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**QUESTION**

Write a concise review on the developmental genetics of the cerebellum and highlight the genetic bases of known cerebellar disorders.

DEVELOPMENTAL GENETICS OF THE CEREBELLUM

Developmental genetics is the study of how genes control growth and development of organisms

In this case however, developmental genetics of the cerebellum is the study of how genes control the growth and development of the cerebellum.

The mature cerebellum has exquisite, stereotypical morphology, foliation, and lamination, which are consistent between individuals and highly conserved across vertebrates. At the cellular level, unlike other regions of the CNS, the cerebellum is composed of very few neuronal types, each with distinct morphology, arranged in discrete lamina, and connected in stereotypical circuits. The cerebellum has essential roles in motor coordination, but is not essential for viability. Thus, compared with other regions of the central nervous system (CNS) the cerebellum has been more amenable to genetic studies since disruptions in development, which lead to abnormal morphology or function, are readily observed in obvious neurological and behavioral phenotypes. Because of this, it has been possible to obtain a precise understanding of cerebellar development.

To discover genes influencing cerebellum development, we conducted a complex trait analysis of variation in the size of the adult mouse cerebellum. We analyzed two sets of recombinant inbred BXD strains and an F2 intercross of the common inbred strains, C57BL/6J and DBA/2J. We measured cerebellar size as the weight or volume of fixed or histologically processed tissue. Among BXD recombinant inbred strains, the cerebellum averages 52 mg (12.4% of the brain) and ranges 18 mg in size. In F2 mice, the cerebellum averages 62 mg (12.9% of the brain) and ranges ∼20 mg in size. Five quantitative trait loci (QTLs) that significantly control variation in cerebellar size were mapped to chromosomes 1 (Cbs1a), 8 (Cbs8a), 14 (Cbs14a), and 19 (Cbs19a,Cbs19b). In combination, these QTLs can shift cerebellar size an appreciable 35% of the observed range. To assess regional genetic control of the cerebellum, we also measured the volume of the cell-rich, internal granule layer (IGL) in a set of BXD strains. The IGL ranges from 34 to 43% of total cerebellar volume. The QTLCbs8a is significantly linked to variation in IGL volume and is suggestively linked to variation in the number of cerebellar folia. The QTLs we have discovered are among the first loci shown to modulate the size and architecture of the adult mouse cerebellum.

The cerebellum is an ideal structure in which to study molecular and genetic networks controlling morphogenesis in the CNS. Its laminar cytoarchitecture and modular structure facilitate the study of developmental compartments (Herrup and Kuemerle, 1997), the effects of single-gene mutations (Goldowitz and Eisenman, 1992; Heintz and Zoghbi, 2000), and normal individual differences (Inouye and Oda, 1980; Neumann et al., 1990; Cooper et al., 1991; Wahlsten and Andison, 1991, 1993). The networks of genes that are critical in cerebellar development are beginning to be identified (Goldowitz and Hamre, 1998; Oberdick et al., 1998). For example, Yang et al. (1999) recently reported a convincing dependence on Zipro1 gene dosage for granule cell number and folial patterning in the mouse cerebellum. Relatively subtle quantitative effects on cerebellar size and foliation have also been well demonstrated for Engrailed-2 null mice (Joyner et al., 1991; Millen et al., 1994; Kuemerle et al., 1997). Null mutant, induced expression, or other transgenic technologies are indispensable for molecular dissection of cerebellar development. It remains unknown whether these (or other) early patterning genes are responsible for normal differences between inbred mice, wild mouse populations, or larger species differences. We might expect early patterning genes to be fixed within species and that later acting genes tolerate the polymorphisms that realize individual differences.

Recent advances in quantitative genetics allow neurogeneticists to systematically identify the polymorphic gene loci that, in concert with environmental influences, generate quantitative variation. This forward genetic approach, i.e., from trait to gene, initially involves associating differences in alleles near defined marker loci with differences in the relevant trait. A strong statistical association indicates the location and effect size of a quantitative trait locus (QTL) (Darvasi, 1998; Williams, 2000). In one of the first studies of gene loci affecting normal strain differences in the mouse CNS, Neumann et al. (1993) reported three loci that have quantitative effects on the number of cerebellar fissures in mice derived from the C57BL/6J and DBA/2J inbred strains. Most recently, in the NZB/BINJ and C57BL/6By inbred strains, LeRoy-Duflos (2001) identified seven loci affecting the presence or absence of the declival and intraculminate cerebellar fissures. Furthermore, LeRoy-Duflos (2001) colocalized one of these gene loci with a measure of motor coordination, illustrating a potentially powerful approach to investigate possible causal relationships between CNS structure and function. By taking advantage of differences between normal strains of mice, we have recently mapped gene loci that modulate numbers of neurons in mouse retina (Strom and Williams, 1998; Williams et al., 1998), olfactory bulb weight (Williams et al., 2001), and hippocampus weight and internal architecture (Lu et al., 2001).

In the present study we have initiated a neurogenetic analysis of the normal adult mouse cerebellum. Using experimental crosses of mice derived from C57BL/6J and DBA/2J inbred strains, we have started our analysis by first considering genes that affect total cerebellar size and the size of the cell-rich internal granule layer. We demonstrate that differences in cerebellar anatomy are generated in part by a set of five QTLs with relatively large effects that map to chromosomes 1, 8, 14, and 19.

