

DISCOVERING INFLUENTIAL VARIABLES: A METHOD OF PARTITIONS

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A trend in all scientific disciplines, based on advances in technology, is the increasing availability of high dimensional data in which are buried important information. A current urgent challenge to statisticians is to develop effective methods of finding the useful information from the vast amounts of messy and noisy data available, most of which are noninformative. This paper presents a general computer intensive approach, based on a method pioneered by Lo and Zheng for detecting which, of many potential explanatory variables, have an influence on a dependent variable Y . This approach is suited to detect influential variables, where causal effects depend on the confluence of values of several variables. It has the advantage of avoiding a difficult direct analysis, involving possibly thousands of variables, by dealing with many randomly selected small subsets from which smaller subsets are selected, guided by a measure of influence I . The main objective is to discover the influential variables, rather than to measure their effects. Once they are detected, the problem of dealing with a much smaller group of influential variables should be vulnerable to appropriate analysis. In a sense, we are confining our attention to locating a few needles in a haystack.

1. Introduction. Lo and Zheng [18, 19] introduced the backward haplotype-transmission association (BHTA) algorithm, an efficient computationally intensive method of detecting important genes involved in complex disorders. This method, using haplotype information on multiple markers for affected subjects and their parents, was applied to Inflammatory Bowel Disease data [19]. In that application, a total of 235 case-parent trios (each family contains an affected child and his/her parents) and 448 markers (variables) are included in the analysis. Because the proposed method efficiently draws information from both joint and marginal effects, interesting and novel scientific results were obtained, some of them intriguing.

In order to accommodate different types of genetic data (such as in case-control designs, for example), the method has been modified recently to

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other genetic approaches using multiple markers [12, 28]. A brief summary of these methods and their results appears in Appendix S1.

In brief outline, the methods consist of subjecting a small randomly selected group of markers to analysis to see which, if any of these, seem to be moderately associated with the disease. A measure I (defined later in Equation 1) that evaluates the amount of influence of this set of markers is used to quantify their associations with the disease. A stepwise elimination process reduces this set to a smaller set of possibly influential variables which are retained. By repeating this process many times on randomly selected groups of markers, a subset of markers is obtained which frequently appear to be associated with the disease, and this subset is regarded as potentially involved in the disorder.

An advantage of this method consists of avoiding a difficult direct analysis involving hundreds or thousands of markers in favor of a simple but effective analysis repeated many times. Another advantage is that, as opposed to other methods depending mainly on marginal information, this method can make use of both marginal and interactive effects to yield more effective detections.

The main idea applies much more generally than to special genetic problems. In this paper, a general version, which we shall call *Partition Retention*, is proposed to deal with the problem of detecting which, of many potentially influential discrete variables X_s , $1 \leq s \leq S$, have an effect on a dependent variable Y using a sample of n observations on, $\mathbf{Z}=(\mathbf{X}, Y)$ where $\mathbf{X}=(X_1, X_2, \dots, X_S)$.

There exists a substantial literature, especially in engineering journals, on feature and variable selection [5, 9, 11, 15, 23], but much of it is directed toward improving techniques in classification. A set of variables that are useful for classification purpose can be potentially very different from the set of influential variables that we seek to identify in this paper. In our text we will make comparisons with Random Forests [5] and some comments on Multifactor Dimensionality Reduction (MDR) [23]. In our discussion, we include some comments on the interesting technique of Koller and Sahami [15].

In the background is the assumption that Y may be slightly or negligibly influenced by each of a few variables X_s , but may be profoundly influenced by the confluence of appropriate values within one or a few small groups of these variables.

At this stage the object is not to measure the overall effects of the influential variables, but to discover them efficiently. Once these variables have been detected, the problem of dealing with a much smaller group of influ-

ential variables should be vulnerable to appropriate analysis. In a sense we are confining our attention to locating a few needles in a haystack.

The object of this paper is to introduce the general approach, and to indicate that there are many important variations of strategies which may be worth exploring in order to increase the effectiveness for finding influential variables and discarding *impostors*.

Section 2 provides a preliminary illustration of the approach with an artificial example. This is followed by Section 3 which gives a formal presentation of terminology.

Sections 4 and 5 address the following major issues and the novel advantages of our method with simple artificial examples in Section 4 and four more substantial ones in Section 5, where two are based on real data.

1. As has been noted by Guyon and Elisseeff [11], while one of a set of influential variables may have no *causal* effect by itself, it may have an *observable* marginal effect. That observable effect might be small or negligible. Many current methods rely heavily on the presence of strong observable marginal effects and are unlikely to succeed if marginal observable effects are weak. Under certain circumstances, some *impostor* variables with no causal influence may seem to have substantial marginal observable effects.
2. The method we present is sensitive to the combined effects of several influential variables when there are many potential influential candidates. When the number of candidates is very large, our original plan may not succeed in observing the combined effects of several influential variables, and it may be necessary to *thin* out the set of candidates with a preliminary stage where all variables are first considered one or two at a time. In a following stage our method may *resuscitate* influential variables that did not show up early.
3. Our method uses a measure of information related to the multiple correlation (or t test in the case of one explanatory variable). It is more sensitive to influence than the correlation when applied to several variables at a time.

Section 6 is a summary which also includes a discussion of the comparison with Random Forests, and describes some aspects of an interesting procedure by Koller and Sabadi [15]. Finally, an appendix contains some derivations and related results. Except for Appendices A1 and A2, the other parts (Appendices S1-S3) are included in the online supplementary file.

2. Preliminary Illustration. We introduce the partition retention (PR) approach and related terminology and issues by considering a small artificial

example.

EXAMPLE 1. Suppose that an observed variable Y is normally distributed with mean X_1X_2 and variance 1, where X_1 and X_2 are two of $S = 6$ observed and potentially *influential* variables which can take on the values 0 and 1. Given the data on Y and $\mathbf{X} = (X_1, \dots, X_6)$, for $n = 200$ subjects, the statistician, who does not know this model, desires to infer which of the six explanatory variables are causally related to Y . In our computation the X_i were selected independently to be 1 with probabilities 0.7, 0.7, 0.5, 0.5, 0.5, 0.5.

The approach is to partition the 200 observations into $2^6 = 64$ *partition elements*, according to the values of $\mathbf{X} = (X_1, X_2, \dots, X_6)$ with n_i observations in the i -th element. We introduce the measure

$$I = n^{-1} \sum_i n_i^2 (\bar{Y}_i - \bar{Y})^2$$

where $\bar{Y} = \sum_i n_i \bar{Y}_i / n$ is the overall average of Y and \bar{Y}_i is the average of Y in the i -th element. We consider I to be a measure of influence based on how well the partition separates the subjects into relatively homogeneous subsets.

To measure the influence of X_1 on I we can repeat this process by using the coarser partition depending on the other 5 variables, in effect pretending that we don't have X_1 available. The difference, D_1 in the two values of I is regarded as a measure of the influence of X_1 on Y in the presence of the other 5 variables. A decrease in I would suggest that X_1 has a substantial influence. We could repeat this process for each of the other 5 variables. Our procedure consists of discarding from consideration the variable for which the D value is least. We repeat this procedure with the remaining 5 variables and continue discarding until we reach a step where all the D values are positive, at which time we *retain* the remaining variables.

We illustrate the method for a particular data set not presented here. First we standardize Y by subtracting the mean and dividing by the standard deviation, a procedure we find convenient but not essential. Then we obtain the value $I = 2.14$ when all 6 variables were considered. Taking turns, eliminating one of these variables at a time gives us, for the remaining five not eliminated, values of I of 1.46, 1.57, 3.25, 3.32, 3.24, and 3.38, with corresponding D values of 0.68, 0.57, -1.11, -1.18, -1.10 and -1.24. Then we discard variable X_6 which led to the smallest value of D , leaving us with a value of $I = 3.38$. Repeating this process on variables X_1 to X_5 leads to discarding variable X_4 with $I = 5.83$. An abbreviated history of this process

is presented in the first two rows of Table 1 which give the successive values of I and the variables discarded at each stage.

The next two rows of Table 1 involve the same procedure applied to the set of five variables X_2, X_3, X_4, X_5, X_6 . The following two rows treat the case where the variable X_2 is originally omitted from the six. Finally the last two rows treat the case where only the last 4 noninfluential variables are considered in the subset analyzed.

TABLE 1
History of the Discarding Procedure for Four Cases.

Initial set: {1, 2, 3, 4, 5, 6}						
I before discarding	2.14	3.38	5.83	10.76	20.32	9.89
variable discarded	6	4	3	5	1	2
Initial set: {2, 3, 4, 5, 6}						
I before discarding	1.46	2.12	3.34	5.49	9.89	
variable discarded	5	6	3	4	2	
Initial set: {1, 3, 4, 5, 6}						
I before discarding	1.57	2.29	3.36	5.49	8.70	
variable discarded	6	3	4	5	1	
Initial set: {3, 4, 5, 6}						
I before discarding	1.00	1.12	1.13	1.01		
variable discarded	6	3	5	4		

When the influential variables X_1 and X_2 are in the subset subject to the process, they end up as the last items to be discarded. When both are present the initial value of I tends to be larger than when only one is present, and when none are present the initial value of I is still smaller. In the first case our plan retains both influential variables. In the next two cases I increases as we discard, and our retention strategy retains only the last variable kept, 2 and 1 respectively. For the case where the discarding process starts with only the “unimportant” $\{X_3, X_4, X_5, X_6\}$, I has the lowest initial value and does not grow much as variables are discarded.

The strategy of retaining all variables when all D values are positive, *i.e.* when I starts to decrease, would lead to retaining variables X_4 and X_5 in the fourth case. With the relatively small initial value of $I = 1.00$, it might be a good idea to retain none of the variables being studied. In other words, our strategy for retaining variables could be reconsidered. In fact, as we shall note later, values of I substantially greater than 1 signify possible influence, and the values of I at the stopping times were 20.32, 9.89, 8.70 and 1.13, in these four situations above. The relatively modest value of I at the stopping time in the fourth case could be regarded as a signal to not retain the remaining variables.

Because we will be dealing with many candidate variables in more realistic problems, our plan is to take small random subsets of the variables under consideration and subject these to a reduction scheme similar to the one described above. If the retention rate for influential variables will be greater than for noninfluential variables, the influential ones will show up more often in many repetitions of this process, and will be discovered by their high retention rates.

Although X_1 has no marginal *causal influence* by itself, the third case shows that it has a marginal *observable effect* which may also be detected by a simple t test. As we shall see, applying the t test on each candidate variable is computationally cheap, and may locate influential variables with a strong marginal observable effect. But, in the case of many candidate variables, it will allow some noninfluential variables to behave as *impostors*. The t test may not be very efficient in detecting observable effects which depend on interactions, and may fail to discriminate against some of the impostors. However, for problems where S , the number of potentially influential variables under consideration, is not very large, we may be able to calculate the value of I for all possible pairs or even all possible triples, as a way of increasing the sensitivity for detecting influential variables, for which the causal effect depends largely on the interactions of groups of variables, while discriminating against impostors.

3. Formulation outline. If we select a subset or group of m binary valued variables from $\mathbf{X} = (X_1, X_2, \dots, X_S)$, this subset defines a partition Π^* of the sample of n observations into $m_1 = 2^m$ subsets which we shall call partition elements, $\{A_1, A_2, \dots, A_{m_1}\}$, corresponding to the possible values of the collection of these m binary variables. For simplicity and without causing confusion, we shall use $\{X_1, X_2, \dots, X_m\}$ to denote the subset of selected variables. Each partition element A_j corresponds to a possibly empty subset of n_j Y values and $\sum n_j = n$. Each nonempty partition element A_j yields a mean value \bar{Y}_j and the overall mean is $\bar{Y} = \sum n_j \bar{Y}_j / n$. Let

$$(1) \quad I_{\Pi^*} = n^{-1} \sum n_j^2 (\bar{Y}_j - \bar{Y})^2.$$

If I_{Π^*} is unduly large, an expression to be explained later, we suspect that some of the m variables may have an influence on Y .

