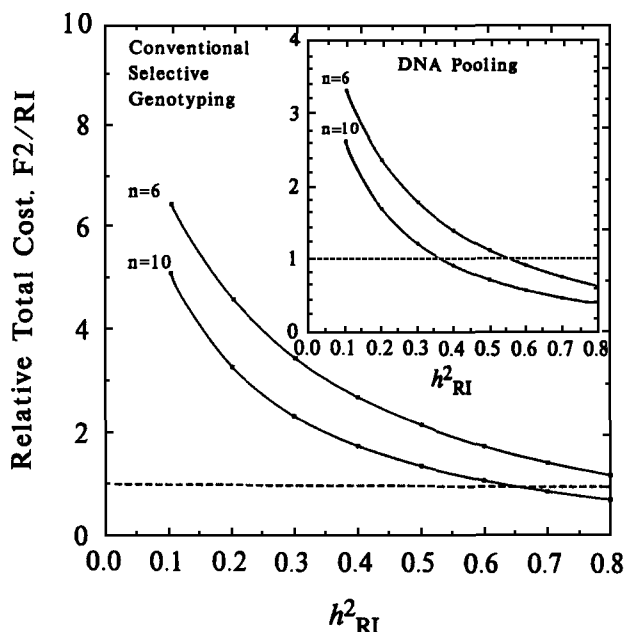


Fig. 4. The relative total cost (RTC = RE × RC) of an F₂ vs. RI experiment yielding the same power to detect additive QTLs based on RE from Fig. 3 and RC from the cost analysis shown in Table I. The results are shown for both conventional selective genotyping (**main body**) and selective genotyping and DNA pooling combined (**inset**). Plotted values assume within strain sample sizes (*n*) of either 6 or 10. The dashed horizontal lines show the equicost conditions (i.e., RTC = 1.0), where both populations have equal power and equal cost. The results shown are typical in our analysis of over 30 behavioral traits but can vary widely from laboratory to laboratory or trait to trait. For example, if genotyping costs are double those shown in Table I, then the plotted relative total cost values (*Y* axis) should be multiplied by 1.65 (conventional selective genotyping) or 1.24 (DNA pooling). In contrast, if genotyping costs are half those shown in Table I, the correction factors are 0.71 and 0.94, respectively. Note that the total costs associated with DNA pooling are relatively little affected by fluctuations in genotyping costs compared to conventional selective genotyping.

will cost relatively more than equivalent RI populations) when either h^2_{RI} or n decreases, and vice versa.

With selective genotyping and DNA pooling, RC declines to 1.09 in our example (Table I). As before, $RTC = RE \times RC$, or RE (from Fig. 3 \times 2.4) \times RC (1.09). The factor of 2.4 reflects the greater number of mice needed to offset the loss in power due to selective genotyping (2.0-fold) and DNA pooling (1.2-fold) used together.

The horizontal dashed line in Fig. 4 shows $RTC = 1.0$, when both populations have the same

cost for the same power. Where this line intersects the curved lines for $n = 6$ or 10 gives the equicost values of h^2_{RI} , where both populations are equal in power and cost. For conventional selective genotyping, this occurs when $h^2_{RI} = 0.90$ or 0.65 for $n = 6$ or 10, respectively. F₂ populations will be more cost-effective (cost less) than RI populations above the equicost h^2_{RI} values and less cost-effective below them. For selective genotyping coupled with DNA pooling, the corresponding “break-even” h^2_{RI} values are 0.55 ($h^2_{F_2} = 0.38$) and 0.37 ($h^2_{RI} = 0.23$) for $n = 6$ and 10, respectively, which fall within the range often seen for behavioral traits. Thus, the question of which population is more cost-effective will depend on observed values of h^2_{RI} for a given n .