MATERIALS AND METHODS

Mice. Three complementary sets of mice were used in this study (n = 514). These included two sets of BXD recombinant inbred (RI) strains and a set of F2 progeny. The RI strains and F2 progeny were generated from the inbred strains C57BL/6J (B for short) and DBA/2J (D for short). RI mice are inbred lines derived from brother–sister matings starting from an F2 intercross. In general, RI strains, when compared with F2 mice, provide lower power but increased mapping resolution for QTL detection. This is because RI sets are small (usually <30 strains) but harbor 3.5–4 times as many recombination events compared with F2 mice. Because they are inbred, RI lines also allow greater precision in phenotyping. The average of several mice can represent the mean value for a strain. The BXD RI strains were generated by Dr. B. A. Taylor (Taylor, 1989; Taylor et al., 1999), and purchased from The Jackson Laboratory (Bar Harbor, ME) from 1994 through 1999 (n = 334). The parental strains used to generate F2 mice were purchased from The Jackson Laboratory. The F2 animals were generated at the University of Tennessee (UT) by intercrossing both BDF1 and DBF1 mice as described in Zhou and Williams (1999). One hundred and five of these F2 mice were BDF2s and 75 were DBF2s. Mice were maintained at 20–24°C on a 14:10 on:off photoperiod in a pathogen-free colony at UT. Most animals were provided with tap water and a 5% fat Agway Prolab 3000 rat and mouse chow ad libitum. The average age of the BXD RI strains was ∼80 d (range, ∼30–300), and that of the F2 mice was 100 d (range, ∼70–160). Both females and males were studied.

Cerebellum. Mice were deeply anesthetized with Avertin (1.25% 2,2,2-tribromoethanol and 0.8% tert-pentyl alcohol in water; 0.8–1.0 ml, i.p.) and weighed to the nearest 0.1 gm. Most mice were perfused transcardially with 0.1 m PBS followed first by 1.25% glutaraldehyde and 1.0% paraformaldehyde in 0.1 m PBS and then by 2.5% glutaraldehyde and 2.0% paraformaldehyde in 0.1 m PBS.

Brains from BXD and F2 mice were dissected from the skull, sectioned free of the spinal cord and cranial nerves, rolled briefly on tissue paper, and weighed to the nearest 0.1 mg. All weights were recorded at room temperature. The cerebellum was dissected free of each brain by inserting a small knife parallel to the dorsal hindbrain surface to cut the peduncles. Two persons dissected cerebellar weights from BXD mice. Technical error was controlled before QTL mapping by using a covariate term (coded as a dummy or indicator variable).

In a second set of BXD mice, brain, cerebellum, and internal granule layer (IGL) volumes were measured from celloidin-embedded Nissl-stained sections available from the Mouse Brain Library. Brain and cerebellum volumes were estimated by summing the areas measured across sections and multiplying the sum by the sampling interval (0.30 mm). This method does not differ significantly from Cavalieri's method for the number of sections per mouse (>10) measured (Rosen and Harry, 1990). Measurements were made on slide images using NIH Image software (developed at the United States National Institutes of Health). The volume of the internal granule layer was measured using the density slice feature of NIH Image. As measured, this volume included the internal granule layer (predominantly the somas of the granule cells) and the adjoining Purkinje cell layer (somas). Cerebellum and IGL volumes were corrected for differences in tissue shrinkage by dividing by the total brain volume and multiplying by the brain volume expected from the known brain weight, assuming a uniform brain density of one milligram per cubic millimeter of fixed tissue.

Genotyping. Genomic DNA from F2 mice was extracted from the spleen using a high salt procedure (Laird et al., 1991). Spleens were removed under anesthesia just before perfusion. A set of 145 microsatellite loci distributed across all autosomes, and the X chromosome was typed in the F2 progeny using a modified protocol ofLove et al. (1990) and Dietrich et al. (1992). Each 10 μl PCR reaction contained 1× PCR buffer, 1.92 mmMgCl2, 0.25 U of Taq DNA polymerase, 0.2 mm of each deoxynucleotide, 132 nm of the primers, and 50 ng of genomic DNA. The microsatellite primers were purchased from Research Genetics (Huntsville, AL). A loading dye (60% sucrose, 1.0 mm cresol red) was added to the reaction before the PCR (Dietrich et al., 1994). PCRs were performed in 96 well microtiter plates. A high-stringency touchdown protocol was used to lower the annealing temperature progressively from 60 to 50°C in 2°C steps over the first six cycles (Don et al., 1991). After 30 cycles, PCR products were run on 2.5% Metaphor agarose gels (FMC Bioproducts, Rockland, ME), stained with ethidium bromide, and photographed. Genotypes were scored, entered into Microsoft Excel 98, formatted, and exported to Map Manager QT and Map Manager QTX (Manly and Olson, 1999). A nonredundant set of ∼320 loci typed for all 35 BXD RI strains was used for analysis, as described in Williams et al. (1998) and Zhou and Williams (1999).

Statistics. Regression analysis was performed to explore and remove (i.e., statistically control) covariance between cerebellum or IGL size and other variables before QTL mapping. The primary purpose of this analysis was to provide protection against claiming that a gene effect is specific to cerebellum or IGL size when the effect is more general, e.g., a whole brain or body effect. Our variables included cerebellum weight (in milligrams) or volume (in cubic millimeters), IGL volume (in cubic millimeters), brain weight (in milligrams) or volume (in cubic millimeters), body weight (grams), sex, and age (days). The regression analysis resulted in a set of new brain phenotypes based on the residuals from linear regression models; these “adjusted” phenotypes were used for gene mapping. A secondary purpose of the regression analysis was to understand relations between brain phenotypes and factors like sex or age. Regression models were computed with Data Desk 6.1. Interaction terms were explored in each model and excluded from final models when not significant. Alternatives to this regression analysis include multiple trait QTL analysis (Jiang and Zeng, 1995) or the use of covariates during single trait QTL analysis (Map Manager QT manual). We chose single trait QTL analysis because we were specifically interested in cerebellum QTLs. We chose to control variance related to factors like brain size, sex, and age before QTL mapping, because these factors are of independent biological interest and potentially confounding.

