#### MIXTURES OF DISTRIBUTIONS

by

RAY LEE MILLER, JR.

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Approved by:

Major Professor

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#### INTRODUCTION

Mixtures of distributions occur when there is an overlapping of two or more distributions so that it is difficult to separate them into their respective components. The main statistical problem is to estimate as accurately as possible the true proportions of overlapping contributed by each distribution and their respective parameters. Mixtures of distributions may also be considered as a form of contagion.

There are many examples of mixtures of distributions or contagious distributions in our environment. Feller (1943) has distinguished two types of these distributions: 1) true contagion and 2) apparent contagion. Student's typing errors, payroll check errors, or bank statement and ledger errors are some of the situations in which the occurrence of one "favorable" event might affect the probability of another event happening. This is true contagion. Heterogeneity of decaying radioactive material or atmospheric data are examples which involve apparent contagion.

The purpose of this paper is to describe mixtures of distributions; to develop their fundamental distributions and interrelationship; and give examples of estimating the parameters of mixtures of exponential distributions by the method of moments and the method of maximum likelihood.

#### HISTORICAL REVIEW

As early as 1894 Karl Pearson had attacked the complicated problem of mixed frequency distributions (Rider, 1961). In particular, he considered dissecting nonnormal populations into normal components. Pearson considered only two distributions in each of which a certain character is distributed normally. The statistical problem reduced to that of estimating the two mean values  $\mathcal{M}_1$ ,  $\mathcal{M}_2$  standard deviations  $\mathcal{T}_1$ ,  $\mathcal{T}_2$  and the proportions of mixture  $\ell$  and  $(1-\ell)$  from the observed frequency distribution. The five parameters were estimated by the method of moments and a solution to the estimates depended upon a suitably chosen root of a nonic (ninth degree equation) constructed from the first five moments of the observed frequency distribution.

In 1920 Greenwood and Yule developed a very general scheme for contagious events through their studies (diseases and accidents) concerned with the nature of frequency distributions representative of multiple happenings. Due to this generality their formulas became too complex for most practical applications (Feller, 1943).

About three years later Polya and Eggenberger were considering a similar problem which later led them to consider a special model of true contagion which is the simplest case of the general Greenwood-Yule scheme. Thus, Feller (1943) warns

l"Dissection" means point estimation of the parameters in a parametric mixture model.

that "in order to decide whether or not there is true contagion, it is not sufficient to consider the distribution of events, but a detailed study of the correlation between various time intervals is necessary."

It wasn't until 1939 when J. Neyman applied contagious distributions to entomology and bacteriology that there was a significant advancement in the study of contagious distributions. Neyman (1939) considered the distribution of larvae in a field which had been divided into plots of equal areas. In the experiments described by Neyman "the attempts to fit the Poisson Law... failed almost invariably with the characteristic feature that, as compared with the Poisson Law, there were too many empty plots and too few plots with only one larva." Thus, this is an excellent example of true contagion because the appearance of one larva in a plot seemed to increase the probability of finding at least one more larva in the same plot. From the related distributions that Neyman derived to fit the experimental data there evolved the generalized Neyman's distribution.

#### GENERAL PROBLEM OF MIXED DISTRIBUTIONS

The problem of mixtures of two distributions is to know exactly with what proportion each distribution contributes in the area where overlapping exists so that each distribution can be reconstructed in that area to learn what effect the second or first distribution had on the other. For example, it may sometimes happen that two normally distributed populations

are mixed with nearly the same mean but different standard deviations. Cassie (1954) illustrated this case by graphing the length of 63 fish meshed in the cod-end of a trawl versus their cumulative frequency and obtained a curve indicating positive kurtosis. It was desirable to know exactly the proportion each distribution contributed in the area where overlapping occurred. Thus, reconstruction in that area would show how each distribution affected the other.

#### CRITERIA OF MIXTURES OF DISTRIBUTIONS

Credit for much of the following is due to Feller (1943) and Conover (1962).

#### Theorem:

If F(x,a) is a distribution function depending on the parameter  $\underline{a}$ , and H(a) is a distribution function, then

$$G(x) = \int F(x,a) dH(a)$$

is a distribution function.

## Proof:

It is sufficient to show that  $G(\mathbf{x})$  satisfies the following three conditions:

- (i) G(x) is monotone non-decreasing, i.e.,  $G(x+b) \ge G(x)$  if b > 0.
- (ii) G(x) is right continuous, i.e.,  $G(x+0) = \lim_{b\to 0} G(x+b) = G(x)$ .

(iii) 
$$\lim_{x\to\infty} G(x) = 1 \qquad \lim_{x\to\infty} G(x) = 0$$

Because F(x,a) is a distribution function and therefore monotone non-decreasing, (i) is true. Thus,

$$G(x+b) = \int F(x+b,a) dH(a) \ge \int F(x,a) dH(a) = G(x)$$
;  $b > 0$ .

(ii) is right continuous since

$$G(x+0) = \int F(x+0,a) dH(a) = \int F(x,a) dH(a) = G(x)$$

where F(x,a) must have been right continuous. The last condition is true because

$$\lim_{x\to\infty} F(x,a) = 1 \qquad \text{and } \lim_{x\to-\infty} F(x,a) = 0;$$

thus.

$$\lim_{x\to\infty} G(x) = \lim_{x\to\infty} \int_{F(x,a)} dH(a) = \int_{x\to\infty} F(x,a) dH(a)$$

$$= 1.dH(a) = 1$$

because H(a) is also a distribution function. Also,

$$\lim_{x \to -\infty} G(x) = \lim_{x \to -\infty} \int_{\mathbb{F}(x,a)} dH(a) = \int_{\mathbb{F}(x,a)} dH(a)$$
$$= \int_{0.0} \int_{\mathbb{F}(x,a)} dH(a) = \int_{0.0} \int_{\mathbb{F}(x,a)} dH(a)$$

Therefore, both F(x,a) and H(a) determine that G(x) is a distribution also.

