Taylor’s power law and the statistical modelling of infectious disease surveillance data

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Abstract

Surveillance data collected on several hundred different infectious organisms over twenty years have revealed striking power relationships between variance and mean in successive time periods. Such patterns are common in ecology, where they are referred to collectively as Taylor’s power law. In this paper, these relationships are investigated in detail, with the aim of exploiting them for the descriptive statistical modelling of infectious disease surveillance data. We confirm the existence of variance to mean power relationships, with exponent typically between 1 and 2. We investigate skewness to mean relationships, which are found broadly to match those expected of Tweedie distributions, and thus confirm the relevance of the Tweedie convergence theorem in this context. We suggest that variance and skewness to mean power laws, when present, should inform statistical modelling of infectious disease surveillance data, notably in descriptive analysis, model building, simulation and interval and threshold estimation, the latter being particularly relevant to outbreak detection.

Key words: exponential dispersion model, infectious disease, power law, surveillance, Taylor’s Law, Tweedie family.

1 Introduction

This paper is devoted to an exploration of variance to mean relationships in surveillance data on infectious diseases, focusing on implications for descriptive statistical modelling. The paper stems from a recent epidemiological study that suggested clear power relationships between mean and variance (Enki et al 2013). The present paper is devoted to a further exploration of these patterns.

Enki et al (2013) undertook a review of the UK statistical surveillance system for outbreak detection, encompassing the 3,303 different organism types reported to the UK LabBase database over the 20 year period 1991 to 2011. Our investigations revealed striking linear relationships between the logarithm of the variance and the
logarithm of the mean of the number of organisms detected in successive six-monthly
time periods, which appears to be present for virtually all of the organism types
for which sufficient data were available. Such variance to mean power laws have
previously been observed for measles and whooping cough (Keeling and Grenfell
1999). Our data suggest they are ubiquitous for infectious diseases, whatever the
mode of transmission.

It is desirable to use statistical models that correctly account for the variance
to mean relationship (see for example McCullagh and Nelder 1989, pages 328-332).
Thus, investigating the variance to mean relationship is a useful preparatory step
for statistical modelling: for example, a Poisson or quasi-Poisson model might be
indicated if the variance is proportional to the mean. In the case of exponential
family models, the variance function largely determines the distribution. In turn,
quantiles of this distribution may be important in some applications, notably for
determining threshold values, as needed for detecting outbreaks of infectious dis-
eeses. These considerations apply to individual data sets; in our case we have access
to data on hundreds of different organisms. The existence of power variance to
mean relationships for virtually all these organisms is of intrinsic interest, may help
in elucidating underlying mechanisms, and should inform the statistical modelling
framework used represent the data.

Power variance to mean relationships are instances of Taylor’s power law, re-
viewed in Kendal (2004). Taylor’s law originally related to the observation that
populations within territories partitioned into quadrats exhibit a variance to mean
power relationship of the form

\[ \text{var}(X) = \phi E(X)^p \]

where \( X \) is the population count within a quadrat (Taylor 1961). Various mecha-
nisms, discussed in Kendal’s review, have been advanced to explain such empirical
power laws, which have been observed in very different contexts. For example, they
have been shown to result from certain types of birth and death processes with im-
migration (Anderson et al 1982, Keeling 2000). Keeling and Grenfell (1999) showed that simulations with their stochastic, pulsed realistic age-structured SEIR model produce power variance to mean relationships at higher incidences. It has also been suggested that Taylor’s law is the empirical manifestation of the asymptotic limiting behaviour of exponential dispersion models, which in turn is linked to the properties of the Tweedie family of distributions (Kendal 2004, Kendal and Jørgensen 2011). Finally, note that the power laws considered here relate to abundances, and should not be confused with power laws governing other features, such as spatial spread (Brockmann et al 2006, Meyer and Held 2014).

In the present paper we do not seek to investigate in any detail the underlying mechanisms which generate power laws in infectious disease surveillance data. We do, however, study the evidence that the data conform to Tweedie distributions, as one might expect if the observed power relationships are the result of Tweedie convergence.

The paper has three aims: first, to investigate in greater detail and with greater rigour the empirical evidence for power laws in observed variance to mean relationships in infectious disease surveillance data; second, to explore whether these data conform to the limiting behaviour of exponential dispersion models; and third, to seek to exploit the observed variance to mean relationships for the purposes of descriptive statistical modelling. In Section 2, we describe the data and present the evidence for a variance to mean power law. In Section 3, we briefly review the Tweedie family of exponential dispersion models. In Section 4 we investigate the evidence that our data conform (in an asymptotic sense) to this family. In Section 5 we discuss the use of the power law and Tweedie models for infectious disease modelling, notably outbreak detection. These methods are applied to surveillance data in Section 6. The paper ends with a discussion of the potential and limitations of this approach in Section 7.
2 Variance-mean power relationships in infectious disease surveillance data

We used weekly counts of infectious organisms reported to Public Health England’s LabBase database between week 1 of 1991 and week 26 of 2011. This is a computerized database of reports of infectious disease organisms identified in biological specimens (taken from samples of blood, faeces, or urine) collected by laboratories in England, Wales, and Northern Ireland. These specimens are obtained primarily for diagnostic purposes, rather than for routine screening. The laboratory identification of causative organisms in the specimens analysed is highly accurate, but the weekly counts generally represent only a small proportion of infections, except perhaps in some outbreaks. The laboratory protocols and classifications used have changed over time, so that for some organisms data are not available for the entire 20 year period. Our data were arranged by week of collection of the specimen from which the infection was identified. There is an overall increasing trend in numbers of isolates and numbers of organisms reported over time; more details of these trends and other aspects of these data may be found in Enki et al (2013). These data form a key resource for outbreak detection in England, Wales, and Northern Ireland. A similar system is operated in Scotland by Public Health Scotland.

2.1 Empirical variance to mean relationships

We explored the variance to mean relationship in successive 6-month periods. First, we de-seasonalised the data where appropriate. This was done by fitting a quasi-Poisson generalized additive model with log link, smooth trend and a seasonal factor with levels $\gamma_j$, $j = 1, \ldots, 12$. The de-seasonalised data are then obtained from the weekly counts $y_t$ as follows:

$$z_t = y_t \exp(\bar{\gamma} - \gamma_{s(i)})$$
where $\bar{\gamma}$ is the average of the $\gamma_j$ and $s(i)$ is the seasonal level for week $i$. We only applied this seasonal adjustment for 1015 organisms with nonzero counts in every season; for other organisms, too sparse to apply a meaningful seasonal adjustment, we used $z_i = y_i$. We then grouped the data in 41 6-monthly periods. Within each calendar year there are two such periods, indexed by $k = 1$ for the period January to June, and $k = 2$ for July to December.