Map Manager QT and QTX programs (Manly and Olson, 1999) were used for QTL mapping. In essence, QTL mapping divides a panel of mice into groups based on their genotypes (e.g., BB orDD in the case of RI strains and BB,BD, and DD in the case of F2 progeny) at defined chromosomal loci and compares these groups using a quantitative phenotype (e.g., cerebellum weight). QTL mapping generally proceeds from analysis at defined loci (single marker analysis), to positions inferred between loci (simple interval mapping), and then to positions inferred between loci but with statistical control for other loci known to affect the trait (composite interval mapping). Interval mapping helps to better localize QTLs between markers. Composite interval mapping helps to understand the independent or incremental effects of QTLs. By controlling for multiple unlinked QTLs, the power to detect secondary QTLs may also be increased substantially. In our genome scans we first mapped our phenotype against genotype data on a locus-by-locus basis to assess linkage. Loci with log of the odds ratio (LOD) scores greater than ∼2.8 were considered suggestive (Lander and Kruglyak, 1995). Chromosomes harboring suggestive loci were analyzed by simple interval mapping. When more than one locus was identified, composite interval mapping was used. Map Manager QT and QTX implement simple and composite interval mapping using methods described by Haley and Knott (1992). Genome-wide (experimentwise) significance probabilities for mapped QTLs were estimated by comparing peak likelihood ratio statistics (LRSs) (LRS = LOD × 4.6; Liu, 1998) of correctly ordered data sets with those computed for 10,000 permutations of the phenotype data (Churchill and Doerge, 1994). Probabilities reported below as “genome-wide” (PG) reflect correction for multiple tests; other reported probabilities are comparisonwise. Confidence of QTL position is given as the 2-LOD support interval that bounds the QTL with ∼95% confidence (Lynch and Walsh, 1998).

RESULTS

Our results are divided into two parts. The first section describes the normative phenotypic measures and their relation to differences in sex, age, body weight, and brain weight. The second section presents evidence for QTLs modulating cerebellar size and structure.

Cerebellar size and variation

Parental cerebella

The cerebella of C57BL/6J mice are on average 18% larger (F(1,58) = 110; p < 0.0001) than those of DBA/2J mice: 59.6 ± 0.52 mg (n = 36; 15 male, 21 female) versus 50.5 ± 0.73 mg (n = 24; 11 male, 13 female). Male and female mice do not differ in cerebellar weight within these strains. Despite the significant 18% difference in absolute size, the cerebellum makes up ∼12% of total brain weight in both strains. This contrast highlights the importance of assessing specificity of gene effects or the need to determine whether loci affect cerebellar size directly or indirectly (e.g., through brain or body size).

BXD cerebella

The random assortment of multiple alleles at different loci will often generate much greater differences among RI strains than between the parental strains (Neumann et al., 1993; Lu et al., 2001; Williams et al., 2001). This is definitely the case for the cerebella of BXD strains. Weights range from a low of 45.9 ± 1.09 mg in BXD12 to a high of 62.5 ± 2.32 mg in BXD5 (Table 1). In a separate sample of BXD mice, cerebellar volumes range from 43.1 ± 1.21 mm3 in BXD23 to 62.9 ± 3.55 mm3 in BXD5. Strain distribution patterns for either measure of cerebellar size do not depart significantly from normality (Kolmogorov tests), arguing that the strain differences are polygenic in origin (cf. Williams et al., 1998). The correlation between strain means for cerebellar weight and volume is high and significant (r = 0.86; p < 0.0001). Using ANOVA with strain as the single factor, strain differences are significant for cerebellar weight (R2 = 67.5%;F(33,144) = 9.1; p < 0.0001) and volume (R2 = 51.9%, F(30,124) = 4.5;p < 0.0001). Using estimates of environmental and genetic influences on cerebellum size from within strain and between strain variances [0.5VA/(0.5VA+ VE)], the heritability is ∼47% and 33% for weight and volume, respectively (Hegmann and Possidente, 1981).

F2 cerebella

The average cerebellar weight of 180 F2 mice is 61.5 ± 0.33 mg. The distribution of cerebellar weight in F2 mice is bell-shaped and is not significantly different from a normal distribution (Kolmogorov test) (Fig. 2). Cerebellar weight in the F1 mice averages 62.1 ± 0.63 mg (n = 46). In a single-factor ANOVA with F1, F2, and parental mice, the parental strains differ from each other (Scheffe post hoc test;p < 0.0001), F1 mice differ from D mice (p < 0.0001) but not B mice (p = 0.10), and F1 and F2 mice do not differ significantly (p = 0.80). Using variances in the isogenic F1 and heterogeneous F2 samples, (VF2 −VF1)/VF2, heritability of cerebellum weight is 23%.