### Definition:

A distribution function  $\theta(x)$  is called a "mixture of distributions" if

(1) 
$$G(x) = \int F(x,a) dH(a)$$

where F(x,a) is an arbitrary non-degnerate cumulative distribution function  $^1$  (c.d.f.), depending on a parameter  $\underline{a}$ , and another c.d.f. H(a). The domain of variation of  $\underline{a}$  determines the range of integration. If H(a) is a step function, then we must define a non-degenerate cumulative distribution  $F(x,a_1)$  whose parameters are finite numbers  $a_1, a_2, \ldots, a_1$ ; let  $a_1 \in A_1, a_2 \in A_2, \ldots, a_1 \in A_1$  where  $A_1, A_2, \ldots, A_1$  are given sets of real numbers that make up the population. Let  $p_1$  be the weight attached to the population  $A_1$  which are mixed at random in proportions  $p_1$ :  $p_2$ : ...:  $p_1$  ( $p_1 \ge 0$  and  $\sum p_1 = 1$ ) where  $p_1$  are real constants. Thus, a family of distribution functions may be obtained by letting the parameters vary over  $A_1$  independently of each other. The function

(2) 
$$G(x) = \sum p_i P(x, a_i)$$

or more simply

(3) 
$$G(x) = \sum p_1 F_1(x)$$

is a cumulative non-degenerate distribution whose components are  $F_1$  (x) and the real values  $P_1$  are weights (Medgyessy, 1961, p. 1).

Teicher (1960, 1961, and 1963) and Robbins (1948 and 1949) give a more rigorous approach to mixtures of distributions.

 $<sup>^{1}\</sup>mathrm{B}(x)$  is said to be a degenerate distribution function if  $\mathrm{B}(x)\!=\!0$  when  $x\!\leq\!s$  (constant) and if  $\mathrm{B}(x)\!=\!1$  when x>s. Its graph consists of a single step of height one at x=s. A probability distribution is degenerate if its members are : one when x=s (integer) and 0 (otherwise); its graph consists of a single point of height one at x=s

#### DEVELOPMENT OF FUNDAMENTAL DISTRIBUTIONS OF MIXTURES

Feller (1943) gives the following definitions of contagion: Definition:

True contagion. Each "favorable" event has a direct effect of increasing or decreasing the probability of some future event happening.

#### Definition:

Apparent contagion. Inhomogeneity in populations where the events are independent of each other.

#### Definition:

A simple Poisson distribution function with parameter  $\underline{\mathbf{a}}$  must be of the form

(4) 
$$\pi(n;a) = e^{-a} \frac{n}{n!}, \quad a > 0, \quad n = 0,1,2, \dots,$$

where a gives the expected number of "events."

#### Definition:

A distribution function  $P(\mathbf{x})$  is a compound Poisson distribution if it has the form

(5) 
$$P(x) = \sum_{n=x} \int_{0}^{\infty} e^{-a} \frac{a^{n}}{n!} dH(a), \quad n = 0,1,2,...,$$

where H(a) is the distribution function of a non-negative random variable and  $\underline{a}$  is distributed according to the cumulative probability law H(a).

From (5) one can go to the probability function, first considered by Greenwood and Yule (1920),

(6) 
$$\pi_n = \int_0^\infty e^{-a} \frac{n}{n!} dH(a) .$$

Thus, (6) is called the compound Poisson distribution.

If H(a) is a step function, then the probability function of the compound Poisson distribution becomes

(7) 
$$\pi_{n} = \frac{1}{n!} \sum_{i=0}^{\infty} e^{-a_{i}} a_{i}^{n} p_{i}.$$

By defining Pearson's type III distribution, two special cases of the Polya-Eggenberger distribution can be developed.

# Definition:

The Pearson type III distribution is the distribution of the probability density function

(8) 
$$h(a) = \frac{1}{\int \frac{1}{2} \left(\frac{1}{d}\right)^{\frac{9}{d}} \left(\frac{9}{d} - 1\right)} e^{-\frac{1}{d}(a-c)}$$
;  $a > c$ ,  $\frac{9}{d} > 0$ ,  $\frac{1}{d} > 0$ 

where  $\underline{a}$ ,  $\frac{9}{d}$ , and  $\frac{1}{d}$  are constants.

To obtain a more condensed form of Pearson's type III distribution, let  $k=\frac{1}{d}$  and  $t=\frac{9}{d}$ , then (8) becomes

(9) 
$$h(a) = \frac{k^{t}}{\Gamma(t)} (\lambda - c)^{t-1} e^{-k(\lambda - c)}$$

If C=0, then h(a) in (9) has the origin of its distribution shifted by the amount (- C) and

(10) 
$$h(a) = H^{\dagger}(a) = \begin{cases} \frac{K^{t}}{\Gamma(t)} & e^{-Ka} \\ O & , a < 0 \end{cases}$$

where  $\underline{a}$  ranges over the desired values, k and t being constants determined by comparing the resulting compound Poisson distribution with the actual observed data.

The Polya-Eggenberger distribution may be obtained as a special case of (6). Substitute (10) into (6).

(11) 
$$T_n = \int_0^\infty e^{-a} \frac{n}{n!} \left[ \frac{t}{K} a e^{-tA} e^{-xA} \right] da$$

(12) 
$$\overline{\mathcal{H}}_{n} = \frac{1}{n!} \frac{K^{t}}{\Gamma(t)} \frac{1}{(1+K)(1+K)^{n+t-1}} \left[ \widehat{a}^{(1+K)} \right]^{n+t-1} e^{-a(1+K)}$$

$$(1+K) da$$

which is of the form

$$\Gamma(t) = \int_{0}^{\infty} x^{t-1} e^{-x} dx$$

Therefore, (12) becomes

(13) 
$$\Pi_{n} = \frac{K^{t}}{\Gamma(t)} \frac{1}{n!} \frac{\Gamma(n+t)}{(1+K)^{n+t}}, K>0, t>0, n \leq 0, 1, 2, \dots$$

which is the Polya-Eggenberger distribution.

