Comparison of Bayesian and frequentist methods for prevalence estimation under misclassification

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Comparison of Bayesian and frequentist methods for prevalence estimation under misclassification

Matthias Flor, Michael Weiß, Thomas Selhorst, Christine Müller-Graf and Matthias Greiner

Abstract

Background: Various methods exist for statistical inference about a prevalence that consider misclassifications due to an imperfect diagnostic test. However, traditional methods are known to suffer from truncation of the prevalence estimate and the confidence intervals constructed around the point estimate, as well as from under-performance of the confidence intervals' coverage.

Methods: In this study, we used simulated data sets to validate a Bayesian prevalence estimation method and compare its performance to frequentist methods, i.e. the Rogan-Gladen estimate for prevalence, RGE, in combination with several methods of confidence interval construction. Our performance measures are (i) error distribution of the point estimate against the simulated true prevalence and (ii) coverage and length of the confidence interval, or credible interval in the case of the Bayesian method.

Results: Across all data sets, the Bayesian point estimate and the RGE produced similar error distributions with slight advantages of the former over the latter. In addition, the Bayesian estimate did not suffer from the RGE’s truncation problem at zero or unity. With respect to coverage performance of the confidence and credible intervals, all of the traditional frequentist methods exhibited strong under-coverage, whereas the Bayesian credible interval as well as a newly developed frequentist method by Lang and Reiczigel performed as desired, with the Bayesian method having a very slight advantage in terms of interval length.

Conclusion: The Bayesian prevalence estimation method should be preferred over traditional frequentist methods. An acceptable alternative is to combine the Rogan-Gladen point estimate with the Lang-Reiczigel confidence interval.

Keywords: prevalence estimation; imperfect diagnostic test; misclassification; Bayesian prevalence estimate; Rogan-Gladen estimate; diagnostic sensitivity; diagnostic specificity

Background

Prevalence estimation is fundamental to a lot of epidemiological studies. However, to obtain an accurate estimation of prevalence, misclassification and measurement errors should be considered as part of bias analysis in epidemiological research [1]. Frequentist and Bayesian methods for bias adjustment of epidemiological risk estimates have been reviewed in Keogh et al. [2] and Shaw et al. [3]. Estimation of prevalence is always based on the application of a diagnostic test to classify samples with respect to the binary trait under investigation. Major sources of uncertainty of prevalence estimates are related to the study design and sampling issues and are usually described using the concepts of bias and precision (statistical parameter...
uncertainty of the estimate). A typical source of (information) bias is diagnostic
misclassification due to imperfect sensitivity and specificity. Rogan and Gladen have
derived an estimator for prevalence with adjustment for diagnostic misclassification
[4]. This approach requires that unbiased estimates of the diagnostic accuracy be
available for the given application [reviewed in 5]. Here, we present a comparison of
Bayesian and frequentist methods for prevalence estimation taking into account all
relevant uncertainties associated with the study- and meta-data, e.g. the diagnostic
test performance.

In the following, $D^+/−$ denotes the individual disease status and $T^+/−$ the result of
a diagnostic test applied to an individual. The test sensitivity is then defined as the
probability that a $D^+$ individual tests positive, $\tilde{Se} = \Pr(T^+|D^+)$, and its specificity
is defined as the probability of a $D^−$ individual testing negative, $\tilde{Sp} = \Pr(T^-|D^-)$.
The disease prevalence, $\tilde{\pi}$, is the proportion of diseased individuals in a population
but can also be thought of as the probability that a randomly sampled individual
is diseased. The probability that a test applied to a random individual from such a
population yields a positive result is called the apparent prevalence, $\tilde{AP}$.

$$\tilde{AP} = \Pr(T^+) = \Pr(T^+|D^+) \Pr(D^+) + \Pr(T^+|D^-) \Pr(D^-)$$
\[= \tilde{Se} \tilde{\pi} + (1 - \tilde{Sp})(1 - \tilde{\pi}).\] (1)

By replacing the true population quantities (denoted by the tilde) in Eq. (1) with
estimates of these quantities (denoted by a circumflex accent or "hat") that are
subject to sampling variability and subsequently solving for the prevalence, the
Rogan-Gladen point estimate [4] is derived:

$$\text{RGE} = \tilde{\pi} = \frac{\tilde{AP} - (1 - \tilde{Sp})}{\tilde{Se} - (1 - \tilde{Sp})}.$$ (2)