If Taylor’s law is deemed to apply to the weekly counts $y_i$ with

$$\log\{\text{var}(y_i)\} = \alpha + p \log\{E(y_i)\}$$

then

$$\log\{\text{var}(z_i)\} = \alpha_s(i) + p \log\{E(z_i)\}$$

with $\alpha_s(i) = \alpha + (2 - p)(\bar{\gamma} - \gamma_s(i))$. Now let $W_{jk}$ denote the set of weeks $i$ within period $k$ of year $j$, and assume that $E(z_i)$ is roughly constant for $i \in W_{jk}$. We fit the model

$$\log\{\text{var}(z_i)\} = \alpha_k + p \log\{E(z_i)\}, \quad i \in W_{jk}$$

which describes a linear relationship with two intercepts, corresponding to $k = 1, 2$. When the data are sparse, and are not de-seasonalised, we just used one intercept.

Perry (1981) showed that simple regression of the logarithm of the empirical variance against the logarithm of the empirical mean (as used in Enki et al 2013) produces a negatively biased estimate of $p$ (that is, $\hat{p}$ is too low), and suggested that a gamma generalised linear model be used instead. This produces the same point estimates as the approach of Jørgensen et al (2011) using unbiased estimating equations.

Accordingly, we applied the gamma model to the 1737 different organism types with sufficient data to estimate the regression parameters, with two intercepts when de-seasonalized, and a single intercept when not.

In all cases, the empirical relationship between log variance and log mean is linear. Figure 1 shows the plots, together with the gamma regressions, for six common
organisms. The full set of 1737 plots is in the Supplementary Materials. Figure 2 shows caterpillar plots of values of \( \hat{p} \), along with approximate 95% confidence intervals. The variances of the estimates \( \hat{p} \) were obtained using the scale parameters for the gamma regression models based on Pearson’s chi-square statistic, which are consistent provided that the \( p \) have been consistently estimated (McCullagh and Nelder 1989, page 296). On the left is the caterpillar plot for all 1737 organisms. The strange appearance of the leftmost tail of the plot is the result of sparse organism counts: when there is a single count in a six-month period, then the estimated mean and variance are identical. In the extreme case where this is true of all six-monthly periods, the dispersion (and hence the width of the confidence interval) is zero. The caterpillar plot on the right of Figure 2 is restricted to the 374 organisms for which the count in every six-month period is at least 2. These caterpillar plots show that most values of \( p \) lie between 1 and 2, and few other than those affected by the sparseness issue just described have 95% confidence intervals entirely located outside this range.

We repeated the analysis with an annual rather than 6-monthly grouping of the data (and consequently a single intercept). The slopes are similar to those obtained with 6-monthly periods, as shown by the scatterplot in Figure 3, except for the organisms with larger slopes. We suspect that residual periodicity may contribute to inflating the variances in the analysis based on annual data. These results confirm the striking linear relationships between log variance and log mean observed by Enki et al (2013). These are strongly suggestive of power variance-to-mean relationships, with exponent typically lying between 1 and 2. These observations lend empirical support for Taylor’s law over an extremely wide range of different micro and macro-organisms.
2.2 Serial dependence

Throughout the paper, we focus on the marginal distribution of the counts, and ignore any correlation between successive weekly counts. Since the data relate to infectious diseases, which are transmitted directly or indirectly over time, this requires some justification. For 1774 organisms with sufficient data to do so (including the 1737 organisms to which we fitted regressions as described above), we obtained roughly stationary residuals by fitting a quasi-Poisson generalized additive model with log link, smooth trend with knots every 26 weeks, and a 12-level seasonal factor for the 1015 organisms with non-zero counts in every season. Let $\mu_i$ denote the expected value in week $i$ from this model; the residual was defined as

$$r_i = (y_i - \hat{\mu}_i)\hat{\mu}_i^{-\frac{1}{2}}.$$
Figure 2: Caterpillar plot of estimated values of slopes and corresponding 95% confidence limits. Left: all organisms; Right: excluding sparse organisms.

We then obtained correlograms for the 1774 organisms; these are in the Supplementary Materials. Similar results were obtained with residuals based on $\log(y_i + 1)$. The correlograms show that for the majority of organisms, there is little serial dependence. The Ljung-Box portmanteau test applied to the first 30 lags gives a $p$-value greater than 0.05 for 1127 (64%) of organisms. Furthermore, owing to the large sample size, autocorrelations are flagged as significant even when close to zero. Figure 4 shows the histogram of the 1774 lag 1 autocorrelations $\rho_1$. Of the 647 organisms with significant serial correlation, 269 (42%) had negative lag 1 autocorrelations. These typically occur for uncommon organisms. Many of the large positive lag 1 autocorrelations correspond to organisms such as Cryptosporidium, Norovirus, Respiratory Syncytial Virus and Rotavirus with substantial seasonality which is incompletely removed by the seasonal factor, often owing to fluctuating amplitude or periodicity. In a few cases, notably Influenza A and B, the autocorrelation is substantial and not due to residual periodicity. However, the great majority of organisms display little serial correlation: for 84% of organisms $-0.1 \leq \rho_1 \leq 0.1$, and for 95% $-0.2 \leq \rho_1 \leq 0.2$. We comment on these observations in the Discussion.
Figure 3: Scatter plot of estimated slopes: bi-annual vs annual grouping.

Figure 5 shows the caterpillar plots of $\hat{p}$ in two subgroups, according to whether serial dependence was or was not significant at the 5% level in the portmanteau test. Power variance to mean relationships with slopes $\hat{p} > 1$ are observed in both groups.

