# THE IDENTITIES HIDDEN IN THE MATCHING LAWS, AND THEIR USES

#### DAVID R. THORNE

#### WALTER REED ARMY INSTITUTE OF RESEARCH

Various theoretical equations have been proposed to predict response rate as a function of the rate of reinforcement. If both the rate and probability of reinforcement are considered, a simple identity, defining equation, or "law" holds. This identity places algebraic constraints on the allowable forms of our mathematical models and can help identify the referents for certain empirical or theoretical coefficients. This identity can be applied to both single and compound schedules of reinforcement, absolute and relative measures, and to local, global and overall rates and probabilities. The rate matching equations of Hernstein and Catania appear to have been approximations to, and to have been evolving toward, one form of this algebraic identity. Estimates of the bias and sensitivity terms in the generalized ratio and logarithmic matching models are here held to be averaging artifacts arising from fitting procedures applied to models that violate or conceal the underlying identities.

Key words: matching, law, compound schedules, concurrent schedules, identity, tautology, behavioral economics, foraging

The success in the late 1960s and early 1970s in predicting relative response rates on concurrent interval schedules led to attempts to predict absolute response rates on schedules of reinforcement in general. For the case of a single response class, Herrnstein (1961) suggested a simple proportionality of the form

$$b = k r \tag{1}$$

where b and r are the mean response and reinforcement rates respectively. Catania (1963) suggested a power function which for the single response case reduced to the form

$$b = k r^c \tag{2}$$

Herrnstein (1970) stated that the simple proportionality did not appear to work well empirically and proposed a hyperbolic function

$$b = \frac{k \, r}{r + r_0} \tag{3}$$

where  $r_0$  was assumed to represent sources of reinforcement other than those specified by the experimenter and k was assumed to be the asymptotic rate attainable for that response in the absence of competition from other sources. Catania (1973) proposed a mathematically similar equation

$$b = \frac{k C r}{C + \sum r_i} \tag{4}$$

Where k was a coefficient of proportionality, C was assumed to represent an inhibitory effect of reinforcement upon responding, and  $\Sigma$   $r_i$  was the total rate of the known sources of reinforcement.

Each of these formulations treats rate of reinforcement as the primary variable influencing responding.

It will be shown below that in both singleresponse and many choice-response situations Equation 1 is the more generally correct; that is, that the exponent is unity and that the obtained response and reinforcement rates (and their ratios) are directly proportional, but that the multiplier is an identifiable variable rather than a hypothetical constant.

# Simple Schedules, Absolute Measures

At the end of a typical operant experiment one has three totals at hand—the duration of the session or schedule component of interest (T), the number of Behavioral responses that occurred therein (B), and the number of Reinforcements delivered (R). From these



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Address correspondence to David R. Thorne, Department of Behavioral Biology, Division of Psychiatry and Neurosciences, Walter Reed Army Institute of Research, 503 Robert Grant Ave., Silver Spring, Maryland, 20910-7500 (email: david.thorne@us.army.mil or davidthorne@verizon.net). doi: 10.1901/jeab.2010.93-247

three raw totals we frequently derive three additional variables (or their reciprocals): response rate b = B/T (or a mean IRT, T/B); rate of reinforcement r = R/T (or a corresponding obtained mean VI value T/R); and probability of reinforcement given a response p = R/B (or a corresponding obtained mean VR value, B/R). We make these conversions in order to generalize across sessions or experiments having different absolute totals. Whether the three original numbers (B, R and T) are viewed as free and independent of one another, the three derived variables (b, r and p) are ratios sharing common terms, hence they are interdependent. If two of the variable pairs are known the third is strictly determined. Thus we can write

Response Rate =

where the denominator is the "conditional" probability of reinforcement given a response and all values are obtained as opposed to programmed.

The word-equation 5a conceals from easy view that it is equivalent to the algebraic identity

$$\frac{B}{T} = \frac{(R/T)}{(R/B)} = \frac{R}{T} \cdot \frac{B}{R} = \frac{B}{T}$$
 (5b)

This identity will of course hold for all values of B, R and T; and for the grand totals on any simple, compound or second-order schedule of reinforcement. It will also hold, and can be evaluated, for the local totals on those compound schedules where one can legitimately determine each component's duration and counts (e.g. chain, multiple, concurrent, but not single-operandum conjunctive or alternative schedules, where it is not possible to assign a response exclusively to one component over the other). It would hold equally well for dubious or inappropriate cases such as noncontingent time schedules or the differential reinforcement of "other" behavior (where the reinforcement probabilities given B and Not-B are equal, or zero and nonzero respectively).

Two rearrangements of this identity are well known. One, often used in the present context, places R/T on the left and is typically

used to derive the rate of reinforcement generated by ratio schedules. The other, often used in the context of behavioral economics, places B/R on the left and is used to derive the response "cost" imposed by interval schedules (where cost equals 1/p). Moving B/T to the left is trivial mathematically but perhaps significant perceptually or interpretively.

One important point this makes is that, because things equal to the same thing are equal to each other, any theoretical equation that accurately describes or purports to describe or predict response rate must also equal Equation 5a. The two equations may then be solved simultaneously to yield additional relationships, or to reveal arithmetic incompatibility.

Equation 5a can be abbreviated to the form

$$b = r/p \tag{5c}$$

which may be compared to Equation 1, or both sides of Equation 5b can be multiplied by T to give an equation in number and probability, rather than rate:

$$B = \frac{R}{R/B} = R/p \tag{6}$$

The identity can also be inverted and rewritten in terms of obtained mean interresponse times, variable-interval values, and variable-ratio values as

$$IRT = VI/VR$$
 (7)

Although this last equation would seldom be useful in practice it can be used to make a somewhat contrived point. In any single experimental session where the experimenter imposes a particular VI (or VR) schedule, there is a post hoc VR (or VI) sequence that would have produced exactly the same recorded result. An uninformed observer, or the subject, would be unable to tell which was in effect. Zeiler (1977) has addressed this issue in the context of control by direct and indirect variables.