BXD and F2 differences

There are significant differences between the F2 and BXD mice: cerebellar weight is on average 10 mg heavier, brain weight is 45 mg heavier, and body weight is 6.7 gm heavier in F2 mice. The age and sex composition of our samples are also somewhat different. The F2 mice are on average 10 d older, but the age range is comparatively narrow (66–156 d vs 30–300 d). The F2 sample is 52% female, whereas the BXD sample is 44% female. The cerebellar weight of F2 cases are still significantly higher (4 mg) after adding covariates for brain weight, body weight, sex, and age (F(1,344) = 53; p < 0.0001). The increased size of F2 mice likely reflects heterosis of the non-inbred F2 progeny and F1 maternal superiority (Lynch and Walsh, 1998).

Cerebellar covariate analysis

Sex is not associated with differences in cerebellar size. When restricted to 19 BXD strains for which both sexes were replicated, no sex difference is evident (F(1,100) = 0.21; p = 0.65). Sex is also not an important predictor in either BXD cerebellar volume (F(1,88) = 0.14; p = 0.71; 18 strains) or F2 cerebellar weight (F(1,178) = 0.03; p = 0.86).

Increased body weight is weakly associated with larger cerebella. In BXD mice, body weight has a correlation of 0.36 with cerebellar weight (df = 162; p < 0.0001) and 0.34 with cerebellar volume (df = 148; p < 0.0001). In the F2 data set, the correlation is 0.19 (df = 177; p = 0.014).

Age is also weakly associated with cerebellar size. In BXD mice, cerebellar weight correlates 0.27 with age (df = 174;p = 0.0003; 30–300 d; log-transformed), and cerebellar volume correlates 0.19 with age (df = 150; p = 0.01). We did not detect any increase in cerebellar weight within the more restricted age sample of the F2 mice.

Brain size is strongly associated with cerebellar size. Cerebellar weight among BXD strains has a correlation of 0.70 with brain weight (df = 176; p < 0.0001); for volume the correlation is 0.53 (df = 153; p < 0.0001). Cerebellar weight in F2 mice has a correlation of 0.72 with brain weight (df = 178; p < 0.0001).

Adjusted cerebellar size

In multiple regression models including sex, body, age, and brain, only brain size is an important predictor of cerebellum for both BXD and F2 mice. Interactions are not significant. For these reasons, simple additive models including only brain weight or volume were used to create a set of adjusted measures for QTL mapping (Table 1 for adjusted BXD measures; Fig. 2 for adjusted F2 measures). Strain differences are still highly significant even after correction for differences in brain weight (R2= 80.3%; F(33,144) = 17.78;p < 0.0001) or brain volume (R2 = 45%;F(33,124) = 3.43; p < 0.0001).

Internal granule layer volume

The volume of the internal granule layer ranges from a low of 14.5 ± 0.57 mm3 in BXD20 to a high of 23.0 ± 0.20 mm3 in BXD5 (Table1). As a percentage of cerebellar volume, the IGL ranges from 33.6 ± 2.2% (BXD14) to 42.9 ± 2.9% (BXD22). Strain differences are again highly significant (R2 = 52%; F(30,124) = 4.42;p < 0.0001). Heritability for IGL volume is ∼31%. The correlation between strain means (an estimate of the genetic correlation) for IGL volume and the remaining cerebellum volume (molecular layer, internal white matter, and deep nuclei) is 0.62 (df = 29; p = 0.0002). This correlation remains significant after variance associated with brain volume is removed (r = 0.41; df = 29; p = 0.02).

IGL covariate analysis

Sex is not associated with differences in IGL volume. When restricted to 18 BXD strains for which both sexes are replicated, no sex difference is evident (F(1,88) = 0.27; p = 0.61). Males and females are also not different in the volume of the rest of the cerebellum (F(1,88) = 0.01; p = 0.91).

Body weight is weakly associated with IGL volume (r = 0.17; df = 148; p = 0.03), but more strongly associated with the rest of the cerebellum (r = 0.36; df = 148; p < 0.0001).

As with body weight, there is a dissociation between the effects of age on IGL volume and the volume of the rest of the cerebellum. Age (log-transformed) does not correlate with IGL volume (r= 0.05; df = 150; p = 0.50) but does correlate with the remaining cerebellar volume (r = 0.23; df = 150; p = 0.004).

Larger brains are positively associated with both greater IGL volume and the volume of the rest of the cerebellum. Brain volume is moderately correlated with IGL volume (r = 0.40; df = 153; p < 0.0001) and with the rest of the cerebellum (r = 0.46; df = 153; p< 0.0001).

Adjusted IGL volume

As was true for cerebellum size, analysis by multiple regression shows that only brain size significantly predicts IGL volume when sex, body, and age covariates are included. Interactions are not important predictors. A simple additive model including only brain volume was again used to create a set of adjusted IGL measures (Table 1). In contrast, the volume of the remainder of the cerebellum is predicted better by two factors: brain volume (t1,147 = 4.69;p < 0.0001) and body weight (t1,147 = 2.50; p = 0.01). Strain differences are significant for adjusted IGL volume (R2 = 48%;F(30,124) = 3.8; p < 0.0001) and marginal for the adjusted cerebellar white matter (R2 = 28%;F(30,119) = 1.6; p = 0.05).

