



Structured latent growth curves for twin data

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We describe methods to fit structured latent growth curves to data from MZ and DZ twins. The well-known Gompertz, logistic and exponential curves may be written as a function of three components – asymptote, initial value, and rate of change. These components are allowed to vary and covary within individuals in a structured latent growth model. Such models are highly economical, requiring a small number of parameters to describe covariation across many occasions of measurement. We extend these methods to analyse longitudinal data from MZ and DZ twins and focus on the estimation of genetic and environmental variation and covariation in each of the asymptote, initial and rate of growth factors. For illustration, the models are fitted to longitudinal Bayley Infant Mental Development Scale data published by McArdle (1986). In these data, all three components of growth appear strongly familial with the majority of variance associated with the shared environment; differences between the models were not great. Occasion-specific residual factors not associated with the curve components account for approximately 40% of variance of which a significant proportion is additive genetic. Though the growth curve model fit less well than some others, they make restrictive, falsifiable predictions about the mean, variance and twin covariance of other (not yet measured) occasions of measurement. *Twin Research* (2000) 3, 165–177.

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Introduction

Growth and decay are essential properties of living organisms. An improved understanding of the mechanisms of development seems fundamental to explaining individual differences in almost all biological science, and behavior genetics is no exception. In this paper, we describe, extend and apply growth curve models which predict changes in mean, variance, and covariance over time. These curve models can predict non-monotonic increases and decreases in means and covariances, and the two types of statistics do not necessarily change in parallel.

Growth curve models are well-suited to data where the number of occasions of measurement is large, because they describe changes in mean, variance and covariance with a limited number of parameters. We begin with a review of the conceptual background to the curves, and then show how they may be used to derive an appropriate set of factor loadings for model-fitting. Our methods are based on those previously published by Browne¹ and Browne and Du Toit.² We extend them to model genetic and environmental components of variation

in growth curves, using data from relatives, and provide Mx scripts to make the methods readily available.

At present, behavior genetic analyses of longitudinal data usually employ one of a few basic models: a) the Cholesky factorization; b) a Markov chain; or c) growth curves. As described by Neale and Cardon,³ the Cholesky factor model, also known as the square root factorization,^{4,5} (see Figure 1a) is a transformation of a covariance matrix. This model, presented as a path diagram in Figure 1a, has the same number of parameters as there are covariances ($m(m + 1)/2$ for m variables), and will always fit perfectly. In a genetic Cholesky model, the additive genetic (A), common (C) or specific (E) environmental covariance matrices are each decomposed into their Cholesky factors. While they describe the covariance of each of the components perfectly, the Cholesky factor model may fail because the within-person covariances are estimated from four different sources in the classical twin model, and these replicate statistics may be unstable. Similarly, the cross-twin cross-variable covariances may be non-symmetric and therefore contribute to lack of fit of the Cholesky. Such statistical fluctuations of the data do not reflect the appropriateness of the Cholesky as a model. Other causes of failure – such as DZ covariances being less than half the corresponding MZ covariances, or phenotypic variance differences between

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MZ and DZ twins – would be informative about the general suitability of the ACE model for the data, but still do not test the number of factors or the triangular pattern of their loadings.

In contrast to its use in the multivariate analysis of a set of variables, the Cholesky decomposition may have a useful conceptual interpretation when applied to longitudinal data which are ordered from time 1 to time m . All factors are constrained to impact variation at current and later – but not earlier – occasions. Thus the first factor ($F1$) may create

variation at all occasions, but $F2$ may influence all occasions *except* the first. Just as the model may predict any change in variation and covariation over time, so it can predict any pattern of change in means. However, this explanatory power can be seen as a disadvantage, because the model is not falsifiable. No pattern of genetic covariances over time exists that could not be accounted for by the Cholesky model. Even worse, it makes no prediction about genetic variation or covariation on future, not yet measured, occasions.