DISCUSSION

Typical values of n reported in the RI behavioral literature are 6–10, and h^2_{RI} is often in the range of 0.3 to 0.6 (reviewed by Belknap *et al.*, 1997a), which is approximately equivalent to $h^2_{F_2}$ of 0.18 to 0.43 from Eq. (B). The approximate equivalent F₂ N conferring the same power for this frequently encountered range is about 50 to 115 mice when 25 RI strains are used. For QTL mapping purposes, this estimated F₂ N is capable of detecting only the very largest QTLs at Lander and Schork (1994) significance levels (Belknap *et al.*, 1997a). To detect smaller effect QTLs reliably, newer strategies have been developed involving RI and additional mapping populations (e.g., Johnson *et al.*, 1992; Plomin and McClearn, 1993; Belknap *et al.*, 1996, 1997a, b; Crabbe *et al.*, 1994). The most common strategy is a two-step approach, where an RI population is used as a preliminary screen of the genome for provisional QTLs at relatively relaxed α levels of (usually) .01 or .05 (Step 1), followed by confirmation testing of each RI-implicated chromosomal region in a large F₂ population derived from the same progenitors (Step 2). The advantages and disadvantages of this two-step approach have been recently reviewed (Belknap *et al.*, 1996, 1997a).

Increasing n increases the power to detect QTLs, but the relative efficiency diminishes rapidly as n exceeds 4, especially at higher values of h^2_{RI} (Figs. 2 and 3). For example, the increase in the estimated equivalent F₂ N as n is increased from 4 to 10 (a 2.5-fold increase) is 1.7-fold (70%) at h^2_{RI}

= 0.1 but is only about 4% when $h_{RI}^2 = 0.8$. When $h_{RI}^2 = 0.3$ to 0.6, as is often the case in the behavioral literature, the increase in equivalent $F_2 N$ is only about 1.1- to 1.3-fold (10–30%) in response to the 2.5-fold (150%) increase in n (and N). Moreover, inspection of Figs. 2 and 3 shows that the added burden of testing 15 or more mice per strain, compared to only 6 or 10, for traits with high h_{RI}^2 (say, >0.5) is probably not economically justified. While increasing n under these conditions is not efficient in terms of animal numbers, perhaps a more important question is, Is it cost-effective? This question is discussed below.

Since QTL mapping is inherently a large-scale enterprise, the costs per trait are high and often beyond the resources of many laboratories. This, in turn, inhibits progress. For this reason, the study of relative costs of one experimental design vs. another for QTL detection is an especially important consideration. As an example of a cost analysis of F_2 vs. RI populations, data from our laboratory are presented in Table I. They roughly follow the cost efficiency analysis explicated by Sokal and Rohlf (1995).

We compared the cost of an individual RI mouse vs. an F_2 mouse for QTL detection using the cost structure shown in Table I. While all genotyped RI mice are also genotyped, this is generally not the case in segregating populations. A common practice to minimize F_2 genotyping costs is to employ conventional selective genotyping, where only individual mice at the extreme ends of the trait distribution are genotyped (Lander and Botstein, 1989; Darvasi and Soller, 1992). While effective in dramatically reducing costs, selective genotyping has several disadvantages that must also be considered. Mapping accuracy and power are somewhat reduced (Darvasi and Soller, 1992; Darvasi, 1997) and the newer and more powerful multiple regression-based QTL analyses, e.g., Jansen (1993), Zeng (1994), Manly and Cudmore (1996) and Basten *et al.* (1996), cannot be used. Also, the assessment of interactions among QTLs is weakened, as is the analysis of linked QTLs (Lin and Ritland, 1966). Finally, QTL results emerging from selective vs nonselective genotyping can, at times, be surprisingly different when the selection fraction is small, as observed, for example, by Gershenfeld *et al.* (1997) using a selection fraction of 0.12 (0.06 in each tail). This raises questions about whether a highly restricted sample is (1) increasing the sam-

pling error to serious levels or (2) magnifying the spurious effects of experimental artifacts that cause extreme scores for reasons unrelated (or poorly related) to genotype (phenocopies). Thus, there are several reasons to avoid selective genotyping, especially when the selection fraction is small.