In practice, the Rogan-Gladen estimate is truncated to the interval $[0, 1]$ in order
to guarantee a proper proportion [6],

$$[\text{RGE}]_0^1 = \min(\max(\text{RGE}, 0), 1).$$ (3)

The potential need for truncation stems from the fact that for a prevalence estimate
to be a valid proportion, $0 \leq \text{RGE} \leq 1$, three conditions must be satisfied [4, 6]
which may not always be the case in practice:

$$1 - \tilde{Sp} < \tilde{Se}$$ (4a)
$$1 - \tilde{Sp} \leq \tilde{AP}$$ (4b)
$$\tilde{AP} \leq \tilde{Sp}$$ (4c)

Condition (4a), $1 - \tilde{Sp} < \tilde{Se}$, should hold for any diagnostic test to meet the basic
requirement that a disease be detected better than by chance alone [4]. Failing to
satisfy the second condition \((4b)\), \(1 - \hat{S}_p \leq \hat{A}P\), results in a negative estimate for the true prevalence, \(RGE < 0\), whereas violating the third requirement \((4c)\), \(\hat{A}P \leq \hat{S}_e\), yields an estimate larger than unity, \(RGE > 1\) [6]. Note that it is possible for degenerate cases to occur where the first condition is violated but the estimate still yields a value between zero and unity.

In order to quantify the precision of an estimate, it is good practice to accompany the point estimate with a 95% confidence interval (CI). In this study, we included CI construction methods by Clopper and Pearson [7], Sterne [8], Blaker2000 [9], Rogan and Gladen [4], and by Lang and Reiczigel [10].

In a Bayesian framework, credible parameter values are described by probability distributions. In order to provide comparability with the frequentist approach, the mean of a distribution may then be used as a point estimate, and the Bayesian analogue to the CI is the credible interval (CrI) that marks a range of values that combine a specified percentage of the distribution’s probability mass. More specifically, we consider the 95% highest density interval (HDI) which is the shortest of all possible 95% CrIs.

The aim of this study is to validate a Bayesian model for prevalence estimation with an imperfect diagnostic test and to compare its performance with traditionally used methods. We use simulated data sets based on simulated true parameter values. Our performance measures are (i) estimation error of point estimates and (ii) confidence interval coverage and length. Of special interest is the performance of the Bayesian point estimates in situations when the Rogan-Gladen estimate must be truncated because it otherwise yields a negative value or a value larger than unity.

Methods

The present validation study consists of four steps: (1) Simulation of parameter sets, including true prevalence values, (2) simulation of data sets, (3) estimation of prevalence for each of the generated data sets, and finally the actual (4) validation of the estimates against the simulated true values. Fig. 1 gives an overview of these steps, and we describe each step in detail in the following sections.

All computations were performed on a workstation running Ubuntu Linux 18.04.3 LTS, using the statistical software R version 3.6.2 [11] and the MCMC software JAGS version 4.3.0 [12]. The R code used as well as all data sets generated and analysed during this study are available via a Zenodo snapshot, DOI: 10.5281/zenodo.3631123.

Parameter set simulation

We distinguish between statistical parameters and design parameters. The statistical parameters are the true prevalence in a population under study, and the true characteristics of the test that is used to diagnose individuals in a sample drawn from such a population, i.e. true sensitivity and true specificity. The statistical parameters may be estimated by a statistical model. Design parameters describe the circumstances under which data is generated, and are typically chosen by scientists conducting a study or by users applying a diagnostic test. In our case there are three design parameters, all of them sample sizes: The size of the sample that a
diagnostic test is applied to, the sample size of a study that has been conducted\(^1\) to validate the sensitivity of the diagnostic test, and likewise the sample size of a validation study of the test’s specificity.

For the purposes of this study we generated 1,000 parameter set realisations\(^4\) consisting of the three statistical and the three design parameters. Simulated true\(^5\) values (indicated by a tilde, \(\tilde{\cdot}\), in this article) for the statistical parameters, prevalence\(^6\), sensitivity, and specificity, were randomly drawn from the ranges specified in Tab. 1. True prevalence (\(\tilde{\pi}\)) can take on any value between 0% and 100%, and true sensitivity (\(\tilde{Se}\)) and specificity (\(\tilde{Sp}\)) values are assumed to be at least 60% up to a maximum of 100%.