Although these equations are simple algebraic tautologies, that does not necessarily mean that they are trivial. Many, but not all, of the equations or laws of physics are equally circular (e.g.  $E = I \cdot R$ ,  $f = m \cdot a$ ) as can be shown by reexpressing them in their basic units and dimensions. Such laws are sometimes called defining equations.

Equations 5 through 7 may be considered descriptive but not explanatory in the colloquial sense. They say more about the particular behavioral measures we have come to use for parsing our subject matter than about behavior per se. Nevertheless, the identity can be useful in a number of ways ranging from the philosophical or heuristic to the practical:

The identity can be interpreted as a reminder that even on the simplest schedule we do not have a single dependent variable but a dependent variable pair. This is most clearly seen in the case of simple ratio or interval schedules (e.g. Equation 7) where the experimenter has complete or effectively complete control over one variable while the subject controls the other two. If it is known that only one of the dependent variables is directly influenced by an experimental manipulation and that the other dependent variable merely follows as it mathematically must, then the common practice of recording and attending to only one of the two variables may be logically justifiable, albeit potentially misleading. In any single isolated experiment the two dependent variables will be inextricably confounded and we may arbitrarily decide to call one of them the relevant consequent and the other an irrelevant concomitant. Across a series of experiments, however, it may be possible to separate the two still-confounded effects based on differential manipulations and findings and/or a preferred criterion of "orderliness" such as linearity, invariance, or goodness-of-fit.

The practice of attending to only one of the two dependent variables becomes less defensible if both variables are directly influenced by an experimental manipulation while simultaneously constrained to obey the numerical identity. We tend to describe such cases as exhibiting strong interaction. Possible analogies here might be simultaneous manipulation of both sensitivity and bias in Signal Detection experiments or of speed/accuracy tradeoffs in reaction time studies—two areas where double dependent measures have proven fruitful. Attending to rate while ignoring probability carries risks.

- 2) The identity suggests a potentially stronger parallel than usually recognized between the phenomena identified with "rate matching" and those historically identified with "probability learning", or with those more recently considered in behavioral economics.
- 3) The identity can also be interpreted as showing that, as independent variables, neither rate of reinforcement nor probability of reinforcement is logically prior to the other. Both may act concurrently. Which if either of the two variables is behaviorally the more potent or pertinent may differ with the situation, but the identity still suggests an inverse relationship.
- 4) The identity can also be used simply for calculating the "missing variable" in an ongoing experiment, an armchair experiment, or from a published report. This is the most frequent use of Ohm's Law, for example.
- 5) The identity can also serve as a useful check on our theoretical models, since they may not contradict it. As an illustration it will be used below, first to ask whether several different matching models are mathematically allowed, disallowed, or forced. Finally, we will return to a single-response case combining the identity with an empirical finding.

## Compound Schedules, Absolute Measures

On a two-ply (two-component) compound schedule we can distinguish at least 10 different absolute rates grouped into three distinct sets: Local rates, e.g.  $R_1/T_1$ ,  $B_2/T_2$ ; global or grand rates, e.g.  $(B_1 + B_2)/(T_1 + T_2)$ ; and what have come to be called overall rates, e.g.  $R_2/(T_1 + T_2)$ . The logically completing fourth set, a kind of "underall rate", e.g.  $(R_1 + R_2)/T_1$ , has not been deemed useful.

Parallel distinctions can be made for at least six different local, global, and overall conditional probabilities of reinforcement, for example,  $(R_1 + R_2)/(B_1 + B_2)$ . Each of these rate and probability measures can be related to the others through appropriate algebraic manipulation. The identity of Equation 5c holds in a straightforward manner for both the local and global measures. That is, the local (global) response rate will equal the local (global) reinforcement rate divided by the local (global) probability of reinforcement. Overall rates and probabilities, being to some

extent unsymmetrical mixed measures, lead to their own mixed identities. Thus absolute overall response rate could be written as either of the following

$$\frac{B_1}{T_1 + T_2} = \frac{(R_1 + R_2)/(T_1 + T_2)}{(R_1 + R_2)/B_1} = \frac{\text{global } r}{\text{underall } p} (8a)$$

$$= \frac{R_1/(T_1 + T_2)}{R_1/B_1} = \frac{\text{overall } r}{\text{local } p}$$
 (8b)

where the latter may have some potential theoretical usefulness.

Compound Schedules, Relative Measures

Having expressions for the various absolute rates in each component of a compound schedule allows us to compute various relative rates as well. It has been traditional practice to use overall rates for such computations, in which case all temporal terms cancel, leaving a dimensionless ratio that resembles a relative number or frequency rather than a relative rate. Thus relative overall response rate becomes

$$\frac{B_1/(T_1+T_2)}{B_1/(T_1+T_2)+B_2/(T_1+T_2)} = \frac{B_1}{B_1+B_2}.$$
 (9)

In the matching literature the term "rate" is still applied, perhaps because all numbers are normalized to a common session duration, hour, minute or other temporal unit. In other contexts and disciplines it might be more appropriate and conventional to treat this quantity as a probability, but the now historical terminology will be followed here.