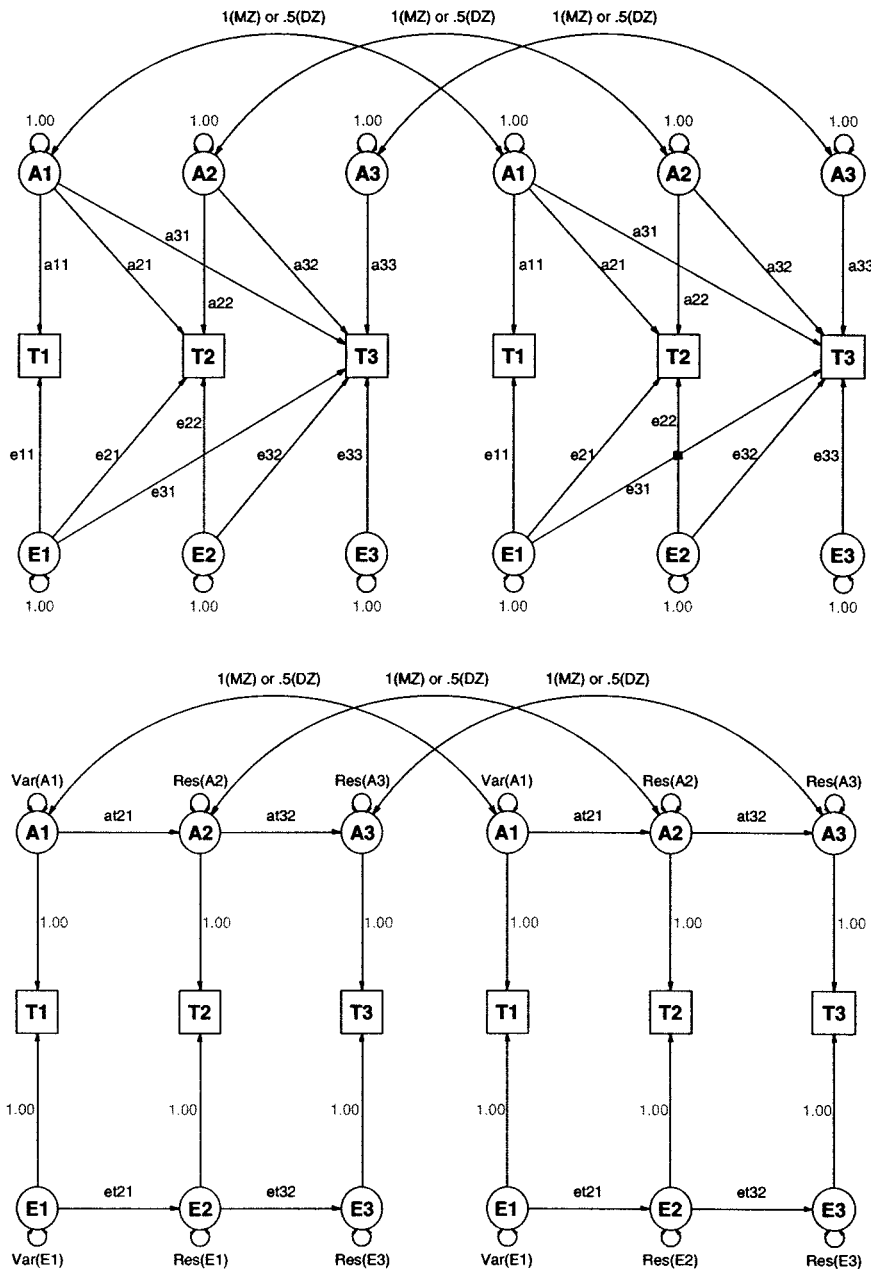


Figure 1 Two path models for longitudinal data from an individual: (top) Cholesky factor; and (bottom) Simplex or Markov

The ‘Simplex’ or ‘Markov chain’ model of Figure 1b provides an alternative account of changes in variance over time. Dolan and colleagues^{6,7} describe a constrained version of this model which specifies a linear relationship between longitudinal changes in variance and concurrent changes in mean. In contrast to the Cholesky factor model, the simplex model makes highly restrictive predictions about covariances and hence is falsifiable with sufficient occasions.⁸ Differences in mean and variance (either at the phenotypic or genetic factor level) may be related in a non-linear fashion, and the linear model may fail. For example, a decrease in variance might be accompanied by an increase in mean on one occasion, while at other times both may increase or decrease. Certain growth curve models predict more complex patterns of relationship between mean, variance and covariance, and they may be empirically tested against simpler models; hence our interest in them.

A popular, modern approach to the study and modeling of change is to use dynamical systems theory.^{1,9} Essentially these methods focus on the rate of change of a variable of interest (ie its slope or partial derivative) as a way to predict the level at a series of points in time. These methods have many applications, including pollution levels, disease epidemics, supermarket queues, population growth, radioactive decay, weather and chaotic systems. In this paper we relate structural equation modeling of genetically informative longitudinal data to dynamical systems.

A growth curve approach to behavioral genetic analysis was introduced by Vandenberg and Falkner¹⁰ who first fitted polynomial growth curves for each subject and then estimated heritabilities of the components. The latent variable growth model version of this idea was later presented by McArdle.¹¹ Here we extend models of this type by incorporating structured factor loadings that represent elementary theories about growth and change. In every case, we model the genetic and environmental sources of variance covariance in the latent growth components. This approach is analogous to using the phenotypic factors (common pathway) model rather than the biometric factors (independent pathway) model.^{3,12} The latter framework, though appealing, cannot be identified for growth curves because there is no predicted mean difference between MZ and DZ twins. In this respect, the growth curve models are quite different from the Simplex and Cholesky which are variants of biometric factor models.

Growth curves

First we present the theoretical background behind three growth curves which display asymptotic

behavior: the exponential, the logistic, and the gompertz. The basic idea behind these curves is that individuals start at some initial point, then grow at a rate which accelerates. This exponential growth characterizes the early stages of development, where the gradient gets steeper and steeper. However, it also seems reasonable to assume that growth does not continue to accelerate forever; some limiting factor takes over, progressively slowing down growth, making the gradient less and less steep until the growth curve asymptotes at an equilibrium point. Different hypotheses about the type of growth and limiting factors lead to the different growth curves, and some of these will be discussed below.

The exponential curve

One of the simplest models for changing growth rate is where current growth rate is proportional to current size (see Malthus (1798) as reported by Murray¹³). If we were to plot size against time, the gradient would get steeper and steeper (see Figure 2). Mathematically, we can write this concept as

$$\frac{dy}{dt} = y(t)c, \tag{1}$$

where c is a constant and $y(t)$ is the size at time t . This differential equation involves both y and the derivative of y over time t . Some differential equations, including this one, can be solved to find an expression for $y(t)$ by itself:

$$y(t) = y(0)e^{ct}. \tag{2}$$

Braun⁹ gives a good account of methods for solving such equations, and symbolic calculus software such as Mathematica¹⁴ may be used to verify results. Note how $y(t)$ increases exponentially over time from the

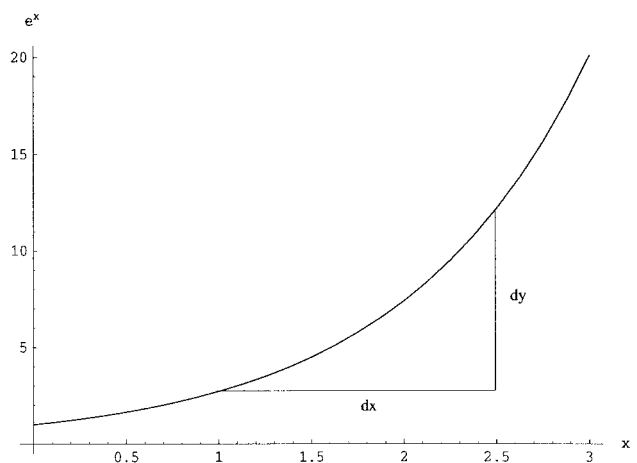


Figure 2 Plot of $y = e^x$ illustrating perpetual acceleration of growth for a system where growth rate is proportional to current size, dy and dx illustrate the derivative of the curve dy/dx

initial period $y(0)$ at time zero. While such growth may be a good approximation during initial stages of many biological systems, it rarely applies to the later ones, so we need to add a 'carrying capacity' to represent an upper limit on size.

One form of asymptomatic behavior arises when the growth rate is proportional to the distance from the maximum height or asymptote ($a = y(\infty)$). In this case the differential may be written:

$$\frac{dy}{dt} = c(a - y(t)), \tag{3}$$

which may be solved to give

$$y(t) = a - (a - y(0))e^{-ct}. \tag{4}$$

Note how this result scales the growth curve to lie between the initial value, $y(0)$, and the asymptote, a . When t is zero, the exponential part gives $e^0 = 1$ so that $a - (a - y(0)) = y(0)$. Conversely, when t is large, $e^{-\infty} = 0$, giving $y(\infty) = a$. This meets the requirement for asymptotic behavior, as illustrated by the solid line in Figure 3.