2.3 Factors associated with the power parameter

Figure 6 shows the estimated power parameters $p$ plotted against the logarithm of the median of the 41 seasonally-adjusted counts; organisms for which the median of the seasonally-adjusted counts was less than or equal to 0.05 were grouped into a single category with median 0.05 (the median value of $p$ for these was 1.07). The loess curve superimposed on the scatterplot suggests that the power $p$ tends to increase slightly with the median adjusted count, though the spread of values is considerable at all medians. At low median adjusted counts $p$ is centrally located a little above 1, corresponding to the quasi-Poisson model. The four outliers with values of $p$ close
Figure 4: Autocorrelations at lag 1 for 1774 organisms.

to or above 2.5 are Rotavirus, Cryptosporidium Sp, Herpes simplex virus untyped
and Escherichia coli untyped; residual periodicity may have contributed to inflating
these values. There is also a single outlier with a negative $p$, corresponding to
Neisseria meningitidis type B (types A and C had values of $p$ close to 1). Table 1
gives the values of $p$ for some specific organisms (in some cases averaged over the
values for distinct subtypes). There is no evident relationship with the mode of
transmission.

Figure 7 shows the power parameters $p$ plotted against the log seasonal amplitude
$A$, defined as

$$A = \max\{\gamma_s : s = 1, \ldots, 12\} - \min\{\gamma_s : s = 1, \ldots, 12\}$$

where the $\gamma_s$ are the seasonal factors used to remove seasonality from the data prior
to analysis. When no seasonal factor was fitted we set $A = 0$. The loess plot shows
some positive dependence between $A$ and $p$. That there should be any relationship
might appear strange, since $A$ measures an effect that has been removed from the
Figure 5: Caterpillar plot of estimated values of slopes and corresponding 95% confidence limits. Left: organisms with significant serial correlation; Right: organisms with no significant serial correlation.

Data prior to analysis. Our interpretation, supported by visual inspection of the residuals, is that it reflects residual periodicity remaining after adjustment. Residual periodicity can occur when the seasonal peaks vary slightly in amplitude and timing, and will tend to inflate the estimated values of \( p \). An alternative interpretation is that highly seasonal organisms also have higher values of \( p \) owing to the mechanism driving the seasonality.

Finally, Figure 8 shows the power parameter \( p \) plotted against the lag 1 autocorrelation \( \rho_1 \). The loess plot shows some positive dependence between \( p \) and \( \rho_1 \), driven by the higher positive values of \( \rho_1 \) outside the main cluster of the data. However, once again the interpretation of the plot is complicated by the residual periodicity previously mentioned, which is related both to \( p \) and to \( \rho_1 \). The overall Spearman correlation between \( p \) and \( \rho_1 \) is 0.189. Splitting the data according to the median log seasonal amplitude \( A = 0.45 \), the correlation in the \( A > 0.45 \) group is 0.325, whereas in the \( A \leq 0.45 \) group it is 0.067. The contrast between the two values is consistent with the view that the correlation is to some degree an artefact induced
by residual periodicity. An alternative interpretation is that serial correlation contributes to higher values of $p$ through the underlying mechanism driving both the spread of infection and the seasonality.

3 Tweedie models, asymptotics and scaling

Exponential dispersion models have densities (or probability mass functions) of the form

$$p(y; \theta, \phi) = c(y; \phi) \exp\{\theta y - \kappa(\theta)\}/\phi,$$

where $\theta$ is the canonical parameter, $\phi > 0$ is the scale parameter, and $\kappa$ is the cumulant function for the underlying probability measure. The expectation of $Y$ is $E(Y) = \mu = \kappa'(\theta)$, and its variance is $\text{var}(Y) = \phi V(\mu)$ where $V(\mu)$ is the unit variance function. Within this class, the Tweedie models are those with power variance function $V(\mu) = \mu^p$ for $p \in (-\infty, 0] \cup [1, \infty)$; there are no such exponential dispersion models with $p \in (0, 1)$. For more details, see Jørgensen (1997).
Table 1: Estimated values of $p$ for selected organisms

<table>
<thead>
<tr>
<th>Organism</th>
<th>Estimate</th>
<th>Organism</th>
<th>Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Bordetella pertussis</em></td>
<td>1.06</td>
<td><em>Mycobacterium tuberculosis</em></td>
<td>1.13</td>
</tr>
<tr>
<td><em>Campylobacter jejuni</em></td>
<td>1.12</td>
<td>Norovirus</td>
<td>1.63</td>
</tr>
<tr>
<td><em>Chlamydia trachomatis</em></td>
<td>1.81</td>
<td><em>Pseudomonas aeruginosa</em></td>
<td>1.65</td>
</tr>
<tr>
<td><em>Clostridium difficile</em></td>
<td>1.24</td>
<td>Rubella virus</td>
<td>1.37</td>
</tr>
<tr>
<td>Cytomegalovirus</td>
<td>1.36</td>
<td><em>Salmonella enteritidis</em></td>
<td>1.27</td>
</tr>
<tr>
<td><em>Escherichia coli</em> O157</td>
<td>1.27</td>
<td><em>Salmonella typhimurium</em></td>
<td>1.21</td>
</tr>
<tr>
<td>Epstein Barr Virus</td>
<td>1.14</td>
<td><em>Staphylococcus aureus</em></td>
<td>1.80</td>
</tr>
<tr>
<td>Herpes Simplex Virus 1</td>
<td>1.36</td>
<td><em>Streptococcus A</em></td>
<td>1.31</td>
</tr>
<tr>
<td>Herpes Simplex Virus 2</td>
<td>1.49</td>
<td><em>Streptococcus B</em></td>
<td>1.61</td>
</tr>
<tr>
<td>Influenza A virus</td>
<td>1.64</td>
<td><em>Taenia</em></td>
<td>1.26</td>
</tr>
<tr>
<td>Influenza B virus</td>
<td>1.81</td>
<td><em>Toxoplasma</em></td>
<td>1.21</td>
</tr>
<tr>
<td>Measles virus</td>
<td>1.11</td>
<td>Varicella Zoster Virus</td>
<td>1.19</td>
</tr>
</tbody>
</table>

Of particular interest to us are the Tweedie models with $p \geq 1$. The exponent $p = 1$ corresponds to (scaled) Poisson models, values $p \in (1, 2)$ to compound Poisson-gamma models, namely Poisson mixtures of gamma densities, which have an atom at 0 and are continuous above 0, while $p = 2$ corresponds to gamma models. The values $p > 2$ correspond to stable distributions, including for instance the inverse Gaussian distribution ($p = 3$). The Tweedie models with $p$ close to 1 are multimodal (Dunn and Smyth 2005).