The three design parameters are denoted by \(n_{Se}\), \(n_{Sp}\), and \(n\). These enable us to simulate the situations where (i) studies are performed to validate the sensitivity and the specificity of a diagnostic test (with sample sizes \(n_{Se}\) and \(n_{Sp}\), respectively), (ii) the actual application of the test takes place (namely, the number of individuals that are tested, \(n\)). The three sample sizes were randomly drawn as specified in Tab. 1, and exemplary parameter set realisations are shown in Tab. 2. No correlations were assumed among these parameters.

Data set simulation

For each simulated set of parameters we generated 1,000 replications of data sets\(^20\) (replications in the sense that the underlying true values for prevalence, sensitivity, and specificity, are the same for such a set of 1,000 data). Data from diagnostic validation studies were simulated by randomly drawing numbers of true positives, \(x_{Se}\), and numbers of true negatives, \(x_{Sp}\), from the binomial distributions\(^1\) given in Equations (5). The number of positive test results, \(x\), when applying the test in a population with true prevalence \(\tilde{\pi}\), was simulated analogously:

\[
\begin{align*}
x_{Se} & \sim B \left( n_{Se}, \tilde{Se} \right) \\
x_{Sp} & \sim B \left( n_{Sp}, \tilde{Sp} \right) \\
x & \sim B \left( n, \tilde{AP} \right),
\end{align*}
\]

where \(\tilde{AP} = \tilde{Se} \tilde{\pi} + (1 - \tilde{Sp}) (1 - \tilde{\pi})\) may be called the ‘true apparent prevalence’. Thus for each data set, the maximum likelihood estimators for sensitivity, specificity, and apparent prevalence would be

\[
\begin{align*}
\tilde{Se} & = \frac{x_{Se}}{n_{Se}} \\
\tilde{Sp} & = \frac{x_{Sp}}{n_{Sp}} \\
\tilde{AP} & = \frac{x}{n}
\end{align*}
\]

In our parameter sets, the true values, \(\tilde{Se}\), \(\tilde{Sp}\), and \(\tilde{AP}\), would (by definition) always meet all of the three conditions (4). However, due to the random sampling procedure\(^3\) the binomial distribution \(B(n, p)\) is the discrete probability distribution of the number of successes in a series of independent trials where parameters \(n\) and \(p\) are the number of trials and the success probability for each trial, respectively.
described above for simulating (and mimicking random processes occurring in the real world) validation studies and application of a diagnostic test, the maximum likelihood estimators, $\hat{Se}$, $\hat{Sp}$, and $\hat{AP}$, will sometimes lead to violation of one or more of the conditions. From the total of 1,000 (parameter set realizations) \times 1,000 (data set replications) = 1,000,000 data sets, we excluded 21 because they failed to meet the first condition, $1 - \hat{Sp} < \hat{Se}$, arguing that a test that has been validated in this manner would not be applied in practice. The data simulation process thus generated a total of 999,979 data sets. Tab. 3 shows exemplary data sets from the generation process described in this section.

### Estimation of prevalence

In the next step, we estimated prevalence adjusted for misclassification for each of the simulated data sets using frequentist and Bayesian methods.

#### Frequentist estimation

For each data set, we calculated the Rogan-Gladen point estimate for prevalence according to Eq. (3), using the maximum likelihood estimates for the sensitivity and specificity values as well as the apparent prevalence. In order to construct a confidence interval (CI) for the Rogan-Gladen estimate, several different methods are used in the literature. Here, we consider the methods proposed by Clopper and Pearson [7], Sterne [8], Blaker [9], and Rogan and Gladen [4]. All of these methods except for the one by Rogan and Gladen assume that sensitivity and specificity are known, and the MLE values were used in this case. Rogan and Gladen [4] do allow for sensitivity and specificity to be estimated from validation studies (thus taking into account uncertainty around the diagnostic test properties) but due to the normal approximation that they suggest, their CI is known to perform poorly [5]. For a more comprehensive description of how these intervals are constructed, we refer to the study by Reiczigel et al. [13]. Indeed, the code we used to calculate these CI’s is based on the code provided in Reiczigel et al. [13] and in Lang and Reiczigel [10].