The logistic curve

Another model for limited growth was proposed by Verhulst in 1838,¹⁵ for changes in population size. He suggested that populations would grow in the exponential fashion of equation 1 above, but would be limited by the square of their size. Thus,

$$\frac{dy}{dt} = ay(t) - by(t)^2 \tag{5}$$

$$= ay(t)(1 - y(t)b) \tag{6}$$

where a is the ultimate size or asymptote, and b is the initial level. The solution of this equation is

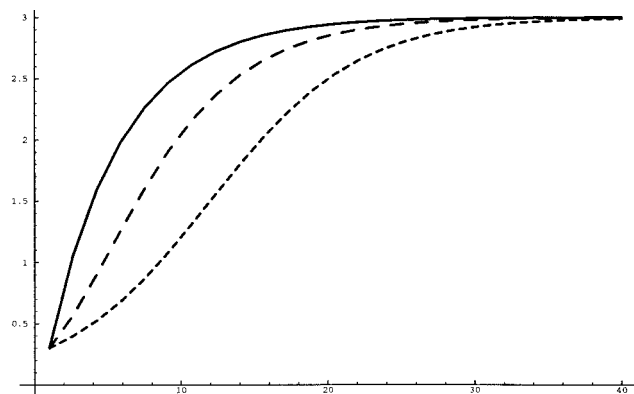


Figure 3 Growth curves from three functions that may display asymptotic behavior: exponential (-----), logistic (— — —), Gompertz (- - - -). Parameter values used to draw the curves were: $a = 3, i = 0.3, r = 0.2$

$$y(t) = \frac{ay(0)}{by(0) + (a - by(0))e^{-at}} \tag{7}$$

which is familiar to many researchers as the logistic curve. The middle line in Figure 3 is an example of a logistic curve. While it was developed for the growth of populations, it has (like many mathematical models) proved useful in a variety of contexts.

The Gompertz curve

A third model for asymptotic growth comes from Gompertz.¹⁶ It is similar to the logistic in that the rate of growth decays as a non-linear function of current height. Instead of the square of the current size, the product of logarithm of current height and the height itself are used. We write the differential equation for this model as

$$\frac{dy}{dt} = ay(t) - b \log[y(t)]y(t), \tag{8}$$

so growth is initially exponential (from $ay(t)$), but as time passes the term on the right may counteract it. This equation can be solved and rearranged to give

$$y(t) = \exp\left[\frac{a - e^{b(c-t)}}{b}\right] \tag{9}$$

The behavior of the Gompertz curve varies according to the parameters; for some values of b the curve does not asymptote but increases exponentially. An example Gompertz curve is the bottom line in Figure 3. While the model was developed for cell reproduction during ontogeny (the growth of chicken hearts), it has been widely used in tumor growth and models of population death rates.^{17,18}

We emphasize that the three curves presented here are only a small sample from the set of possibilities. However, they provide a foundation for a bridge between dynamical systems theory widely used in mathematical biology and structural equation modeling of genetically informative data.

Modeling phenotypic data

Now that we have mathematical equations for growth curves, we need to find a way to generate appropriate predictions for population means, variances and covariances across time. Each curve has parameters which control the initial level, the growth rate, and the asymptote. Suppose that there are latent factors which represent individual differences in these components, and that observed scores at different time points are a linear combination of these components and some random error. This can be described as a factor model with three latent

factors, as shown in Figure 4. Algebraically, we can write

$$\mathbf{p} = \mathbf{F}\mathbf{f} + \mathbf{E}\mathbf{e}, \quad (10)$$

where \mathbf{p} , \mathbf{f} and \mathbf{e} are respectively vectors of phenotypes, factors and residual errors, \mathbf{F} is the (full) matrix of factor loadings, and \mathbf{E} is the diagonal matrix of loadings on the residual components. The rules of path analysis^{19–22} or multivariate path analysis²³ may be useful to derive predicted covariances from this model as

$$\Sigma = \mathbf{R}\mathbf{F}\mathbf{F}' + \mathbf{E}\mathbf{E}' \quad (11)$$

where \mathbf{R} is the covariance matrix of the latent factors \mathbf{f} . Population variation in the latent factors is assumed to be symmetric, so the predicted mean for occasion t equals the height of growth curve at that time point. We can write this model using factor means,

$$\mu_t = \mathbf{F}\mu_f, \quad (12)$$

where μ_t is the $n_t \times 1$ vector of means, n_t is the number of timepoints, and $\mu_f = (a, i, 0)'$ is the 1×3 vector of factor means. As we later show, within a computer program such as Mx, the expression for the growth curve (Equations 4, 7 and 9) can be specified directly for the predicted means.

Our next task is to find expressions for the elements of \mathbf{F} , the factor loadings, that generate predicted covariances according to the growth curve model in use. We follow Browne¹ by using a first-

order Taylor series to reproduce the growth curve. In brief, this involves use of the partial derivatives of the growth curve function (Equation 4 or 7 or 9 above) with respect to the parameters a , b and c . For the vector of factor loadings from the asymptote factor under the exponential model we use the partial derivatives of Equation 4 with respect to a . These partial derivatives involve t , so the loadings are different from the different occasions $t = 1, 2, \dots, T$. They form the columns of the matrix \mathbf{F} in Equation 10. While these expressions have been published elsewhere, we reproduce them in Table 1 for convenience. We could directly follow the equations for the three curves derived in equations 4, 7 and 9 above, but a slight reparameterization due to Browne¹ makes the estimates easier to interpret, with a being the final asymptote and i being the value at time $t = 0$ for all three curves.