Suppose that we now introduce another binary variable from the original set of S potentially influential variables, which we shall call X_0 for notational convenience. This leads to a more refined partition $\Pi = \{A_{jk} : 1 \leq j \leq 2^m, k = 0, 1\}$ where A_{j0} corresponds to that part of A_j with $X_0 = 0$ and A_{j1} corresponds to that part of A_j with $X_0 = 1$. Now let \bar{Y}_{jk} be the mean of the

n_{jk} elements in A_{jk} and hence, $n_j = n_{j0} + n_{j1}$, and $n_j \bar{Y}_j = n_{j0} \bar{Y}_{j0} + n_{j1} \bar{Y}_{j1}$. We refer to Π^* and Π as the *coarse* and *refined* partitions respectively. The measure I_{Π^*} is now replaced by

$$(2) \quad I_{\Pi} = n^{-1} \sum n_{jk}^2 (\bar{Y}_{jk} - \bar{Y})^2,$$

and

$$(3) \quad D_I = \frac{1}{2} (I_{\Pi} - I_{\Pi^*})$$

can be regarded as a measure of how much X_0 contributes in influence on Y in the presence of $\mathbf{X} = (X_1, X_2, \dots, X_m)$. It is easy to see that

$$(4) \quad D_I = -n^{-1} \sum n_{j0} n_{j1} (\bar{Y}_{j1} - \bar{Y})(\bar{Y}_{j0} - \bar{Y}).$$

Thus, D_I tends to be negative when both means in the refined partition elements tend to be on the same side of \bar{Y} as in the coarse partition element from which the refined elements came. We would expect that if the new variable contributes influence on Y , then D_I would tend to be positive.

In Appendix A1, we calculate the expectation of D_I conditional on the partition sample sizes, in a more general framework described at the end of this section. This expectation consists of the difference of two positive quantities plus one which is relatively small and can be estimated. Neglecting this small term, we see that if the new variable has no influence on Y , other than random noise, the expectation of D_I will be nonpositive, and strictly negative if there are some influential variables in the selected subset $\{X_1, \dots, X_m\}$. On the other hand, if the new variable X_0 contributes influence on Y and the old ones don't, then the expectation of D_I will be positive.

Our policy is not that of adding new variables to our group of m variables, but one of deleting variables from an initial group. Thus if we start with $m + 1$ variables, we consider the effect, *i.e.*, D_I , of using the coarser partition obtained by eliminating one of the $m + 1$ variables. The one with the smallest D_I is then eliminated, and we repeat this procedure on the remaining m variables. We may continue eliminating until we are satisfied by some criterion (*e.g.* when all the remaining D_I are positive), that most of the remaining variables are *good* candidates for being influential and should be retained.

The set of $m + 1$ variables will be selected at random from the original set of S variables. The retention procedure is to be carried out many times. We can observe which of the original variables is retained with an unusually high frequency among those retained, and use these for further analysis.

Because we expect to repeat this procedure many times, the approach is computationally intensive. It may be possible sometimes to use the initial value of I_{Π} to decide whether a randomly selected group of m variables is worth pursuing with the elimination scheme, thereby avoiding the calculations required for the successive eliminations. Sometimes, it may be sensible to stop after the first step in the elimination process and select the variables which lead to large values of D_I .

While our discussion was confined to binary valued explanatory variables, there is no such essential limitation. In fact the applications of Zheng et al. (2006) [28] used SNP genotypes, which assume three possible values, as explanatory variables. We could easily partition based on discrete valued explanatory variables. Then, if X_0 assumes a finite set of values, say 1 to r , the equation for D must be adjusted to give

$$D_I = -n^{-1} \sum_i \sum_{j < k} n_{ij} n_{ik} (\bar{Y}_{ij} - \bar{Y})(\bar{Y}_{ik} - \bar{Y}).$$

where the partition element A_{ij} is that subset of A_i where $X_0 = j$ and has n_{ij} elements averaging \bar{Y}_{ij} . If we define W_{ij} as the sum of all Y variables in the partition element A_{ij} and W as the sum of all the Y values, then we may write

$$D_I = -n^{-1} \sum_i \sum_{j < k} (W_{ij} - (n_{ij}/n)W)(W_{ik} - (n_{ik}/n)W).$$

In those cases where the explanatory variables are continuous, the investigator could select cutoff points to separate the possible values into a few discrete subgroups. This process might involve subjective decisions. Such subjective decisions could also be applied to a more complicated case where a pair of discrete or continuous variables may be assigned to a small number of discrete values depending on the expert opinions of the investigator.

In Appendix A1, we deal with two models. In the first, the *random* Y model, we assume that, the distribution of Y depends on \mathbf{X} which may be random or may be selected in advance as part of an experimental design. In the second, the *specified* Y model, the values of Y are selected in advance. For example in case-control experiments we select the number of cases and controls and examine the related values of \mathbf{X} .

An alternative measure of influence, one more aligned with standard analysis of variance calculations, is given by

$$(5) \quad J_{\Pi} = n^{-1} \sum n_{jk} (\bar{Y}_{jk} - \bar{Y})^2.$$

The use of J to compare two sets of $m > 1$ variables for influence is the same as using the multiple correlation of Y on these variables. For $m = 1$ using J , the squared correlation coefficient and the absolute value of Student's t will give almost the same comparisons when $m = 1$ and $r = 2$. The same could be said for the chi-square statistic and J when the variable X_i assumes more than 2 values or $m \geq 2$.

4. General Comments. Our object is to locate influential variables. Whatever method we use there is always the possibility that, among the ones we characterize as influential, there will be some that are impostors. When S is large, it may be necessary to go through several stages of an elimination process, eliminating many of the candidate variables from consideration at each stage.

Our major method is most effective when the subgroup of variables randomly selected has a reasonable probability of containing more than one of the interacting influential variables. But for this to be the case the size of the randomly selected group of variables m , should be a substantial portion of S , the number of candidate variables. When the sample size n is not very large, m has to be modest for our approach to be effective, for otherwise, there will be many partition elements that are empty or have only one subject. For example if n is 200, we would like to have no more than 50 to 100 partition elements. In the case of binary valued X values, that means that we should consider subgroups of 6 or 7 variables at a time. But if S is 1,000, it is rarely the case that a randomly selected subgroup of 7 variables will contain more than one of a small number of interacting influential variables. In that case, we have to thin out the set of competitive variables before we can hope to have the advantage anticipated when our subgroup frequently has more than one of the interacting influential variables.

One way to thin out the candidates is to apply I or the t test to one explanatory variable at a time, and to concentrate energy on those which indicate strong marginal observable effects. If S is not too huge, we may even consider all possible pairs and concentrate on those variables which appear in many *high ranking* pairs.

In the examples of Section 5 we will show how the partition retention method applied after thinning can *resuscitate* influential variables with mediocre ranking by marginal considerations.

An issue of importance is the relative powers of using $m = 1, 2$ and 7 . Another issue is whether a variable, which is an impostor using one of these methods is likely to fail by some of the others. If that is the case, then we can hope to weed out impostors by combining the various techniques.

Finally, when our analysis points to some likely candidates for being influential, we would like to have some way of deciding how plausible our results are. One way would be to add randomly selected additional variables which should have no relationship to the dependent variable, and see how their presence affects the various statistics used. This approach does not seem to be as reliable as simply permuting the values of the observed dependent variable Y . This latter approach does not upset the relationships among the S explanatory variables, whereas the first proposal would require assuming independence or some arbitrarily selected correlations.

We have used the word impostor to suggest that some explanatory variables which are not causally related to the dependent variable tend to be easily confused with influential variables. The following simple artificial example provides some insight on the circumstances that can lead to impostors.

EXAMPLE 2. There are three independent explanatory variables X_1, X_2, X_3 which take on the values 0 and 1. Let $Y = X_1X_2$, and in a sample of n cases, $n_{ijk} = np_{ijk}$ is the number of cases where $X_1 = i, X_2 = j, X_3 = k$. We use the subscript d to replace the conventional dot to indicate summation over the corresponding index. For example $p_{11d} = \sum_k p_{11k} = p_{110} + p_{111}$. (In small print the symbol d is easier to read than a dot.) We also relate the p values with the probabilities they estimate and so we may write $p_{11d} = p(X_1 = 1, X_2 = 1) = p(X_1X_2 = 1) = p(Y = 1)$.

Using the partitions based on X_1 we obtain

$$(6) \quad I_{X_1} = 2n(p_{11d}p_{0dd})^2$$

and using the noninfluential X_3 we have

$$(7) \quad I_{X_3} = 2n(p_{11d}(p_{111}/p_{11d} - p_{dd1}))^2.$$

The ratio I_{X_3}/I_{X_1} depends on the ratio of $p(X_3 = 1|X_1X_2 = 1) - p(X_3 = 1)$ to $p(X_1 = 0)$. In Appendix S2, we prove that, assuming independence of X_3 and X_1X_2 , the asymptotic distribution of the first term of this ratio has mean 0 and variance $n^{-1}P(X_3 = 1)P(X_3 = 0)P(X_1X_2 = 0)/P(X_1X_2 = 1)$. Thus the probability that the random noninfluential variable X_3 will act as an impostor is small if n is large. However when S is very large and n is modest, there may be several impostors.

Note that in most real problems the dependent variable is typically not completely determined by the causal variables. There is usually some random variation and the signal to noise ratio is of consequence. In our example there

was no noise, and the signal to noise ratio is effectively infinite. Even so it is possible to have impostors.

The following is an artificial example where two explanatory variables determine the dependent variable but neither one has a marginal observable effect.

EXAMPLE 3. This is a variation of Example 2, where the data consist of (Y, X_1, X_2) for n observations, and $Y = X_1X_2 + (1 - X_1)(1 - X_2)$. Then

$$(8) \quad I_{X_1} = 2n(p_{11}p_{01} - p_{10}p_{00})^2.$$

If (X_1, X_2) takes on the values $(1,1)$, $(1,0)$, $(0,1)$ and $(0,0)$ with probabilities q_1, q_0, q_0, q_1 , then the expression $p_{11}p_{01} - p_{10}p_{00}$ is asymptotically normal with mean 0 and variance $2q_0q_1/n$. In effect, the marginal observable effects of X_1 and of X_2 are negligible even though both variables are influential. This lack of marginal observable effect depends on a certain amount of symmetry in the causal mechanism and on the distribution of explanatory variables.

It is useful to observe that if a group of noninfluential variables are distributed independently of a dependent variable Y which is standardized to have sample variance 1, then I will be distributed roughly like a weighted sum of independent chi-squares with one degree of freedom. Also the distribution of J conditional on m' , the number of nonempty partition elements, will have the approximate distribution of a chi-square with m' degrees of freedom divided by n . More precise statements and derivations are presented in Appendix A2. These results provide a clue as to when a group of variables are likely to contain some influential ones.

Some experimental results, one listed in Appendix S3, suggest that both of these measures tend to have roughly the same ability to detect influential variables when the number of partition elements is small and of comparable sizes. However, for a special alternative to independence, the ratio of sensitivities of I and J depend heavily on $m' \sum (n_i/n)^2$, which attains a minimum of 1 when all the partition elements have equal sample sizes, and a maximum close to m when most of the observations are concentrated in one partition element. The advantage of I over J depends on the variance of the frequencies n_i (see Appendix S3 for detail).

At this time, we hesitate to present a specific program to carry out our aim of detecting influential variables. Each applied problem has special needs which may call for variations on the procedures we described.

5. Examples. In this section we present four examples. One is an extension of Example 3 of the last section and involves 10 influential variables.