Neyman's type A distribution can be obtained if  $\underline{a}$  takes on the values be only, where c>0 is a constant and b=0,1, ..., and if  $\underline{a}$  is distributed according to the Poisson law (Feller, 1943)

(14) 
$$p_i = \text{Prob}\left\{a = bo\right\} = e^{-\lambda} \frac{\lambda^b}{b!}$$
,  $\lambda > 0$ ,

then (7) becomes

$$\Pi_{n} = \sum_{b=0}^{\infty} \left[ e^{-bc} \frac{(bc)^{n}}{n!} \right] e^{-\lambda} \frac{\lambda^{b}}{b!}$$

or

(15) 
$$T_n = \frac{c^n}{n!} e^{-\lambda} \sum_{b=0}^{\infty} \frac{b^n}{b!} (e^{-c} \lambda)^b$$

which is Neyman's contagious distribution of type A.

RELATIONSHIPS BETWEEN DISTRIBUTIONS OF MIXTURES

Wilks (1961) stated that the gamma distribution is a Pearson type III distribution when the given distribution is a probability density function (p.d.f.) of the form

$$h(a) = \frac{a^{u-1}e^{-a}}{\Gamma(u)}$$

where M is the mean and corresponds to t in equation (10).

According to Feller (1957), p. 131, the Polya-Eggenberger distribution can be linked to the negative binomial distribution by going through numerous limiting processes on the Polya-Eggenberger distribution. Also, Conover (1962) showed that the

negative binomial distribution is a compound Poisson distribution. Gurland (1957) further related the negative binomial to other distributions not mentioned in this paper.

It was shown above that Neyman's contagious distribution of type A can be obtained by letting the parameter  $\underline{a}$  in the compound Poisson step function equal bc only where  $\underline{a}$  is distributed according to the Poisson law.

# APPLICATION TO EXPONENTIAL DISTRIBUTIONS AND ESTIMATION OF PARAMETERS BY THE METHOD OF MOMENTS

Rider (1961) has stated that life characteristics of certain types of electronic components (resistors, capacitors, vacuum tubes, etc.) and complex systems of highspeed digital computers are very well described by exponential distributions. In experiments in life testing a probability density function of the following form may be assumed:

(16) 
$$f(t) = \frac{1}{\theta} e^{-t/\theta} ; \theta > 0, 0 \le t < \infty$$

where  $\Theta$  is the mean lifetime between failures.

Suppose two populations of type (16) have been mixed in unknown proportions p and q =(1-p) with parameters  $\theta_1$  and  $\theta_2$  respectively. Then,

(17) 
$$f(t) = p \theta_1^{-1} e^{-t/\theta_1} + (1-p) \theta_2^{-1} e^{-t/\theta_2}$$

At this point, the method of moments can be applied to estimate the parameters  $\theta_1$ ,  $\theta_2$ , and p.

Let  $\mathbf{m}_{\mathbf{r}}^{i}$  be the  $\mathbf{r}\underline{\mathbf{th}}$  sample moment about zero. In particular,  $\mathbf{m}_{\mathbf{r}}^{i}$ ,  $\mathbf{m}_{\mathbf{r}}^{i}$ , and  $\mathbf{m}_{\mathbf{r}}^{i}$  are the moments of a random sample from (16). The estimators of p,  $\theta$ <sub>1</sub>, and  $\theta$ <sub>2</sub> will be obtained by method of moments and denoted p\*,  $\theta$ <sub>2</sub>, and  $\theta$ <sub>3</sub>. Thus,

(18) 
$$p^{\#} \theta_{1}^{\#} + (1-p^{\#}) \theta_{2}^{\#} = m_{1}^{\prime}$$
,

(19) 
$$p^* \theta_1^{*2} + (1-p^*) \theta_2^* = \frac{1}{2} \pi_2'$$

(20) 
$$p^* \quad \theta_2^{*3} + (1-p^*) \quad \theta_2^{*3} = \frac{1}{6} \, m_3'$$

From (18)

(21) 
$$p^* = (m_1' - \theta_2^*)/(\theta_1^* - \theta_2^*)$$

Upon substituting (21) into (19) and (20), the following equations result:

(22) 
$$(m_1' - \theta_2^*) (\theta_1^* + \theta_2^*) = \frac{1}{2}m_2' - \theta_2^{*2}$$

(23) 
$$(m_1' - \theta_2^*) (\theta_1^{*2} + \theta_1^* \theta_2^* + \theta_2^{*2}) = \frac{1}{6}m_3' - \theta_2^{*2}.$$

Equation (22) may be solved for  $\theta_1^{\text{S}}$  (1 = 1 or 2), the solution being

(24) 
$$\theta_1^{\#} = (\frac{1}{2} m_2^{\prime} - m_1^{\prime} \theta_1^{\#}) / (m_1^{\prime} - \theta_1^{\#})$$

where j=2 or 1 according as i=1 or 2. If equation (24) is substituted into (23), upon simplification we obtain  $\theta_j^s$ .

(25) 
$$6(2m_1^2 - m_2^2) \theta_1^{*2} + 2(m_3^2 - 3m_1^2 m_2^2) \theta_1^* + 3m_2^2 - 2m_1^2 m_3^2 = 0.$$

Solving this quadratic equation gives two roots  $\theta_1^*$  and  $\theta_2^*$ . Which root is designated  $\theta_1^*$  and which  $\theta_2^*$  is immaterial.  $p^*$  can be obtained by substituting  $\theta_1^*$  and  $\theta_2^*$  into (21) and will refer to the component  $\theta_1$ ; whereas,  $1 - p^*$  will refer to  $\theta_2$ .

The roots of (25) may not always be positive or even real. If every observation in a sample were equal to some constant k>0, then it follows that  $m_1^1=k$ ,  $m_2^1=k^2$  and  $m_3^1=k^3$  and (25) is reduced to

(26) 
$$k^2 \left[ 6(\theta - \frac{1}{3}k)^2 + \frac{1}{3}k^2 \right] = 0$$
,

whose roots are imaginary. This may occur provided we have positive probability as seen from continuaity considerations.