Using the above definitions and identities, various expressions for relative overall response rates can now be derived. Beginning with Equation 6, and using subscripts to denote the two components, we have  $B_1 = R_1/p_1$  and  $B_2 = R_2/p_2$ , hence

$$\frac{B_1}{B_1 + B_2} = \frac{R_1/p_1}{(R_1/p_1) + (R_2/p_2)}.$$
 (10)

Multiplying both numerators and denominators by  $(p_1 p_2)$  simplifies to

$$\frac{B_1}{B_1 + B_2} = \frac{R_1 p_2}{R_1 p_2 + R_2 p_1}. (11)$$

Herrnstein (1970) used a formally equivalent equation in discussing matching under a concurrent ratio schedule, though the equation is a general identity true of all compound schedules. It is not identical to Herrnstein's (1961) matching relationship:

$$\frac{B_1}{B_1 + B_2} = \frac{R_1}{R_1 + R_2} \tag{12}$$

but reduces to it as a special case if and when  $p_1 = p_2$ , as noted by Revusky (1963). Thus matching is mathematically allowed but not forced. If relative rate matching does occur then local and global reinforcement probabilities will be equal, and vice versa. The two phenomena (rate matching and equal probabilities) are confounded correlates, of equal potential significance, with neither necessarily being the cause of the other. It also follows from Equation 11 that when relative rate matching does *not* occur (as in undermatching on concurrent schedules or behavioral contrast on multiple schedules) it must necessarily be due to differences in the two probabilities of reinforcement.

Equation 11 does not reduce to a general equation of the form

$$\frac{B_1}{B_1 + B_2} = \frac{k R_1}{R_1 + R_2} \tag{13}$$

although for any isolated schedule pair,  $R_1$   $R_2$ , one can always find some value for k that forces numerical equality. However, by dividing the numerator and denominator of Equation 11 by  $p_2$  we get the somewhat more familiar form

$$\frac{B_1}{B_1 + B_2} = \frac{R_1}{R_1 + m R_2} \tag{14}$$

where  $m = p_1/p_2$ . This too is a general identity that must always hold.

For the case of a three-ply compound schedule the corresponding identity can be variously written as

$$\begin{split} \frac{B_1}{B_1 + B_2 + B_3} &= \frac{R_1/p_1}{(R_1/p_1) + (R_2/p_2) + (R_3/p_3)} \\ &= \frac{R_1p_2p_3}{R_1p_2p_3 + R_2p_1p_3 + R_3p_1p_2} \\ &= \frac{a\ R_1}{a\ R_1 + b\ R_2 + c\ R_3} \\ &= \frac{R_1}{R_1 + m_2\ R_2 + m_3\ R_3} \end{split} \tag{15}$$

where  $m_j = p_1/p_j$ .

Equations 14 and 15 are written for the relative rates in component number 1, but corresponding equations will hold for components 2, 3, or n. For notational exactitude m should be doubly subscripted as  $m_{ij} = p_i/p_j$  where i is the component of interest selected for the numerator. Relative overall response rates for the generalized case of any n-ply compound schedule (multiple, concurrent or other) can then be written as either of the following identities

$$\frac{B_i}{\sum B_j} = \frac{R_i}{p_i \sum (R_j/p_j)}$$
 (16a)

$$= \frac{R_i}{\sum m_{ij} R_j}.$$
 (16b)

Application to Classic Rate Models

It will be convenient for what follows to rewrite the last identity as

$$\frac{B_1}{\sum B_j} = \frac{R_1}{R_1 + \sum m_{1j} R_j}$$
 (17)

where the right-hand summation is over all components *except* component number 1. This awkwardness could easily be avoided by assigning an m term of unity to the  $R_{\rm I}$  term in the denominator and then moving both back within the summation, but it is left in the less orthodox form to facilitate comparisons with Herrnstein's (1970) theoretical equation

$$B_1 = \frac{k R_1}{R_1 + m R_2 + R_0} \tag{18}$$

and with Catania's (1973) theoretical Equation 19 (before C k became a new k)

$$B_1 = \frac{C \ k \ R_1}{C + \sum m_i \ R_i}.$$
 (19)

Although there are obvious differences between the last three equations, the similarities are the more interesting.

Herrnstein (1970) viewed m as an interaction term between components in multiple schedules, having a value between zero and one, to be estimated from the data. The present general account provides an explicit definition for m, having a value between zero and infinity, which can be calculated directly from the raw totals.

In the two-ply case Herrnstein (1970) began with the reasonable first assumption that the m terms were the same for both components, whereas in the present account  $m_{12}$  and  $m_{21}$  are seen to be reciprocally related—becoming the same only when the two probabilities are equal, and their ratios are unity. When unequal, the m terms could be viewed as representing a differential preference or bias toward one schedule, stimulus, or response of a pair.

The terms B and k in Equations 18 and 19 carry the dimensions of "responses" or "responses per unit time" (Catania, 1973; Herrnstein, 1974) whereas Equations 14–17 are written as dimensionless proportions or relative rates. The simplest resolution of this apparent difference is to assume that the term k represents the quantity  $(B_1 + B_2)$ , or more generally  $\sum B_j$ . Transposing k to the left-hand denominator makes the similarities among Equations 17–19 more obvious.

In matching experiments where the total rate of reinforcement is held constant the observed response total also appears to remain relatively constant, or to vary unsystematically (see Catania, 1963, Figure 2). A constant response total can also result if the organism responds at a fixed local rate that remains invariant with changes in schedule values or relative durations, as has been reported with pigeons (Blough, 1963; Stubbs & Pliskoff, 1969). The constancy of k has been assumed in several theoretical accounts, and these particular effects would tend to produce it, but it is not required for  $\sum B_i$  in the identities, and many researchers have reported conditions under which it is not (see review by McSweeney, Melville & Whipple, 1983).