Another is a more realistic one featuring two small groups of influential variables. Two are based on a real data set for Rheumatoid Arthritis. A major advantage of the artificial ones is that *truth* is known and the properties of the methods can be evaluated for those examples. By simulation we can see how our methods respond as parameters of the model in the example vary. We have the opportunity to compare the results with those of Random Forests (RF), a method pioneered by Breiman [5]. For examples based on real data, we have to rely on supplementary information to determine the reliability of our conclusions.

Our simulations give rise to a great deal of data. For the sake of this presentation, devoted to introducing the partition retention approach, we will occasionally omit some useful information in an attempt to avoid overwhelming the reader. In particular, we depend heavily on ranking the influential variables among all the candidates, and rarely present the measures used for the ranking. Thus the reader will seldom see those situations where there is a precipitous drop in the measure as one goes from one variable to the one ranked next.

Our comparisons often will involve rankings of variables based on $|t|$, I_1 , I_2 , I_{2f} , I_7 and RF. Here t is the Student's t test statistic and behaves very much like I_1 , which is the marginal measure I based on $m = 1$. For I_2 , we rank the $S(S - 1)/2$ pairs of variables using I based on $m = 2$. There is no unique way to rank the importance or influence of the individual variables given this ranking of pairs. Two alternatives suggest themselves. The first, somewhat ambiguously labeled I_2 , is measured by the number of variables that have appeared at least once in the ranking of the pairs before the candidate appears. The potential trouble with this method is the possibility that a very strong candidate in one group of influential variables will carry some noninfluential variables with it before we see indications from influential variables in another group. An alternative ranking I_{2f} , depends on the number of times a candidate variable appears in the n_r most highly ranked pairs where n_r is a substantial portion of the number of pairs.

Given a data set, $|t|$, I_1 , I_2 and I_{2f} are determined. The partition retention method with $m = 7$, yielding I_7 , is random since it depends on the random sample of n_s subsets of m variables. Good choices of n_s would depend on how far apart are the frequencies of retention of influential and noninfluential variables. While real problems could use sequential methods to help select n_s , we have generally settled almost arbitrarily on $n_s = 30,000$ or $20,000$ for many of the experiments presented here. Similarly, in comparisons with random forests, we have taken the number of variables sampled at each node, m_t , to be 7 and the number of trees n_t to be 20,000. Results for random

forests seem to be insensitive to variations in these parameters. The rankings for random forests is given by RF.

Example 4 is an artificial example, representing an extension of Example 3 to deal with 10 well balanced influential variables in a set of 500 variables with 400 observations on each. We shall see that marginal methods give poor results.

EXAMPLE 4. The $S = 500$ variables X_s are binary valued with values 0 and 1. The first 10 are influential. The number of ones among these, R , is uniformly distributed from 1 to 9. The subset of R of these 10 variables to be equal to one is chosen at random with equal probability from among all such subsets. The remaining 490 variables are independent and each is chosen to have its probability of one to be uniformly distributed from 0.4 to 0.6. Given R , the dependent variable, Y is normally distributed with mean and variance equal to $4(R(R - 1) + (10 - R)(9 - R))$. Here, the sample size $n = 400$.

The ranks of the influential variables are listed in Table 2 when the methods $|t|$, I_1 , I_2 , I_7 and RF are applied.

TABLE 2
Ranks of Influential Variables Using $|t|$, I_1 , I_2 , I_7 and RF. Notation “ r ” is short for rank.

vars	1	2	3	4	5	6	7	8	9	10
$r t $	162	281	363	69	370	52	493	337	183	290
rI_1	159	279	361	65	369	50	493	335	183	288
rI_2	8	9	3	1	6	5	4	7	2	18
rI_7	39	123	144	35	130	33	154	161	67	45
rRF	127	363	213	51	208	48	221	220	186	266

In summary, $|t|$ and I_1 are in close agreement and the ranks they give are essentially those of a random sample of 1 to 500. There are no first order observable effects, which is to be expected given the construction. On the other hand I_2 is almost perfect in identifying the influential variables. Only the tenth is superseded by 8 impostors. The I_7 and RF methods did not do as well as I_2 , but better than I_1 . Furthermore, in this example I_7 seems to do considerably better than RF, suggesting that RF is more dependent than I_7 on strong marginal observable effects. The same calculations were done on the data sets consisting of the first 100 and the first 200 observations. The results for I_2 deteriorate slightly as the sample size decreases, allowing 13 impostors. For I_7 and RF we still seem to do better than chance, but not by very much.

One way of testing for influence is to introduce noisy variables and see what effect these have on I_1 . Another is to make comparisons with the methods applied after Y is randomly permuted a number of times. Since we know truth in this case, these methods are not required, but we demonstrate a couple of exercises. In Table 3 we apply I_1 and I_2 to the variables 11 to 500 and see how the noisy variables 11 to 20 are ranked. In Table 4 we apply I_1 and I_2 to variables 1 to 500 and see how variables 1 to 10 are ranked after subjecting Y to a random permutation.

TABLE 3
Ranks of Noninfluential Variables 11 to 20 by I_1 and I_2 .

vars	11	12	13	14	15	16	17	18	19	20
rI_1	207	290	345	88	190	374	466	251	158	321
rI_2	370	247	348	258	33	152	63	386	28	343

TABLE 4
Ranks of Variables 1 to 10 under I_1 and I_2 after Y is Randomly Permuted.

vars	1	2	3	4	5	6	7	8	9	10
rI_1	251	374	485	283	392	338	333	430	265	465
rI_2	306	433	412	241	311	293	57	217	277	340

The comparison between the results in Tables 3 and 4 with those of I_2 in Table 2 is striking. The comparison with those of I_7 and RF are less striking but apparent. In a real data problem, a number of such randomized variations of the original data set can be used to estimate the false discovery rate when this method is applied to real data. Such an application appears later in Example 7.

We will now introduce Example 5 which has two small groups of 3 and 4 influential variables among 1,000 candidate binary valued variables. We consider two major aspects. First we examine the average behavior of some of the methods, as sample size and signal strength change. Here we find that average ranks are not very informative, since one case with a large rank will hide the fact that most of the time the rank is small. Thus we report both the average and median ranks and values of I . Second, on the assumption that a better understanding of the intrinsic variabilities due to the underlying model and due to the analysis would come from looking at a few examples in detail, we also study five data sets generated by one of the models. Here we explore the ability, by using I_{2f} or a variation of I_7 , to resuscitate influential variables previously neglected

EXAMPLE 5. The vector \mathbf{X} has 1,000 components which assume the values 0 and 1. The first 7 consist of two sets of influential variables which

interact slightly. The dependent variable Y is normally distributed with mean μ and standard deviation σ where

$$\mu = \max(\mu_1, \mu_2) + 0.1(\mu_1 + \mu_2)$$

and

$$\sigma = \max(\sigma_1, \sigma_2)$$

with $\mu_1 = \mu_0 X_1 X_2 X_3$, $\mu_2 = 1.5\mu_0 X_4 X_5 X_6 X_7$, $\sigma_1 = 1 + X_1 X_2 X_3$ and $\sigma_2 = 1 + 2X_4 X_5 X_6 X_7$. The binary valued explanatory variables are independent of each other and take on the value of 1 with probabilities .4, .5, .6, .35, .45, .55 and .65 for the seven influential variables. The probabilities for the remaining 993 variables are randomly uniformly selected in the range of 0.4 to 0.6. In this example, there is a slight interaction between the rare dual effects of the two groups of influential variables. After the data set is observed, Y is normalized to have sample mean 0 and variance 1.

First we will describe results based on 400 simulations of 4 *conditions*. Then we will explore in depth 5 cases for one of these conditions. The four conditions involve sample sizes 200 and 400, and the values 4 and 6 for μ_0 . For the four conditions we had subsamples of $m = 7$. We start with the results using the marginal methods based on one or two variables

Table 5 presents, for each influential variable, the rank it gets among the 1,000 variables when each is subjected to the t test and when each is evaluated by I_1 . This table is based on 400 data sets corresponding to each of 4 conditions.

Note that the first group of influential variables gets better average results than the second group, and within each group the variables with lower probabilities of 1 tend to do better. Given that the first group is influential about one eighth of the time while the second is influential about half as often, it is natural to expect that elements of the first group will be easier to detect in spite of the somewhat weaker mean signal (mean 4 instead of 6 when $\mu_0 = 4$, and 6 instead of 9 when $\mu_0 = 6$). If we think of the other variables in a group as providing support to a designated variable being tested, the variables which show up less frequently are being more strongly supported by the more prominent members of its group. That is a heuristic explanation for the second phenomenon.

The two methods, using the t test and I_1 give comparable results. This partially supports our claim that the use of I is expected to be preferred to J when there are many unevenly occupied partition elements, but not otherwise. It seems that the increase of μ_0 from 4 to 6 has less effect than the doubling of the sample size, n . Even if the mean grows very large, there is a

TABLE 5

Ranks of the influential variables using $|t|$ and I_1 based on a single variable. Means and Medians of the ranks using 400 data sets with $S = 1,000$. Four cases involve $n = 200$ and 400 and $\mu_0 = 4$ and 6.

variable	1	2	3	4	5	6	7	n	μ_0	statistic
mean	14.95	37.74	87.53	61.46	110.33	168.91	232.51	200	4	$ t $
median	2.00	5.00	22.50	8.00	24.00	62.00	134.50			
mean	15.57	37.00	90.25	67.06	110.11	168.63	245.70			I_1
median	2.00	4.00	25.00	9.50	24.00	62.50	154.00			
mean	5.61	18.95	55.76	43.21	81.55	130.26	200.07	6		$ t $
median	1.00	3.00	13.00	5.00	17.00	47.00	98.00			
mean	5.88	18.37	57.92	47.88	81.33	129.88	212.23			I_1
median	1.00	3.00	14.00	7.00	17.00	47.00	109.00			
mean	1.82	4.99	13.55	7.44	23.54	51.12	116.78	400	4	$ t $
median	1.00	3.00	4.00	3.00	6.00	15.00	37.50			
mean	1.84	4.86	14.19	8.60	23.32	50.92	125.17			I_1
median	1.00	2.00	4.50	3.00	6.00	15.00	46.00			
mean	1.64	3.97	8.40	4.86	12.93	28.37	89.84	6		$ t $
median	1.00	3.00	4.00	3.00	5.00	9.00	29.00			
mean	1.66	3.83	8.78	5.52	12.73	28.15	97.81			I_1
median	1.00	2.00	4.00	3.00	5.00	9.00	33.00			

limited range of improvement for a fixed sample size. In a sense, the experiment where the dependent variable is a two valued deterministic function of the influential variables, corresponds to a problem of our type where the standard deviation of Y given X is zero or where the mean is effectively infinite. But even there we can not eliminate impostors with a finite sample size.

We present some of the results from calculating the value of I_2 for pairs of variables in Table 6. More precisely, we calculate I_2 for all 499,500 pairs, and rank them in descending order. The value of I_2 and the rank r for a given pair of influential variables, say 4 and 7, are obtained. Finally, we determine the rank, by which all of the influential variables have shown up, and the corresponding value of I_2 . We carry out these calculations, calculating means and medians for 400 simulated data sets for each of the four conditions, $n = 200$ and 400, and $\mu_0 = 4$ and 6. For simplicity and to save space we show the results for only 7 of the 21 pairs of influential variables in Table 6. For comparison purposes, keep in mind that the distribution of I for a set of noninfluential variables is approximately that of a weighted sum of independent chi-square variables with one df, and hence has mean about 1 and variance about $\sum 2(n_i/n)^2$.