If  $\theta_1 \neq \theta_2$ , then our proposed estimators are consistent and the probability that  $\theta_1^* > 0$ ,  $\theta_2^* > 0$ ,  $0 \le p^* \le 1$  approaches 1 as n tends to infinity. This happens because the estimators which are thought of as functions of  $(m_1', m_2', m_3)$ , are continuous at the point  $(\mathcal{M}_1', \mathcal{M}_2', \mathcal{M}_3')$ , where  $\mathcal{M}_1$  are the population moments, and  $\theta_1^* > 0$ ,  $0 \le p^* \le 1$  if  $(m_1', m_2', m_3')$  is sufficiently close to  $(\mathcal{M}_1', \mathcal{M}_2', \mathcal{M}_3')$ .

If  $\theta_1 = \theta_2 = \theta$ , then the behavior of the estimators change radically. Thus,  $\mathcal{M}_1 = \theta$ ,  $\mathcal{M}_2 = 2\theta^2$ ,  $\mathcal{M}_3 = 6\theta^3$ ; and therefore,  $2 \mathcal{M}_1 = \mathcal{M}_2 = \mathcal{M}_3 =$ 

Thus, the coefficients in the quadratic equation (25), multiplied by  $n^{\frac{1}{2}}$ , are normally distributed in the limit as n

approaches infinity, with zero means and finite and positive variances. This implies that  $\theta_1^*$  and  $\theta_2^*$  have no constant limits in probability and their imaginary parts do not become negligibly small as n increases. In particular, the estimators are not consistent in this case.

The reliability of the estimators  $\theta_1^*$  and  $\theta_2^*$  can be tested if it is temporarily assumed p is known. Otherwise, the calculation of the variances of the three estimators is difficult. With p known, only two sample moments are needed to estimate  $\theta_1$  and  $\theta_2$ . Using equations (18) and (19),

(27) 
$$\theta_{1}^{*} = m_{1}^{1} + \left(\frac{9}{2p}\right)^{1/2} \left(m_{2}^{2} - 2m_{1}^{2}\right)^{1/2}$$

where q = (1 - p). If  $\theta_1 \geq \theta_2$ , use the upper sign; otherwise, use the lower sign if  $\theta_1 \leq \theta_2$ . One can see the shortcoming of the methods of moments because it is the knowledge about  $\theta_1$  and  $\theta_2$  that we desire. Knowing whether  $\theta_1$  is  $> or < \theta_2$  will tell us which pair of estimators is consistent.

Cramer (1958), p. 354, gives a theorem for the asymptotic variance of  $\theta_1^{\#}$  :

(28) 
$$\operatorname{Var} \theta_{1}^{\#} = \mathcal{U}_{2}(m_{2}^{\prime}) \left( \frac{\partial \theta_{1}^{\prime \dagger}}{\partial m_{1}^{\prime}} \right)^{2} + 2 \mathcal{U}_{i_{1}}(m_{1}^{\prime}, m_{2}^{\prime}) \frac{\partial \theta_{1}^{\dagger}}{\partial m_{1}^{\prime}} \frac{\partial \theta_{1}^{\prime \dagger}}{\partial m_{2}^{\prime}} + \mathcal{U}_{2}(m_{2}^{\prime}) \left( \frac{\partial \theta_{1}^{\prime \dagger}}{\partial m_{2}^{\prime}} \right)^{2};$$

Here  $\mathcal{M}_2(\mathfrak{m}_1^i)$  and  $\mathcal{M}_2(\mathfrak{m}_2^i)$  are the variances of  $\mathfrak{m}_1^i$  and  $\mathfrak{m}_2^i$  respectively with  $\mathcal{M}_{11}$   $(\mathfrak{m}_1^i, \mathfrak{m}_2^i)$  the covariance of these two moments. The partial derivatives are to be evaluated at the point

(29) 
$$m_1' = p \theta_1 + q \theta_2$$
 ,  $m_2' = 2(p \theta_1^2 + q \theta_2^2)$ 

Rider (1961) gives formulas for finding the values of the coefficients of the partial derivatives in (29).

(30) 
$$\mathcal{U}_{2}(m_{i}') = n^{-1} \left[ (2p - p^{2}) \theta_{i}^{4} - 2pq\theta_{i}^{2} \theta_{2}^{2} + (2q - q^{2}) \theta_{2}^{4} \right]$$

(31) 
$$\mathcal{U}_{\eta}(m_1', m_2') = 2\eta^{-1} \left[ (3p-p^2)\theta_1^3 - pq\theta_1^2\theta_2 - pq\theta_1\theta_2^2 + (3q-q^2)\theta_2^3 \right]$$

(32) 
$$\mathcal{M}_{2}(m_{2}^{\prime}) = 4n^{-1} \left[ (6p - p^{2}) \theta_{1}^{4} - 2pq \theta_{1}^{2} \theta_{2}^{2} + (6q - q^{2}) \theta_{2}^{4} \right]$$

If  $\theta_1 > \theta_2$ , the partial derivatives needed are

(33) 
$$\frac{\partial \theta_1^*}{\partial m_1'} = 1 - \frac{2^{1/2} \ 9^{1/2} \ m_1'}{p^{1/2} (m_2' - 2 m_1')^2}$$

(34) 
$$\frac{\partial \theta_{1}^{*}}{\partial m_{2}^{\prime}} = \frac{g^{h_{2}}}{2^{3h_{2}} p^{h_{2}} (m_{2}^{\prime} - 2m_{1}^{\prime 2})^{h_{2}}}$$

At the point (29), these derivatives have values

(35) 
$$\frac{\partial \theta_1^*}{\partial m_1'} = \frac{-\theta_2}{p(\theta_1 - \theta_2)} , \frac{\partial \theta_2^*}{\partial m_2'} = \frac{1}{4 p(\theta_1 - \theta_2)}$$

If  $\theta_1 < \theta_2$ , change the signs of the fractions on the righthand sides of (33) and (34), and again (35) is obtained.