QTL mapping

QTLs affecting cerebellar weight in BXD mice

Single marker analysis shows that variation in adjusted cerebellar weight is associated with the pattern of B and Dalleles at several microsatellite markers, but linkages to mid-distal chromosome (Chr) 8 and distal Chr 1 appear strongest. Simple interval mapping of Chr 8 shows the tightest linkage (LRS = 22.5) to be just distal of D8Mit312 at 45 cM. At this locus, interval mapping estimates 47% of the variance in cerebellar weight is explained, and each B allele is associated with a decrease of 2.5 mg (Fig. 3A). The genome-wide significance for this linkage isPG = 0.0006. With control forD8Mit312, composite interval mapping reveals a secondary QTL near D1Mit150 at 100 cM on Chr 1 (LRS = 15;PG = 0.06). At this locus, 18% of the variance in cerebellar weight is independently explained, and eachB allele is associated with a decrease of 1.3 mg (Fig.3C). Collectively, these QTLs control 60% of the variance in adjusted cerebellar weight among BXD strains. A 6.4 mg difference in weight characterizes BXD strains with BB (n= 9) or DD (n = 9) alleles at both loci

DISCUSSION

Synopsis

We have discovered five new QTLs that are the among first normal gene variants known to modulate the size of the vertebrate cerebellum. We have named these QTLs Cbs1a, Cbs8a,Cbs14a, Cbs19a, and Cbs19b. Among BXD recombinant inbred strains, the cerebellum averages 52 mg (12.4% of the brain) and ranges 18 mg in size. The two quantitative trait loci on Chr 1 (2-LOD support interval 89–102 cM) and Chr 8 (44–53 cM), are responsible for ∼6 mg of this range, a 33% effect size. In C57BL/6J × DBA/2J F2 mice, the cerebellum averages 62 mg (12.9% of the brain) and ranges in size ∼20 mg. Four QTLs on Chrs 1 (81–92 cM), 14 (5–16 cM), and 19 (28–56 cM, 4–21 cM) are responsible for ∼7 mg of this range, a 35% effect size. In mapping these QTLs, the effects of brain size, sex, age, and body weight on cerebellar size were examined before mapping and statistically controlled when necessary. Apart from brain size, which accounted for ∼50% of cerebellar weight, we detected no discernable, independent effects on total cerebellar size for sex, age, or body weight.

Regional cerebellar effects of Cbs8a

Regional effects of cerebellar size QTLs were explored in BXD strains by examining the volume of the cell-rich internal granule layer versus the remaining volume. Our measurement of the IGL contained the somas of two of the main cell types in the cerebellum, the Purkinje, and the granule neurons. The remaining cerebellar volume included the molecular layer where granule and Purkinje cells synapse, the internal white matter, and the deep cerebellar nuclei. Although the IGL and remaining cerebellum are genetically correlated, we found that age and body weight correlated differently with these parts of the cerebellum, and we found a difference in the strength of linkage to the Chr 8 locus Cbs8a. Age and body weight correlated more strongly with the aggregate volume of the molecular layer, internal white matter, and deep nuclei than with the volume of the internal granule layer. BXD strain differences in internal granule layer volume are more strongly associated with genotypes at Cbs8a(PG = 0.08) than the volume of the remaining cerebellum (PG = 0.53). This suggests that the main effect of Cbs8a is on cell population size. This is consistent with evidence for linkage betweenCbs8a and the number of fissures, in which Dalleles that are associated with increased IGL volume are also associated with greater numbers of fissures. Although mechanisms of fissure formation are not well understood, a role for the proliferation of granule cells in fissure formation has been suggested (Mares and Lodin, 1970), and it has been noted that overall size of the cerebellum accounts significantly for fissure number (Wahlsten and Andison, 1991). Over expression of Zipro1 results in increased granule cell proliferation and increased numbers of intralobular fissures (Yang et al., 1999). Although Cbs8a is not Zipro1 (which resides on Chr 5), it is tempting to postulate that it may be a downstream target of Zipro1 or part of a network in whichZipro1 is involved.

Specificity of QTL effects

A difficult problem in correlational analyses is specificity of causation. For instance, if we mapped cerebellum size without considering the effects of total brain size, we could map genes that do not specifically affect the cerebellum. In the present study we used residuals from linear regression models that included brain size covariates. This improves our ability to map genes with selective effects on the cerebellum. Another approach to the problem of specificity is to gather many phenotypes from the same mice. For instance, in two companion studies we report QTLs in BXD and BDF2 mice for olfactory bulb (Williams et al., 2001) and hippocampus (Lu et al., 2001). QTLs for olfactory bulb weight were discovered on chromosomes 4, 6, 11, and 17. QTLs for hippocampus weight were discovered on chromosomes 1 and 5. Bulb4a on Chr 4 and Hipp1aon Chr 1 map within the same 2-LOD support intervals as the putative linkage of cerebellar white matter volume to Chr 4 and the linkage ofCbs1a to Chr 1, respectively. The linkage ofCbs1a distal Chr 1 reported here is independent of that reported for hippocampus at Hipp1a by Lu et al. (2001); a linear model predicting F2 weight of the cerebellum fromD1Mit57 remains significant with hippocampus weight used as a covariate (data not shown). Yet another approach to the problem of specificity is to gather more refined phenotypes related to the primary phenotype, as Lu et al. (2001) did for hippocampal volume, and as we have done for cerebellar volume. Lu et al. (2001) reported thatHipp1a exhibits shared effects on the volume of the hippocampal complex, the hippocampus proper, the pyramidal cell layer, and the granule cell layer. Hipp1a was also observed to overlap a previously mapped locus affecting the volume of the mossy fiber projection to CA3 from granule cells (Lassalle et al., 1999). In a similar way, by remapping cerebellar folial pattern in BXD strains (Neumann et al., 1993) with a more complete marker set, not only did we see the linkages previously reported by Neumann et al. (1993) on Chrs 5, 7, and 11, but we also discovered thatCbs8a is linked to the number of fissures (LRS = 8.3;p = 0.005), with D alleles adding fissures. Thus, DBA/2J alleles at Cbs8a increase cerebellar weight, cerebellar volume, internal granule layer volume, and the number of cerebellar folia in BXD mice. These observations illustrate one of the advantages of recombinant inbred strains: data are cumulative over time and across laboratories. However, they also illustrate the problem of specificity of QTL action and highlight a need for improved mapping resources and methods. QTL fine mapping (Darvasi, 1998), perhaps in combination with multiple trait QTL mapping methods (Jiang and Zeng, 1995), may help significantly in this regard.