#### Bayesian estimation

In the Bayesian prevalence estimation model, prior knowledge or belief about the true prevalence as well as the sensitivity and specificity of the diagnostic test is expressed in terms of probability distributions. Prior knowledge in the present context derives from validation studies of the diagnostic test properties but in other situations may also stem from expert opinion. Through data from the application of the test to diagnose samples from a population the model updates the probability distributions which afterwards describe the posterior knowledge about the true prevalence as well as about the sensitivity and specificity of the test. Posterior probability distributione must often be numerically approximated by random sampling algorithms referred to as Markov Chain Monte Carlo (MCMC) methods. To ensure that values are sampled from a stationary distribution, so-called convergence diagnostics are used.

Our prevalence estimation model was implemented using the JAGS software [12], a Gibbs sampler for MCMC simulations that uses a dialect of the BUGS modeling...
The model, expressed in the BUGS language, is basically a description of the process that provides parameters for data generation:

```
model {
  pi ~ dbeta(1, 1) # prevalence
  se ~ dbeta(x_se + 1, n_se - x_se + 1) # sensitivity
  sp ~ dbeta(x_sp + 1, n_sp - x_sp + 1) # specificity
  ap <- pi * se + (1 - pi) * (1 - sp) # apparent prevalence
  x ~ dbin(ap, n) # number of positive tests
}
```

By definition, the variables prevalence, sensitivity, and specificity are all proportions and thus appropriately modeled as Beta probability density distributions \(\text{dbeta}\) in the BUGS language; the tilde symbol, \(\sim\), denotes drawing a random variable from a distribution whereas the left arrow, \(<-\), implies a deterministic relationship. Importantly, this guarantees that all probability mass for each of these variables is restricted to the domain \([0, 1]\) thus eliminating the problem of prevalence estimates that are negative or larger than unity, and the same conveniently holds for limits of credible intervals. For the prevalence we use the uniform \(\text{Beta}(1, 1)\) as a non-informative prior distribution \([2]\). In contrast, prior information on the sensitivity and specificity of the diagnostic test is available from their respective validation studies. E.g., if the sensitivity of the test has been validated against a gold standard in a study of size \(n_{\text{Se}}\), and \(x_{\text{Se}}\) of the truly positive samples yielded a positive test result, then our knowledge about the true sensitivity can be expressed as \(\text{Beta}(x_{\text{Se}} + 1, n_{\text{Se}} - x_{\text{Se}} + 1)\). As with all MCMC simulation techniques, using the JAGS software requires that convergence of the Markov chains to a stationary distribution be checked. In order to realize the validation study presented here, i.e. applying our model to approximately 1,000,000 data sets, we made use of the R package \texttt{runjags} \([15]\) which provides an interface to JAGS with capabilities for automated calculation of convergence \([16]\) and of appropriate sample length \([17]\) via an \texttt{autorun} function. All chains were initialized explicitly to ensure that convergence can be evaluated appropriately. For all simulations, we used three chains, 1,000 adaptive iterations, a burn-in length of 4,000, and a minimal sample length of 20,000. This yielded an effective sample size of approximately 10,000 on which the prevalence estimates are based. When the Bayesian estimation model was applied to each of the data sets, the Gelman and Rubin’s statistic as used by the \texttt{autorun} function indicated convergence in all cases. The mean of the posterior distribution –approximated by the three combined chains– is a minimum mean square error estimator and provides a Bayesian point estimate for the true prevalence. The broader the probability distribution, the less certain our knowledge about the true prevalence is. A 95% credible interval (CrI) denotes a range of prevalence estimates that together account for 95% of the probability mass of the distribution. The 95% highest density interval (HDI) is the shortest 95% CrI, such that any value outside the HDI is considered less plausible. More accurately, the \(\text{Beta}(1, 1)\) prior is weakly informative, as it implicitly considers all possible values to be equally likely. However, for the sake of simplicity we will use the term non-informative throughout this article.
than the values inside of it. Therefore, the 95% HDI constitutes a natural measure of uncertainty for the estimate. Note that the Bayesian model also provides updated knowledge on the sensitivity and the specificity of the test, but in this study we focus on the performance of the prevalence estimates.

Validation

In the final validation step of our study, we calculated two performance metrics and their distributions in order to compare the various estimation methods: (i) the estimation error of the point estimates and (ii) the coverage of the confidence or credible intervals.