At this point we have rationalized and redefined the terms m and k in order to make the theoretical Equations 18 and 19 most easily consistent with what is algebraically required. The same approach also leads to questioning the necessity of the inhibitory term C in Equation 19 and the reinforcement term  $R_0$  in Equation 18.

In Equation 19 the C term in the denominator should have the dimensions associated with reinforcement—to prevent adding apples to oranges, and to cancel the corresponding dimensions of R in the numerator. But if C in the numerator also has these same dimensions then cancellation no longer occurs and we

ultimately end up with the statement that a number equals a dimension. If C in the numerator does have the dimensions of reinforcement, then k cannot be a dimensionless coefficient, nor have the dimensions of responses or responses per time as in the Herrnstein (1970) account (and the above account), but must have the dimensions of responses per reinforcement (which, interestingly, is equivalent to 1/p in the above). Both Catania (1973) and Williams and Wixted (1994) discussed this dimensional necessity. It would seem less awkward and more parsimonious to discard C as unnecessary. Since Equation 19 already contains an m term for each R term within the summation, it then becomes algebraically and dimensionally compatible with the required identity of Equation

The term  $R_0$  in Equation 18 is assumed to represent unmeasured hypothetical reinforcers beyond those arranged by the experimenter. If there are cases where such intrinsic or extraneous sources of reinforcement must be considered, the formal equations require an additional  $m_{10}$  term on the right and a corresponding  $B_0$  term on the left, paralleling the third components in Equation 15. Identifying the physical referents for such terms and quantifying their values presents a number of problems and dangers one would hope to avoid. The most cautious and parsimonious approach might be to consider the possibility that  $R_0$  has served as a floating parameter to account for inaccuracies in the estimates of m and/or inconstancies in the values of k. Having explicit definitions for these terms makes it easier to evaluate this possibility.

If  $R_0$  were assumed to exist anyway, then the three required theoretical terms  $R_0$ ,  $m_{10}$ , and  $B_0$  represent only two degrees of freedom since they too must fulfill the identity  $B_0 = R_0$  $p_0$ . In the most general case, there will be no way of estimating  $p_0$  independently of  $R_0$  and  $B_0$ . However, in the special case where overt matching occurs and probabilities are equal, then  $p_0$  should equal the empirically measurable value of  $p_1$ ,  $p_2$ , etc. Unfortunately, this still leaves an infinite number of unmeasurable  $R_0/B_0$  pairs that would fulfill the necessary relationship. This problem cannot be unequivocally resolved even by the, perhaps tempting, assumption of a constant k greater than the observed  $\Sigma B_i$ , such that  $B_0 = k - \Sigma B_i$ . Why?

Any given n-ply schedule can be expressed as having only (n-1) plies by discarding one set of real terms, or as having (n+1) plies by adding appropriately selected imaginary terms, and in either case the equations will balance. Any value of k can be made to suffice. The existence of  $R_0$  cannot be formally confirmed or disconfirmed within this closed system.

It would be interesting to reexamine those studies where archival data permit comparing the assumed or computed m values with the obtained reinforcement probabilities, and the computed k values with the actual response sums. A computed k value less than any one of its contributing sums would seem fatal to the  $R_0$  hypothesis since it would indicate a negative number of  $B_0$  responses—a notion difficult to fathom or defend. (McSweeney et al., 1983, reported a number of cases where the computed  $R_0$  was negative, which equates to the same thing.)

The above derivations and analysis suggest that Herrnstein's (1970) and Catania's (1973) formulations in Equations 18 and 19 are incorrect in their details or parametric interpretations but are close approximations to the general identity of Equation 16b. That these investigators came so close, out of an unlimited number of alternative possibilities, is a tribute to their hypothetico-deductive skills.

## Application to Ratio Models

In the above it has been shown how identities stemming from Equation 5 determine or constrain expressions relating responding and reinforcement, using relative overall rates as an example. An additional example will show how it applies to the alternative measures of response and reinforcement ratios.

In concurrent schedules the subject determines the time allocations  $T_1$  and  $T_2$ , which usually differ, and the use of relative overall rates is conventional (following the discovery of matching). In multiple schedules the experimenter determines  $T_1$  and  $T_2$ , which are usually equated rather than "matched" to the schedule values, and the use of relative local rates is conventional. (In a typical multiple schedule the overall rate  $B_1/(T_1 + T_2)$  would equal  $B_1/(2T_1)$ , or half the local rate.) However, by using rate *ratios* the temporal terms can be made to cancel in both

cases allowing a common formulation (although possibly concealing an important underlying distinction).

For multiple schedules, Lander and Irwin (1968) proposed a nonmultiplicative ratio power function which Baum (1974) generalized to concurrent schedules and matching as

$$\frac{B_1}{B_2} = a \left(\frac{R_1}{R_2}\right)^c \tag{20}$$

where a represents any differential bias and c represents sensitivity ('censitivity') to reinforcement. Equation 5b, on the other hand, leads to the identity

$$\frac{B_1}{B_2} = \frac{R_1/p_1}{R_2/p_2}$$

$$\frac{B_1}{B_2} = \frac{p_2}{p_1} \left( \frac{R_1}{R_2} \right) \tag{21}$$

with an R ratio exponent of unity and a "bias" multiplier equivalent to  $m_{2,1}$  in the above. Many investigators have found cases where Equation 20 fits data more accurately when the exponent c differs from unity. These cases typically involve numerical fits computed across several schedule combinations of  $R_1$   $R_2$  pairs. There are at least two ways that this result can arise.