With the sole exception of the case $p = 1$, Tweedie models are continuous above 0, and hence are not suitable for direct modelling of infectious disease counts. They are nonetheless highly relevant owing to the Tweedie convergence theorem. This states that a random variable $cY$ where $Y$ belongs to an exponential dispersion family whose variance function $V(\mu) \sim c_0 \mu^p$ as $\mu$ tends to zero (or infinity), converges in distribution to the Tweedie model with power parameter $p$ as $c$ tends to 0 (or infinity) (Jørgensen 1997, pages 148-149). Thus, a wide range of exponential...
dispersion models may be approximated by Tweedie models. It has been suggested that this theorem provides the theoretical basis underpinning the empirical evidence for Taylor’s law (Kendall 2004). This contention will be discussed briefly in Section 7.

Tweedie models are attractive also by their scaling properties: it may be shown that the Tweedie models correspond precisely to those exponential dispersion models with $V(1) = 1$ which are closed under scale transformations, that is, for which $V(c\mu) = g(c)V(\mu)$ for some function $g$, for all $c > 0$. Such a property is natural for counts of infectious diseases: if $Y$ is the number of cases per time unit, the statistical properties of $Y$ should not depend in fundamental respects on the time unit chosen. Note that this scaling invariance only applies to the power parameter $p$: unsurprisingly, the scale parameter $\phi$ is scale-dependent.

Let $\text{T}_p(\mu, \phi)$ denote the Tweedie model of power $p$ with mean $\mu$ and scale parameter $\phi$. If $Y \sim \text{T}_p(\mu, \phi)$ then $\kappa_2 = \text{var}(Y) = \phi \mu^p$. We shall also be interested
in its skewness. The third central moment of $Y$ is $\kappa_3 = p\phi^2 \mu^{2p-1}$ and its skewness is

$$\rho_3 = \frac{\kappa_3}{\kappa_2^{3/2}} = p\phi \mu^{p-1}.$$

These relationships are preserved when iid variables are aggregated. Thus, suppose that $Y_i \sim Tw_p(\mu, \phi)$, $i = 1, \ldots, n$, are independent and let $Z = Y_1 + Y_2 + \cdots + Y_n$. Then $Z \sim Tw_p(\mu_Z, \phi_Z)$ where $\mu_Z = n\mu$ and $\phi_Z = n^{1-p}\phi$. For example, the skewness of $Z$ is $p\phi_Z^{1/2} \mu_Z^{p/2-1}$. Thus, aggregating independent counts of disease in successive equal time periods will preserve the key power relationships. As noted previously, successive counts can often reasonably be considered to be independent, or only weakly dependent. The practical implication is that power relationships may validly be examined by aggregating data over several time units (in our case, weeks).

Figure 8: Scatter plot of estimated powers $p$ against lag 1 autocorrelation. The full line is a loess curve.
4 Evidence for convergence to Tweedie models

In this section we examine further evidence that Tweedie models provide an adequate description of count data for infectious diseases, in an asymptotic sense. Clearly, Tweedie models other than the Poisson (corresponding to \( p = 1 \)) can only be approximate, since they are continuous above zero. It has already been established in Section 2 that our infectious disease data exhibit the variance to mean relationship typically expected of Tweedie models. Here we consider the empirical skewness to mean relationship, to examine whether it displays the Tweedie power relationship elucidated in Section 3.

4.1 Empirical skewness to mean relationships

We shall use the following sample skewness coefficient (Joanes and Gill 1998) for a sample of size \( n \):

\[
\hat{\rho}_3 = \frac{\sqrt{n(n-1)} \left( \frac{1}{n} \sum_{i=1}^{n} (x_i - \hat{\mu})^3 \right)}{(n-2) \left\{ \frac{1}{n} \sum_{i=1}^{n} (x_i - \hat{\mu})^2 \right\}^{3/2}}.
\]

If in a period of \( n \) weeks there is just a single weekly count of 1, the others being zero, then \( \hat{\mu} = n^{-1} \) and \( \hat{\rho}_3 = n^{1/2} \). Thus, for rare organisms in which the count is 0 or 1 in every six-month period, the Tweedie skewness to variance relationship is exactly satisfied with \( p = 1 \) and \( \phi = 1 \). If Taylor’s law is deemed to apply to the weekly de-seasonalised data \( z_i \), then

\[
\log\{\rho_3(z_i)\} = \alpha_{s(i)} + \left( \frac{p}{2} - 1 \right) \log\{E(z_i)\}
\]

where \( s(i) \) is the seasonal level for week \( i \). Accordingly we fitted a normal errors generalised linear model to the empirical skewnesses, with logarithmic link and the linear predictor just described. When the data were not de-seasonalised (owing to sparseness) we just fitted a single intercept.

There is considerable scatter in the estimated skewness coefficients, making an assessment of the skewness to mean relationships more difficult than for the variance.
to mean relationships, a difficulty compounded by the curvilinear shape of the theoretical functional dependence. Nevertheless, for many organisms the asymptotic relationship implied by the Tweedie model does appear to hold, as exemplified by the six organisms displayed in Figure 9. The full set of 1724 plots (comprising the organisms for which sufficient data were available) is in the Supplementary Materials. Figure 10 shows caterpillar plots for the values of $p$ estimated from the skewness to

![Graphs of six organisms showing empirical relationship between skewness and log mean.](image)

**Figure 9:** Empirical relationship between skewness and log mean for six organisms (in 6-monthly periods): observed values (dots) and regression curves (with different intercepts for different seasons).

mean relationships, with approximate 95% confidence limits. These values typically lie between 0.5 and 2.5. The confidence intervals are quite wide, however, and are affected by the sparseness of the data for some organisms.
Figure 10: Caterpillar plot of estimated values of \( p \) and corresponding 95\% confidence limits based on log(mean) vs skewness. Left: all organisms; Right: excluding sparse organisms.

4.2 Bias correction

Sample skewness evaluated from small samples may be prone to bias. Accordingly, we repeated the analyses with intervals of 1 year, rather than 6 months, so as to increase the typical sample size from which the sample skewness is calculated. We also applied a first-order bias correction to the skewness coefficient (Pewsey 2005):

\[
\rho_3 = \sqrt{\frac{n(n-1)}{n-2}} \left\{ \beta_1 + \frac{3}{8n} \left( \beta_1 (7 + 5\beta_2) - 4\beta_3 \right) \right\} + o(n^{-1}),
\]

where

\[
\beta_k = \frac{\mu_{k+2}}{\mu^2}, \quad \mu_k = \frac{1}{n} \sum_{i=1}^{n} (x_i - \mu)^k.
\]

Figure 11 shows the corresponding caterpillar plots for \( p \) with annual groupings, with and without bias correction. Increasing the sample size narrows the confidence intervals and brings most values \( p < 1 \) closer to 1. Applying the skewness correction produces a marked increase in the estimated values of \( p \).