For each data set, the estimation error of the two point estimates (Rogan-Gladen estimate, Bayesian mean) was calculated as the difference to the simulated true value, estimation error = \( \hat{\pi} - \tilde{\pi} \). Note that a consistent estimator should exhibit a symmetric distribution (across all data sets) of estimation errors that is centered at zero. To investigate which estimator performs better, we carried out a regression of the estimation errors of all Bayesian estimates on the corresponding Rogan-Gladen estimate's estimation errors. Because obviously there are errors in both estimators, we used a Deming regression and assumed equal error variances for the two distributions. This means that the regression minimizes the sum of squared orthogonal distances to the regression line.

For each parameter set, coverage of the various frequentist 95% CI’s [4, 7–10] and of the Bayesian 95% HDI was computed as the percentage of the 1,000 replication data sets for which the true prevalence value was contained in the respective interval. Note that this implies assessing the frequentist behaviour of the Bayesian HDI. A well-behaved 95% confidence (or credible) interval is expected to have a symmetric distribution (across all parameter sets) of coverage values that is centered at its nominal value of 95%. Additionally, we calculated the lengths of the CI’s for all data sets. If one compares two well-behaved CI methods then the one producing shorter intervals is the better one.

Results

In general, the Rogan-Gladen point estimate as well as the Bayesian estimator (i.e. the MCMC mean) yield a consistent point estimate for the prevalence. More specifically, estimation error distributions of the two methods look very similar when compared across the entirety of all simulated data sets (Fig. 2a, top left). This property of being consistent also holds true for data subsets in which the Rogan-Gladen estimate yields a value between zero and unity (Fig. 2a, top right). However, for subsets of the data in which the non-truncated Rogan-Gladen estimate is negative (Fig. 2a, bottom left) or larger than unity (Fig. 2a, bottom right), the Bayesian estimator is more consistent and exhibits a distribution approximately symmetrical to an estimation error of zero. Its asymmetrical error distribution indicates a deficiency of the Rogan-Gladen estimator.

In order to compare the two estimators in more detail, we plotted their estimation error values against one another for all of the 999,979 data sets (see Fig. 2b). To deal with overplotting, we used hexagonal binning; the darker the shade of a hexbin the more data sets were registered within that bin. In general, the hexbins...
are distributed along the diagonal showing that the estimation errors of the two es-
1timators behave similarly. The correlation coefficient for the two estimation errors is
2r = 0.921, and the orthogonal regression has a slope of 0.939. The confidence inter-
3val for the slope estimate, (0.938, 0.941), does not include the diagonal (slope 1)
4thus revealing that overall the Bayesian estimator performs slightly better than the
5Rogan-Gladen estimate. A linear regression analysis for both estimators suggests
6that the residual estimation error after adjusting prevalence estimates for diagnostic
7sensitivity and specificity does depend on the application scenario as represented by
8our design variables for sample sizes, true prevalence, true sensitivity and true speci-
9ficity, but that the effects are only very minor for the whole model, the individual
10parameters, as well as their first-order interactions (see Additional file 1).

11The coverage of the Bayesian HDI credible intervals and of the Lang-Reiczigel CI
12is in good accordance with their nominal value of 95%, as can be seen in Fig. 3. This
13is in stark contrast to traditional confidence intervals which exhibit coverage much
14lower than 95%. In fact, the nominal 95% are not even included in the interquartile
15ranges of coverage values for these traditional methods.

16Coverage is one performance metric for a confidence (or credible) interval, length
17of the interval is the other. Fig. 4a shows that traditional CI’s (Clopper-Pearson,
18Blaker, Sterne, Wald-Rogan-Gladen) are much shorter than the Lang-Reiczigel CI
19and the Bayesian HDI, the two of which have very similar lengths. This pattern is
20even more pronounced for the data subsets in which the Rogan-Gladen estimate
21has to be truncated at zero (case 2; Fig. 4a bottom left) or at unity (case 3; Fig. 4a
22bottom right).

23Fig. 4b shows how the lengths of the Bayesian HDI and of the best performing (in
24terms of coverage) conventional confidence interval, the Lang-Reiczigel CI, relate to
25one another. The correlation coefficient is, r = 0.975. The dashed black line shows
26an orthogonal Deming regression, again under the assumption that both interval
27lengths have equal variance. The slope of the regression is 0.955 (0.954, 0.957) show-
28ing that the Bayesian HDI tends to be slightly narrower than the Lang-Reiczigel
29CI.