Table 6 shows that sample size has a large effect on uncovering influence,

TABLE 6

Means and medians, based on 400 data sets, of values of I_2 and the ranks, r , for a few or the 21 pairs of influential variables.

pairs	(1,2)	(1,3)	(2,3)	(1,4)	(4,5)	(5,7)	(6,7)	final
$n = 200, \mu_0 = 4$								
mean I_2	8.34	7.74	6.25	6.43	4.40	3.18	2.68	4.85
med. I_2	8.06	7.32	6.05	6.31	4.11	2.88	2.48	4.63
mean r	758.45	921.87	4897.03	392.86	13178.51	42790.07	53546.21	1034.57
med. r	3.00	6.00	50.00	46.50	1215.00	5697.50	11065.50	514.00
$n = 200, \mu_0 = 6$								
mean I_2	9.79	8.91	7.46	7.57	5.39	3.87	3.15	5.60
med. I_2	9.51	8.57	7.09	7.60	5.22	3.49	3.01	5.44
mean r	226.14	278.58	2074.82	170.89	7688.84	23337.26	37028.30	507.66
med. r	2.00	3.00	25.00	16.00	460.00	2923.00	5688.00	274.00
$n = 400, \mu_0 = 4$								
mean I_2	15.55	14.23	12.20	12.20	8.58	6.26	5.22	8.68
med. I_2	15.44	14.08	12.07	12.11	8.37	5.85	4.86	8.47
mean r	16.84	48.54	212.85	22.03	1197.38	44268.99	9317.81	177.03
med. r	2.00	2.00	4.00	4.00	101.00	1207.50	2167.00	70.50
$n = 400, \mu_0 = 6$								
mean I_2	18.77	17.32	14.54	14.67	10.23	7.41	5.95	10.21
med. I_2	18.51	17.06	14.30	14.57	10.11	7.22	5.59	9.93
mean r	14.01	27.80	87.34	7.61	841.75	3315.74	5092.48	141.41
med. r	1.00	2.00	4.00	4.00	67.50	1096.50	2124.00	47.50

The column “final” refers to the rank at which all 7 influential variables have appeared at least once.

and signal to noise ratio has a relatively small effect. For these conditions, many pairs have precedence over the ones where all the influential variables have finally shown up. This table indicating that we need a meaningful way of using pairs to rank single variables suggested our use of I_{2f} to supplement I_2 . In Table 7, we apply I_{2f} to 200 data sets for the condition $n = 400$ and $\mu_0 = 4$ with the two values of $n_r = 2,000$ and $4,000$. The results using I_{2f} are substantially better than those using I_1 and I_2 , and a little better than for I_7 . In other applications with 1,000 variables, we tend to use $n_r = 5,000$ more or less arbitrarily. Presumably there may be a rational way of selecting an appropriate value of n_r , but this question has not yet been examined.

Next the partition retention method described in the early sections of this report was applied for the case where $n = 400$ and $\mu_0 = 4$. This was applied with 20,000 random subsets of 7, for each of 200 data sets. For each data set the influential variables were ranked according to how many times they were retained. The means and medians of these ranks are presented in Table 8.

These result seem to be worse than those using the t test for the first 3 variables but better for the latter 4. The reason for this is not clear, but the

TABLE 7

Means and Medians of the ranks of influential variables, using frequency of appearance in the n_r most high ranked pairs for 200 data sets. In this case $n = 400$ and $\mu_0 = 4$.

variable	$n_r = 2,000$		$n_r = 4,000$	
	mean	median	mean	median
1	1.86	1.00	1.64	1.00
2	4.29	2.00	6.32	2.00
3	10.62	5.00	11.20	4.00
4	6.60	3.00	10.91	4.00
5	15.04	6.00	27.89	6.00
6	16.74	8.00	39.51	12.00
7	24.84	12.50	51.52	28.50

TABLE 8

Means and medians, based on 200 data sets, of the ranks of each of 7 influential variables, ranked according to the number of retentions in 20,000 samples of 7 by the Partition Retention Scheme when $n = 400$, $\mu_0 = 4$ and $m = 7$.

variable	1	2	3	4	5	6	7
mean	19.42	39.94	86.02	17.73	6.81	8.60	22.92
median	5.00	8.00	16.50	7.00	3.00	3.50	6.00

difference suggests that, used in tandem, the two approaches will have some effect in detecting and eliminating impostors.

A key question concerns how to take advantage of the partial information gained from the marginal observable effects. One way is to reduce the number of plausible candidate variables, so that our methods can apply higher order interactions to help detect influential variables. Another way is to use likely candidates to resuscitate influential variables that have not yet shown up well. To investigate these possibilities we will concentrate on a few data sets. For each of these we will apply various techniques to see how well these methods work.

First, Table 9 provides a list of the 30 most favored candidates by I_1 , I_2 , I_{2f} , I_7 and RF, using $m = 1$, $m = 2$, $m = 2$ with $n_r = 5,000$, $m = 7$ with $n_s = 20,000$, and $m_t = 7$ with $n_t = 20,000$. Note that 30 was selected mainly to facilitate presentation, and that in most comparable problems a larger number would usually be more appropriate at this thinning stage.

One of the methods of using a reduced list consists of applying the partition retention method to a sample of variables, 3 of which are selected from a reduced list of the top 10 candidates and 4 of which are selected from the variables not in the list. This method increases the probability of getting two influential variables in the sample, one of which may not yet be in the

TABLE 9
30 most highly ranked variables from five methods for each of 5 data sets.

data set 1					data set 2					data set 3					data set 4					data set 5				
I_1	I_2	I_{2f}	I_7	RF	I_1	I_2	I_{2f}	I_7	RF	I_1	I_2	I_{2f}	I_7	RF	I_1	I_2	I_{2f}	I_7	RF	I_1	I_2	I_{2f}	I_7	RF
6	4	5	3	4	4	4	1	1	4	1	1	1	2	1	2	1	1	2	4	1	1	1	4	1
5	5	6	4	5	1	7	4	4	1	2	2	2	5	4	4	2	2	6	2	3	2	3	1	5
4	6	4	5	6	8	1	7	3	8	4	3	4	1	2	472	3	4	676	1	5	3	5	5	4
3	7	3	870	1	7	5	8	5	3	5	4	5	4	5	1	218	472	472	472	2	593	2	163	3
2	1	2	913	3	3	584	3	628	7	984	915	984	3	984	3	4	3	1	5	4	888	4	999	2
870	268	870	288	2	628	8	628	8	628	3	676	3	314	314	677	472	677	462	677	593	5	593	3	593
1	182	1	809	106	2	6	2	469	2	874	7	314	759	207	6	462	6	4	3	163	4	163	2	163
106	673	346	2	623	674	3	542	794	5	314	97	874	984	358	676	22	676	668	814	462	7	462	593	48
660	3	106	6	870	469	2	674	2	690	759	609	358	874	874	5	962	5	3	668	873	462	999	462	873
789	106	660	106	660	690	614	690	542	6	358	790	759	358	175	462	6	462	677	6	999	732	873	661	999
346	341	623	251	789	542	269	469	7	611	730	984	730	376	702	668	853	668	5	676	143	999	143	240	462
623	251	789	623	403	233	628	6	674	63	427	42	929	427	957	100	627	100	956	100	48	941	48	873	79
288	687	288	403	520	6	961	233	611	661	251	571	427	966	730	571	682	248	571	462	661	163	888	721	798
800	2	520	673	195	611	818	5	813	542	376	5	717	657	759	713	362	956	713	178	888	306	661	893	143
573	820	573	789	288	603	674	603	661	469	929	168	207	730	3	956	916	571	276	742	610	38	233	233	467
520	243	800	962	346	5	562	152	233	794	717	909	251	499	337	248	540	703	703	224	233	976	287	499	610
809	913	962	660	809	661	153	611	603	750	207	771	657	929	391	178	204	713	937	713	79	136	721	287	888
962	454	809	7	573	262	469	63	614	233	175	592	775	308	987	937	629	853	100	248	941	869	79	34	941
944	463	165	346	944	405	872	262	858	674	775	767	376	853	929	742	874	742	814	853	499	679	610	48	233
913	563	403	944	800	63	690	794	690	603	957	522	175	280	775	703	677	606	886	909	346	79	6	143	661
165	800	913	628	48	152	294	661	767	423	455	376	455	455	975	924	590	924	853	924	929	143	346	346	470
403	288	944	1	407	813	418	750	6	143	660	864	957	207	280	596	5	596	178	956	128	312	893	732	137
182	870	182	800	905	794	913	143	262	262	657	856	592	649	717	853	124	178	596	937	240	315	929	79	929
195	287	7	520	82	143	405	858	152	858	68	649	14	957	251	551	571	737	737	276	287	605	941	919	308
251	369	195	799	962	353	143	351	760	119	280	726	975	175	376	814	862	814	28	703	893	800	128	128	476
827	702	673	573	165	717	93	405	693	601	975	874	34	661	524	662	741	937	51	422	470	721	470	758	302
140	799	140	287	913	750	750	460	119	152	337	150	499	251	880	909	676	456	456	662	721	893	240	136	499
7	140	827	165	893	858	323	717	750	405	928	401	660	767	447	737	34	224	741	571	136	75	283	929	55
673	216	251	82	827	601	252	767	63	813	499	459	308	775	502	224	889	227	256	92	283	817	312	983	893
628	573	144	195	976	802	687	693	351	534	702	581	380	126	455	456	956	869	498	728	137	815	137	137	283

reduced list. We use the reduced list of the 10 top variables. These will show up in 3/10 of the samples, while the ones not in the list will only appear in about 4 out of 1,000 times. While 20,000 seemed reasonably large for applying I_7 using $m = 7$, our candidates for resuscitation will only show up 80 times, and will be paired with a given variable of the reduced list only about 24 times. Of course, if the reduced list has two variables of a group, then a member of the group not in the reduced list may pair up with one of the two more often. However, 24 plus or minus about 5, is not a good basis for discriminating between influential variables and impostors. We have used 100,000 trials to get more opportunities for observing interactions, although both the 20,000 in the first application and 100,000 here seem a bit modest.

TABLE 10

Five new candidates for each of 5 methods on five data sets derived from the use of I_7 on the top 10 candidates.

data set 1					data set 2					data set 3					data set 4					data set 5				
I_1	I_2	I_{2f}	I_7	RF	I_1	I_2	I_{2f}	I_7	RF	I_1	I_2	I_{2f}	I_7	RF	I_1	I_2	I_{2f}	I_7	RF	I_1	I_2	I_{2f}	I_7	RF
7	2	7	7	7	5	690	5	7	413	7	5	7	7	7	7	676	7	703	462	128	535	128	869	128
881	520	944	1	513	985	674	985	985	782	369	314	369	369	3	703	677	703	100	7	299	177	558	7	299
332	623	623	660	789	915	628	915	674	56	154	874	154	326	759	294	956	100	853	676	7	999	869	505	505
520	972	789	520	543	405	639	603	590	690	308	984	918	126	154	100	853	853	294	853	869	505	299	299	7
543	288	968	623	322	687	794	405	6	628	326	812	126	154	369	853	5	294	834	956	38	55	7	977	869

The 10 members of the reduced list are sampled frequently and are bound to be retained often. But some are retained so markedly less often than others that they deserve to be eliminated. For this presentation, we hesitate to do so in order to avoid unnecessary complications. Instead, we will simply adjoin the five next most frequently retained ones to the list of 10. These five appear in Table 10. Because they were at a disadvantage in the first resuscitation step, we eliminate that disadvantage in the next step by selecting 3 of the 15 in addition to 4 of the remaining 985 variables for each sample of 7. After this step there may be a rearrangement of the top 15, and a few new contenders may appear after these 15.