RI strains, F2 intercrosses, and advanced intercrosses

In the present study we used recombinant inbred strains derived from the parental strains C57BL/6J and DBA/2J as well as the F2 intercross progeny. Because both RI and F2 mice used in this study were derived from the same parental strains, the same alleles are assumed to be segregating in each mapping panel. However, we detected significant differences between the 34 BXD strains and our F2 progeny. The F2 progeny had heavier bodies, brains, and cerebella. Such effects probably reflect heterosis (hybrid vigor). Possibly related to these phenotypic differences, two of the five QTLs we discovered, those on Chr 8 and Chr 14, were not detected in both RI and F2 mapping panels. However, Cbs8a finds support in two separate samples of BXD mice measured for both cerebellar weight and volume. Loci that are not replicated may also reflect the random assortment of alleles in relatively small samples or a mutation that has occurred in either the C57BL/6J or DBA/2J genomes since creation of the BXD strains in the 1970s and 1980s. We are currently investigating a third mapping panel derived from C57BL/6J and DBA/2J strains. This is an advanced intercross, created by outbreeding F2 mice. Examining cerebellar size and IGL volume in this panel will allow an additional opportunity to test the QTLs we have reported here and should reduce significantly the 2-LOD support intervals of confirmed QTLs (Darvasi, 1998). Fine mapping of Cbs8a can alternatively proceed using a recombinant inbred segregation test (Darvasi, 1998).

Candidate genes

The goal of our QTL mapping is to identify genes influencing cerebellar development. Once linkage is established, further study can proceed in three directions that complement the main goal of complex trait analysis (Moisan, 1999; Williams, 2000). First, mapping can continue with panels designed to reduce the interval containing the QTL, such as with advanced intercross or congenic lines (high-resolution mapping). Second, the effects of variants in identified genes near the QTL can be assessed (candidate gene analysis). Third, the phenotype associated with a QTL can be more rigorously defined or examined during development. For example, in a developmental analysis Strom and Williams (1998) established that a QTL that modulates retinal ganglion cell number acts through proliferation rather than cell death, thereby effectively eliminating the need to examine candidate genes that engage apoptotic mechanisms. The 2-LOD support intervals of the QTLs we report here are not yet sufficiently precise for effective candidate gene analysis, but we highlight several genes with identified effects on the cerebellum lie within these intervals.

On chromosome 1, the dreher mutation (dr; 88.2 cM) is known to have drastic effects on the cerebellum (Sekiguchi et al., 1992). Recent work reveals dr to be a mutation of the LIM homeodoman protein Lmx1a, which disrupts development of the roof plate, resulting in part in the loss of granule neurons in the cerebellum (Millonig et al., 2000). Also on Chr 1 is the retinoid × receptor gene (Rxrg; 88.1 cM). Retinoic acid receptors are expressed in the cerebellum of the rat, and granule cell loss results from teratogenic effects of the ligand (Yamamoto et al., 1999).

On chromosome 8, four of the cadherin (Cdh) genes reside within the 2-LOD support interval for Cbs8a(Cdh5, 51 cM; Cdh8, 46.5 cM; Cdh11, 46.5 cM; Cdh16, 50.0 cM). Cadherins are cell adhesion molecules with widespread morphogenetic effects and are thought to be important in regionalization of the CNS. Cadherin 8 is expressed widely in the developing mouse brain and shows a patterned distribution in the cerebellum (Korematsu and Redies, 1997; Korematsu et al., 1998). Also on chromosome 8, Cerebellin1 (Kavety et al., 1994), is highly enriched in the cerebellum (Urade et al., 1991; Pang et al., 2000), and its levels are decreased in mice that exhibit a loss of cerebellar granule neurons (Slemmon et al., 1988).

On chromosome 14, both the gene for bone morphogenetic protein 4 (Bmp4; 14 cM) and for bone morphogenetic protein receptor 1A (Bmpr1a; 13 cM) reside within the 2-LOD interval forCbs14a. Work by Alder et al. (1999) suggests that bone morphogenetic proteins can initiate granule cell genesis, and work byIantosca et al. (1999) suggests that Bmp4 may regulate granule cell number through inhibition of apoptosis.

Finally, on chromosome 19, both genes for fibroblast growth factor 8 (Fgf8; 45 cM) and Pax2 (43 cM) reside within the 2-LOD interval for Cbs19a. Both genes are critical to the early development of the cerebellum (Goldowitz and Hamre, 1998). Favor et al. (1996) and Urbanek et al. (1997) showed that inactivation ofPax2 can lead to loss of the cerebellum and posterior midbrain. Liu et al. (1999) have more recently showed thatFgf8 regulates the cascade of genes that transform the hindbrain into cerebellar structures.