Discussion

In this simulation study we evaluated the validity of a Bayesian method to estimate
true prevalence based on the results of imperfect diagnostic tests. The Bayesian
point estimate of the true prevalence performed slightly better than the conventional
Rogan-Gladen estimate. Our study demonstrated that the traditional confidence
intervals, Clopper-Pearson, Blaker, Sterne, and Wald-Rogan-Gladen exhibit consi-
derable under-coverage and should be considered unfit for prevalence estimation
under misclassification. In contrast, both the Lang-Reiczigel CI and the Bayesian
HDI can be considered fit for use as they provide coverage close to the level of 95%.

The fact that the traditional methods generally provide much narrower CI’s than
the Lang-Reiczigel CI or the Bayesian HDI does obviously not ameliorate this short-
coming but rather gives an impression of certainty that is unwarranted. It could be
said that the Lang-Reiczigel CI evenly errs slightly to the safe side (see inset in
Fig 3).

A basic assumption underlying the present study is that the sensitivity and spec-
icity of the diagnostic test have been validated in a manner appropriate for the
1 given application of the test. This refers to the concept of “operational parameters”\(^1\) and requires that the panel selection (of truly positives and truly negatives) for the \(^2\) validation of sensitivity and specificity of the test is representative for its application \(^3\) in the field. This assumption, however, needs to be made independent of whether \(^4\) one uses the conventional Rogan-Gladen estimate or a Bayesian method. \(^5\)

Violations of the conditions (4b) or (4c) can indicate that the prior information \(^6\) about test accuracy is inconsistent with the application data. In this case it may be \(^7\) a better option to refrain from adjustment for misclassification until reliability and \(^8\) relevance of the diagnostic’s parameters are clarified.

Our simulations resulted in calculated Rogan-Gladen estimates outside the possible range of \([0, 1]\) in 4% of the cases. This percentage results from our choices \(^1\) of simulation scenarios and cannot be extrapolated to real applications. However, \(^12\) bias adjustment has led to “impossible” results which does occur in practice. E.g., \(^13\) Moujaber et al. \([18]\) compared prevalences of *Helicobacter pylori* infection in Australia among different age groups but only reported apparent prevalences. Had they \(^15\) corrected for sensitivity and specificity of the ELISA test used, prevalence estimates \(^6\) for two age groups would have been negative. This becomes obvious also in other \(^17\) statistical models, for example when bias adjustment results in negative count data. \(^18\) Lash et al. \([1]\) noted that encountering impossible adjusted data is often of substan-

\(^{19}\) tive importance, as it may represent a fundamental disconnect between the priors \(^20\) and the data or data model, and may signal poor prior information, poor data mod-

\(^{21}\) eling or unrecognized data problems. We believe that this reflects well the situation \(^22\) of truncated Rogan-Gladen estimates. As pointed out elsewhere \([5]\), an adjustment \(^23\) for sensitivity and specificity can be very misleading if those accuracy parameters \(^24\) are not valid for the given prevalence study.

For the construction of their confidence interval, Lang and Reiczigel –as we did \(26\) in our study– use non-informative \(Beta(1, 1)\) priors for sensitivity and specificity \(^27\) and update the probability distributions according to the results of the validation \(^28\) studies and thus gain increased performance by adopting Bayesian concepts. An \(^29\) advantage of our model may be seen in the opportunity to easily incorporate expert \(^30\) judgment on sensitivity and specificity instead of validation studies. This might be \(^31\) particularly useful in situations where a lot of practical experience is available but \(^32\) the existence of a reliable gold standard is questionable.

Latent class models for estimating sensitivity and specificity without gold stan-

\(^{35}\) dard \([19]\) are related to our approach since in this context, prevalence estimation \(^36\) adjusted for sensitivity and specificity occurs as a “by-product” of estimating ac-

\(^{37}\) curacy parameters. Only four out of 64 empirical studies on human populations \(^37\) reported prevalence estimation as primary goal of latent class modelling \([20]\). \(^38\)

There is a recent tendency in health risk assessment studies to replace the tradi-

\(^{40}\) tional Monte Carlo methods by Bayesian network analysis \([21]\). The Bayesian \(^41\) method for the estimation of true prevalence validated in our study can easily be \(^42\) incorporated into a Bayesian network analysis by using the BUGS language (e.g. by \(^42\) using JAGS) and thus provides a single software solution for the complex task of \(^43\) risk assessment in scenarios with a high level of uncertainty.