If one substituted (30), (31), (32), (33) in (27), and simplified, the variance of the asymptotic distribution of  $\theta_1^{\phi}$  is

(36) 
$$\frac{1}{4n p^{2}(\theta_{1}-\theta_{2})} \left[ p(6-p)\theta_{1}^{4} - 4p(3-p)\theta_{1}^{3}\theta_{2} + 2p(5-3p)\theta_{1}^{2}\theta_{2}^{2} - 4p(1-p)\theta_{1}\theta_{3}^{3} + (1-p^{2})\theta_{2}^{4} \right].$$

The variance of the asymptotic distribution of  $\theta_2^*$  may be obtained by replacing p by q and interchanging  $\theta_1$  and  $\theta_2$  in (36).

Rider (1961) stated that data should not be assumed to have come from a mixed exponential distribution until it has been determined that they have not come from a single exponential distribution  $\frac{1}{\theta} e^{-t/\theta}$ . That is,  $\theta$  of this distribution should be estimated and a chi-square test made to see whether the data conforms to this distribution. Should the hypothesis be rejected, then a mixed exponential population may be assumed.

It is possible that the chi-square test could give a wrong conclusion; whereupon, it would be impossible for the method of moments to estimate  $\theta$  according to Rider (1961). Should the population be mixed and  $\theta_1$  nearly equal to  $\theta_2$ , it still may be difficult to obtain valid estimates of them. Therefore, for practical purposes, estimators are not recommended until further research reveals some way to correct their shortcomings.

APPLICATION TO EXPONENTIAL DISTRIBUTIONS AND ESTIMATION OF PARAMETERS BY THE METHOD OF MAXIMUM LIKELIHOOD ESTIMATORS

Mendenhall and Hader (1958) attacked the problem of estimating the parameters of a population that was obtained by mixing two exponential failure time distributions in unknown proportions, the population model being based upon censored sampling. After a predetermined length of time had elapsed or after a predetermined number of units had failed, the life test was concluded. It was assumed that each unit of the population conceptually contained a tag indicating the subpopulation to which the unit belonged. Of course, this unit or tag of information is only available after failure has occurred.

The estimation of population parameters will be by the method of maximum likelihood estimators. Consider a population composed of s=2 subpopulations representing failure types, mixed in proportions p and q=(1-p), where  $0 \le p \le 1$ . The test termination time, T, is in units of size T at which time r units have failed,  $r_1$  from subpopulation (i) and  $r=r_1+r_2$ . The time of failure of the jth unit from subpopulation (i),  $t_{ij}$  is observed. It is assumed that j always ranges from j=1 to  $r_1$  when not specified. The (n-r) are the units which have not failed and yield no information as to the subpopulation they were drawn from. The cumulative failure probability distribution from equation (1) assuming G(x) = constant = 1s

$$F(t) = 1 - e \qquad , \quad 0 \le t < \infty$$

or to be more general

$$F_i(t) = 1 - e^{-t/\theta_i}$$
,  $0 \le t < \infty$ 

where 1 = 1 or 2.

Now let x = t/T and  $\beta_i = \theta_1/T$ , then

(37) 
$$F_{i}(x) = 1 - e^{-x/\beta_{i}}$$
,  $0 \le x < \infty$ 

If p is the proportion of units belonging to subpopulation i=1, then the cumulative distribution function for the population is

(38) 
$$F(t) = pF_1(t) + qF_2(t)$$

and the density function,

(39) 
$$f(t) = p f_1(t) + q f_2(t)$$

Also let

(40) 
$$H_i(t) = 1 - F_i(t)$$

and

(41) 
$$H(t) = 1 - F(t)$$

where H(t) is the probability that a unit will survive to time +.

The probability of  $r_1$  units failing due to cause (1) and  $r_2$  units failing due to cause (2), and (n-r) units surviving at t-1, given a random sample of n units, the multinomial

(42) 
$$P\left\{r_{i}, r_{2}, (n-r) \mid n\right\} = \frac{n!}{r_{i}! r_{2}! (n-r)!} \left[PF_{i}(1)\right]^{n} \left[QF_{2}(1)\right]^{r_{2}} \left[H(1)\right]^{(n-r)}$$

The conditional density of obtaining ordered observations,

$$(x_{11}, x_{12}, \dots, x_{1r_1} | r_1)$$
 and  $x_{1j} \le 1$  is
$$r_i \mid \frac{r_i}{l}, f_i(X_{ij})$$

(43) 
$$P\left\{X_{ii}, X_{i2}, \dots, X_{ir_{i}} \mid r_{i}, X_{ij} \leq 1\right\} = \frac{r_{i}! \prod_{j=1}^{i} f_{i}(X_{ij})}{\left[F_{i}(I)\right]^{r_{i}}}$$

From (42) and (43) the likelihood for this sample becomes

(44) 
$$L = \frac{n!}{(n-r)!} H(1)^{(n-r)} p^{r_1} q^{r_2} \prod_{j=1}^{r_2} f_i(X_{1j}) \prod_{j=1}^{r_2} f_2(X_{2j})$$

(45) 
$$\frac{\partial \ln L}{\partial \beta_1} = \frac{K(N-r)}{\beta_1^2} - \frac{r_1}{\beta_1} + \frac{r_1 \overline{\chi}_1}{\beta_1^2}$$

(46) 
$$\frac{\partial \ln L}{\partial \beta_2} = \frac{(I-K)(N-r)}{\beta_2^2} - \frac{r_2}{\beta_2} + \frac{r_2}{\beta_2^2} \frac{\overline{\chi}_2}{\beta_2^2}$$

(47) 
$$\frac{\partial \ln L}{\partial p} = \frac{K(n-r) + r_1}{p} - \frac{(1-K)(n-r) + r_2}{q}$$

where

$$K = \frac{p e^{-1/\beta_1}}{p e^{-1/\beta_2} + q e^{-1/\beta_2}} = \frac{1}{1 + \left(\frac{q}{p}\right) e^{\left(\frac{1}{\beta_1} - \frac{1}{\beta_2}\right)}}$$

At time t, the subpopulations would be mixed in the proportions p(t) and 1 - p(t).