Note that if the sample skewness has relative bias that is constant as the mean \( \mu \) varies, then the estimate of \( p \) obtained by regressing the sample skewness against
log(\(\mu\)) will be largely unaffected, owing to the log-linear relationship between them; a constant relative bias will be reflected in the estimated intercept. Thus, what matters primarily for our purposes is the dependence of the relative bias on the mean, rather than the magnitude of this bias.

To investigate these effects in more detail, and their possible impact on our observations, we undertook simulations of the sample skewness \(\hat{\rho}_3\) from Tweedie distributions, using sample sizes \(n = 26\) and \(n = 52\), corresponding to typical six monthly and annual groupings, respectively. We also investigated the impact of the first-order bias correction to the skewness coefficient. Each value was obtained from 100 000 runs, using the R package `tweedie` (Dunn 2014). The relative biases in \(\hat{\rho}_3\) for \(\phi = 1\) are in Table 2; a broadly similar dependence of relative bias on the mean was observed with \(\phi = 0.5\) and \(\phi = 2\) (not shown). The simulation results show that skewness is generally estimated with negative bias, as suggested by Joanes and Gill (1998). As expected, the relative bias is less in absolute value with greater sample size, and (generally but not universally) with bias correction. However, for
the uncorrected skewness, the bias becomes more negative as $\mu$ increases for $p$ close to 1. Thus, for values of $p$ close to 1, the trend in the plot of skewness against $\log(\mu)$ will tend to be too steeply negative, resulting in an underestimate of $p$. For larger values of $p$, applying a bias correction increases the dependence of the relative bias on the mean (while reducing its absolute value). Thus, applying a skewness correction is likely to reduce the bias in $p$ for $p$ close to 1 (thus increasing the estimated $\hat{p}$) but will make matters worse for larger $p$. The magnitude of the bias for the estimated variance is very much smaller than for the skewness (results not shown).

Figure 12 shows a scatterplot of the values of $p$ estimated from the variance to mean relationship against those estimated from the skewness to mean relationship, using 6-monthly groupings. The Spearman correlation is 0.84. Similar results were obtained with annual groupings (correlation 0.85). The insights provided by the

Figure 12: Scatter plot of estimated slopes: variance-based vs skewness-based, with 6-monthly groupings.

simulations help explain the appearance of Figure 12, and suggest that the imperfect correspondence of the values of $p$ estimated from the variance and from the skewness
are due, at least in part, to mean-dependent bias in the estimation of the skewness, especially for low $\mu$. For less common organisms, a further contributing factor may be that the asymptotic conditions for convergence to Tweedie distributions are not met.

Overall, these results lend some support to the view that the empirical variance to mean power relationship reflects asymptotic convergence to Tweedie distributions, at least insofar as third moments are concerned. This would suggest, in turn, that Tweedie distributions may perhaps be used to approximate the distributions of infectious disease counts for the purpose of estimating quantiles.

### 4.3 Birth, death and immigration processes and the Tweedie distribution

For human infections for which a specific birth and death process, for example an SIR or SIS model, is believed to apply, a more powerful analysis may be undertaken with a model representing that process, rather than the more empirical Tweedie models described above. This is particularly true when the data exhibit substantial positive autocorrelation attributable to the transmission process, which our modelling approach ignores. However, convergence to a Tweedie distribution does not necessarily follow from observance of a power relationship between variance and mean.

We illustrate this point with a specific branching process model with immigration, in a simulation suggested to us by a referee. Applications of the model may be found in Held et al (2005), Held et al (2006) and Paul et al (2008). A simple version of the model for a count $y_i$ in week $i$ is

$$y_i \sim \text{Poisson}(\lambda y_{i-1} + \nu).$$

The model is stationary when the offspring mean $\lambda$ (which is equal to the lag 1 autocorrelation) is $0 \leq \lambda < 1$; the parameter $\nu$ is the immigration rate. We simulated
data from one such model, chosen to obtain the wide range of means that we observed in our data. We fixed the immigration rate at \( \nu = 5 \), but allowed \( \lambda \) to vary between runs, taking values whose frequencies are represented by samples from the beta density \( \lambda \sim \text{Beta}(1, 4) \). 1000 runs of 1000 values from each sample were obtained. This generates an approximate power variance to mean relationship with power parameter \( p = 1.5 \), as shown in Figure 13.

![Figure 13](image.png)

Figure 13: Left: scatterplot of log variance against log mean, with regression line of slope 1.5. Right: scatterplot of skewness against log mean, and line corresponding to Tweedie model with parameter 1.5.

However, the corresponding Tweedie model with \( p = 1.5 \) bears no relationship to the scatterplot of skewness against log mean. (As the immigration rate \( \nu \), and hence the process mean, is increased, the model converges to the Tweedie model with \( p = 1.5 \), in keeping with asymptotic theory.) Thus, it is perfectly possible for a birth and death process model to produce a power variance to mean relationship that does not fit the Tweedie framework which we have found useful for our data. Empirical investigations of variance and skewness to mean relationships can help to decide which modelling approach is most appropriate.
5 Power laws and Tweedie models for infectious disease modelling of surveillance data

In this and the next section, we focus on the descriptive statistical analysis of infectious diseases from surveillance systems. We ignore any serial dependence, which may be due to the transmission process or, as is often the case, to extraneous factors such as the surveillance system itself. These methods are particularly relevant for infections whose spread in human populations is not primarily governed by SIR and SIS models, but by factors such as food contamination and breakdown of health and safety precautions. Typically, descriptive analyses of counts of infectious diseases from surveillance systems have been based on the Poisson or negative binomial distributions, the latter resulting from gamma mixtures of Poisson distributions to produce overdispersion models in which the variance is proportional to the mean (McCullagh and Nelder 1989, page 199). Where surveillance of relatively common infections is involved, overdispersion relative to the Poisson is typical, perhaps owing to variations in reporting rates from different locations. It is convenient in such circumstances to use a quasi-Poisson model, with dispersion estimated from the data.