Even small improvements in the precision of prevalence estimations are likely to \(^45\) contribute considerably to ameliorate the quality of risk analysis studies because in
these studies information about prevalence is incorporated into larger systems of dependencies such that biases display a tendency to be enlarged in the process of end point estimation.

Conclusions

Prevalence estimations can easily be adjusted for diagnostic misclassification if the diagnostic performance of the test or instrument has been characterized in terms of sensitivity and specificity in a validation study. Furthermore, the use of a Bayesian model is a flexible approach for quantifying the combined uncertainties of all model parameters as it can be informed by empirical data as well as by expert opinion. In a validation study on simulated data sets, Bayesian estimates of the true prevalence and the inherent uncertainty proved superior to traditional frequentist methods, exhibiting less bias than the Rogan-Gladen estimate and better coverage than conventional methods.

Abbreviations

| Abbreviation | Definition |
|--------------|------------|
| AP           | Apparent Prevalence |
| B            | Binomial distribution |
| BUGS         | Bayesian inference Using Gibbs Sampling; language for specifying Bayesian models |
| CI           | Confidence Interval |
| CrI          | Credible Interval |
| D+/-         | Disease status |
| HDI          | Highest Density Interval |
| JAGS         | Just Another Gibbs Sampler; software for simulation of Bayesian models using MCMC |
| MCMC         | Markov Chain Monte Carlo |
| MLE          | Maximum Likelihood Estimator |
| π            | prevalence |
| Pr           | Probability |
| RGE          | Rogan-Gladen Estimator for prevalence |
| S+           | Sensitivity |
| Sp           | Specificity |
| T+/-         | Test result |

Declarations

Ethics approval and consent to participate
Not applicable

Consent for publication
Not applicable

Availability of data and materials

The R code used in this study is available on GitHub, github.com/BfRstats/bayespem-validation-code. From this repository, a Zenodo snapshot was created that includes the code as well as all data sets generated and analysed during this study, DOI: 10.5281/zenodo.3631123.

Competing interests

The authors declare that they have no competing interests.
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Authors' contributions

Conceptualization: MF, CMG, MG; Methodology: MF, MW, MG; Data curation, Software, Formal analysis, Validation, and Visualization: MF; Writing - original draft preparation: MF; Writing - review and editing: MF, MW, TS, CMG, MG; Supervision: TS, CMG, MG; Funding acquisition: CMG, MG.

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Figure 3  Confidence interval coverage. Coverage by parameter set for several methods of 95% confidence interval (CI) construction. The nominal coverage of 95% is marked by a solid gray line, and the dashed lines mark coverage values of 90% and 100%. The traditional CI’s (Clopper-Pearson, Blaker, Sterne, and Wald-Rogan-Gladen) all exhibit significant undercoverage. The bottom left inset shows the Lang-Reiczigel CI’s and the Bayesian HDI’s coverage distributions in more detail. It reveals the the Lang-Reiczigel CI tends to have some over coverage and the Bayesian HDI’s coverage appears to be more more symmetrical around the nominal 95% value.

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Table 1 Parameter set simulation. For each parameter set realisation, true values for the statistical parameters were simulated by drawing from continuous uniform distributions, $U(\min, \max)$. Values for the design parameters were simulated by randomly drawing from fixed sets of values (in the case of sample sizes for simulated validation studies of a diagnostic test) or from a discrete uniform distribution (in the case of the sample size for an application of the test).

| Parameter | Description                          | Values sampled from |
|-----------|--------------------------------------|---------------------|
| $\bar{S}_n$ | True sensitivity                     | $U(0.6, 1)$         |
| $\bar{S}_p$ | True specificity                     | $U(0.6, 1)$         |
| $\tilde{\pi}$ | True prevalence                     | $U(0, 1)$          |
| $n_{\bar{S}_n}$ | Sample size for a sensitivity validation study | $\{50, 100, 200, 500, 1000, 2000, 5000\}$ |
| $n_{\bar{S}_p}$ | Sample size for a specificity validation study | $\{50, 100, 200, 500, 1000, 2000, 5000\}$ |
| $n$ | Sample size for a test application | $U(50, 2000)$         |
Table 2  Exemplary parameter sets. The first ten out of the total of 1,000 parameter sets.