TABLE 11

Ranks of influential variables for five methods applied to 5 data sets. Initial ranks, rI and rRF , and ranks $ud1$ and $ud2$ after resuscitations.

var	data set 1							data set 2							data set 3							data set 4							data set 5						
	1	2	3	4	5	6	7	1	2	3	4	5	6	7	1	2	3	4	5	6	7	1	2	3	4	5	6	7	1	2	3	4	5	6	7
rI_1	7	5	4	3	2	1	28	2	7	5	1	16	13	4	1	2	6	3	4	389	50	4	1	5	2	9	7	191	1	4	2	5	3	37	351
ud1	4	6	5	3	1	2	11	2	6	3	1	11	20	4	2	1	5	3	4	199	11	2	1	4	3	10	6	11	2	3	1	5	4	48	13
ud2	6	5	4	3	2	1	9	2	6	5	1	8	16	3	2	1	5	3	4	85	7	2	1	5	3	10	6	15	1	3	2	5	4	25	15
rI_2	5	14	9	1	2	3	4	3	9	8	1	4	7	2	1	2	3	4	14	748	7	1	2	3	5	22	10	73	1	2	3	7	6	49	8
ud1	6	11	5	1	3	2	4	2	6	3	1	7	8	4	1	2	3	4	11	269	5	2	1	4	3	14	6	74	2	3	1	5	4	18	8
ud2	6	7	5	3	2	1	5	2	6	4	1	7	11	3	1	2	5	3	4	190	6	2	1	4	3	10	6	18	2	3	1	5	4	31	10
rI_{2f}	7	5	4	3	1	2	24	1	7	5	2	14	12	3	1	2	6	3	4	42	163	1	2	5	3	9	7	44	1	4	2	5	3	20	199
ud1	5	6	4	2	3	1	11	2	6	3	1	11	55	5	2	1	5	3	4	113	11	1	2	4	3	10	6	11	1	3	2	5	4	32	15
ud2	6	5	4	3	2	1	7	2	6	5	1	8	16	4	1	2	5	3	4	148	8	2	1	5	3	10	6	15	1	3	2	5	4	23	15
rI_7	22	8	1	2	3	9	18	1	9	3	2	4	22	11	3	1	5	4	2	842	44	5	1	9	7	11	2	270	2	7	6	1	3	41	135
ud1	12	5	4	2	3	1	11	2	5	3	1	7	15	11	1	2	5	3	4	204	11	2	1	4	3	18	6	190	2	3	1	5	4	24	11
ud2	7	6	4	3	2	1	5	2	6	4	1	7	10	3	1	2	5	3	4	40	8	2	1	5	3	16	6	215	2	3	1	5	4	20	15
rRF	4	6	5	1	2	3	53	2	7	4	1	8	10	5	1	3	15	2	4	982	454	3	2	7	1	5	10	882	1	5	4	3	2	63	1000
ud1	2	1	4	3	8	6	12	2	6	3	1	7	8	4	1	2	12	3	4	320	11	2	1	4	3	8	6	12	2	3	1	5	4	20	14
ud2	2	1	5	3	4	6	14	2	6	5	1	7	10	3	1	2	5	3	4	59	8	2	1	5	3	4	6	14	1	3	2	5	4	20	15

In Table 11 we list the rankings of the influential variables for each of the five methods, I_1 , I_2 , I_{2f} , I_7 and RF, and for each of the five data sets. These rankings are followed by ud1 and ud2 which give the rankings after each of the two resuscitation steps. In these examples, the resuscitation methods, almost always seems to improve the rankings of influential variables and often succeed in making prominent those influential variables that were poorly regarded in the first approach. Note that it is virtually impossible for variables with rank greater than 10 to achieve a rank less than 11 in the first resuscitation. Also, because we did not discard poor performers in that first step, it is unlikely that a variable with rank greater than 15 will achieve a rank less than 16 in the second step. Nevertheless, these resuscitations provide an opportunity to reorder the candidates.

An alternative approach to discarding impostors and resuscitating poorly ranked influential variables is to take a relatively large number of prospects and submit those to the I_{2f} approach. This alternative approach can not resuscitate a variable which fails to appear in the list of prospects, and so it would pay to use a relatively large number of such prospects for this reduction stage, which can then be repeated with a smaller list. This method, when applied to the modest list of 30 prospects, provided considerable improvement on the relative rankings of the influential variables in the list, but failed to resuscitate variables not on the list of 30 for the data sets 3, 4 and 5. The results appear in Table 12.

TABLE 12

Ranks of Influential Variables for Five Methods Applied to Five Data Sets. Initial Ranks rI_1 , rI_2 , rI_{2f} , rI_7 , rRF before and after Resuscitation with I_{2f} Based on the Top 30 Ranked Variables.

	data set 1	data set 2	data set 3	data set 4	data set 5
rI_1	7 5 4 3 2 1 28	2 7 5 1 16 13 4	1 2 6 3 4 389 50	4 1 5 2 9 7 191	1 4 2 5 3 37 351
ud	7 5 1 2 3 4 12	1 8 2 3 12 10 4	1 2 6 3 4 - -	1 2 7 3 10 5 -	1 2 3 4 5 - -
rI_2	5 14 9 1 2 3 4	3 9 8 1 4 7 2	1 2 3 4 14 748 7	1 2 3 5 22 10 73	1 2 3 7 6 49 8
ud	9 6 4 1 2 3 10	1 8 5 2 9 10 3	1 2 6 3 4 - 17	1 2 5 3 9 7 -	1 2 3 5 4 - 21
rI_{2f}	7 5 4 3 1 2 24	1 7 5 2 14 12 3	1 2 6 3 4 42 163	1 2 5 3 9 7 44	1 4 2 5 3 20 199
ud	7 5 1 2 3 4 12	1 7 5 2 12 8 3	1 2 6 3 4 - -	1 2 6 3 10 7 -	1 2 3 4 5 17 -
rI_7	22 8 1 2 3 9 18	1 9 3 2 4 22 11	3 1 5 4 2 842 44	5 1 9 7 11 2 270	2 7 6 1 3 41 135
ud	7 5 1 2 3 4 12	1 8 2 3 14 9 4	1 2 5 3 4 - -	1 2 7 3 10 5 -	1 2 3 4 5 - -
rRF	4 6 5 1 2 3 53	2 7 4 1 8 10 5	1 3 15 2 4 982 454	3 2 7 1 5 10 882	1 5 4 3 2 63 1000
ud	7 5 1 2 3 4 -	1 7 5 2 12 11 3	1 2 7 3 4 - -	1 2 6 3 10 7 -	1 2 3 5 4 - -

“-” is used to represent ranks not observed.

5.1. *Applications to Rheumatoid Arthritis.* This section describes an application of the methods of this paper to a real data set on Rheumatoid Arthritis in two *examples*. The first is a brief summary and expansion of work in which some of us participated [10], and applies I_2 to thin the large set of available SNPs, I_8 on the reduced set, and random permutations of the dependent variable to estimate false discovery rates. In the second we apply the ideas of resuscitation to obtain some additional results.

EXAMPLE 6. Rheumatoid arthritis (RA, MIM 180300) is known as a common disorder with complex genetic etiology. In [10] the Illumina genome scan on RA, originally studied by Amos et al. [1], was analyzed as part of the Genetic Analysis Workshop 15 [6]. The Illumina genome scan consists of 5,407 Single Nucleotide Polymorphism (SNPs) genotyped from 642 Caucasian families. For the analysis, 349 unaffected individuals were selected as “controls” and 474 RA patients as “cases”. The analysis was carried out in two stages. It should be noted that in dealing with SNPs, we have explanatory variables which can assume three possible values. Also, as pointed out in Appendix S1, the BGTA method used in [10] is equivalent to that of the partition retention method using I .

Because there was a large number of three valued explanatory variables, and I_1 seemed to be nonproductive, the first stage consisted of using I_2 to select the 707 SNPs which appeared in the 1,000 top ranking pairs. Then the partition retention scheme I_8 was applied to these 707 SNPs using 70,000 randomly selected subsets of 8 SNPs. Each subgroup retained was assigned the value of I at the stopping time. This process yields a sample of 70,000 values of I . We plan to select the elements of those retained subgroups for which the value of I is above a certain threshold. To determine that threshold, we applied 50 permutations to the case-control labels, repeating the process described above for the original data each time. This yields 3,500,000 values of I . For each value of I , there are a number of selected subsets from the original data that have a larger stopping value I , say, m_1 . At the same value of I , we also calculated, for each permutation b , the proportion of the 70,000 permuted I values that are greater than the given value of I , say, $p_0^{(b)}$. The false discovery rate (FDR) at this given value I is then estimated as

$$\text{fdr}(I) = \frac{\text{median} \left(p_0^{(b)} \right)}{m_1/70,000}$$

[4, 27]. We used the value of I when the FDR estimate reaches 30% as the selection threshold. Elements of subsets with stopping I values which exceed

this threshold were selected. These consisted of 50 SNP's which are located within 39 distinct genes. We shall call these SNP's *qualified* since one can not claim, that they are truly related to RA and not impostors, without additional evidence from biological or other studies.

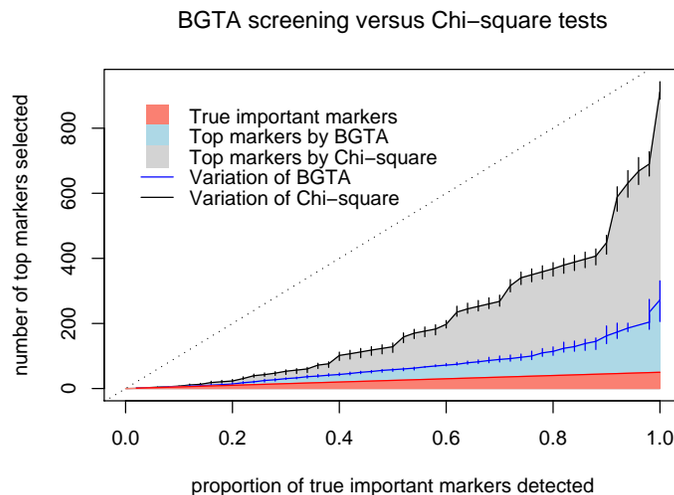


FIG 1. Screening performance of BGTA and χ^2 tests. The black and blue curves are average (out of five simulated sets) number of top markers required to be selected in order to attain a specific proportion of important markers. Vertical bars on these curves indicate the maximum and minimum number of markers required for the five simulations, which reflect the variability of the retention method.

In this paper, an additional procedure was carried out to determine how well these qualified SNPs are ranked by I_7 in the presence of noise, and how well this ranking compares with that of using the marginal χ^2 -test. As illustrated in Appendix S1, the I_7 approach is equivalent to the BGTA method studied in [28]. For this procedure the 50 qualified SNPs are augmented by 950 additional SNPs selected at random from the remaining 5,357. For these additional SNPs the case-control designation was permuted, while it was not for the 50. For example if the permutation moves the case row 16 to control row 35, our new row 35 will have the designation of case and the 50 SNPs will correspond to those of the original row 16, while the remaining 950 will correspond to those of row 35. In this way the structure of the values of the unqualified SNPs is maintained while their relation with the dependent variable is destroyed. This procedure was repeated 5 times with the SNPs ranked by retention frequency using I_7 or BGTA (500,000 screenings for each data set) and by the χ^2 -test. In Figure 1 we plot the average proportion of SNP's ranked above a given value against the average number of qualified

SNPs ranked above that value, for both methods. For example, by the time we retained 40 or 80% of the qualified SNPs, we will have retained 100 by I_7 (BGTA) and 368 by the χ^2 -test. Each of these methods does substantially better than pure chance in recognizing qualified SNPs. If the qualified SNPs represented true effects, this figure would provide an indication of an optimal cut off ranking, given the relative costs of false discovery and of missing true relations using these methods.

EXAMPLE 7. In this example we apply resuscitation analysis on the real data on Rheumatoid Arthritis (RA). We apply I_{2f} with $n_r = 50,000$ to the almost 15 million of the pairs of the 5,407 SNPs. In other words we evaluated I for every pair, and ranked the individual SNPs according to how often they showed up in the top 50,000 pairs. The 25 SNPs with the top ranks were selected as the first reduced list.

To resuscitate influential SNPs not in this short list, the partition retention method I_7 was applied to 2 million subsets with 3 members from the 25 and 4 from the remaining SNPs. The top ranking 50 SNP's from this stage, was then used for a second stage of I_7 with 2 million subsets with 3 from the 50 and 4 from the remaining 5,357. Table 13 displays the top 75 SNP's in this final resuscitated list, 38 of which are within 10 Mb of previously identified RA susceptibility loci.

