

Computing the distribution of the maximum in balls-and-boxes problems with application to clusters of disease cases

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We present a rapid method for the exact calculation of the cumulative distribution function of the maximum of multinomially distributed random variables. The method runs in time $O(mn)$, where m is the desired maximum and n is the number of variables. We apply the method to the analysis of two situations in which an apparent clustering of cases of a disease in some locality has raised epidemiological concerns, and these concerns have been discussed in the recent literature. We conclude that one of these clusters may be explained on purely random grounds, namely the leukemia cluster in Niles, IL, in 1956–1960; whereas the other, a leukemia cluster in Fallon, NV, in 1999–2001, may not.

algorithm | multinomial

It happens from time to time that cases of a disease will cluster both geographically and in time in a manner that seems not to be random and that invites further epidemiological study.

Of course, mathematics alone cannot answer serious questions of public health, but it can provide guidelines about what sort of clustering should be regarded as unusual and what sort is to be expected. In particular, the calculation of a P value is required for an objective assessment of any observed event. In this paper we provide a rapid and exact P value calculation for the standard “balls-in-boxes” model appropriate to a disease-clustering situation.

The Model

Suppose that during a certain time period, a number r of cases of some disease arise randomly in some large population, such as that of the United States. Let N be the size of that population and N_0 be the population of the community in which the seemingly large number of cases has occurred.

We think of the entire country as consisting of $n = N/N_0$ identical communities, or cells, each containing N_0 people, and we ask

If r cases occur randomly in the populations of n communities of the same size, what is the probability that no community gets more than m cases of the disease?

The standard calculation required to answer this question involves the “balls-in-boxes” model, discussed below. If, for example, it turns out that it is extremely likely that *some* community of equivalent size to that in which the seemingly large number of cases occurred would have that many cases purely by chance, we could conclude that the observed cluster would not be a cause for further suspicion of communicability of the disease or the existence of environmental causative factors. Likewise, if it turns out that it is extremely unlikely that, by chance, *any* community of that size would have the observed number of cases of the disease, then support would be given to the possibility of a public health hazard.

The Mathematics

Mathematically speaking, we have r “balls” (the disease cases) being dropped randomly into n labeled “boxes” (the communities). The relevant calculation thus concerns the P value associated with the box (or boxes) having the largest number of balls in it. It is well known that the distribution function of the maximum of a number of random variables changes sharply near the mean of the maximum, so that an exact rather than an approximate calculation is needed to find this P value. We provide this exact calculation in this paper.

The P value associated with an observed value m of cases of the disease in the community of interest is the probability that the maximum number of balls in any box is m or more. We find this probability by first finding the probability that no box contains more than m balls. Denote this probability by $P(r, n, m)$.

Now, the probability that there are r_1 balls in box 1, and r_2 in box 2, and \dots , and r_n in box n , is given by the well known multinomial distribution,

$$\Pr(r_1, r_2, \dots, r_n) = \frac{1}{n^r} \frac{r!}{r_1! r_2! \dots r_n!} \cdot (r = r_1 + \dots + r_n) \quad [1]$$

The probability that no box contains more than m balls (i.e., the cumulative distribution function of the maximum of the r_i , evaluated at m) is

$$\begin{aligned} P(r, n, m) &\stackrel{\text{def}}{=} \Pr(\text{all } r_i \text{ are } \leq m) \\ &= \sum_{\substack{0 \leq r_1, r_2, \dots, r_n \leq m \\ r_1 + r_2 + \dots + r_n = r}} \frac{1}{n^r} \frac{r!}{r_1! r_2! \dots r_n!} \cdot \quad [2] \end{aligned}$$

The Computation

At first sight, the expression of Eq. 2 seems appallingly complicated for exact computation, if r and n are large. Various approximations, such as the Poisson approximation, have been used by researchers to avoid the apparently tedious computation in Eq. 2.

However the exact calculation can be completely tamed by two steps. First, we introduce the function

$$e_m(x) = 1 + x + \frac{x^2}{2!} + \frac{x^3}{3!} + \dots + \frac{x^m}{m!},$$

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Table 2. The Fallon, NV, computation

<i>m</i>	<i>P</i> (8000, 12000, <i>m</i>)	<i>P</i> value
4	0.000472	1.000000
5	0.436361	0.999528
6	0.925122	0.563639
7	0.993604	0.074878
8	0.999528	0.006396

Further Comments on the *P* Value

The *P* value corresponding to any value of *m* in the balls-in-boxes case can in principle be calculated exactly by using standard inclusion/exclusion formulae. In practice, this seems extremely difficult, because the alternating signs can cause catastrophic loss of significant digits. A Poisson approximation is also possible but may be inaccurate, particularly around the tails of the distribution. Our exact method, described in Eq. 4, is fast and does not suffer from any of those problems.