| Parameter set | \( \tilde{\mathcal{S}}_{e} \) | \( \tilde{\mathcal{S}}_{p} \) | \( \tilde{\pi} \) | \( n \) | \( n_{e} \) | \( n_{p} \) | \( n \) |
|---------------|-----------------|-----------------|----------------|------|------|------|------|
| 1             | 0.7479478       | 0.9332674       | 0.4459405      | 100  | 200  | 323  |
| 2             | 0.7768875       | 0.6758782       | 0.3946503      | 50   | 500  | 1285 |
| 3             | 0.9816547       | 0.6470824       | 0.4837289      | 500  | 100  | 1820 |
| 4             | 0.9410975       | 0.8721922       | 0.9188760      | 1000 | 100  | 1524 |
| 5             | 0.8979315       | 0.6941977       | 0.8438814      | 500  | 200  | 933  |
| 6             | 0.6628104       | 0.9243710       | 0.5173496      | 5000 | 1000 | 427  |
| 7             | 0.9812932       | 0.9842258       | 0.4371250      | 5000 | 2000 | 1624 |
| 8             | 0.7668501       | 0.9065918       | 0.3431982      | 1000 | 50   | 1667 |
| 9             | 0.9812932       | 0.9842258       | 0.4371250      | 5000 | 2000 | 1624 |
| 10            | 0.6628104       | 0.9243710       | 0.5173496      | 5000 | 1000 | 427  |

Table 3  Exemplary data sets. The first ten of the replicate data sets for the first parameter set.

| Data set | \( \tilde{\mathcal{S}}_{e} \) | \( \tilde{\mathcal{S}}_{p} \) | \( \tilde{\pi} \) | \( n \) | \( n_{e} \) | \( n_{p} \) | \( n \) | \( x_{e} \) | \( x_{p} \) | \( x \) | \( \tilde{\mathcal{S}}_{e} \) | \( \tilde{\mathcal{S}}_{p} \) | \( \tilde{\pi} \) | \( \tilde{\mathcal{X}} \) |
|----------|-----------------|-----------------|----------------|------|------|------|------|------|------|------|-----------------|-----------------|----------------|----------------|
| 1        | 0.7479478       | 0.9332674       | 0.4459405      | 100  | 200  | 323  | 74   | 193  | 129  | 0.74 | 0.965 | 0.3993808 |
| 2        | 0.7479478       | 0.9332674       | 0.4459405      | 100  | 200  | 323  | 70   | 189  | 125  | 0.70 | 0.945 | 0.3869969 |
| 3        | 0.7479478       | 0.9332674       | 0.4459405      | 100  | 200  | 323  | 78   | 189  | 124  | 0.78 | 0.945 | 0.3839009 |
| 4        | 0.7479478       | 0.9332674       | 0.4459405      | 100  | 200  | 323  | 73   | 191  | 109  | 0.73 | 0.955 | 0.3374613 |
| 5        | 0.7479478       | 0.9332674       | 0.4459405      | 100  | 200  | 323  | 76   | 184  | 117  | 0.76 | 0.920 | 0.3622291 |
| 6        | 0.7479478       | 0.9332674       | 0.4459405      | 100  | 200  | 323  | 66   | 188  | 132  | 0.66 | 0.940 | 0.4086687 |
| 7        | 0.7479478       | 0.9332674       | 0.4459405      | 100  | 200  | 323  | 76   | 179  | 121  | 0.76 | 0.895 | 0.3746130 |
| 8        | 0.7479478       | 0.9332674       | 0.4459405      | 100  | 200  | 323  | 75   | 188  | 118  | 0.75 | 0.940 | 0.3653251 |
| 9        | 0.7479478       | 0.9332674       | 0.4459405      | 100  | 200  | 323  | 74   | 187  | 127  | 0.74 | 0.955 | 0.3931889 |
| 10       | 0.7479478       | 0.9332674       | 0.4459405      | 100  | 200  | 323  | 75   | 191  | 127  | 0.75 | 0.955 | 0.3931889 |
Figures

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Supplementary Files

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