Table 1 Exponential, logistic and Gompertz growth curve functions and their partial derivatives

Exponential

$$F_E = a - (a - i) \exp[-(t - 1)r] \quad (13)$$

$$\frac{dF_E}{da} = 1 - \exp[-(t - 1)r] \quad (14)$$

$$\frac{dF_E}{di} = \exp[-(t - 1)r] \quad (15)$$

$$\frac{dF_E}{dr} = (a - i)(t - 1) \exp[-(t - 1)r] \quad (16)$$

Logistic

$$F_L = \frac{ai}{i + (a - i) \exp[-(t - 1)r]} \quad (17)$$

$$\frac{dF_L}{da} = \frac{i - \exp[-(t - 1)r]F_L}{i + (a - i) \exp[-(t - 1)r]} \quad (18)$$

$$\frac{dF_L}{di} = \frac{a - (1 - \exp[-(t - 1)r])F_L}{i + (a - i) \exp[-(t - 1)r]} \quad (19)$$

$$\frac{dF_L}{dr} = \frac{(a - i)(t - 1) \exp[-(t - 1)r]F_L}{i + (a - i) \exp[-(t - 1)r]} \quad (20)$$

Gompertz

$$F_G = a \exp \left[\log \left[\frac{i}{a} \right] \exp[-(t - 1)r] \right] \quad (21)$$

$$\frac{dF_G}{da} = [1 - \exp[-(t - 1)r]] \exp \left[\log \left[\frac{i}{a} \right] \exp[-(t - 1)r] \right] \quad (22)$$

$$\frac{dF_G}{di} = \frac{a}{i} \exp \left[-(t - 1)r + \log \left[\frac{i}{a} \right] \exp[-(t - 1)r] \right] \quad (23)$$

$$\frac{dF_G}{dr} = -a \log \left[\frac{i}{a} \right] (t - 1) \exp \left[-(t - 1)r + \log \left[\frac{i}{a} \right] \exp[-(t - 1)r] \right] \quad (24)$$

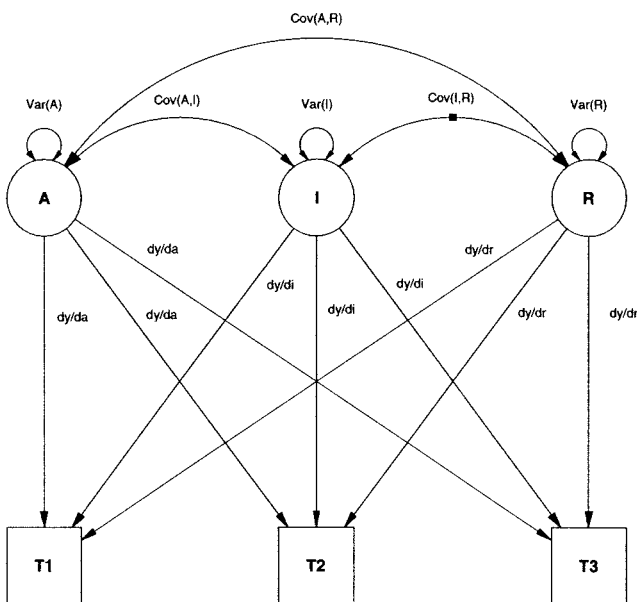


Figure 4 Three-factor model which represents variation in asymptote, initial level, and growth rate. All factors cause phenotypic variation on all occasions (T1, T2, T3), but the factor loadings are constrained according to the predictions of the model. Residual genetic and environmental components that influence only one time each are not shown

Thus the only differences between the growth curve models are the constraints on the factor loadings in the matrix **F**. Figure 5 plots the factor loadings for the Gompertz curve for time points $t = 1, \dots, 10$. As would be expected, the loadings on the asymptote factor increase parallel to the height of the curve itself. Loadings on the initial factor are highest at the start of the process, whereas those on the rate factor are highest when growth is most rapid.

The model allows for both variation within, and covariation between the latent growth factors. So, for example, within a population, initial value might be correlated with asymptote, giving rise to one of the three possible covariances between the components of growth. Second, the model is specified for discrete, evenly spaced time periods, but it is not limited to data of this type. If we had data assessed at ages 1, 2, 4 and 7, we could create or select the rows of **L** accordingly.

Modeling data from relatives

It is a relatively direct matter to extend the phenotypic growth models to cover data from relatives such as twins, but there are several important options. McArdle,¹¹ following Vandenberg and Falkner²⁴ presented a model of the genetic decomposition of the growth parameters. In this case, we have three covarying latent factors (asymptote, initial and rate) and uncorrelated residual variance. When there are data from relatives, we can model the familial resemblance for both the latent factors and the residual components. A multivariate path diagram²³ of such a model is shown in Figure 6. In moving from Figure 4, we have substituted the vector of latent variables **G** for the three growth factors a , i and r , and **P_i** represents the n_t observed measures. The covari-

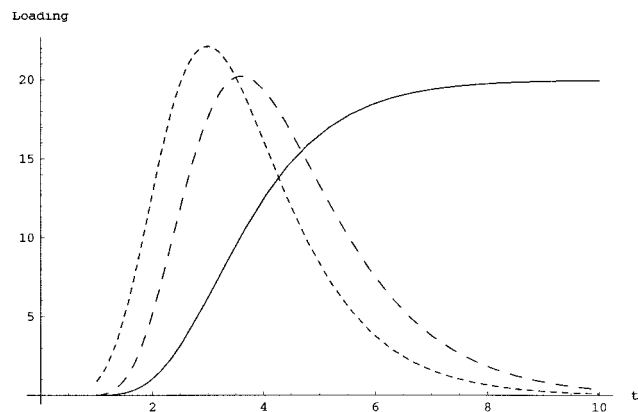


Figure 5 Plot of factor loadings according to partial derivatives of the Gompertz curve for time points $t = 1, \dots, 10$. Key: - - - - asymptote, a ; - - - - initial, i ; - · - · rate, r

ance between the factors, **A**, **I** and **R** in Figure 4, is now modelled in the **A**, **C** and **E** matrices. Likewise, we have substituted **A_s**, **C_s** and **E_s** for the residual variance. The model gives predicted covariance among twins which may be written:

$$\Sigma = (\mathbf{I} \otimes \mathbf{F}) \begin{pmatrix} \mathbf{A}' + \mathbf{C}' + \mathbf{E}' & \alpha \mathbf{A}' + \mathbf{C}' \\ \alpha \mathbf{A}' + \mathbf{C}' & \mathbf{A}' + \mathbf{C}' + \mathbf{E}' \end{pmatrix} (\mathbf{I} \otimes \mathbf{F})' + \begin{pmatrix} \mathbf{A}_s' + \mathbf{C}_s' + \mathbf{E}_s' & \alpha \mathbf{A}_s' + \mathbf{C}_s' \\ \alpha \mathbf{A}_s' + \mathbf{C}_s' & \mathbf{A}_s' + \mathbf{C}_s' + \mathbf{E}_s' \end{pmatrix}$$

where $\alpha = 1$ for MZ and 0.5 for DZ twins, **I** is a 2×2 identity matrix, **F** is the factor loading matrix, and \otimes denotes right kronecker product. The structure of the matrices in this model may be chosen from a variety of identified forms. All the usual tools of multivariate genetic analysis may be applied to the three covarying latent growth factors. We could, for example, use a Cholesky decomposition of the additive genetic, common and specific environmental factors if MZ and DZ twin data are available, in which case the matrices **A**, **C** and **E** would be lower triangular. The matrices could be restricted to represent single-factor or independent sources of variation for the