Table 13: 75 SNPs selected after two rounds of resuscitation. Ranks from I_{2f} and the two resuscitations ud1 and ud2.

SNP	locus	ud2	ud1	rI_{2f}	previously identified locus and reported significance (within 10 Mb)
rs7534363	1p36.3	19	31	213	[20]: p=0.003; [7]: p=0.0035; [25]: p=0.00585;
rs2817594	1p36.2	33	32.5	236	—
rs235256	1p36.2	28	32.5	151.5	—
rs569668	1q42	2	2	3	[13]:p=0.003; [20]:0.04;
rs1389622	1q44	54	104	29	—
rs300739	2p25	51	52	32	[25]: note 1
rs2685263	2p25	20	16	12	—
rs6547142*	2p12	60	57	246	[7]: p=0.041;
rs1473357*	2p12	58	61	151.5	—
rs921423	2q11	16	11	10	
rs7561232	2q21	10	27	205	
rs1402810	2q21-22	9	13	8	
rs970595	2q33	1	1	1	[20]: p=0.03; [7]: p=0.024;
rs1921789	2q33	8	6	15	—
rs3821280*	2q37	55	58.5	100	[7]: p=0.0043;

Continued on Next Page...

Table 13 – Continued

SNP	locus	ud2	ud1	rI_{2f}	previously identified locus and reported significance (within 10 Mb)
rs164466	3p26	11	9	9	
rs1385654	3p12	40	23	25	
rs4572747	3q27	63	55	273.5	[7]: p=0.046;
rs2067078	3q27	45	39	258	—
rs881641	4p16	62	86	71	[20]: p=0.01;
rs1424903	5q11.2	46	19	17	
rs1004531	5q22-23	3	5	4	
rs1560657*	5q32-33	52	162	27	[7]: p=0.033
rs190129	6p25	25	45	125	
rs910516	6p21	31	14	13	[20]: p=6e-5; [25]: p=0.00127; [13]: p=e-12; [14]: p=4e-5;
rs2277123	6p21	13	41	97	—
rs508557	6q13	72	67	252	[13]: p=0.0028;
rs6915493	6q13-14	21	29	213	[13]: p=0.0028; [14]: p=0.006;
rs2296412	6q14-15	49	34	291	
rs6934871	6q15	73	80	265	
rs1873219	6q15-16	44	49	100	[13]: p = 0.01;
rs4302647	6q16	14	28	241	—
rs3827786	6q16	36	40	230.5	—
rs2151913	6q24-25	12	8	11	[7]: p=0.036;
rs1852210	7p13	50	20	24	
rs691183	7q32	7	10	7	
rs1531381	7q36	67	70	79	
rs2442567	8p23	26	35	176.5	[7]: p=0.040
rs766811	8q24.2	41	22	19	
rs751279	9q22	15	30	165	
rs715846	9q22	59	64	187	
rs2298033	10p13	56	51	176.5	
rs224136	10q21	71	68	95.5	[13]: p=0.0002;
rs1649183	10q26	64	54	213	
rs4077638	11q13	42	38	181.5	
rs2276189	11q24	74	74	241	
rs6590098	11q24	53	56	192.5	
rs1558507	12p13	35	25	20	[7]: p=0.0077;
rs1517815	12q21	37	21	23	[7]: p=0.0067;
rs2070628	12q24.3	70	62	252	
rs866781	12q24.3	38	46	213	
rs4758930	12q24.3	39	36	291	[14]: p=0.05;
rs1318725	13q22	17	17	22	[7]: p=0.039, [14]: p=0.03;
rs3811310	14q11.2	29	43	187	
rs1570342	14q11.2	4	3	5	
rs1889387	14q12-13	18	12	14	
rs7149108	14q12-13	27	42	213	
rs4904723	14q31-32	48	47	236	
rs8005578	14q32	34	48	213	
rs7159412	14q32	75	79	213	

Continued on Next Page...

Table 13 – Continued

SNP	locus	ud2	ud1	rI_{2f}	previously identified locus and reported significance (within 10 Mb)
rs1365591	15q12	61	60	265	
rs1565863*	15q14-15	32	15	18	[25]: p=0.01634;
rs3093291	16p12-11.2	5	4	2	[7]: p=0.0080;
rs7190151	16q23	69	58.5	143.5	[7]: p=0.038;
rs723919	16q23	22	18	16	—
rs116719	17q25	57	66	221.5	
rs4479277	17q25	66	84	187	
rs6416862	17q25	24	44	173	
rs879588	18p11.2	68	97	165	[2]: note 2; [20]: p=0.05;
rs1661965	19q13.3-.4	23	26	192.5	[16]: p=0.019; note 3.
rs241605	20p13	43	24	21	[20]: p=1e-4; [7]: p=0.030;
rs761319	20p12	6	7	6	—
rs1389157	21q21	47	50	265	
rs6517799	21q21	30	37	199.5	
rs5994180	22q11.1-11.2	65	53	273.5	[2]: note 2; [20]: p=4e-2; [7]: p=0.019; [21]: note 4.

note 1: Juvenile RA, LOD 6.0, stratified based on HLA-DRB1 presence.
note 2: The mapping is based on chromosomal rearrangements in Danish population on Juvenile RA.
note 3: Association to RA observed only for those that do not carry HLA-DRB1.
note 4: A southern blot experiment revealed a gene IGLV8 being absent in RA patients.
*: identified SNP is < 20 Mb from previously reported locus.

In Table 13 the SNPs are arranged according to the position of the locus in the genome. Thirty eight of these 75 SNPs appear in 22 regions previously referenced in a publication. These regions are grouped by horizontal lines in the table. The remaining 37 SNPs have no reference in the literature. Eleven of them end up among the top 25, and 13 among the second 25. We find that the resuscitation has sent 9 from I_{2f} rankings ranging from 97 to 241 to ud2 rankings in the top 25.

To spare the reader from extensive tables, we have often presented ranks and neglected frequencies and values of I . A careful reading of such extended tables would have made some things now obscure more obvious. In particular, there is almost no chance that any of the first 25 SNPs would have a higher rank than 25 after the first resuscitation, even if one shows up very poorly compared to the others in the first 25. The second resuscitation gives a chance for those in the second 25 to push out some in the first 25. This happens for nine SNPs. There is a possibility of using the results from nearby SNPs to give support to a given locus, but we have not done so here.

In this paragraph we relate some of our results to the biological literature. We use the term locus to represent a region of the genome that has

been identified as relevant to RA in the literature and may contain several genes. Table 13 has 26 such loci, about 12 of which were highly ranked by I_{2f} . These include 1q44, 2p25, 2q33, 6p21, 6q24-25, 12p13, 12q21, 13q22, 15q14-15, 16p12-11.2, 16q23 and 20p13. Some of these contain genes that are considered important in the biological literature. For example, 6p21 contains HLA-DRB considered the most important RA gene identified to date. Also 2q33 contains the important genes CTLA4, CD28 and STAT4, while 12pter-12p12 (centered at 12p13) contains CD4. The locus 1p36 harbors an important RA susceptibility gene PADI4. At this locus we identified 3 SNPs which required resuscitation to appear in the top 50 at ranks 19, 33, and 28. The gene LILR at 19q13.3-13.4 is known to be associated with RA susceptibility among patients who are not HLA-DRB carriers[16]. This may explain why this locus was discovered by resuscitation and not in the initial I_{2f} screening.

6. Summary. We address a problem, expected in medical cases of complex diseases, of a dependent variable influenced by one or a few small groups of explanatory variables, when data is available on many such variables. Our object is to detect these influential variables. Lo and Zheng pioneered a method, generalized here under the name Partition Retention. This method samples m of the S variables many times and uses a reduction process to retain a few of the m variables. Those variables that are retained most frequently are considered to be good candidates for being influential. The reduction process uses a statistic I which is considered to be a measure of information or influence for the set of m variables and n is the sample size. On the null assumption that the subset has no influential variables, the distribution of I is approximately that of a weighted sum of independent chi-squares with one degree of freedom.

When S , the number of variables, is large, the method is unlikely, in its original form, to detect any variable that has a negligible marginal observable effect. The fact that an influential variable has no marginal causal effect does not prevent it from having a marginal observable effect. However, such effects can also be detected by other first order methods. For example a simple t test will detect such an effect, as will I_1 based on $m = 1$. On the other hand, if S is not too huge, it is possible to consider second order interactions by considering I_2 based on all pairs of variables. An alternative to I_2 would be to evaluate the multiple correlations of the dependent variable Y on the pair of variables. There is some evidence, that J or the multiple correlation are as effective as the use of I to rank influence when m is small. But when m is large, we are likely to have a large number of partition elements, many of

which are empty or have few members, and in that case the use of I is more sensitive to detect marginal observable effects .

The rankings of pairs can be used to rank variables in several ways. One is to see how early influential variables are recognized when I_2 is used to rank all pairs. An alternative which we prefer is to rank all variables on how often they appear in the n_r most highly ranked pairs, where n_r is a substantial fraction of approximately $S^2/2$ pairs evaluated.

Assuming that S is too large to consider all possible third order interactions, we now have 4 methods. The methods we label I_1 , I_2 , I_{2f} , and I_m , based on one variable, two based on two variables, and one based on m variables, plus a few others similar to these but using correlations. However, if S is large, each of these methods may pick out impostors among the plausible candidates for influential variables. Part of our task is to discriminate against as many of the impostors as possible. One approach may be to see how these various techniques agree, The assumption is that these methods provide tests to determine influential variables, and insofar as the methods are different, they will, in combination, provide a more difficult test for an impostor than for a truly influential variable.

Another approach is that of using higher order methods on the relatively few plausible candidates. We have used I_{2f} , with the S variables replaced by 30 candidates from each of the four procedures on 5 distinct data sets with $S = 1,000$. We have also used a variation of I_7 where 3 variables are selected from the top 10 candidates and 4 from the remaining $S - 10$ variables. The former method does not make it possible to resuscitate influential variables not among the 30 selected. The latter method does make it possible.

For these second stage methods it would be feasible and sensible to take a longer string of candidates to increase the probability of not omitting influential variables. In fact we used much longer strings of plausible candidates in Example 7 on RA. It would also be feasible to apply I_3 to a list of candidates, relatively small compared with the original S . We have not done so here, nor have we carried out another stage of reductions.

Our application of the partition retention system has been relatively crude. The desire to keep the presentation simple, without making clever use of our knowledge of truth, led us to select numbers like 10 and 15 in our resuscitation scheme and other numbers almost arbitrarily, without an attempt to show off the methods to advantage. In the RA problem with real data, some necessary reasonable flexibility in the choices was applied. It is worth while investigating various strategies based on the use of I . It may save computing time if subsets with initially small values of I are ignored and not subjected to the retention scheme. One alternative is to retain all

those variables which show a large positive value of D_I on the first step, and not bother with the rest of the reductions. Another is to stop eliminating only if all D_I values exceed a number depending on the number of reductions that have taken place. In fact one of the weaknesses of the current method is that only one variable is retained when I increases with each stage of the reduction. This sometimes permits a variable with a very strong signal to overwhelm other influential variables that happen to be there.

Another valuable strategy adopted in our recent work [17] which led to the discovery of interactions between various breast cancer genes, was the use of the ratio of I_2 for a pair of genes to the maximum of the values of I_1 for each of those genes compared to a function of the maximum derived from the data.

The Partition Retention (PR) method has some similarities with Multifactor Dimensionality Reduction (MDR) [23] and Random Forests (RF) [5]. MDR uses what we called partition elements, but requires the dependent variable to be two valued. It does such an intensive multifactor analysis on all possible partitions that it is limited to problems with few explanatory variables, about 20. It uses an error rate criterion, which we conjecture might be improved by using the weighting implicit in I .