(48) 
$$\therefore p(t) = \frac{pH_{i}(t)}{H(t)} , p(0) = p .$$

Therefore, k = p(1), the conditional mixture proportion at the test termination time, x = 1.

When (43), (44), and (45) are equated to zero, the estimating equations are

$$(49) \qquad \stackrel{\wedge}{p} = \frac{r}{n} + \stackrel{\wedge}{K} \frac{(n-r)}{n} \qquad ,$$

$$\beta_1 = \overline{\chi}_1 + \frac{\Lambda}{K} \frac{(n-r)}{r_1}$$

(51) 
$$\beta_2 = \overline{\chi}_2 + \underline{(I-\hat{\chi}(N-r))}_2,$$

where

(52) 
$$\overline{X}_{i} = \frac{\sum_{j=1}^{r_{i}} X_{ij}}{r_{i}} , \quad \hat{K} = \frac{1}{1 + \left(\frac{\hat{G}}{\hat{F}}\right)} e^{(\hat{F}_{i}, -\hat{F}_{i})}$$

Now solve (49), (50), (51) and (52) simultaneously to obtain estimates of  $\beta_1$ ,  $\beta_2$ , and p. If (49), (50) and (51) are substituted into (52) and (52) is solved for k, a single equation is obtained

$$\hat{K} = h(\hat{K})$$
 ,  $(o \le \hat{K} \le 1)$ 

where h(k) is a function of k. k can be obtained by plotting h(k) - k versus k and obtaining the solution where h(k) - k = 0. When k = 0, h(k) - k will be positive or zero.

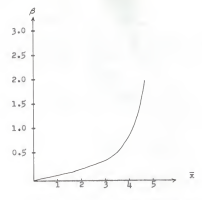


Figure 1. Maximum likelihood estimator of  $\rho$  as a function of  $\bar{x}$  based on a sample from a truncated exponential distribution. Measurements expressed in units of truncation time T.

Mendenhall and Hader (1958) give a procedure to find a good approximation of k for samples drawn from a single truncated exponential distribution. Actually, it is obtained by making a modification of the maximum likelihood estimate. Where the exponential distribution is assumed to be truncated at time T, then the maximum likelihood estimate of  $\beta_1$  is a solution of

(53) 
$$(\beta_i - \bar{x}_i)(e^{-\frac{1}{\beta_i}} - 1) = 1$$
.

Thus, solutions of  $\hat{\beta}_1$  are obtained graphically from figure 1 where  $\hat{\beta}_1$  is given as a function of  $\bar{x}_1$ . Subpopulation (1) is chosen as the smaller  $\bar{x}$  and  $\hat{\beta}_{10}$  is the corresponding value  $\bar{x}$  obtained from figure 1. Now substitute into (50) and solve for  $\hat{k}_0$ .

(54) 
$$\hat{\beta}_{io} = \bar{X}_i + \hat{K}_o \frac{(n-r)}{r_i}$$

Now the quantity  $A_0 = g(\overset{\circ}{k}_0) - \overset{\circ}{k}$  can be solved. Thus, a solution to equations (49), (50), (51) and (52) is possible if A = 0. Since  $g(0) \ge 0$ , then the value of  $\overset{\circ}{k}$  which will satisfy A = 0, must be  $\overset{\circ}{k} < k_0$  or  $\overset{\circ}{k} > k_0$ . It all depends upon whether  $A_0$  is negative or positive.

Let 
$$W = \frac{\hat{q}}{\hat{\beta}} \in \frac{(/\hat{\beta}_1 - /\hat{\beta}_2)}{\hat{\beta}}$$

(56) 
$$d\hat{K} = \frac{-dA}{1 + g(\hat{K})^2 \left(\frac{dW}{d\hat{K}}\right)}$$

where

(57) 
$$\frac{dW}{d\hat{K}} = -W(n-r) \left[ \frac{1}{\hat{q}} + \frac{1}{r_1} \hat{\beta}_1^2 + \frac{1}{r_2} \hat{\beta}_2^2 \right]$$

Choosing

Therefore, this iterative process may be repeated until one achieves the degree of accuracy desired. Then the solution for k obtained can be substituted into the estimating equations (49), (50) and (51) to find the estimates p,  $\beta_1$ , and  $\beta_2$ .

Should it happen that  $\mathbf{r}_1=0$ , then there could be no estimates of  $\boldsymbol{\beta}_1$  obtained from the estimating equations. Really, this is not a problem in the practical sense because it can reasonably be concluded that  $\boldsymbol{\beta}_1$  must be very large or else  $\mathbf{p}_1=0$ . So, let us adopt the convention that when  $\hat{\boldsymbol{\beta}}_1=\infty$ , we shall mean that  $\boldsymbol{\beta}_1$  is very large when  $\mathbf{r}_1=0$ . In experimental work it is desired to choose T and n large enough that the probability that  $\mathbf{r}_1=0$  or  $\mathbf{r}_2=0$  is very small. Actually, one could not expect to get any information on failure parameters if he is not willing enough to test until some failures are observed.

The maximum likelihood procedure appears to give satisfactory results under the following conditions:

 when the sample size is large. Tukey (1960), p. 463, stated that a statistical problem is usually called a

- large sample problem when 1/n is so small everything else can be neglected.
- 2) when the test termination time, T, is large relative to  $\theta_1 \text{ and } \theta_2 \text{ because } \beta_1 = \theta_1/\text{T must be very small to}$  have an efficient estimation  $E(\mathbf{r_i})$  proportional to  $\beta_1$ . When n and T are small, the estimates are badly biased and have large variances. Thus, it would seem desirable to investigate other estimators having better small sample properties.

Rao (1948) stated that for higher efficiency the estimates of the parameters must be found by the method of maximum likelihood because this gives rise to the "best" estimates in large samples. Thus, in practice one would not use the method of moments with large samples since it is not as efficient as the method of maximum likelihood.