The preceding sections indicate that such models may not be appropriate for infectious disease counts in surveillance data, if examination of the variance to mean relationship suggests that the variance is proportional to a power $p$ of the mean with $p > 1$. Tweedie models with $p > 1$ are not discrete, and hence not directly appropriate. One option is to use discrete counterparts of the Tweedie model, such as Tweedie-Poisson mixtures or Hinde-Demétrio models (Jørgensen 1997, pages 165-170, Kokonendji et al 2004), for which the unit variance function is approximately proportional to $\mu^p$ for large $\mu$. A simpler option is to adopt a quasi-Tweedie modelling approach, in which the variance is (at least approximately) proportional to a power of the mean with exponent $p \geq 1$, but without explicitly specifying the likelihood. Quasi-Tweedie models can be fitted within the quasi-likelihood generalised
linear modelling framework, using the following quasi-deviance contribution for the count in week \(i\) (McCullagh and Nelder 1989, page 327) when \(p \neq 1, 2\):

\[
d_i = 2 \left\{ \frac{1}{1-p} \left( y_i^{2-p} - y_i \mu_i^{1-p} \right) + \frac{1}{p-2} \left( y_i^{2-p} - \mu_i^{2-p} \right) \right\}.
\]

This requires the power parameter \(p\) to be fixed. In a first stage, \(p\) is estimated using the gamma regression method described in Section 2. If the data are too sparse to allow this, or if \(\hat{p} < 1\), then take \(p = 1\) and fit a quasi-Poisson model; otherwise, set \(p = \hat{p}\).

Dunn and Smyth (2005) argue that assuming that \(p\) is fixed has little impact on inferences for the Tweedie model, owing to the fact that \(p\) is orthogonal to \(\mu\) and \(\phi\). We show that this is also true of quasi-Tweedie models, in which \(p\) is taken as fixed.

First, note that the quasi-loglikelihood contribution for a single observation \(y_i\) is

\[
ql_i = \frac{1}{\phi} \left\{ \frac{1}{1-p} \left( y_i \mu_i^{1-p} - y_i^{2-p} \right) + \frac{1}{p-2} \left( \mu_i^{2-p} - y_i^{2-p} \right) \right\}.
\]

Let \(\beta\) denote the regression parameters for the \(\mu_i\). The quasi-score function contribution for \(\beta\) is the first derivative of \(ql_i\) with respect to \(\beta\):

\[
U_i(\beta|p) = \frac{1}{\phi} (y_i - \mu_i) \mu_i^{-p} \frac{\partial \mu_i}{\partial \beta}.
\]

Suppose now that \(p\) is consistently estimated by some estimator \(\hat{p}\) (in the present context, \(\hat{p}\) is the slope of the gamma regression described in Section 2). The parameters \(\beta\) are then estimated by solving \(U(\hat{\beta}|\hat{p}) = 0\), where \(U(\beta|p) = \sum_{i=1}^{n} U_i(\beta|p)\).

Expanding in Taylor series to first order around the true values \(\beta\) and \(p\) gives

\[
0 = U(\hat{\beta}|\hat{p}) = U(\beta|p) - I(\beta|p)(\hat{\beta} - \beta) + \frac{\partial U(\beta|p)}{\partial p}(\hat{p} - p) + O_p(1),
\]

where \(I(\beta|p) = E\{ - \partial^2 U(\beta|p)/\partial \beta^2 \}\). Rearranging, we obtain

\[
\hat{\beta} = \beta + I(\beta|p)^{-1} U(\beta|p) + I(\beta|p)^{-1} \frac{\partial U(\beta|p)}{\partial p}(\hat{p} - p) + O_p(n^{-1}).
\]

As \(E\{U_i(\beta|p)\} = 0\), the second term on the right of this equation is \(O_p(n^{-1/2})\). Since

\[
\frac{\partial U_i(\beta|p)}{\partial p} = -\frac{1}{\phi} (y_i - \mu_i) \mu_i^{-p} \log(\mu_i) \frac{\partial \mu_i}{\partial \beta},
\]

we have

\[
\frac{\partial U_i(\beta|p)}{\partial p} = -\frac{1}{\phi} (y_i - \mu_i) \mu_i^{-p} \log(\mu_i) \frac{\partial \mu_i}{\partial \beta}.
\]

As \(E\{U_i(\beta|p)\} = 0\), the second term on the right of this equation is \(O_p(n^{-1/2})\). Since
it follows that $E\{\partial U_i(\beta|p)/\partial p\} = 0$. Hence $\partial U(\beta|p)/\partial p$ is $O_p(n^{1/2})$. Now $(\hat{p} - p)$ is $o_p(1)$ by assumption, so the third term in the expansion of $\hat{\beta}$ is $o_p(n^{-1/2})$. In other words, estimation of $p$ only influences $\hat{\beta}$ by a factor $o_p(n^{-1/2})$, rather than $O_p(n^{-1/2})$, and thus has relatively little bearing on the estimation of $\beta$.

Tweedie models are likely to be useful in other ways relevant to infectious disease surveillance data. For example, the evaluation of statistical techniques for infectious disease surveillance is often based on simulations using Poisson or negative binomial distributions. A more realistic option, especially for more common infections, may be to simulate data from Tweedie distributions, rounding the results to the nearest integer to obtain counts, or from their discrete counterparts.

Finally, a key advantage of the quasi-Tweedie approach is that the true but unknown distribution, though not Tweedie, can nevertheless be approximated by a Tweedie density, by virtue of the Tweedie convergence theorem, at least when the means $\mu$ are large, which is the setting in which the approach is likely to be most fruitful (if counts are low, the appropriate limit when $\mu$ tends to zero is typically Poisson). The evaluation of Tweedie densities is discussed by Dunn and Smyth (2005, 2008). This circumvents a disadvantage of the quasi-likelihood approach: no distribution is presumed, which makes it difficult to obtain reliable prediction intervals for individual observations. This convergence property is likely to be particularly useful in regression-based outbreak detection, where the upper threshold estimated under the null hypothesis that no outbreak is occurring is obtained using a prediction interval. For example, the outbreak detection system in use since the early 1990s at Public Health England, and applied to the LabBase data described above, uses a quasi-Poisson regression method, with the upper prediction limits based on a normal approximation or the quantiles of the negative binomial distribution (Farrington et al 1996, Noufaily et al 2013). These limits are compared to those obtained with the asymptotic Tweedie distribution in the next section.
6 Application to infectious disease surveillance data

We compared the upper prediction limits obtained for several organisms using (a) normal approximations, (b) negative binomial quantiles, and (c) Tweedie approximations. The organisms were selected to obtain a range of values of the exponent $p$, which was estimated using the gamma model of Section 2 applied to adjacent six-monthly periods. We used upper 99.5% prediction limits: organisms with counts above these thresholds are deemed aberrant and undergo further investigations. The algorithm is designed to detect the following types of aberrations: sudden changes in level, sudden changes in trend, and sudden changes in the phase or amplitude of seasonal fluctuations.