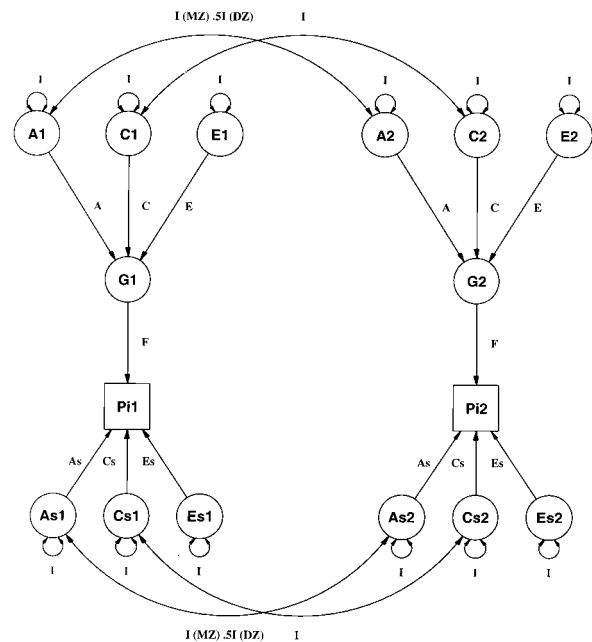


Figure 6 Multivariate path diagram of resemblance between MZ or DZ twins under a growth curve model. Additive genetic, common and specific environmental components (**G**, **C** and **E**) act additively to produce variation and covariation in growth curve factors **G**, which in turn cause variation and covariation in the phenotypes (**P**) over time. Residual components, not explained by individual differences in growth curve factors, may also be partitioned into genetic (**A_s**), and environmental (**C_s** and **E_s**) components

components by making them 3×1 or 3×3 diagonal, respectively.

We can take similar liberties with modeling the residual variation not explained by the growth curve. One approach, used here, is to allow time-specific A, C and E factors, which requires \mathbf{A}_s , \mathbf{C}_s and \mathbf{E}_s to be diagonal. In factor models, modeling of residual variation is often limited to such simple structure of the residuals because of the large number of parameters used to fit the factor structure, which can explain almost any covariance structure. However, growth curve models are highly efficient, requiring only nine parameters, *regardless of the number of occasions*. Many options for exploring residual covariation not explained by the growth curve are therefore available, including conventional general factors or simplex models, or combinations thereof. In this article we use only the time-specific approach to modeling the residual covariances.

Note that the method described here is a single-step analysis. This is in contrast to the approach used by Baker *et al.*²⁵ which uses two steps. First, individual growth curve parameters are estimated from each subject in the sample. Second, genetic and environmental parameters are estimated from the individual parameters.

Application to the Bayley Infant Mental Development Scale

Data to illustrate the approach come from the Louisville Twin Study²⁶ and were published and analyzed by McArdle.¹¹ The Bayley Infant Mental Development scale was administered to twins aged from 6 months to 2 years, at regular 6-month intervals. As described in McArdle's paper, means, standard deviations and correlation matrices for MZ and DZ pairs were computed from the data based on unequal sample sizes ranging from 72 to 105. Scores were computed as 100 times raw score divided by the maximum BIMD of 163, and are thus interpreted as a 'percentage correct' metric. For analysis in this article, covariance matrices and means are used. These statistics are less than optimal for the analysis of incomplete data, especially since Mx is capable of analyzing raw data via maximum likelihood.²⁷⁻³⁰ However, these summary statistics are sufficient for the purposes of illustration.

An Mx script for fitting the genetic growth curve model is given in Appendix 1. It is relatively simple. The first group is used to compute the partial derivative vectors for each of the time points; these form the columns of matrix \mathbf{F} , the factor loadings. The second group sets up the \mathbf{A} , \mathbf{C} and \mathbf{E} components of covariance between the asymptote, initial and rate factors. The third and fourth groups fit the model to

the MZ and DZ data, respectively. Several sub-models are then fitted using the multiple fit option.

As a starting point for comparison, we note that McArdle¹¹ fitted a latent growth curve model with:

- a) an initial level
- b) unrestricted loadings on a second component; and
- c) common and specific biometric components.

This model achieved a fit of $\chi^2 = 169$ on 21 degrees of freedom, compared with a saturated baseline model. We hope to improve on this fit here.

Results

Table 2 shows the parameter estimates for all three growth curve genetic models. The estimates of the means are illustrated graphically in Figure 7, which also shows the estimated exponential, logistic and Gompertz growth curves, based on the parameters a , i and r for the four models. Evidently, the models fit the means well, as would be expected since these statistics have small standard errors relative to the covariances. The figure suggests that additional measurement occasions would help discriminate between the curves, as their closely parallel forms diverge after the two-year final measurement.

In Table 2, there are 3×3 matrices reported for each of A_f , C_f and E_f . Each matrix contains estimates of the A, C or E contribution to the covariance matrix among the three growth components: the asymptote, initial and rate. The top-left element in each matrix represents the variance in the asymptote factor, which is much greater for the C component than for A or E in the exponential model (1756.19 vs 4.76 or 4.90). The same is true, to a lesser extent, for the logistic and Gompertz models. It is also the case that the C component is larger than the A or E for the variance of the initial factor (10.4 vs 1.14 or 0.16). There is little evidence of variation in the rate parameter; it is estimated to be near zero across all three models. This boundary condition may be one reason that the growth curve models do not fit as well as the Cholesky decomposition for these data. It suggests that the pattern of change in mean and variance does not match the partial derivative of the growth curve with respect to the rate parameter r .