A further comment about *P* values is more wide-ranging. Many diseases might come to our attention because of an apparent clustering in some location in some time period. In addition, many different time periods might be potentially observed. An overall *P* value calculation taking these matters into consideration would be desirable but in practice would probably be impossibly difficult, since no precise value can be attached to “the number of diseases that might come to our attention” or, possibly, to the number of time periods that we might have considered.

A Disclaimer

Mathematics cannot prove or disprove the communicability or environmental origins of a disease process. It can only help to define the word “unusual.” The benchmark given above seems like an appropriate one to use when investigating an outbreak that is localized spatially, temporally, or both. By this benchmark, the clustering of leukemia cases in Niles, IL, between 1956 and 1960 was not unusual. In fact, some collection of that number of cases in some community the size of Niles in a 5-year period of keeping records was to be expected with high probability. On the other hand, the Churchill County data seem extremely significant.

Some Related Work

The problem of finding the distribution of the maximum occupancy in a balls-and-cells problem is very old. Already in the work of Barton and David (5) one finds the first of our two observations, namely that the desired probability is a certain coefficient in a power of a given power series. In ref. 6, this observation of Barton and David is cited and is said to be “not in a form convenient for computing,” which is true absent our second step (in Eq. 3) of vastly accelerating the computation of the high power of the given series.

Freeman’s algorithm in ref. 6 sought to economize the computation by grouping together vectors of occupancy numbers that, as unordered multisets, were the same. Hence, he listed partitions with given largest size part and counted the occupancies of that subset of all partitions. This approach requires considerably more labor than our method above.

Likewise the recurrence (Eq. 3) for computing powers of power series has a long history. Although we have followed ref. 1 in our presentation, the recurrence method was certainly not invented by the authors of ref. 1, because this method is described in several earlier works. Nonetheless, the concatenation of the two methods in connection with finding the distribution of the maximum cell occupancy seems to be new.

Finally, we mention some very recent work (7, 8) on a different problem but one that presents a similar challenge. This problem is the normalized maximum likelihood distribution, which arises in connection with finding the shortest possible encoding of a given data set. The problem concerns the rapid computation of

$$R(n, k) = \frac{1}{n^n} \sum_{r_1+r_2+\dots+r_k=n} \frac{n!}{r_1!r_2!\dots r_k!} r_1^{r_1}r_2^{r_2}\dots r_k^{r_k}. \quad [5]$$

This formula, aside from the multiplicative factor, is evidently the coefficient of x^n in $B(x)^k$, where $B(x) = \sum_n n^n x^n/n!$. Our algorithm (Eq. 3) clearly applies here. In ref. 8, the authors discovered that the elegant recurrence

$$R(n, k) = R(n, k - 1) + \frac{n}{k - 2} R(n, k - 2)$$

holds, owing to special properties of the function $B(x)$, and this yields an algorithm that runs in time $O(n + k)$, which is faster than our general algorithm (Eq. 3) when specialized to this case. However, if, for a fixed *k*, we want a table of $R(n, k)$ for all $n = 1, 2, \dots, N$, then our algorithm (Eq. 3) will compute all *N* of those numbers at an average cost of $O(N)$ computations per number computed, which is about the same as the method of ref. 8.

Appendix 1: Powers of a Power Series

Suppose we have a power series $f = \sum_r a_r x^r$. Then, when we raise the series *f* to the *n*th power, we obtain

$$\begin{aligned} f^n &= \left(\sum_r a_r x^r \right)^n \\ &= \left(\sum_{r_1} a_{r_1} x^{r_1} \right) \left(\sum_{r_2} a_{r_2} x^{r_2} \right) \dots \left(\sum_{r_n} a_{r_n} x^{r_n} \right) \\ &= \sum_{r_1, r_2, \dots, r_n} a_{r_1} a_{r_2} \dots a_{r_n} x^{r_1+r_2+\dots+r_n}. \end{aligned}$$

The coefficient of x^r in the above is evidently obtained by requiring that $r_1 + r_2 + \dots + r_n = r$, and therefore it is

$$\sum_{r_1+r_2+\dots+r_n=r} a_{r_1} a_{r_2} \dots a_{r_n}. \quad [6]$$

Next we specialize this expression to the case for which the *f* series is the *m*th section of the exponential series. This means that we are taking $a_j = 1/j!$, for $j \leq m$, and $a_j = 0$ otherwise. The general expression (Eq. 6) then becomes exactly the cumulative multinomial probability (Eq. 2), aside from the factor $r!/n^r$, as claimed. For more information about power series generating functions, see, for example, ref. 9.

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