First, if the investigator manipulates  $R_1/R_2$  in order to observe its effects upon  $B_1/B_2$  and c then, ideally, a should be held constant throughout. If it is not known that a represents the measurable quantity  $p_2/p_1$  then there is no way of detecting its variation. Instead, one might assume and impose invariance by fiat. The resultant fit will therefore be inexact and it can be improved upon by a power transformation, a multiplicative correction factor, an added constant, or any other operation that introduces one or more free parameters. A nonunitary c implies a nonconstant a.

Second, if the experimental procedure allows or causes  $p_2/p_1$  to become a direct or indirect function of  $R_1/R_2$ , then Equations 20 and 21 can hold simultaneously. As an example of an indirect function, assume that the organism "probability matches", i.e. that it apportions its responses in the same manner as the obtained probabilities of reinforcement:

Note that this is not an identity but a hypothetical empirical finding. Substituting for  $p_2/p_1$  in Equation 20 yields

$$\frac{B_1}{B_2} = \frac{B_2}{B_1} \left( \frac{R_1}{R_2} \right). \tag{23}$$

Transposing the response terms on the right to the left gives

$$\left(\frac{B_1}{B_2}\right)^2 = \left(\frac{R_1}{R_2}\right) \tag{24}$$

hence

$$\frac{B_1}{B_2} = \left(\frac{R_1}{R_2}\right)^{1/2} \tag{25}$$

Equation 25 is still equal to Equation 21, but the reinforcement probabilities have become indirect (and hidden) functions of the reinforcement rates. This can be revealed by rewriting Equation 25 as

$$\frac{B_1}{B_2} = \left(\frac{R_2}{R_1}\right)^{1/2} \left(\frac{R_1}{R_2}\right)^1. \tag{26}$$

If, instead, the reinforcement probabilities were equal to, or proportional to, some other power than 1/2, for example  $p_1 = k R_1^C$  and  $p_2 = k' R_2^C$  then the  $(R_1/R_2)$  exponent would become 1-c with a = k'/k. Many other p = f(R) relationships are of course possible, yielding different  $(R_1/R_2)$  exponent values and nonunitary as. Identifying such relationships will require attending to measured probabilities as well as rates.

The Matching Law "Constants" as Averaging Artifacts

If an experiment employs, say, five different  $R_1$   $R_2$  combinations, then Equation 21 will hold for each individual set of triple pairs. However, in most cases each different  $R_1/R_2$  pair will generate a different  $B_1/B_2$  pair and a different  $p_2/p_1$  pair, hence a different local a value. That is, there is no forced invariance, constancy or even continuity across the five conditions. All can change.

Consider an experiment in which a subject exhibits bias but neither under- nor overmatching and c = 1. If one fits and estimates the best constant a across all five pairs then the value of this a will usually fall somewhere between the obtained minimum and maximum values of the five  $p_2/p_1$  pairs. That is, it will be some average, though not necessarily

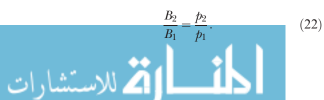


Table 1				
Data from Baum, Schwendiman and Bell (199	9) for Bird 973 on 12 concurrent VI VI schedules.			
Key 1 Rich	Key 1 Lean			

Key 1 Rich			Key 1 Lean				
$B_1$	$B_2$	$R_1$	$R_2$	$B_1$	$B_2$	$R_1$	$R_2$
14217	2646	793	87	7051	10847	168	552
29241	3345	1853	67	3190	52281	35	2365
138469	325	7985	15	766	38959	15	1825
28572	9272	2244	316	7121	12042	247	1033
61808	1907	3585	15	889	48641	19	2541
77265	557	3667	13	1513	50724	16	2704
		raw	log 2			raw	log 2
$p_1$	$p_2$	$p_2/p_1$	$p_2/p_1$	$p_1$	$p_2$	$p_2/p_1$	$p_2/p_1$
0.0558	0.0329	0.5895	-0.7625	0.0238	0.0509	2.1359	1.0948
0.0634	0.0200	0.3161	-1.6616	0.0110	0.0452	4.1230	2.0437
0.0577	0.0462	0.8004	-0.3213	0.0196	0.0468	2.3922	1.2583
0.0785	0.0341	0.4339	-1.2044	0.0347	0.0858	2.4731	1.3063
0.0580	0.0079	0.1356	-2.8824	0.0214	0.0522	2.4443	1.2894
0.0475	0.0233	0.4918	-1.0240	0.0106	0.0533	5.0409	2.3337
Geometric Mean = 0.4035			Geometric Mean =		2.9371		
Arithmetic Mean = $2^{(-1.3094)}$ =		0.4035	-1.3094	Arithmetic Mean = $2^{(1.5544)}$ =		2.9371	1.5544

*Note.* Top portion is the obtained number of responses  $(B_i)$  and reinforcers  $(R_i)$ , separated by whether key-1 was the Rich or Lean schedule of each pair. Middle portion shows the obtained probabilities of reinforcement  $(p_i)$  on each key (as  $R_i/B_i$ ) and their absolute and  $\log_2$  ratios. Bottom portion shows that means of the latter give the same bias and  $\log(\text{bias})$  values estimated by the authors' using a least squares fit on the  $\log_2(R)$  and  $\log_2(B)$  ratios, when matching was assumed.

their arithmetic mean. It can be shown that if one fits Equation 20 using a least squares fitting procedure on the obtained values, then the fitted a will equal the geometric mean of the obtained  $p_2/p_1$  values. If instead of the ratio form of the matching law one fits Baum's (1974) logarithmic form

$$\log(B_1/B_2) = \log a + c \log(R_1/R_2)$$
 (27)

using least squares, then the best fitting  $\log a$  value will equal the arithmetic mean of the five  $\log (p_2/p_1)$  pairs. When unlogged, this necessarily gives the same value for a as the geometric mean.