With only four occasions, there is little information to assess individual differences in rates of growth. Likewise, there is little power to estimate the covariances between the additive genetic factors that contribute to variance in asymptote, initial and rate; the correlations between these components are estimated at unity so a single factor would suffice. The same is true of the specific environmental factors in

Table 2 Parameter estimates for three growth curve models fitted to Bayley Infant Mental Development data on MZ and DZ twins assessed on four occasions

Parameter	Model											
	Exponential				Logistic				Gompertz			
<i>a</i>	140.05				97.29				108.01			
<i>i</i>	39.88				40.17				40.00			
<i>r</i>	0.17				0.64				0.41			
<i>Gf</i>		<i>a</i>	<i>i</i>	<i>r</i>		<i>a</i>	<i>i</i>	<i>r</i>		<i>a</i>	<i>i</i>	<i>r</i>
	<i>a</i>	4.76	-1.00	-1.00	<i>a</i>	5.74	1.00	1.00	<i>a</i>	2.88	1.00	1.00
	<i>i</i>	-2.32	1.14	1.00	<i>i</i>	2.80	1.37	1.00	<i>i</i>	1.89	1.25	1.00
<i>Cf</i>		<i>a</i>	<i>i</i>	<i>r</i>		<i>a</i>	<i>i</i>	<i>r</i>		<i>a</i>	<i>i</i>	<i>r</i>
	<i>a</i>	1756.19	-0.16	-1.00	<i>a</i>	88.66	0.06	-0.90	<i>a</i>	223.12	-0.04	-0.99
	<i>i</i>	-21.49	10.40	0.16	<i>i</i>	1.67	9.46	-0.49	<i>i</i>	-1.72	9.92	-0.08
<i>Ef</i>		<i>a</i>	<i>i</i>	<i>r</i>		<i>a</i>	<i>i</i>	<i>r</i>		<i>a</i>	<i>i</i>	<i>r</i>
	<i>a</i>	146.94	1.00	-1.00	<i>a</i>	8.96	1.00	-1.00	<i>a</i>	21.32	1.00	-1.00
	<i>i</i>	4.90	0.16	-1.00	<i>i</i>	1.18	0.16	-1.00	<i>i</i>	1.86	0.16	-1.00
<i>A_s</i>	<i>t₁</i>	<i>t₂</i>	<i>t₃</i>	<i>t₄</i>	<i>t₁</i>	<i>t₂</i>	<i>t₃</i>	<i>t₄</i>	<i>t₁</i>	<i>t₂</i>	<i>t₃</i>	<i>t₄</i>
<i>C_s</i>	2.21	-1.19	-2.09	0.00	2.21	1.19	2.10	0.00	2.21	-1.18	2.11	0.00
<i>E_s</i>	-1.10	0.00	0.00	0.00	-1.33	0.00	0.43	0.00	1.23	0.00	0.00	0.00
	2.07	1.96	1.79	1.94	2.06	1.95	1.79	1.94	2.06	1.96	1.79	1.94

Variance in the growth curve components *a*, *i* and *r* due to genetic (*G_f*), common environment (*C_f*) and specific environment (*E_f*) factors are shown on the diagonal of matrices, with covariances below and correlations above. Variance due to occasion-specific genetic and environmental factors *A_s*, *C_s* and *E_s* are shown for the four occasions *t₁* to *t₄*.

growth. For shared environmental effects, which have a much greater effect on phenotypic variation, the correlations are large between initial and rate for all models. This would suggest that shared environmental factors that contribute to initial level are very similar to those that influence rate of growth. However, variability in rate is small, and therefore these high correlations are of little consequence.

The relative contribution of the factors to the phenotypic variance changes across occasion. This is because the factor loadings are strict functions of the parameters *a*, *i* and *r* and the time point *t*. The

proportion of variance accounted for by each component could be computed for all four occasions, and for all three models. In all three models, the growth curve accounts for approximately 64%, 88%, 74% and 100% of the variance in occasions one to four respectively.

Approximate χ^2 fit statistics vary from 129 to 138 across the different growth curve models, despite their having the same number of degrees of freedom. This suggests that, even with these few occasions, information in the pattern of variances and covariances over time can effect some discrimination between the models, and that these differences would be enhanced with more occasions of measurement. A further indication of model quality is the reasonableness of the parameter estimates. The logistic and Gompertz curves have asymptote parameters quite close to the theoretical maximum (100) for these test scores, whilst the exponential suggests a less plausible maximum of 139.

The full models demonstrate the identification and feasibility of fitting genetic growth curves, and that some fit the data better than others. To test some specific hypotheses about individual differences in growth, we can compare the fit of submodels. A series of comparative fit statistics for the three growth curves is shown in Table 3. On the whole, the changes in fit from a model to a submodel are similar across the three types of curve, so we focus on one curve, the logistic. The submodels address the following hypotheses:

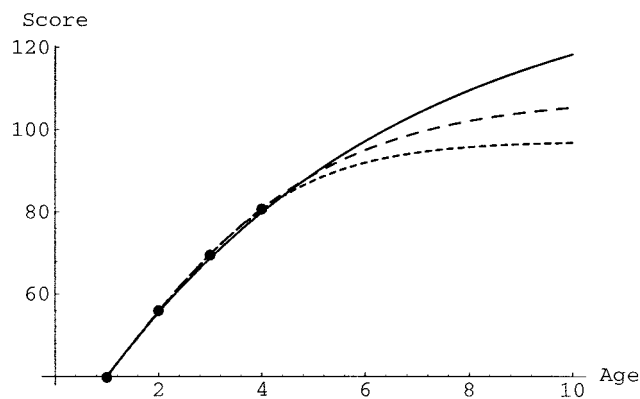


Figure 7 Plots of the exponential (---), logistic (-.-) and Gompertz (....) curves with parameters estimates set to the solution of fitting the growth curve models to the MZ and DZ data on Bayley Infant Mental Development scales

1. The asymptote, initial and rate factors covary;
2. Genetic factors contribute to factor variance and covariance;
3. Environmental factors contribute to factor variance and covariance; and
4. Residual variance not due to the latent growth curve does not covary between family members.

The full models do not fit very well by the χ^2 criterion. This may be due to non-normality in the data or the use of covariance matrices as summary statistics. As we shall see below, this problem is also present for the Cholesky model, which suggests that the loss of fit is due to inconsistencies of the four within-person covariance matrices (MZ and DZ twin 1 and twin 2). A likely consequence of this poor fit is that all χ^2 difference statistics (the likelihood ratio tests we use to test the difference between models) are inflated (Kendall and Stuart p.230).³¹ Evidence for significant loss of fit would be weaker, and evidence for non-significantly poorer fit would be stronger than usual.