The relative magnitudes of  $\theta_1$  and  $\theta_2$  will be known in most experimental situations; hence, the estimating procedure may be modified in a simple way when this is true. This modification will reduce the bias and variances of the estimates.

The disadvantage of maximum likelihood estimates is that the equations leading to them are usually non linear and thus difficult to solve.

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APPENDIX

#### Numerical Example

Since the method of maximum likelihood estimation is more efficient than the method of moments where the sample size n is very large, the method of maximum likelihood has been chosen to illustrate a numerical problem (Mendenhall and Hader, 1958) to utilize the theory developed in this paper.

In this numerical example it was the policy to remove all ARC-1 VHF communication transmitter-receivers which had operated for 630 hours from the aircraft. Thus, T = 630 hours was the fixed time at which the sample was censored. Units which failed were removed from the aircraft for maintenance. The failure times for the ith population are assumed to have a cumulative failure probability distribution defined by

$$F_1(t) = 1 - \exp -t / \Theta_1$$
 ,  $0 \le t < \infty$ .

In some cases the apparent failure of the units were not confirmed and upon arrival at the maintenance center were found to operate satisfactorily. This makes it desirable to estimate that fraction of unconfirmed failures in the population. Thus, the sample of failures may be subdivided into two subpopulations, 1) confirmed failures and 2) unconfirmed failures (shown in tables 1 and 2 respectively).

To identify the two subpopulations let the unconfirmed failures be called subpopulation (1) and the confirmed failures as subpopulation (2). Using tables 1 and 2, one obtains the

Table 1. Confirmed failures. Hours to failure for ARC-1 VHF radio transmitter receivers\*

16 392 408 304 256 488 60 360 168 96 616 328 80 80 85 272 72 1184 152	224 576 384 16 194 216 120 208 232 72 168 224 464 72 168 288 268 268 268 208	16 128 256 722 136 168 208 440 644 1140 408 424 1328 400 168 160	80 56 246 824 184 32 104 112 40 80 32 616 608 264 2480 152 352 176	128 112 184 88 32 144 1128 128 120 480 192 164 125 128 140 192 164 125 128 140 125 128 140 125 128 129 129 129 129 129 129 129 129 129 129	168 160 160 160 224 288 384 32 152 416 112 126 112 126 408 276 480 276 584	144 384 480 488 488 336 488 336 488 390 390 490 490 490 490 490 490 490 490 490 4	176 600 104 168 120 304 256 360 280 560 104 288 296 560 256 96 96 96 96 176	176 40 168 80 320 40 40 80 104 328 144 122 248 3248 3248 3248 3248 3248 326 360	568 416 408 518 160 296 168 192 203 336 120 552 248 184	

Table 2. Unconfirmed failures. Hours to failure for ARC-1 VHF radio transmitter receivers\*

136 246 168 112 72 208 112 272	512 72 120 56 168 114 96 320	136 80 616 184 288 480 64	472 312 24 40 112 114 360 440	96 24 176 256 80 392 136 224	144 128 16 160 584 120 168 280	112 304 24 456 368 48 176	104 16 32 48 272 104 256	104 320 232 24 208 272 112 216	
272 256	320 104	104	440	224 304	280 240	88	56 248	216 472	
88 184	400	392	168	72 152	184	88	176	216	
	136 246 168 112 72 208 112 272 256 88 184	136 512 246 72 168 120 112 56 72 168 208 114 112 96 272 320 256 104 88 200 184 400	136 512 136 246 72 80 168 120 616 112 56 184 72 168 288 208 114 480 112 96 64 272 320 8 256 104 104 88 200 392 184 400 424	136 512 136 472 216 72 80 312 168 120 616 21 112 56 181 40 72 168 288 112 208 111 480 111 112 96 61 360 272 320 8 440 256 104 104 8 88 200 392 168 184 400 424 88	136 512 136 472 96 216 72 80 312 24 168 120 616 24 176 112 56 184 40 256 72 168 288 112 80 208 114 480 114 392 112 96 64 360 136 272 320 8 440 224 256 104 104 8 304 88 200 392 168 72 184 400 424 88 152	136   512   136   172   96   144     246   72   80   312   24   128     168   120   616   24   176   16     112   56   184   40   256   160     72   168   288   112   80   584     208   114   480   114   392   120     112   96   64   360   136   168     272   320   8   440   224   280     256   104   104   8   304   240     280   280   392   168   72   40     184   400   424   88   152   184	136   512   136   172   96   144   112     246   72   80   312   24   128   304     168   120   616   24   176   16   24     112   56   184   40   256   160   456     72   168   268   112   80   584   368     208   114   480   114   392   120   48     112   96   64   360   136   168   176     272   320   8   440   224   280   8     256   104   104   8   304   240   88     88   200   392   168   72   40   88     184   400   424   68   152   184   -	136   512   136   172   96   114   112   104     214   72   80   312   24   128   304   16     168   120   616   24   176   16   24   32     112   56   184   40   256   160   456   18     72   168   288   112   80   584   368   272     208   114   1480   114   392   120   48   104     112   96   64   360   136   168   176   256     272   320   8   140   224   280   8   56     256   104   104   8   304   240   88   248     88   200   392   168   72   40   88   248     184   400   424   88   152   184	136   512   136   172   96   114   112   104   104     246   72   80   312   24   128   304   16   320     168   120   616   24   176   16   24   32   232     112   56   184   40   256   160   456   48   24     72   168   268   112   80   584   368   272   208     208   114   148   114   392   120   48   104   272     112   96   64   360   136   168   176   256   112     272   320   8   140   224   280   8   56   216     256   104   104   8   304   240   88   248   472     88   200   392   168   72   40   88   176   216     184   400   124   88   152   184

<sup>\*</sup>Tables reproduced from Biometrika 1958, 45(3 and 4): 509.

following data:

where r is the number of units failed in each subpopulation and n is the random sample size.

$$n-r = 369 - 325 = 444$$
  $\bar{x}_1 = \frac{\bar{t}_1}{\bar{T}} = 0.3034862$   $\bar{x}_2 = \frac{\bar{t}_2}{\bar{T}} = 0.3644677$ 

t is the time the unit survived which is a proportional part of the fixed time T at which the life of a unit is censored.