The same thing will result using a dataset where over- or undermatching occurs if one forces c to equal 1 and then refits. This will be demonstrated using data from Baum, Schwendiman and Bell (1999). This study exposed 4 pigeons to Conc VI VI schedules having a range of reinforcement rate ratios counterbalanced across two keys. When the data were conventionally plotted as log response ratios versus log reinforcement ratios (that is, of log  $[B_{LEFT}/B_{RIGHT}]$  versus log  $[R_{LEFT}/R_{RIGHT}]$ ) the fitted lines showed minimal bias (log  $a \approx 0$ )

but noticeable undermatching ( $c < 1 \approx 0.8$ ), as most typically seen in other studies. When the ratios were instead dichotomized (subscripted) by preferred and nonpreferred schedule values, independent of key position, a different picture emerged with negative log bias and near matching ( $c \approx 1$ ). For 1 of their 4 pigeons they replotted the "Left/Right" data showing the function not as one continuous line of reduced slope but as two separate lines of near-unit slope and opposing bias. They then forced the two slopes to be exactly equal (at c = 1) and solved for the "Lean and Rich" bias values separately, getting log(a)values of 1.55 and -1.31 respectively. Table 1 shows how these same values can be obtained by averaging the raw or logged data values without an explicit least squares fitting procedure.

The same thing will result using a dataset where the empirically obtained  $\epsilon$  is unity or near unity without being "forced". Such data sets are rather rare but Table 2 presents one example where the raw data values are publicly available. This example is derived from the Appendix and Table 2 of Elliffe and Alsop



Table 2					
Data from Elliffe and Alsop (1996) for Bird 131 on five concurrent VI VI schedules where near matching occurred ( $c = 1.01$ ).					

$B_1/B_2$	$R_1/R_2$	p <sub>2</sub> /p <sub>1</sub>	$\log B_1/B_2$	$\log R_1/R_2$	$\log p_2/p_1$
0.370768	0.845238	0.438656	-0.4309	-0.07302	-0.35788
0.06206	0.084337	0.735853	-1.20719	-1.07398	-0.13321
2.417017	3.394737	0.711989	0.38328	0.530806	-0.14753
0.199578	0.24812	0.804358	-0.69989	-0.60534	-0.09455
7.953125	9.388889	0.847078	0.900538	0.972614	-0.07208
	Geometric Means			Arithmetic Means	
0.615416	0.891695	0.690164	-0.21083	-0.04978	-0.16105

*Note.* Top portion lists the absolute and  $\log_{10}$  ratios of the obtained responses, reinforcers, and (inverse) probabilities of reinforcement. Means of the p ratios give the same bias and log bias values obtained by the authors using a least squares fit on the  $\log B$  and  $\log R$  ratios.

(1996) for Bird 131 under one reinforcement/min, where the reported log bias value was -0.16 with a c of 1.01 (close enough to unity for the present purpose).

Again, it is seen that the fitted log bias value equals the arithmetic mean of the log  $p_2/p_1$  values.

Finally, consider the more common case of inexact matching. If one interprets the fitted a value as a true constant then, since it usually isn't, the fit will not be optimal and a better fit will result by introducing a free parameter c and estimating both parameters simultaneously. In this case the two fitted parameters will seldom be identifiable simple means of any of the raw data values. Table 3 gives an example derived from the appendix of Davison and Jones (1995) for Bird 21where the reported c was 0.54 and log a was 0.00 (a concurrent VI Extinction condition has been omitted to avoid division by zero).

In this case the arithmetic mean of the log  $p_2/p_1$  values is not 0.00 (actually 0.0004 before rounding) but -0.095.

Interestingly, there is still some hidden underlying order. Equation 21 holds for each row of ratio triplets taken individually, as it must, but also applies to the means of the entire dataset. That is, the geometric mean of the n  $B_1/B_2$  ratios equals the geometric mean of the n  $p_2/p_1$  ratios times the geometric mean of the n  $R_1/R_2$  ratios. Similarly, the arithmetic mean of the log  $B_1/B_2$  ratios equals the sum of the arithmetic means of the two corresponding log ratios. The reader can verify this with the data in Tables 2 and 3. Neither of these mean calculations involves the parameter c. This shows that the behavioral identity underlies this result, that it is independent of the theoretical matching equation, and that it is not due to the fitting procedure, be it least squares or other.

Table 3 Data from Davison and Jones (1995) for Bird 21 on nine concurrent VI VI schedules where marked undermatching occurred (c = 0.54).

$B_1/B_2$	$R_1/R_2$	$p_2/p_1$	$Log B_1/B_2$	$Log R_1/R_2$	$\mathit{Log}\ p_2/p_1$
3.05813	10.11111	0.302452	0.485456	1.004799	-0.51934
0.163176	0.069519	2.34722	-0.78734	-1.1579	0.370554
6.782114	65.66667	0.103281	0.831365	1.817345	-0.98598
0.068611	0.005025	13.65352	-1.16361	-2.29885	1.135245
14.23062	99.00000	0.143744	1.153224	1.995635	-0.84241
0.857934	0.709402	1.209376	-0.06655	-0.14911	0.082561
2.782693	5.451613	0.510435	0.444465	0.736525	-0.29206
0.26971	0.081081	3.326429	-0.5691	-1.09108	0.521978
4.776889	10.11111	0.47244	0.679145	1.004799	-0.32565
	Geometric Means			Arithmetic Means	
1.2939	1.6103	0.8035	0.1119	0.2069	-0.0950

*Note.* Format is as in Table 2. The mean of the log p ratios does not give the same log bias value obtained by the authors (0.00), but other order is still present (see text).