RF uses random subsets and is not limited to discrete explanatory variables. Where PR is a backward recursion method which gets rid of the worst candidates first, RF generates trees in a forward system, that is vulnerable to confusion if the first choice is a poor one. In other words, if the decision on the best first choice is not very good, it is likely that future splits will not be useful. In PR, if the first choice for deletion is not the least informative, the process is not likely to be ruined.

In our Example 5, we compared RF results with those of the other techniques for the five special data sets. In those data sets which exhibited strong first order observational effects, RF was comparable to the other methods. Where the first order effects were not too strong, RF seemed a little weaker. Resuscitation by I_7 and I_{2f} worked well on RF, but not quite as well as for the other methods. As far as we know, RF does not exploit the concept of resuscitating variables that previously looked poor, but interact strongly with some of those that looked good.

One of the referees brought the paper by Koller and Sahami [15] to our attention. It has some interesting parallels to this manuscript. It uses Kullback-Leibler information, for which J is a first order approximation, as a measure of influence, which we consider sensible, but has two shortcomings. It requires that Y be discrete, and it lacks some of the advantages of I over J . The application of this information is designed to attack the problem of

causal variables without marginal effect, by considering the effect of pairs. These pairs are employed in an interesting way using so called “Markov blankets.” However, insofar as that method depends on those pairs, it is like I_2 and I_{2f} , deterministic, and does not allow for the resuscitation of influential variables which require higher order interactions to be observed.

Acknowledgement. The research is partially supported by NSF grant DMS 0714669 and NIH Grant R01 GM070789. We would like to dedicate this to T.W. Anderson, a pioneer in Multivariate Analysis in honor of his 90th birthday. We wish to thank the editors and referees for useful comments and references.

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APPENDIX A1: $E(D_I)$

We derive expressions for the conditional expectation of D_I given \mathbf{n} for the random Y model and the expectation of D_I for the specified Y model.

Random- Y model: The partition element A_{ij} yields n_{ij} independent observations on Y , with mean μ_{ij} and variance σ_{ij}^2 , summing to W_{ij} . Let $\tilde{\mu} = n^{-1} \sum n_{ij} \mu_{ij}$ and $\tilde{\sigma}^2 = n^{-1} \sum n_{ij} \sigma_{ij}^2$. We use the tildes over the Greek letters to remind ourselves that these depend on \mathbf{n} and are not true parameters.

We calculate $E(W_{ij}W_{ik}|\mathbf{n}) = n_{ij}n_{ik}\mu_{ij}\mu_{ik}$ for $j \neq k$, $E(W_{ij}W|\mathbf{n}) = n_{ij}\sigma_{ij}^2 + n_{ij}\mu_{ij}n\tilde{\mu}$, and $E(W^2|\mathbf{n}) = n\tilde{\sigma}^2 + n^2\tilde{\mu}^2$. Combining these expectations, we have

$$E(D_I|\mathbf{n}) = -n^{-1} \sum_i \sum_{j < k} n_{ij}n_{ik}[(\mu_{ij} - \tilde{\mu})(\mu_{ik} - \tilde{\mu}) + n^{-1}(\tilde{\sigma}^2 - \sigma_{ij}^2 - \sigma_{ik}^2)].$$

The term involving the variances, is relatively small and can be estimated. We will neglect it in this discussion. The main term can be rewritten. We replace the sum for $j < k$ by the sum for $j \neq k$ and introduce $\mu_{ij} = \tilde{\mu}_i + \varepsilon_{ij}$ where $\tilde{\mu}_i = n_i^{-1} \sum n_{ij} \mu_{ij}$. Then

$$\begin{aligned} -2E(D_I|\mathbf{n}) &= n^{-1} \sum_i \sum_{j \neq k} [n_{ij}(\tilde{\mu}_i - \tilde{\mu} + \varepsilon_{ij})n_{ik}(\tilde{\mu}_i - \tilde{\mu} + \varepsilon_{ik})] \\ &= H_1 - H_2 \end{aligned}$$

where $H_1 = n^{-1} \sum_i [n_i(\tilde{\mu}_i - \tilde{\mu})]^2$ and $H_2 = n^{-1} \sum_i \sum_j [n_{ij}(\tilde{\mu}_i - \tilde{\mu} + \varepsilon_{ij})]^2$.

We may regard $\tilde{\mu}_i - \tilde{\mu}$ as representing the effect of (X_1, X_2, \dots, X_m) and ε_{ij} as representing the effect of X_0 in the presence of (X_1, X_2, \dots, X_m) . Thus if the m variables have no effect H_1 will be zero, and $E(D_I|\mathbf{n})$ will be positive. If X_0 has no effect in the presence of the other variables, ε_{ij} will be zero and $E(D_I|\mathbf{n})$ reaches a minimal value which is non-positive. The greater the effect of X_0 in the presence of (X_1, X_2, \dots, X_m) , the more positive D_I tends to be. The presence of influence in the m variables tends to diminish the effect of influence, if any, of X_0 .

Specified-Y model. Given that Y assumes the values y_1, y_2, \dots, y_R with frequencies given by $n^{(1)}, n^{(2)}, \dots, n^{(R)}$, the partition elements A_{ij} have $n_{ij}^{(r)}$ members equal to y_r , where the $\mathbf{n}^{(r)} = \{n_{ij}^{(r)}\}$ are independent for $1 \leq r \leq R$ with multinomial distributions $Mn(\mathbf{n}^{(r)}, \mathbf{p}^{(r)})$, and $\mathbf{p}^{(r)} = \{p_{ij}^{(r)}\}$. The number and probability for the partition elements A_i are similarly labeled $n_i^{(r)}$ and $p_i^{(r)}$. After normalization $W = \sum n^{(r)} y_r = 0$ and $I_{\Pi} = n^{-1} \sum W_{ij}^2$ and $I_{\Pi^*} = n^{-1} \sum W_i^2$.

We may write $W_i = \sum n_i^{(r)} y_r$ and $W_{ij} = \sum n_{ij}^{(r)} y_r$ and these have expectations $E(W_i) = \sum n^{(r)} p_i^{(r)} y_r$ and $E(W_{ij}) = \sum n^{(r)} p_{ij}^{(r)} y_r$. To calculate $E(D_I)$, we need $E(W_i^2)$ and $E(W_{ij}^2)$ which involve the variances. We have

$$E(W_i^2) = (EW_i)^2 + \sum n_i^{(r)} p_i^{(r)} (1 - p_i^{(r)}) y_r^2$$

and

$$E(W_{ij}^2) = (EW_{ij})^2 + \sum n_{ij}^{(r)} p_{ij}^{(r)} (1 - p_{ij}^{(r)}) y_r^2.$$

Since the sums in the two expressions above are of order n and those of the squared expectations are of order n^2 , we may approximate $E(D_I) = E(I_{\Pi}) - E(I_{\Pi^*})$ by

$$E(D_I) \approx n^{-1} \sum_i \left(\sum_j (EW_{ij})^2 - (EW_i)^2 \right).$$

Let $p_{ij} = \sum n^{(r)} p_{ij}^{(r)} / \sum n^{(r)}$ and $p_i = \sum n^{(r)} p_i^{(r)} / \sum n^{(r)}$. let $e_{ij}^{(r)} = p_{ij}^{(r)} - p_{ij}$ and $e_i^{(r)} = p_i^{(r)} - p_i$. Then

$$\begin{aligned} \sum_i (\text{EW}_i^2) &= \sum_i \sum_{r,s} n^{(r)} y_r n^{(s)} y_s \left(p_i^2 + p_i e_i^{(s)} + p_i e_i^{(r)} + e_i^{(r)} e_i^{(s)} \right) \\ &= \sum_i \sum_{r,s} n^{(r)} y_r n^{(s)} y_s e_i^{(r)} e_i^{(s)} \\ &= \sum_i \left(\sum_r n^{(r)} y_r e_i^{(r)} \right)^2. \end{aligned}$$

Also

$$\begin{aligned} \sum_{ij} (\text{EW}_{ij})^2 &= \sum_{i,j} \sum_{r,s} n^{(r)} y_r n^{(s)} y_s \left(p_{ij}^2 + p_{ij} e_{ij}^{(s)} + p_{ij} e_{ij}^{(r)} + e_{ij}^{(r)} e_{ij}^{(s)} \right) \\ &= \sum_{i,j} \sum_{r,s} n^{(r)} y_r n^{(s)} y_s e_{ij}^{(r)} e_{ij}^{(s)} \\ &= \sum_{i,j} \left(\sum_r n^{(r)} y_r e_{ij}^{(r)} \right)^2. \end{aligned}$$

We have expressed $E(D)$ as approximately the difference of two positive expressions, one of which involves $e_i^{(r)}$ which relates to the influence of X_1, X_2, \dots, X_m on Y , and the other which involves $e_{ij}^{(r)}$ which relates to the combined influence of X_0, X_1, \dots, X_m .

APPENDIX A2: NULL DISTRIBUTION OF I AND J .

Consider the null distribution of I for the random Y model. If the explanatory variables have no influence on \mathbf{Y} , we observe n independent identically distributed observations on Y with $n_i = np_i$ allocated to partition element A_i . Then $I = n^{-1} \sum [n_i(\bar{Y}_i - \bar{Y})]^2$ and $J = n^{-1} \sum n_i(\bar{Y}_i - \bar{Y})^2$. Suppose Y has mean μ and variance σ^2 . The mean has no effect on the distribution of I or J , and σ^2 has only a multiplicative effect. Thus there is no loss of generality in assuming that $\mu = 0$ and $\sigma = 1$.

Naively assuming that $\bar{Y} = 0$, we have the approximations that $I = \sum p_i(n_i \bar{Y}_i^2)$ is distributed like $\sum p_i V_i$ where the V_i are independent with approximately the chi-square distribution with 1 d.f. as $n \rightarrow \infty$. A similar argument would have the distribution of J approach that of chi-square with n' degrees of freedom where n' is the number of nonempty partition elements.

A more precise derivation takes \bar{Y} into account, but assumes that all the p_i are bounded away from 0 and 1 as n gets large. Let $Z_i = n^{-1/2} n_i(\bar{Y}_i - \bar{Y})$.

Conditioning on $\mathbf{n} = \{n_i\}$, the asymptotic distribution of \mathbf{Z} is $N(0, A)$ where $A = D(\mathbf{p}) - \mathbf{p}\mathbf{p}^T$ and $D(\mathbf{p})$ is the diagonal matrix with elements p_i .

Since $I = \mathbf{Z}^T \mathbf{Z}$, the limiting distribution of I is that of $\sum \lambda_i V_i$ where the λ_i are the eigenvalues of A . This singular matrix has one zero eigenvalue, but the others are nonnegative and add up to the trace of A which is $1 - \sum p_i^2$. In most of our ordinary applications $\sum p_i^2$ tends to be relatively small and the naive approximation is a good fit. The correction for J corresponds to the loss of one df.

In our applications we typically normalize Y so that it has sample mean 0 and $n^{-1} \sum (Y_i^2) = 1$. This normalization corresponds, asymptotically, to replacing σ by one. On the other hand our implicit assumption that all the n_i are large is really inappropriate for many of our applications where the partition elements have a good number which are empty or singletons. Nevertheless, it is easy to see that $E(I|\mathbf{n}) = 1 - \sum p_i^2$, and it seems clear that a more sophisticated theorem will apply for these applications.

If we deal with the null distribution for the specified Y model, then the values of Y in a given partition element with n_i entries corresponds to a sample without replacement of n_i observations from a finite population of n elements with sum 0 and sum of squares equal to n after normalization. But then the sum W_i of the Y values in partition element i has $E(W_i) = 0$ and $E(W_i^2) = n_i(1 - (n_i - 1)/(n - 1)) \approx np_i(1 - p_i)$. Moreover, for $i \neq j$, the covariance, $E(W_i W_j) = -n_i n_j / (n - 1)$. Since $I = \mathbf{W}^T \mathbf{W} / n$, the application of the Central Limit theorem for sampling from finite populations repeats the analysis for the random Y model. Once again, it is easy to see that $E(I|\mathbf{n}) = 1 - \sum p_i^2$. We have proved

THEOREM 1. *Conditioned on \mathbf{n} , the null distribution of I when Y is normalized, is asymptotically that of a weighted sum of independent chi-square variables, with nonnegative coefficients adding up to $1 - \sum p_i^2$.*

This applies to the null random Y and the null specified Y models, under the standard conditions for the applicability of the Central Limit Theorem and the assumption that the elements of \mathbf{p} are bounded away from 0.