The confidence intervals for the QTLs we have discovered are homologous to the following chromosomal regions in humans: Cbs1a: 1q23–43, Cbs8a: 16q12–16q22, Cbs14a: 10q11–23,Cbs19a: 9q13-q24 and 11q12-q13, Cbs19b: 10q23-qter. The conserved sequence similarity among mammals makes the likelihood high that identification of genes that influence cerebellar development in mice will aid discovery of homologous genes in humans.

GENETIC BASIS OF KNOWN CEREBELLAR DISORDERS

Cerebellar malformations are now widely diagnosed during pregnancy and associated with significant morbidity and mortality in the newborn period and throughout life. Given the broad range in prognosis and associated medical concerns, a working knowledge of these disorders and their causes is essential for obstetricians, perinatologists and neonatologists. For clinicians, it is most relevant to organize cerebellar malformations by their clinical and imaging features, which then directs additional diagnostic testing, medical monitoring for associated complications, and counseling about prognosis, treatment and recurrence risk. Distinguishing genetic disorders from similar conditions caused by extrinsic factors, such as infection, stroke, or prematurity, is particularly important to provide quality patient care. Cerebellar malformations may be classified as predominantly involving the cerebellum or involving both the cerebellum and brainstem. They may occur in isolation or as part of broader syndromes involving multiple systems. Though the cerebellum has long been recognized for its role in motor co-ordination, it also shapes the functions of other brain regions, especially cognition and affect, by processing external sensory and internally generated information to influence neocortical circuit refinement. Thus, not surprisingly, most cerebellar malformations are associated with neurodevelopmental issues affecting multiple domains: motor, communication, cognition, emotional regulation, and executive function. Human cerebellar development begins around the ninth gestational week and continues beyond birth. This protracted developmental timeline makes the human cerebellum particularly vulnerable to insult, especially during 24–40 weeks of gestation, when considerable neurogenesis in the external granule cell layer results in a five-fold increase in cerebellar size. Malformations that arise early in development typically affect both cerebellum and brainstem, whereas, later in development, cerebellar malformations have less effect on the pons. Here we present some of the most frequently occurring and best understood human cerebellar malformations and their genetic causes.

**Specific malformations with known genetic causes**

Predominantly cerebellar malformations

A malformed cerebellum may be abnormally small, dysplastic, or unusually large. The vermis and both hemispheres may be equally or disproportionately affected. Primary malformations of the pons, midbrain, and supratentorial structures are also seen in a substantial subset of patients. The wide range in morphological presentations results from the diversity of causes, including chromosomal abnormalities, specific genetic syndromes, and extrinsic factors. Dandy–Walker malformation Dandy–Walker malformation (MIM 220200) is a heterogeneous disorder defined by a hypoplastic, upwardly rotated vermis, an enlarged fourth ventricle, and an enlarged posterior fossa with an elevated confluence of sinuses. Typically, the cerebellar hemispheres are less affected than the vermis, and the brainstem is normal to moderately hypoplastic. DWM can occur with additional brain abnormalities including agenesis of the corpus callosum (ACC) and hydrocephalus, but more often it occurs as an isolated brain-imaging finding. The clinical features and developmental outcomes vary widely. Patients may exhibit symptoms ranging from intellectual disability to autism or they may be completely unaware of any deficits until diagnosed as adults for unrelated reasons. The recurrence risk in isolated DWM is low at an estimated 1–5%, suggesting de-novo, somatic mosaic, or complex genetic causes.

Few genes have been implicated in rare cases of DWM, including genomic imbalances that are part of a congenital syndrome and rare single gene disorders [5–8]. FOXC1-related DWM is associated with multiple congenital anomalies, especially eye malformations consistent with Axenfeld–Rieger syndrome [6]. Congenital anomalies associated with FOXC1-related DWM in severely affected patients overlap with Ritscher–Schinzel, or 3C (cranio-cerebello-cardiac) syndrome. Recently, mutations in CCDC22 were found in X-linked cases of 3C syndrome, suggesting that CCDC22 mutations may be a new cause of DWM. Though these patients were noted to have DWM, limited neuroimaging data were reported to substantiate this diagnosis. ZIC1/4-related DWM is also associated with multiple congenital anomalies, including dysmorphic facial features and abnormal development of the eyelids. Recently, exome sequencing identified autosomal dominant mutations in LAMC1 and NID1 as the cause of DWM with encephalocele in two families. Despite these genetic advances, the genetic cause remains unknown in the majority of DWM patients.

Case reports and small case-series also suggest that extrinsic factors may contribute to DWM. For example, serial fetal prenatal neuroimaging identified evidence of prenatal hemorrhage above the cerebellum in a patient postnatally diagnosed with DWM. Two comprehensive population-based studies suggest that clomiphene citrate exposure and twinning may be additional non-genetic risk factors for DWM. Additional rigorous studies are needed to assess prenatal risk factors for DWM.

Cerebellar hypoplasia

Cerebellar hypoplasia (CH) refers to an underdevelopment of the cerebellum. This category of cerebellar malformation is distinct from DWM in that it does not involve a concurrent enlargement of the posterior fossa, and almost all individuals exhibit cognitive and motor impairments. CH is a feature of many different disorders and it is often a non-specific feature associated with genomic imbalances. CH is frequently associated with additional brain abnormalities, including lissencephaly, cortical dysplasia, microcephaly and heterotopia, pointing to specific genetic causes.