Table 3 Fit statistics obtained for growth curve models and submodels applied to Bayley Infant Mental Development data on MZ and DZ twins

Model	Fit statistic			Difference χ^2		
	χ^2	d.f.	AIC	χ^2	d.f.	P
Exponential						
Full	129.06	55	19.06	–	–	–
Orthogonal	143.20	64	15.20	14.14	9	0.12
No A	140.78	61	18.78	11.72	6	0.07
No C	169.27	61	47.27	40.21	6	0.00
No E	134.15	61	12.15	5.09	6	0.53
No A _s , C _s	166.86	63	40.86	37.80	8	0.00
No A _s	136.88	59	18.89	7.82	4	0.10
No C _s	129.21	59	11.21	0.15	4	0.99
Logistic						
Full	137.89	55	27.89	–	–	–
Orthogonal	148.73	64	20.73	10.84	9	0.29
No A	149.49	61	27.49	11.60	6	0.07
No C	177.04	61	55.04	39.15	6	0.00
No E	142.74	61	20.74	4.85	6	0.56
No A _s , C _s	183.91	63	57.91	46.02	8	0.00
No A _s	145.83	59	27.83	7.94	4	0.09
No C _s	138.30	59	20.30	0.41	4	0.98
Gompertz						
Full	131.93	55	21.93	–	–	–
Orthogonal	144.13	64	16.13	12.20	9	0.20
No A	143.54	61	21.54	11.60	6	0.07
No C	171.72	61	49.72	39.79	6	0.00
No E	137.00	61	15.00	5.07	6	0.54
No A _s , C _s	172.76	63	46.76	40.83	8	0.00
No A _s	139.81	59	21.81	7.88	4	0.01
No C _s	132.21	59	14.21	0.27	4	0.99
Cholesky and Simplex						
Full Cholesky	117.75	50	17.75	–	–	–
Simplex	134.59	63	8.59	–	–	–
Simplex G Means	135.64	65	5.64	–	–	–

The orthogonal model does not fit significantly worse than the full model, so there is no evidence that the growth curve components covary. There is little evidence for either genetic or random environmental variation in the growth curve components, but significant variation is associated with the shared environment. The picture for the residual variance is quite different; familial resemblance appears to be additive genetic rather than shared environmental in origin. Thus of our four hypotheses above, 1, 2 and 4 are rejected and 3 is not.

Comparison with Cholesky and Simplex

To provide a comparative framework for the fit of the growth curve models, we fitted two standard behavior genetic models for longitudinal data, the Cholesky and the Simplex. The A, C, E Cholesky (Figure 1a) gave a fit of 117.75, with 50 df which yields Akaike's Information Criterion, AIC;³² = 17.75. Several of the growth curve submodels fitted better by AIC, so the poor fit (significant χ^2) to these data does not seem to be a feature of growth curves models *per se*.

We followed Dolan's⁷ treatment and specified separate A, C and E factor means which were constrained to equal a constant times the variance of each factor. The simplex model has a lower AIC (is more parsimonious) than most of the growth curves ($\chi^2 = 134.59$, df = 63, AIC = 8.59). It allows for A, C and E transmission and innovation components as shown in Figure 1b. Further improvement in AIC was found by eliminating the means on the common and random environment components ($\chi^2 = 135.64$, df = 65, AIC = 4.64), as they had only minor effects on the goodness of fit χ^2 . Though this 'genetic means' model fits well, there seems no theoretical justification to expect that genetic factors affect the means whereas environmental factors – especially shared environmental which are not due to measurement error – do not.

Discussion

We have reviewed growth curve theory from a dynamical systems perspective. In addition to its didactic value, it opens the door to genetic modeling of more complex dynamical systems within the structural equation modeling framework. Both balanced and incomplete data can be analyzed with this method.

The illustrative application to the published BIMD data has some novel substantive conclusions. The variance growth curve parameters are largely under shared environmental control is in agreement with McArdle's¹¹ results for linear latent growth models.

We have expanded on this initial treatment both by specifying different functional forms for growth (logistic etc.) and by partitioning the residual variance into A, C and E components. This latter partition is quite informative; in all models we found evidence for genetic factors in the residual variance, which suggests several possibilities. First, we may be measuring cognitive development at too few and too widely spaced intervals of time. Perhaps what we are ascribing to occasion-specific variance encompasses other growth processes which are partly under genetic control. Alternatively, it might be that the response of the individual to testing on that particular occasion (the 'person \times situation interaction') is genetically mediated. It seems difficult to discriminate between these possibilities without additional data.

Inferences we draw from the application should be viewed in the light of several potentially important limitations. First, the BIMD scores were treated as raw percentage scores, and this may not be appropriate for a scale that was not developed with, for example, Rasch³³ scaling.^{34,35} Second, we would expect similar answers if the raw data were analyzed instead of the means and covariances used here, but differences might occur if the data are not missing completely at random. It would be interesting to see how robust the findings are with raw data. Third, we have ignored the effect of sex in these analyses. Substantial sex differences in growth parameters might have inflated our estimates of common environmental variance. Future analyses separating males and females would be valuable, although quite large sample sizes would be required to detect sex differences in the model parameters.

Possible extensions to the methods presented here are many. First, it would be very simple to apply them to data collected from larger pedigrees than the pairs used here. Such datasets could include irregular family structures, with varying ages. A practical limitation of such studies – especially of cognitive development – would be finding parents who had been assessed in their early childhood. However, with appropriately scaled tests it should be possible to analyse data across wide age ranges. It may not be necessary to assess parents and their children at comparable ages, so long as there is sufficient overlap between the two samples. Indeed non-overlapping samples might suffice if, as in this study, the sources of variability appear to be entirely familial in origin, but we would advocate against designing studies which cannot assess the lasting impact of specific environmental factors. Measurement of different-aged relatives would be a poor substitute for longitudinal data, whose cost to collect can be reduced with cohort-sequential designs.³⁶ While parents and offspring may be the most

obvious addition to a twin study, other collateral relative groups – such as husbands and wives or full and half-siblings may be used in the short term to test assumptions of the twin study.

Second, the methods could be combined with those of standard behavior genetic analysis to enable testing of many types of hypothesis. For example, we could test for sibling interaction in development by modeling differences in MZ and DZ variance in the growth factors. We might test for $G \times E$ interaction by testing for heterogeneity of the growth curve parameters – or even types of growth curve – between groups subdivided according to some environmental factor. Multivariate extensions are readily incorporated, where we could allow for growth factors which influence several different variables simultaneously, or feedback mechanisms between the growth factors of different variables. Similarly, quantitative trait loci for the growth factors could be sought with appropriate data on DNA markers.^{37,38} Future work on these methods has great potential for understanding individual differences in change which could be applied to many areas of human development.