Generally, whether RI populations are more cost-effective (less costly) than a comparable F_2 will depend on the relative efficiency ratio, or RE (which depends on h_{RI}^2 , n , and the selection fraction) and the relative cost ratio per mouse, or RC (which depends on the costs of genotyping relative to the other costs, and the selection fraction), all of which can vary widely from trait to trait and laboratory to laboratory. Darvasi and Soller (1992) discuss the cost implications of selective genotyping in the general case. For the cost data shown in Table I, the equicost value of h_{RI}^2 , when RI and F_2 populations of equal power are also equal in cost, was 0.65 for $n = 10$ and 0.90 for $n = 6$, when optimal (for cost) conventional selective genotyping was practiced (Fig. 4). Since most behavioral traits will have h_{RI}^2 values less than the equicost value, RI populations generally will be more cost-effective than F_2 populations with similar RC values to our example, even when selective genotyping is optimized to reduce cost. The cost advantage of RI over segregating populations is severalfold at low h_{RI}^2 and disappears as h_{RI}^2 reaches the equicost value. When $h_{RI}^2 = 0.4$ ($h_{F_2}^2 = 0.25$) and $n = 10$, for example, typical values in our experience, the cost advantage is just under twofold from Fig. 4 (the F_2 population is almost twice as costly) under the cost conditions shown in Table I, a major difference. This difference is even larger with smaller n ; for example, it is almost threefold when $n = 6$.

When selective genotyping and DNA pooling of each tail is used, equicost h_{RI}^2 is 0.55 ($h_{F_2}^2 = 0.38$) for $n = 6$ and 0.37 ($h_{F_2}^2 = 0.23$) for $n = 10$, values which fall in the range typically seen for behavioral traits. Thus, when DNA pooling is used under the cost conditions of our example (Table I), F_2 compared to RI populations will be more cost-effective (cost less) for traits with heritabilities above this equicost value and less cost-effective below them.

The conclusion drawn above concerning relative total cost strictly hold only for traits with a relative cost ratio per mouse (RC) similar to that used in our example (Table I). However, our example is typical of our experience with over 30 traits subjected to QTL analyses. Actual costs from

other laboratories can easily be substituted for the values shown in Table I to obtain more accurate cost evaluations for a particular experiment. (A MathCad worksheet is available from the author for this purpose.)

Throughout this paper, dominance variation in the F_2 has been ignored, since this source does not exist in the RIs. However, dominance provides another source of QTL information that can increase power to detect QTLs showing dominance, thus increasing F_2 power and cost-effectiveness. On the other hand, the opposite can occur for QTLs showing no dominance, because the assessment of both additive and dominance effects in the QTL analysis (e.g., MapMaker QTL) requires a twofold more stringent p value as the threshold for statistical significance than do additive effects alone (Lander and Schork, 1994), which effectively reduces the power and F_2 cost-effectiveness.

Overall, the use of RI populations to gain QTL information in behavioral studies has much to recommend it for cost as well as other reasons noted elsewhere (Bailey, 1981; Belknap *et al.*, 1996, 1997a; Plomin and McClearn, 1993). This is especially true if the within-strain sample size, n , is reasonably adjusted for the expected heritability (i.e., using smaller n when h^2_{RI} is high, and vice versa), which can greatly reduce RI costs relative to F_2 conferring equal power. However, of all the costs considered, those of genotyping are likely to be most affected by advances in technology, which will likely make F_2 and other segregating populations more attractive economically than they are at present.