The lissencephaly spectrum of brain malformations is caused by defects in either the reelin pathway or microtubule formation and function. Patients with autosomal recessive mutations in RELN have pachygyria and an extremely hypoplastic cerebellum with very little foliation and disproportionate effects on the vermis. Additional clinical features include profound developmental disability, microcephaly, sloping forehead, seizures, and congenital lymphedema. In contrast, patients with autosomal recessive mutations in VLDLR, a reelin receptor, exhibit mild pachygyria and a mildly small cerebellum that retains some foliation. Reelin is transiently expressed by neurons within superficial layers in both the cerebral cortex and cerebellum and regulates radial neuronal migration.

Mutations in the alpha- and beta-tubulins are a major cause of brain malformations, especially lissencephaly, pachygyria, and polymicrogyria, collectively referred to as tubulinopathies. Mutations in TUBA1A, TUBA8, TUBB2A, TUBB2B, TUBB3, TUBB4A, and TUBB are associated with a range of clinical features from isolated congenital fibrosis of the extraocular muscles to severe intellectual disability, quadriplegic cerebral palsy, seizures, cranial neuropathies and hydrocephalus. Neuroimaging features include cortical dysgenesis (lissencephaly or polymicrogyria), malformation of cranial nerves, and basal ganglia dysplasia, often with cerebellar and pontine hypoplasia, and defects in the corpus callosum, anterior commissure and internal capsule. Dysmorphic features are infrequently reported, and, surprisingly, other organ systems are not affected. Most occurrences are sporadic and due to de-novo mutations, but rare recurrences have been reported due to germline mosaicism and autosomal recessive inheritance.

Additional genes and extrinsic exposures have been associated with CH. Mutations in CHD7 account for the majority of patients with CHARGE syndrome (MIM-214800), characterized by Coloboma, Heart defects, choanal Atresia, Retardation of growth and development, Genital and Ear abnormalities. Neuroimaging reveals that more than a third of these patients have a slightly rotated, mildly to moderately hypoplastic cerebellar vermis, a mildly enlarged posterior fossa, and abnormal cerebellar hemisphere foliation. CHD7 loss-of-function disrupts critical gene expression in the isthmus organizer, a transient embryonic structure that directs early cerebellar development. Mutations in OPHN1 are associated with CH, ventriculomegaly, intellectual disability, seizures and mildly dysmorphic facial features that are characteristic in males and occasionally in females. Retinoic acid exposure during early pregnancy has been associated with severe cerebellar vermis hypoplasia with variable effects on the hemispheres. Additional clinical features include microtia, micrognathia, cleft palate, conotruncal heart defects, thymic defects, and retinal abnormalities.

Cerebellar hyperplasia

Cerebellar hyperplasia, or macrocerebellum, is a rare neuroimaging finding that occurs in isolation, or is coincident with a variety of neurodevelopmental disorders, including genomic imbalances and specific overgrowth syndromes, including megalencephaly–capillary malformation (MCAP) and megalencephaly–polydactyly–polymicrogyria–hydrocephalus (MPPH) (reviewed by Poretti et al. [19]). Recently, significant progress has been made in understanding the pathophysiology that leads to brain overgrowth in MCAP/MPPH. Clinical features of MCAP/MPPH include seizures, capillary malformations, macrocephaly, and polydactyly, and diagnosis is often based on cerebral cortical overgrowth and polymicrogyria. Though cerebellar size is normal at birth, patients often develop a macrocerebellum with normal posterior fossa size. This may progress into cerebellar ectopia/Chiari I malformation, causing clinically associated symptoms (posterior headache, dysphagia, stridor) and hydrocephalus. It is unknown whether the cerebellar overgrowth is a feature of generalized brain overgrowth, or whether there are distinct mechanisms that specifically influence cerebellar overgrowth. Most MCAP and MPPH patients have activating de-novo mutations in PIK3R2 and PIK3CA, respectively, that result in increased cell growt. Mutations affecting the PI3K–AKT–mTOR pathway are found in a variety of cancers, opening the possibility of using drugs in development for cancer treatment to reduce brain overgrowth and neurological issues in patients with MCAP/MPPH.

 Cerebellar dysplasia

Any part of the cerebellum can be dysplastic, from small focal regions within one hemisphere to abnormal foliation throughout the cerebellum. Hypoplastic cerebella are frequently also dysmorphic, as observed in tubulinopathies and cobblestone malformations. Chudley–McCullough syndrome (CMS; MIM 604213) is an autosomal recessive disorder in which patients have striking disorganization of the inferior cerebellar hemisphere folia and additional brain abnormalities, including frontal polymicrogyria with subcortical heterotopia, corpus callosum hypogenesis, and arachnoid cysts. Clinical presentation includes severe neonatal sensorineural hearing loss and hydrocephalus that may require shunting in some patients. Surprisingly, these patients typically are not dysmorphic, lack additional congenital anomalies, and have relatively mild developmental problems. CMS is caused by biallelic truncating mutations in GPSM2 that encodes a GTPase regulator required for correct orientation of stem cell divisions in multiple tissues. The cerebellar dysplasia present in CMS is likely due to abnormal cell division, but the precise mechanism remains unknown. Poretti–Boltshauser syndrome (PBS; MIM 150320) is characterized by cerebellar dysplasia, cysts, and vermis hypoplasia with and without retinal dystrophy and is caused by mutations in LAMA1. The superior cerebellar peduncles are long in some patients, though their appearance differs from the classic molar-tooth appearance that defines JS. The clinical features of PBS include motor and speech delay with variable cognitive impact. Finally, cerebellar dysplasia with cysts is a prominent feature in cobblestone malformations and GPR56-related brain malformations.

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