The fitting procedure will, however, influence the value of c and a. Except for the case of strict matching, the identity of Equation 21 does not predict what the fitted values of c and a will be. That will be determined by the particular set of numbers generated by the subject and experimenter, and by the fitting procedure itself. Davison and Elliffe (2009) have shown that the least squares procedure underestimates the sensitivity parameter c. This is due to the variance, unequal variances, asymmetrically skewed distributions among the response and reinforcement measures. The logarithmic transformation ameliorates some of these distortions and is preferable to the raw ratio fits, as Baum (1974, 1979, 1983), too, has discussed, but it does not remove them. A "best" fitting procedure is not yet known or agreed upon.

Because the generalized matching law of Equation 20 describes behavior well, the identity of Equation 21 implies that so might the equation

$$\frac{B_1}{B_2} = a' \left(\frac{p_2}{p_1}\right)^d \tag{28}$$

expressing the response ratios as a function of the probabilities rather than the rates of reinforcement, but with different bias and sensitivity values. Figure 1 compares plots of Equations 20 and 28 using the data in Table 3.

Graphs for the remaining 5 pigeons in the study showed this alternative relationship to be linear in log coordinates, with negative slopes, but with R-squares consistently smaller than with the conventional model (group mean R-square = 0.83 versus 0.95). However, the slope values were consistently closer to "matching" (i.e. to -1.0). Slopes for the conventional model varied from 0.36 to 0.60 with a mean of 0.49. Slopes for the alternative model varied from -0.54 to -1.23 with a mean of -0.92. Bias values were small in both cases.

This finding would give a mixed answer to a simple question such as, "Which variable—rate or probability—is the more important, influential, operative, or causal?" It would appear that both variables are acting simultaneously but in different ways. Clarifying these differences would require a more molecular examination of the underlying factors leading to the particular obtained reinforcement rates and probabilities, which is not the main purpose of this article.

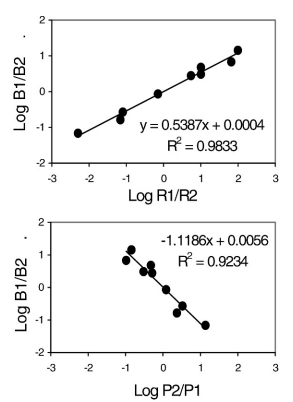


Fig. 1. Data from Table 3 plotting log  $B_I/B_2$  as a function of log  $R_I/R_2$  (top) and as a function of log  $p_2/p_I$  (bottom).

Application to Single Schedules and an Empirical Finding

Catania and Reynolds (1968) exposed pigeons to a number of single VI schedules ranging in value from 10.5 to 420 seconds. They plotted responses per min against reinforcements per hour and obtained monotonically increasing negatively accelerated functions. The identity of Equation 5c states that for each single VI value, rate of response b will equal rate of reinforcement r divided by the probability of reinforcement p. It does not make a specific prediction of how p varies with r across conditions if b also varies across conditions—linearly, nonlinearly or even independently of r. If b is free to vary, this relationship is not forced, but it can be determined empirically. Catania and Reynolds's data were reanalyzed after converting both r and b to rates per min so that their quotient would give the obtained p. The results are plotted on the left side of Figure 2.

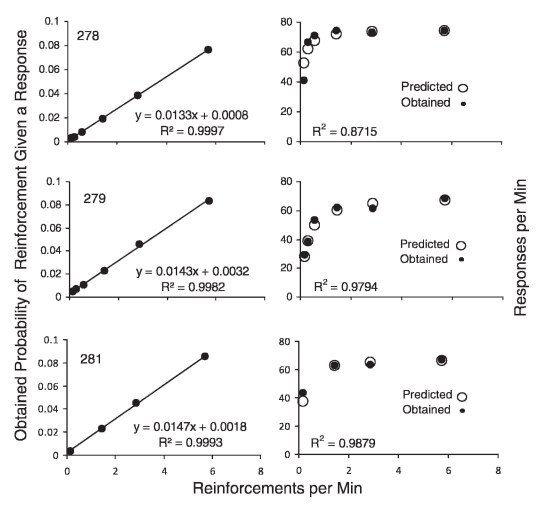


Fig. 2. Data reanalyzed and replotted from Catania and Reynolds (1968) for the last 3 pigeons in their Figure 1. Left axis is the number of reinforcements per min divided by the number of responses per min. Intercept and slope constants from the linear regression lines were used to predict responses per min vs. reinforcements per min in the right column, using an hyperbola based on an identity described in the text.

The data were highly linear in these coordinates and over this range, hence they were fitted with a straight line of the form  $p = \alpha + \beta$  r. Substituting this expression for p into the identity gives the hyperbola.

$$b = \frac{r}{\alpha + \beta r}. (29)$$

Using the fitted intercept and slope constants, the predicted values for b are plotted with the obtained values for b on the right side of Figure 2.

Dividing both the numerator and denominator of this last equation by  $\beta$  gives

$$b = \frac{(1/\beta)r}{\alpha/\beta + r} = \frac{k r}{r + \alpha/\beta} = \frac{k r}{r + \alpha k}$$
 (30)

which can be compared to Herrnstein's theoretical model in Equation 3. Herrnstein's  $r_0$  becomes  $\alpha/\beta$  while  $1/\beta$  closely approximates the asymptotic response rate. For example, in the top two panels of Figure 2 the asymptote appears to be around 75 responses per min and the fitted value for  $\beta$  is 0.0133, whose reciprocal is 75.2. The more exact predicted asymptote is found by evaluating the linear equation when probability p reaches one, and is  $(1 - \alpha)/\beta$  or 75.1



responses per min. Substituting this correction complicates the equations somewhat while making little quantitative difference. With the units used here the absolute values for  $\alpha$  are small. However,  $\alpha$  cannot be zero, or the animal would always respond at its asymptotic response rate regardless of reinforcement rate.