For methods (a) and (b) we applied the algorithm of Noufaily et al (2013). In brief, to obtain the upper prediction limit at week $t$ a quasi-Poisson generalised linear model (with log link, linear time effect and factorial seasonal effect) is applied to the observed counts in recent years, but excluding the most recent six months, lest these counts are contaminated by a current outbreak. This gives estimates of the expected value $\mu_t$ at time $t$ and of the dispersion $\phi$, from which the limits (a) and (b) are derived from the 0.995-quantiles of the normal (after a suitable transformation) and negative binomial distributions. See Noufaily et al (2013) for details. The algorithm is implemented in the R package surveillance (Höhle 2007).

For method (c), we replaced the quasi-Poisson model with a quasi-Tweedie generalised linear model (with pre-estimated $p$) to obtain the expected value $\mu_t$ at $t$ and the dispersion $\phi$ (which is different from that obtained for methods (a) and (b)). The quasi-Tweedie model was fitted using the function glm within R (R development Core Team 2014); suitable code is provided in the Appendix. The upper prediction limit was then obtained as the 0.995 quantile of the Tweedie $\text{T}_p(\mu_t, \phi)$ density, using the function qtweedie within R package Tweedie (Dunn 2014).

For all methods, we switched off the reweighting mechanism in the standard version of the outbreak detection algorithm. This mechanism downweights high
baseline values, in order to reduce the impact of past outbreaks. It was switched off because it would need to be modified for the quasi-Tweedie model, thus complicating the comparison of the three methods.

For all organisms, the expected values obtained using the quasi-Poisson and quasi-Tweedie models are virtually identical. The upper prediction limits obtained using the normal approximation for the quasi-Poisson and negative binomial quantiles were also very close.

For organisms with $p$ close to 1, the upper prediction limits based on Tweedie quantiles were very similar to those obtained using the other two methods. Figure 14 shows the results for Acinetobacter baumanii, with $p = 1.05$, and Varicella zoster virus, with $p = 1.19$. Figure 15 shows the results for two organisms with intermediate values of $p$: Cytomegalovirus ($p = 1.36$) and Chlamydia trachomatis ($p = 1.82$). Here, the Tweedie upper prediction limits are generally higher than those for the quasi-Poisson, the difference increasing with the mean frequency and with $p$. For Cytomegalovirus, note that the system detects the abrupt change in level around

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**Figure 14:** Weekly count (dots), expected values and upper thresholds for quasi-Tweedie (full lines) and quasi-Poisson (dotted lines). Left: Acinetobacter baumanii; right: Varicella zoster virus.
week 875, after which performance is degraded until the algorithm adjusts to the
new level. Figure 16 shows the results for two organisms with high values of $p$:

![Graph showing the results for two organisms with high values of $p$.]

**Figure 15:** Weekly count (dots), expected values and upper thresholds for quasi-
Tweedie (full lines) and quasi-Poisson (dotted lines). Left: Cytomegalovirus; right:
Chlamydia trachomatis.

*Bacteroides fragilis* ($p = 2.14$) and *Cryptosporidium* sp ($p = 2.49$). Here, the
Tweedie upper prediction limits can be very different from those for the quasi-
Poisson. Typically they are higher, except in the troughs of the seasonal cycles. The
practical consequence of these observations is that, when $p$ is high, the false positive
rate (that is, the proportion of non-outbreak weeks in which organism counts are
flagged as aberrant) will tend to be too large using the standard outbreak detection
algorithm, if as suggested in Section 4, the distribution of the organism of interest is
well approximated by a Tweedie distribution. However, organisms with $p$ less than
about 1.4 should not be unduly affected unless the mean frequency is very high.

The Tweedie thresholds in Figures 14 to 16 are plug-in estimates, obtained as-
suming that $p$, in particular, is known. Each is the 0.995-quantile of the Tweedie
density with parameters $\tilde{p}$, $\hat{\mu}$ and $\hat{\phi}$. We investigated the impact of estimating $p$
on these thresholds by a Monte Carlo method based on the approach of Lawless
Figure 16: Weekly count (dots), expected values and upper thresholds for quasi-Tweedie (full lines) and quasi-Poisson (dotted lines). Left: Bacteroides fragilis; right: Cryptosporidium sp.

and Fredette (2005). First, 1000 bootstrap estimates $p_b$, $b = 1, ..., 1000$ of the power parameter $p$ were obtained by bootstrapping the gamma regression described in Section 2. For each value of $p_b$ the model was refitted, yielding estimates $\mu_b$ and $\phi_b$. Then 1000 random variables $y_{br}$, $r = 1, ..., 1000$ were generated from the Tweedie distribution with parameters $\tilde{p}$, $\tilde{\mu}$ and $\tilde{\phi}$ (the original estimated values, as used in the plug-in method) and the values $u_{br} = F(y_{br}; p_b, \mu_b, \phi_b)$ were calculated, where $F$ is the Tweedie cumulative distribution function. The 0.995-quantile $v$ of the $u_{br}$ was obtained, and the adjusted threshold was taken as $Q(v; \tilde{p}, \tilde{\mu}, \tilde{\phi})$ where $Q$ is the Tweedie quantile function. All calculations were undertaken in R using the package *tweedie* (Dunn 2014). This procedure is time-consuming, because a large simulation sample is required to estimate high quantiles accurately. We applied it to the most recent week for each of the six organisms in in Figures 14 to 16. The relative difference between the thresholds obtained using the plug-in method and the method allowing for estimation of $p$ was always less than 0.8% (on average it was 0.2%). Since estimating $p$ has so little impact on the thresholds, we recommend the
much simpler and quicker plug-in method.