Online Supplements

APPENDIX S1: EXAMPLES WHERE THE PARTITION RETENTION METHOD WAS USED.

Several versions of the Partition Retention approach have been used successfully in two past studies devoted to Inflammatory Bowel Disease (IBD) [19] and Rheumatoid Arthritis (RA) [10]. We shall describe briefly the method Backward Genotype Trait Association (BGTA) used in the latter, to show its relation to the recursive approach.

In the IBD study, data supplied by Rioux et al. [22] were subjected to Backward Haplotype Transmission Association (BHTA) analysis. The genetic data used 407 markers on 235 trios of parents and affected child. Forty-eight markers were identified which overlap two previously reported important loci. Four novel markers gave the strongest signal. A subsequent study by Barmada et al [3] support one of these four. Studies by Van Heel et al [26], Stoll et al [24], and Daly et al [8] support some of the others of the 48.

For the RA study, the data consisted of 5,407 SNP's from 642 families, using 349 unaffected individuals as "controls" and 474 patients as "cases". Here 39 loci were identified, 19 of which were previously reported in the literature. Based on the selected SNP's, an association network was constructed, showing several hubs of "interaction".

In the effort to identify genes responsible for complex human disorders, one method is to compare cases of the disease with suitable controls at a large number of available loci, where a given chromosome is known to take on one of two possible values. These are called SNP's and for a given subject with two chromosomes, the result will be labeled aa,ab,or bb, yielding 3 possible outcomes for each SNP. When a set of m SNP's are studied, there will be 3^m possible values for each subject, each representing a partition element of Π . In the RA study with cases and controls, the recursive method was applied to the Genotype Trait Distorion (GTD) statistic defined

$$(9) \quad GTD = \sum_{i=1}^{3^m} \left(\frac{n_{d,i}}{n_d} - \frac{n_{u,i}}{n_u} \right)^2,$$

where $n_{d,i}$ and $n_{u,i}$ are counts of cases and controls in genotype (element) i , and n_d and n_u are the total number of cases and controls under study. Therefore, the sample size $n = n_d + n_u$.

Let Y be the disease status (1 for cases and 0 for controls). Then, for a genotype partition Π , the score discussed in this paper can be naturally

defined as:

$$\begin{aligned}
I_{\Pi} &= n^{-1} \sum_{j \in \Pi} n_j^2 (\bar{Y}_j - \bar{Y})^2 \\
&= n^{-1} \sum_{i=1}^{3^m} (n_{d,i} + n_{u,i})^2 \left(\frac{n_{d,i}}{n_{d,i} + n_{u,i}} - \frac{n_d}{n_d + n_u} \right)^2 \\
(10) \quad &= \frac{n_d^2 n_u^2}{n_d + n_u} \sum_{i=1}^{3^m} \left(\frac{n_{d,i}}{n_d} - \frac{n_{u,i}}{n_u} \right)^2.
\end{aligned}$$

As shown in (10), GTD is indeed a special case of the influence measure I proposed in this paper.

APPENDIX S2: ASYMPTOTIC DISTRIBUTION OF

$$P(X_3 = 1 | X_1 X_2 = 1) - P(X_3 = 1).$$

With a relabeling of the variables, it suffices to establish the asymptotic distribution of $T = p(X_2 = 1 | X_1 = 1) - p(X_2 = 1)$. Then

$$T = \frac{n_{11}}{n_{11} + n_{10}} - \frac{n_{11} + n_{01}}{n}$$

and $\mathbf{n}^T = (n_{11}, n_{10}, n_{01}, n_{00})$ has the multinomial distribution $Mn(n, \mathbf{r})$ where $\mathbf{r}^T = (P_1 P_2, P_1 Q_2, Q_1 P_2, Q_1 Q_2)$, $P_1 = 1 - Q_1 = P(X_1 = 1)$ and $P_2 = 1 - Q_2 = P(X_2 = 1)$.

Let $\mathbf{Z} = n^{-1/2}(\mathbf{n} - n\mathbf{r})$ and thus $\mathbf{n} = n^{1/2}\mathbf{Z} + n\mathbf{r}$. By the Central Limit Theorem, the asymptotic distribution of \mathbf{Z} is $N(0, A)$ where the covariance matrix is given by $A = D(\mathbf{r}) - \mathbf{r}\mathbf{r}^T$ and $D(\mathbf{r})$ is the diagonal matrix formed by the elements of \mathbf{r} .

Writing T in terms of Z we have

$$\begin{aligned}
T &= \frac{r_1 + n^{-1/2}Z_1}{(r_1 + r_2) + n^{-1/2}(Z_1 + Z_2)} - [(r_1 + r_3) + n^{-1/2}(Z_1 + Z_3)] \\
&= \frac{r_1 + n^{-1/2}Z_1}{r_1 + r_2} \left[1 - \frac{n^{-1/2}(Z_1 + Z_2)}{r_1 + r_2} \right] - (r_1 + r_3) - n^{-1/2}(Z_1 + Z_3) + o_p(n^{-1/2}) \\
&= \frac{r_1}{r_1 + r_2} + n^{-1/2} \frac{Z_1}{r_1 + r_2} - n^{-1/2} \frac{r_1(Z_1 + Z_2)}{(r_1 + r_2)^2} - (r_1 + r_3) - n^{-1/2}(Z_1 + Z_3) + o_p(n^{-1/2}) \\
&= n^{-1/2} [Z_1(1/P_1 - P_2/P_1 - 1) - Z_2(P_2/P_1) - Z_3] + o_p(n^{-1/2}).
\end{aligned}$$

It follows that the limiting distribution of $n^{1/2}T$ is Normal with mean 0 and variance

$$\sigma^2 = \mathbf{b}^T \mathbf{A} \mathbf{b} = \sum b_i^2 r_i - \left(\sum b_i r_i \right)^2$$

where $\mathbf{b}^T = (1/P_1 - P_2/P_1 - 1, -P_2/P_1, -1, 0)$,

$$\sum b_i^2 r_i = (1 - P_2 - P_1)^2 P_1^{-1} P_2 + P_1^{-1} P_2^2 Q_2 + Q_1 P_2$$

and

$$\sum b_i r_i = (1 - P_2 - P_1) P_2 - P_2 Q_2 - Q_1 P_2 = -P_2.$$

Thus

$$\begin{aligned} \sigma^2 &= P_2 P_1^{-1} [(1 - P_1 - P_2)^2 + P_2 Q_2 + P_1 Q_1 - P_1 P_2] \\ &= P_2 Q_2 Q_1 / P_1. \end{aligned}$$

APPENDIX S3: RELATIVE EFFICIENCY OF I VERSUS J

We compare the expectations of I and J for the following special model of influence. Suppose that the i -th partition element has mean $\bar{Y}_i = W_i + \epsilon_i$ where W_i has mean 0 and variance $1/n_i$ and is independent of ϵ_i which assumes the values $\delta, -\delta, 0$ with probabilities $\lambda, \lambda, 1 - 2\lambda$. Then

$$E(I|\mathbf{n}) = 1 + 2n\lambda\delta^2 \sum (n_i/n)^2$$

$$E(J|\mathbf{n}) = m' + 2n\lambda\delta^2$$

The ratio of the relative increases due to $\lambda\delta > 0$ is approximately $m' \sum (n_i/n)^2$ which depends mainly on the variability of the sizes of the partition elements.

As an informal test of the theory that I is preferable to J for sensitivity to detect influence, we construct the following example.

EXAMPLE 8. A set of r probabilities p_i , summing to one, and proportional to $(1, 2, 4, \dots, 2^{r-1})$ are selected. We select n normal random variables with mean μ and standard deviation 1, and assign them to r partition elements with probabilities p_i . We select another n such variables with mean 0 and standard deviation 1, and assign them to another r partition elements with probabilities p_i . We construct $I_\mu = n^{-1} \sum n_i^2 (\bar{Y}_i - \bar{Y})^2$ and $J_\mu = n^{-1} \sum n_i (\bar{Y}_i - \bar{Y})^2$.

For several values of μ including 0, we construct $nit = 10,000$ samples of I and J , thereby estimating the distributions $F_I(x, \mu)$ and $F_J(x, \mu)$. To measure how sensitive I is for detecting a nonzero value of μ we see how many values of I_0 exceed a given quantile of I_μ . The greater the sensitivity the smaller that number would be. In effect, we are interested in $(1 - F_I(x_q, 0))$ where $F_I(x_q, \mu) = q$ or

$$G_I(q, \mu) = 1 - F_I[F_I^{-1}(q, \mu), 0].$$

If I is more sensitive than J we should expect to have

$$G_I(q, \mu) < G_J(q, \mu)$$

Our conjecture is that if r is substantial, then the sample sizes in the partition elements will vary considerably with many singletons and empty elements and I should do better than J . Figure 2 corresponds to several values of μ as q varies from 0.5 to 1.0. Notice that each of these curves goes from the origin to an end point corresponding to $q = 0.5$. As μ increases the curves get shorter, because there are very few values of I_0 and J_0 that exceed the large median values of I_μ and J_μ when μ is large. While Figure 2 corresponds to $r = 5$ representing 10 partition elements, Figure 3 corresponds to $r = 2$ where there are only 4 partition elements.

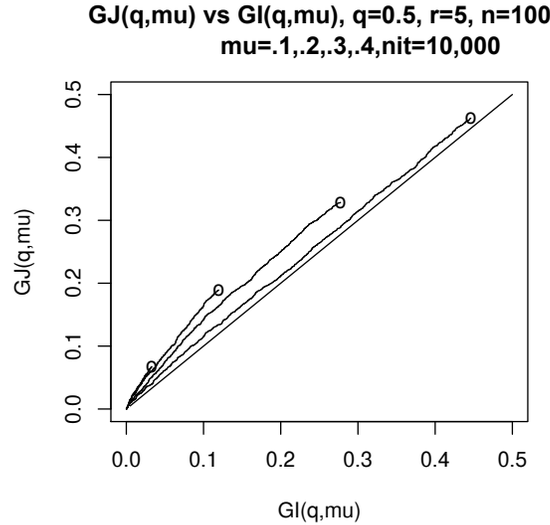


FIG 2. *Relative sensitivity of I and J for r = 5.*

The figures clearly show that I is more sensitive than J when $r = 5$ and that there is not much difference when $r = 2$.

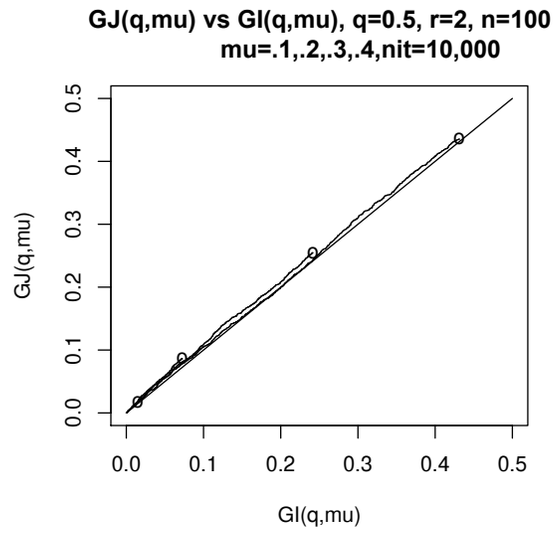


FIG 3. *Relative sensitivity of I and J for r = 2.*