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REFERENCES

- Bailey, D. W. (1981). Recombinant inbred strains and bilineal congenic strains. In Foster, H. L., Small, J. D., and Fox, J. G. (eds.), *The Mouse in Biomedical Research, Vol. I*, Academic Press, New York, pp. 223–239.
- Basten, C. J., Weir, B. S., and Zeng, Z.-B. (1996). *QTL Cartographer: A Reference Manual and Tutorial for QTL Mapping*, Department of Statistics, North Carolina State University, Raleigh.
- Belknap, J. K., Mitchell, S. R., O'Toole, L. A., Helms, M. L., and Crabbe, J. C. (1996). Type I and Type II error rates for quantitative trait loci (QTL) mapping studies using recombinant inbred mouse strains. *Behav. Genet.* **26**:149–160.
- Belknap, J. K., Dubay, C., Crabbe, J. C., and Buck, K. J. (1997a). Mapping quantitative trait loci for behavioral traits in the mouse. In Blum, K., and Noble, E. P. (eds.), *Handbook of Psychiatric Genetics*, CRC Press, Boca Raton, FL, Chap. 25.
- Belknap, J. K., Richards, S. P., O'Toole, L. A., Helms, M. L., and Phillips, T. J. (1997b). Short-term selective breeding as a tool for QTL mapping: Alcohol preference drinking in mice. *Behav. Genet.* **27**:55–66.
- Crabbe, J. C., Belknap, J. K., and Buck, K. J. (1994). Genetic animal models of alcohol and drug abuse. *Science* **264**:1715–1723.
- Darvasi, A. (1997). The effect of selective genotyping on QTL mapping accuracy. *Mammal. Genome* **8**:67–68.
- Darvasi, A., and Soller, M. (1992). Selective genotyping for determination of linkage between a marker locus and a quantitative trait locus. *Theor. Appl. Genet.* **85**:353–359.
- Darvasi, A., and Soller, M. (1994). Selective DNA pooling for determination of linkage between a molecular marker and a quantitative trait locus. *Genetics* **138**:1365–1373.
- DeFries, J. C., Wilson, J. R., Erwin, V. G., and Petersen, D. R. (1989). LS \times SS recombinant inbred strains of mice: initial characterization. *Alc. Clin. Exp. Res.* **13**:196–200.
- Falconer, D. S., and MacKay, T. F. C. (1996). *Introduction to Quantitative Genetics*, 4th ed., Longman, Essex, UK.
- Gershenfeld, H. K., Neumann, P. E., Mathis, C., Crawley, J. N., Li, X., and Paul, S. M. (1997). Mapping quantitative trait loci for open-field behavior in mice. *Behav. Genet.* **27**:201–210.
- Hegmann, J., and Possidente, B. (1981). Estimating genetic correlations from inbred strains. *Behav. Genet.* **11**:103–114.
- Hyde, J. S. (1973). Genetic homeostasis and behavior: Analysis, data and theory. *Behav. Genet.* **3**:233–245.
- Jansen, R. C. (1993). Interval mapping of multiple quantitative trait loci. *Genetics* **135**:205–211.
- Johnson, T. E., DeFries, J. C., and Markel, P. (1992). Mapping quantitative trait loci for behavioral traits in the mouse. *Behav. Genet.* **22**:635–653.
- Kearsey, M. J., and Pooni, H. S. (1996). *The Genetical Analysis of Quantitative Traits*, Chapman and Hall, London.
- Knapp, S. J., and Bridges, W. C. (1990). Using molecular markers to estimate quantitative trait locus parameters: Power and genetic variances for unreplicated and replicated progeny. *Genetics* **126**:769–777.
- Lander, E. S., and Schork, N. J. (1994). Genetic dissection of complex traits. *Science* **265**:2037–2048.
- Lander, E. S., and Botstein, D. (1989). Mapping mendelian factors underlying quantitative traits using RFLP linkage maps. *Genetics* **121**: 185–199.
- Lin, J.-Z., and Ritland, K. (1996). The effects of selective genotyping on estimates of proportion of recombination between linked quantitative trait loci. *Theor. Appl. Genet.* **93**:1261–1266.
- Manly, K. E., and Cudmore, R. (1996). *Map Manager QT: A Program for Genetic Mapping*, Roswell Park Cancer Institute, Buffalo, NY.
- Plomin, R., and McClearn, G. E. (1993). Quantitative trait loci analysis and alcohol-related behaviors. *Behav. Genet.* **23**: 197–212.
- Sokal, R. R., and Rohlf, F. J. (1995). *Biometry*, Freeman, San Francisco.

- Soller, M., and Beckmann, J. S. (1990). Marker-based mapping of quantitative trait loci using replicated progenies. *Theor. Appl. Genet.* **80**:205–208.
- Taylor, B. A. (1995). Recombinant inbred strains. In Lyon, M. F., and Searle, A. G. (eds.), *Genetic Variants and Strains of the Laboratory Mouse*, 3rd ed., Oxford University Press, Oxford, UK.
- Taylor, B. A., and Phillips, S. J. (1996). Detection of obesity QTLs on mouse chromosomes 1 and 7 by selective DNA pooling. *Genomics* **34**:389–398.
- Zeng, Z.-B. (1994). Precision mapping of quantitative trait loci. *Genetics* **136**:1457–1468.

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