Extensions to ordinal or binary data require additional considerations. We could impose thresholds at various heights on the curve and via numerical integration compute the predicted proportion that would lie between each pair of consecutive thresholds. However, for two time points and twins we have four variables, requiring four-dimensional integration to compute the joint likelihood. While this is feasible with current computer hardware and software, studies with more than six occasions of measurement per subject (or 12 per family when larger pedigrees are considered) would prove difficult to analyze. Polychoric correlations and their associated weight matrices (ADF methods³⁹) can be a useful way to summarize ordinal data, and they would capture changes in covariance over time. However, they would lack the information on changes in mean and variance which provide much of the information to fit growth curves. Perhaps joint analysis of polychorics and thresholds and their overall weight matrix would be a good approach, although the large sample sizes required for ADF methods are presumably even larger when the thresholds are included.

Finally, we recognize the limitations to dynamical systems; many differential equations cannot be solved, and that even those that can may not be identified when implemented as a structural model. For example, the general sine wave function $a \sin[b + ct]$ has partial derivatives which are linearly dependent, so the three factor model is not identified. Nevertheless, the methods offer much promise for modeling change over time. While time is the

natural dimension with which to study change, other dimensions such as physical distance might be substituted to study genetic and environmental variation in other phenomena such as EEG waves or cellular chemical concentrations. Future research on these methods could grow in many directions.

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Appendix 1 Mx script to fit logistic growth curve to twin data

```
!  
! Mx script to fit logistic growth curve to data  
collected from  
! twins on nocc occasions of measurement  
!  
#define nt 4 ! Number of time points  
#define nt × 8 ! Twice number of time points
```

```

Group 1: compute factor loading vectors ! Title of
group 1
Calculation Ngroups = 5 ! Type of group & total
No. of groups
Begin Matrices; ! declare matrices
A full 1 1 free ! asymptote parameter
I full 1 1 free ! initial parameter
R full 1 1 free ! rate parameter
T full nt 1 ! vector of occasions t = 1,...nt
U unit nt 1
End Matrices
! A few starting values
Matrix T 1 2 3 4 5 6 7 8 9 10 ! extend if nt > 10
Start 100 a 1 1
Start 10 i 1 1
Start 1 r 1 1
! Boundaries for the parameters
Bound -500 500 all
!
! Now we use matrix algebra to compute the
vectors and paste them together
! into a single matrix of factor loadings, F
!
Begin Algebra;
J = i@U + (a-i)@( \exp(-(t-u)@r)); ! Logistic func-
tion itself
K = (i@u-( \exp(-(t-u)@r)).(((a*i)@U)%J)) % J;
!dJ/da
L = ((a@u) - (u-( \exp(-(t-u)@r))). (((a*i)@U)%J)) %
J ; ! dJ/di
M = ((a-i)@(t-u)). ( \exp(-(t-u)@r)). (((a*i)@U)%J)
% J ; ! dJ/dr
F = K | L | M ;
End Algebra;
End Group
Group 2 Compute the MZ and DZ twin pairs'
predicted factor covariances
Calculation
Begin Matrices
H full 1 1 ! to put 0.5 in for DZ twin genetic
covariance
X Lower 3 3 free ! lower triangular decomposition
used initially
Y Lower 3 3 free
Z Lower 3 3 free
End Matrices;
Matrix H .5
Start 2 X 1 1 Y 1 1 Z 1 1
Start 0.1 X 2 2 X 3 3 Y 2 2 Y 3 3 Z 2 2 Z 3 3
Begin Algebra;
A = X*X' ;
C = Y*Y' ;
E = Z*Z' ;
M = A + C + E | A + C —
A + C | A + C + E ;
D = A + C + E | h@A + C —
h@A + C | A + C + E /

```

```

End Algebra;
End Group
Group 3 fit the model to the MZ data
Data Ninput = ntx2 nob = 75
CMatrix full file = bayleymz.cov ! observed cov
matrix
Mean file = bayleymz.mean ! observed means
Matrices
F Computed = F1 ! Factor loading matrix
R Computed = M2 ! MZ factor covariances
A diag nt nt free ! Residual genetic factors
(occasion specific)
C diag nt nt free ! Residual C factors
E diag nt nt free ! Residual E factors
I iden 2 2
U unit 1 2
V unit 2 2
X full 3 1
End Matrices
Specify X 1 2 0
Start 3 E 1 1 to E 4 4
Bound 0.5 20 E 1 1 to E 4 4
Means U@(F*X)' /
Covariance (I@F) & R + I@(E*E') + V@(C-
*C' + A*A') /
Option rs
End Group
Group 4 fit the model to the DZ data
Data ninput = ntx2 nob = 75
CMatrix full file = bayleydz.cov
Means file = bayleydz.mean
Begin Matrices;
F Computed = F1 ! Factor loading matrix
R Computed + D2 ! DZ factor covariances
A diag nt nt = A3 ! Residual genetic factors
(occasion specific)
C diag nt nt = C3 ! Residual C factors
E diag nt nt = E3 ! Residual E factors
H Full 1 1 = H2
I Iden 2 2
U unit 1 2
V unit 2 2
X full 3 1 = X3
End Matrices;
Means U@(F*X)' /
Covariance (I@F)&R + I@(E*E') + V@(C*C') + (H-
@V + I)@(A*A') /
Option rs ! Print residuals
Option Multiple ! Fit submodels next
End Group
!
! Save current solution then fit seven submodels
from above solution
!
save baylog.mxs
!

```

```
! 1. make asymptote, initial and rate factors
orthogonal
!
drop 17 19 20
drop 5 7 8
drop 11 13 14
end
! 2. delete all genetic effects on factors
get baylog.mxs
drop 4 5 6 7 8 9
end
! 3. delete all C effects on factors
get baylog.mxs
drop 10 11 12 13 14 15
end
! 4. delete all E effects on factors
```

```
get baylog.mxs
drop 16 17 18 19 20 21
end
! 5. delete G parts of residuals
get baylog.mxs
drop 22 23 24 25
end
! 6. delete C parts of residuals
get baylog.mxs
drop 26 27 28 29
end
! 7. delete G & C parts of residuals
get baylog.mxs
drop 22 23 24 25 26 27 28 29
end
```