7 Discussion

Analysis of extensive surveillance data on a large number of distinct organisms provides strong evidence that the variance to mean relationship may be described by power laws in virtually all instances, the power parameter $p$ typically lying between 1 and 2, with some exceptions. Further investigation of skewness to mean relationships provides some evidence to suggest that they conform asymptotically to Tweedie distributions, as predicted by the Tweedie convergence theorem. The strength of evidence is somewhat reduced by the problem of mean-dependent bias in estimating sample skewness. Nevertheless, we tentatively conclude that Tweedie distributions can be used to approximate distributions of counts.

From an epidemiological standpoint, these power relationships are of intrinsic interest, in that they potentially provide insight into infectious disease dynamics. It would be interesting to know whether the power relationships we have observed are also present in infectious disease surveillance data from other countries. Contrasting data sets obtained in different contexts may help to clarify whether the observed patterns stem from underlying mechanisms, the specificities of different reporting systems, or purely stochastic effects.

Our focus has been entirely on exploiting variance to mean and skewness to mean relationships for descriptive statistical modelling purposes. We have not sought to explore the underlying mechanisms which produce such power relationships. It is known that birth and death processes produce such relationships (Anderson et al 1982, Keeling 2000), while Faddy (1997) explicitly constructed a family of statistical models for count data based on birth processes, which exhibits a range of variance to mean behaviours. We have instead focused on the suggestion of Kendal (2004) and Kendal and Jørgensen (2011) that the Tweedie convergence theorem provides a purely mathematical explanation for such observed patterns, in terms of asymptotic
limiting behaviour. Our data appear to provide some support for this contention.

From a statistical standpoint, our findings potentially provide a more accurate
descriptive representation of the variability associated with surveillance data on
infectious diseases. We advocate the explicit use of observed variance and skew-
ness to mean relationships, and the Tweedie convergence theorem, for four related
purposes in infectious disease modelling. In the first instance, an examination of
these relationships (at least if data are plentiful), after removing seasonal variation,
can provide useful information and ought to become a routine element of the de-
scriptive statistical investigation of infectious disease surveillance data. This should
also inform model choice. Second, quasi-Tweedie generalised linear models (the
term quasi-Tweedie solely denoting a power variance to mean relationship) should
perhaps be more commonly used for descriptive modelling purposes. Third, model
evaluation using simulations based on Poisson and negative binomial models is likely
to underestimate the extent of random variation and skewness in infectious disease
counts; simulations using discretised Tweedie distributions or their discrete coun-
terparts might be more realistic. And finally, the calculation of thresholds and
prediction intervals, notably in outbreak detection systems, may be more accurate
if based on asymptotic Tweedie distributions, at least when \( p \) is appreciably greater
than 1.

For infections with \( p \) close to 1, Poisson, quasi-Poisson or negative binomial
models are likely to be adequate. However, for organisms with higher values of
\( p \), we have shown in Section 6 that there are benefits in adopting the methods
we propose. While these may be impractical for the purposes of large multiple
routine surveillance systems, more focused organism-specific surveillance systems
may benefit from making use of the empirical relationships suggested by the data.
It would also be useful to explore suitable discrete models, rather than rely on
quasi-Tweedie methods.

We have focused entirely on marginal variance and skewness to mean relation-
ships, ignoring any serial dependence in the data. In our surveillance data, such
dependencies were generally rather weak, and for this reason we feel our approach is justified for many infections. Many of the instances of significant positive serial dependence are most likely unrelated to the transmission mechanism, resulting for example from residual periodicity. In infectious diseases, seasonality is often subject to variation in amplitude and period. The surveillance system may also induce regular spikes and troughs. These effects are not wholly removed using fixed seasonal cycles: more flexible methods of seasonal adjustment are required. Beyond such spurious effects, the empirical finding that autocorrelations are often quite close to zero (or negative) can be attributed to two main factors: the modes of transmission, and the nature of the surveillance data. First, very many organisms are not primarily transmitted directly from person to person by close contact. For example, many are primarily food, water or airborne from zoonotic or environmental reservoirs, as for *Campylobacter*, *Escherichia coli*, *Listeria monocytogenes*, *Legionella*, and most salmonellas. The temporal autocorrelation for such infections in humans is weak, cases arising primarily sporadically, or occasionally in point and common source outbreaks with perhaps only a limited element of subsequent person-to-person transmission. However, we also observe low autocorrelations for organisms that are directly transmitted by close contact, such as Varicella zoster virus, *Bordetella pertussis* and Measles virus. The reason is most likely that our data include only cases of disease of sufficient clinical severity to warrant a biological specimen being collected for identification: these cases thus represent only a small proportion of actual infections, and selection effects may erase the underlying serial dependence.

There were, however, some exceptions, notably Influenza A and B, for which the autocorrelation is significantly positive in line with known transmission dynamics. For these infections, ignoring the transmission mechanism, while generally producing consistent estimates, may be inefficient. Furthermore, our finding that serial dependence is generally weak may not apply to other data on infectious diseases, obtained from more complete reporting systems. If, in an investigation focused on a particular infection, substantial non artefactual serial correlation is observed which
is related to the mode of transmission, it ought to be incorporated in statistical models. For example the models of Paul et al (2008) and Finkenst dt and Grenfell (2000) may perhaps be adapted, if required, to encompass greater flexibility in the power variance to mean and skewness to mean relationships they generate. It may also be beneficial in some circumstances to incorporate further complexities, such as cross-correlations between distinct infections of specific interest.

**Acknowledgments**

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**References**


Kendal W.S. (2004) Taylor’s ecological power law as a consequence of scale in-


The following R code may be used for fitting the quasi-Tweedie model, for a specified numerical value of \( p \). First, set up the following list to specify the variance function.

```r
varp<-list(
  varfun=function(mu){mu^p},
  validmu=function(mu){all(mu>0)},
  dev.resids=function(y,mu,wt){
    2*wt*((1/(1-p))*(y^(2-p)-y*mu^(1-p))+(1/(p-2))*(y^(2-p)-mu^(2-p)))},
  initialize=expression(mustart<-ifelse(y==0,0.001,y)
  )
)
```

The model is then fitted with a call to R function `glm`, using

```r
family=quasi(link="log",variance=varp).
```
Table 2: Relative bias for sample skewness (uncorrected and bias-corrected) for different values of $\mu$ and $p$ for sample sizes $n = 26$ and $n = 52$, with $\phi = 1$

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<td>bias corrected</td>
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