Now one can form the estimating equations by making use of equations (49), (50) and (51).

(60) 
$$p = 0.2900 + 0.1192 k$$

(61) 
$$\beta_1 = 0.3035 + 0.4112 \text{ k}$$

(62) 
$$\hat{\beta}_2 = 0.5663 - 0.2018 \text{ k}$$

One can simplify the process of obtaining an iterative solution by making use of table 3.

The first step of the procedure is to use  $\bar{\mathbf{x}}=0.303$  and enter figure 1. Thus, the corresponding estimate of  $\beta_1$  is  $\hat{\beta}_{10}=0.380$ ; and the corresponding value of  $\hat{\mathbf{k}}$ ,  $\hat{\mathbf{k}}_0=0.186$  is obtained by using the estimating equation (61). Knowing  $\mathbf{k}_0$ ,  $\hat{\beta}_{20}$  and  $\hat{\mathbf{p}}_0$ , can be easily obtained. Table 3 shows these values in row  $\mathbf{v}=\mathbf{0}$ .

Table 3. Record of iterations

٧	ř.	Â <sub>lv</sub>	β <sub>2▼</sub>	p <sub>V</sub>	W	g(k <sub>w</sub> )	A
0	0.186	0.380	0.529	0.312	4.622	0.1779	-0.0081
2	.167	.3721	.5326 .5330	·3099 ·3097	5.002	.1666 .1654	0001

Next we compute

(63) 
$$g(\hat{k}_0) = \frac{1}{1 + \left(\frac{\hat{q}_0}{\hat{q}_0}\right) \exp\left[\frac{1}{\hat{\beta}_{10}} - \frac{1}{\hat{\beta}_{20}}\right]}$$

and  $A_0 = g(\hat{k}_0) - \hat{k}_0 = -0.0081$  follows (Mendenhall and Hader, 1958). The value of  $\hat{k}$  will be obtained whenever there exists a solution to the maximum likelihood equations, i.e., A = 0. Since A can be either positive or zero when  $\hat{k} = 0$  and negative when  $\hat{k} = 0.186$ , then there exists a solution for  $\hat{k}$  when  $0 < \hat{k} < 0.186$ . Therefore the value  $\hat{k}$  has when  $\hat{i} = 1$  must be less than 0.186. By use of equation (56) we can calculate the change in  $\hat{k}$ ,  $d\hat{k}_0$ .

(6\(\beta\)) 
$$\frac{d\hat{k}_{o}}{1 + g(\hat{k}_{o})^{2} \left(\frac{dw_{o}}{d\hat{k}_{o}}\right)} = \frac{(-0.0081)}{1 + (0.1779)^{2} (-19.0\(\beta\))}$$
$$= -0.02.$$

Thus, 
$$\hat{k}_1 = \hat{k}_0 + d\hat{k}_0 = 0.186 - 0.02 = 0.166$$
,  $\hat{\beta}_{11} = 0.3718$ ,  $\hat{\beta}_{21} = 0.5328$ ,  $\hat{p}_1 = 0.3098$ .

Since the values of these estimating parameters make  $A_1=0.0000$ , then essentially these estimates are the maximum likelihood estimates of the parameters. If a bound is desired on the iteration error, then it can be obtained by calculating A for  $\hat{k}_2=0.167$  and  $\hat{k}_3=0.165$ . If the solution for  $\hat{k}$  is taken to be 0.166 and since  $A_2=-0.0004$  is negative and  $A_3=0.0004$  is positive, then the absolute value of the iterative error for  $\hat{k}$  is less than 0.001.

The estimate of the unconfirmed failures is  $\hat{p}$  = 0.3098 and one can find their estimated average life to be

$$\hat{\Theta}_1 = \hat{\beta}_1 T = (0.3718)(630) = 234.2 \text{ hours.}$$

The estimate of the average life for the confirmed failures is

$$\hat{\theta}_2 = \hat{\beta}_{2^{\text{T}}} = (0.5328)(630) = 335.7 \text{ hours.}$$

Although the accuracy above was good for the estimation equations when only one iteration was used, it may be that one or two iterations on the maximum likelihood equations may not be sufficient. Therefore, one should put bounds on the iterative error as was done in this numerical example.

It can be seen that the estimates of the average life of units from two subpopulations,  $\theta_1$  and  $\theta_2$ , can be quite useful in maintaining excellent service in the communications field as one can anticipate maintenance requirements before any actual break down in service results. Thus, one can see the importance and use of estimation theory.

#### MIXTURES OF DISTRIBUTIONS

by

RAY LEE MILLER, JR.

B. S., University of Missouri, 1957

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Department of Statistics

KANSAS STATE UNIVERSITY Manhattan, Kansas

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Mixtures of distributions occur when there is overlapping of two or more distributions so that it is difficult to separate them into their respective components. Feller has distinguished two types of these distributions, true contagion and apparent contagion.

Karl Pearson attacked the problem of mixed frequency distributions in the early 20th century. About 25 years later Greenwood and Yule developed a very general scheme for contagious events but this proved to be too complex for most practical applications. Several years later Polya and Eggenberger considered a special model of true contagion which was the simplest case of the general Greenwood-Yule scheme. The next significant advancement in the study of contagious distributions was in 1939 when Neyman applied them to entomology and bacteriology. Since that time Neyman's distributions have been generalized by several workers among whom were Feller and Gurland.

Several definitions of distributions have been cited and developed. Some of these were the compound Poisson, Neyman's contagious distribution of type A, Polya-Eggenberger, and Pearson's type III distributions. Also, a theorem of mixtures of distributions was stated and proved.

Applications to exponential distributions and estimation of parameters were done by methods of moments and methods of maximum likelihood. Since the method of maximum likelihood estimation is more efficient than the method of moments where

the sample size n is very large, a numerical example illustrating the use of the method of maximum likelihood was applied to an exponential distribution.