The quantity  $\alpha/\beta$  necessarily has the same dimensions as r, and in this example evaluates to 0.06 reinforcements per min, or 3.61 per hour. This is compatible with Herrnstein's theoretical hypothesis but I leave its interpretation to the reader.

Equations 29 and 30 suggest exploring which classes of experimental variables or "subject strategies" affect  $\alpha$  and  $\beta$  differentially. Experimental variables might include reinforcement magnitude, deprivation level, or response effort. Subject strategies might include proportional changes in the postreinforcement pause, or changes in running rates, within-bout response rates or in the number and durations of between-bout pauses. On VI the subject has minimal control over r but through these can manipulate p.

The linearity between *p* and *r*, and the proportion of variance accounted for suggest the possibility that the obtained probability of reinforcement may be at least as effective a controlling variable as rate of reinforcement even on interval schedules, and allows the hypothesis that the hyperbolic relationship between response rate and reinforcement rate might be more derivative than primary.

### Limitations and Extensions

The identity of Equation 5 is formulated only for the obtained values of the response rate, reinforcement rate, and the conditional probability of reinforcement; not for any of the many other variables known to characterize or influence behavior. Furthermore, it is based only on their totals or arithmetic means, and says nothing about their instantaneous, sequential or distributional characteristics.

This identity and those in Equations 14, 16 and 21 are purely quantitative, atheoretical molar descriptions of how the obtained values of these variables are related. They do not explain, for example, behavioral contrast or choice or why matching occurs, when it does. They do not address, and I will not here address, the vast literature on molecular theoretical models (momentary, local, or

global maximizing, melioration, etc.) proposed as explanations for these phenomena. Neither do I here address the literature of behavioral economics where the same identities apply, with the variables usually inverted and renamed to represent the cost of responding or perhaps the cost of waiting (Collier, Johnson, & Mathis, 2002). However, the identities can serve as checks and guidelines for these molecular theoretical models, and perhaps aid in the interpretation of findings. The algebraic identities dictate that:

- 1) Any quantitative theoretical model that predicts an absolute or local response rate *b* must also equal *r/p*. The two equations can then be solved simultaneously to yield other relationships.
- 2) Any theoretical model that predicts a single  $B_1/B_2$  ratio pair must also equal  $(p_2/p_1)^*(R_1/R_2)$ . If multiple pairs are considered simultaneously the result is more complex but a similar relationship holds for their geometric means.
  - Any theoretical model that predicts a relative response rate  $B_1/(B_1 + B_2)$  must also equal  $(R_1 p_2)/(R_1 p_2 + R_2 p_1)$ , or equivalently  $R_1/(R_1 + m_{12} R_2)$ . If three (or n) components are involved then the summation forms of Equations 15 and 16 apply. However, any one of the components can be temporarily or permanently discarded or ignored and the relationship will hold for the other two (or n-1) components. That is, one can drop  $R_3$  (or  $R_0$ , as is done in the ratio formulation) and the identities will still hold. It is mostly for the single response case that the empirically unmeasurable  $R_0$  has been considered theoretically appealing as a hypothetical variable.

Historically, most operant quantitative models regarded response rate as primarily a function of reinforcement rate (e.g., Herrnstein, Catania, Baum as cited herein). Some more recent models have considered various explicit or implicit reinforcement probabilities (e.g. the Stay/Switch model of MacDonall, 2009). The position taken here is that we need to consider and measure *both* variables. Behavior is presumed to be simultaneously sensitive to both and affects both through interaction and continuous feedback, at least until a stable state is reached.

Although Equation 5 is an equation that must hold, it is only one of many possible equations that may also hold. The identity should become increasingly useful as we discover or devise and incorporate additional equations relating its three variables to themselves or other variables. These additional equations may be empirical, theoretical, or mathematically definitional; and one may anticipate that some of them will be situation-specific rather than universal. Even in the absence of such additional equations the identity has conceptual and computational utility beyond the examples given here.

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APPENDIX

VI data from Catania and Reynolds (1968) showing reinforcement rates, response rates, and the obtained probability of reinforcement p (as their ratio).

Bird	R/hr	R/min	B/min	Þ	$r/(\alpha + \beta r)$
278	8.6	0.143	41	0.0035	52.96
	17	0.283	66.84	0.0042	62.02
	34	0.567	71.15	0.0080	67.97
	86	1.433	74.35	0.0193	72.16
	171	2.850	73.28	0.0389	73.63
	343	5.717	74.69	0.0765	74.41
279	9	0.150	29.11	0.0052	28.06
	17	0.283	38.7	0.0073	39.07
	34	0.567	53.3	0.0106	50.13
	86	1.433	62.3	0.0230	60.49
	171	2.850	61.46	0.0464	64.84
	343	5.717	68.3	0.0837	67.30
281	9	0.150	43.68	0.0034	37.45
	86	1.433	62.39	0.0230	62.67
	86	1.433	63.2	0.0227	62.67
	171	2.850	63.25	0.0451	65.22
	343	5.717	67.05	0.0853	66.60

Note. Right column is the predicted response rate, where r is reinforcement rate and  $\alpha$  and  $\beta$  are the intercept and slope constants shown in Figure 2.

