An FSKX compliant source attribution model for salmonellosis and a look at its major hidden pitfalls

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Abstract

To reduce the burden of human society that is caused by zoonotic diseases, it is important to attribute sources to human illnesses. One powerful approach in supporting any intervention decision is mathematical modelling. This paper presents a source attribution model which considers five sources (broilers, laying hens, pigs, turkeys) for salmonellosis and uses two datasets from Germany collected over two time periods; one from 2004 to 2007 and one from 2010 to 2011. The model uses a Bayesian modelling approach derived from the so-called Hald model and is based on microbial subtyping. In this case, Salmonella isolates from humans and animals were subtyped with respect to serovar and phage type. Based on that typing, the model estimates how many human salmonellosis cases can be attributed to each of the considered sources. A reference description of the model is available under DOI: 10.1111/zph.12645. Here, we present this model as a ready-to-use resource in the Food Safety Knowledge Exchange (FSKX) format. This open information exchange format allows to re-use, modify, and further develop the model and uses model metadata and controlled vocabulary to harmonise the annotation. In addition to the model, we discuss some technical pitfalls that might occur when running this Bayesian model based on Markov chain Monte Carlo calculations. As source attribution of zoonotic
disease is one useful tool for the One Health approach, our work facilitates the exchange, adjustment, and re-usage of this source attribution model by the international and multi-sectoral community.

Keywords

Salmonella, R programming language, mathematical modelling, Bayesian model, Markov chain Monte Carlo method, Food Safety Knowledge Exchange (FSKX) format

Introduction

Zoonotic diseases are a major burden for human society. The burden relates to two categories: 1) human health burden in form of mortality and morbidity (Taylor et al. 2001) and 2) economical burden, e.g., in form of losses due to health care costs (The World Bank 2012). In the European Union, over 320,000 human cases of zoonotic disease were reported in 2019 (European Food Safety Authority and European Centre for Disease Prevention and Control 2021).

Salmonella is the second most common zoonotic disease in Europe with a stable number of salmonellosis cases during 2014–2018 (European Food Safety Authority and European Centre for Disease Prevention and Control 2021). Although the salmonellosis burden is stagnating, the contribution of Salmonella serovars differs in prevalence and the source of human infection (European Food Safety Authority and European Centre for Disease Prevention and Control 2021, Jabin et al. 2019).

To reduce the human cases of zoonoses, it is important to understand the relationship of potential sources and human illness (Batz et al. 2005). In order to reduce consumer exposure and to optimize intervention measures, it is required to identify the different zoonotic sources of human infections and quantify their relative contribution (Batz et al. 2005, Pires et al. 2009). Both can be supported by source attribution methods. A powerful method is mathematical modelling. One approach for attributing foodborne illnesses is based on microbial subtyping. This approach includes various methods to distinguish bacterial and viral isolates from one another (Pires et al. 2009). A widely used microbial subtyping approach combines serotyping with phage typing that is based on phenotypic methods. In recent years further subtyping approaches based on molecular methods like plasmid analysis or whole-genome sequencing have been used (Boysen et al. 2014, Mather et al. 2015, Munck et al. 2020, Arnold et al. 2021). Whichever method is used, the data which describe the distribution of different subtypes in different sources can be used to do source attribution based on mathematical modelling.

One modelling approach for source attribution that is based on microbial subtyping is the Bayesian model. In the context of food safety, the models developed by Hald et al. (2004) and David et al. (2013) are of special interest. Jabin et al. (2019) used the approach of David et al. (2013) to attribute human salmonellosis to potential food sources in Germany.
using two datasets from 2004–2007 and 2010/2011. Although the mathematical model has been published and described in detail in Jabin et al. (2019), it is not available in a ready-to-use format. Here, we present the Bayesian model referred as Bayes data-based model in Jabin et al. 2019 in the Food Safety Knowledge Exchange (FSKX) format. This open information exchange format uses model metadata and controlled vocabulary to harmonize annotations of risk assessment models (Haberbeck et al. 2018). Together with the model script, the visualization script, and simulation settings, the metadata are the key components of the format (de Alba Aparicio et al. 2018). Thus, FSKX format facilitates the model usage and re-usage.

The two datasets and the mathematical model by Jabin et al. (2019) are incorporated into a ready-to-use source attribution model which can be executed, developed further, and easily adapted to new data by the international risk assessment community. With our work, we facilitate the exchange, adjustment, and re-usage of this source attribution model.

**Model metadata**

The model metadata are part of the FSKX-file (see Suppl. material 1 for the FSKX-file). For details about the metadata schema and the used definitions see Haberbeck et al. (2018) and available on https://foodrisklabs.bfr.bund.de/rakip-harmonization-resources/ (we use the metadata schema Version 1.04).

**General metadata**

**Source:** PUBLISHED SCIENTIFIC STUDIES

**Identifier:** SourceAttributionBfRBayesDB

**Rights:** Creative Commons Attribution 4.0 (CC BY 4.0)

**Availability:** Open access

**Language:** English

**Software:** FSK-Lab 1.9.0

**Language Written In:** R 3

**Objective:** The model attributes human cases of the salmonellosis caused by various serovars of *Salmonella* from various sources (namely, broilers, laying hens, pigs, turkeys, and unknown). The model is parameterized using data from Germany.

**Product/matrix**

**Name:** Broilers

**Description:** Tons of broiler meat consumed
Unit: Tons

Origin Country: Germany

Name: Laying hens

Description: Number of eggs consumed

Unit: Number of eggs

Origin Country: Germany

Name: Pigs

Description: Tons of pork consumed

Unit: Tons

Origin Country: Germany

Name: Turkeys

Description: Tons of turkey meat consumed

Unit: Tons

Origin Country: Germany

Hazard

- **Type**: Microorganisms; **Name**: Salmonella Enteritidis; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for Salmonella enterica subsp. enterica ser. Enteritidis (Salmonella Enteritidis or S.E. for short) which were further typed according to their phage type (Phage types: S.E. PT 1, S.E. PT 11, S.E. PT 14b, S.E. PT 19, S.E. PT 2, S.E. PT 21, S.E. PT 21c, S.E. PT 25, S.E. PT 35, S.E. PT 4, S.E. PT 4a, S.E. PT 4b, S.E. PT 5a, S.E. PT 6, S.E. PT 6a, S.E. PT 7, S.E. PT 7a, S.E. PT 8, other); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: Salmonella Typhimurium; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for Salmonella enterica subsp. enterica ser. Typhimurium (Salmonella Typhimurium or S.T. for short) which were further typed according to their phage type (Phage types: S.T. DT001, S.T. DT007, S.T. DT008, S.T. DT009, S.T. DT012, S.T. DT017, S.T. DT040, S.T. DT041, S.T. D066, S.T. DT099, S.T. DT104, S.T. DT120, S.T. DT126, S.T. DT139, S.T. DT195, S.T. DT208, S.T. U302, S.T. U310, other); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.
• **Type:** Microorganisms; **Name:** *Salmonella Agama*; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Agama (*Salmonella Agama* for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

• **Type:** Microorganisms; **Name:** *Salmonella Agona*; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Agona (*Salmonella Agona* for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

• **Type:** Microorganisms; **Name:** *Salmonella Anatum*; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Anatum (*Salmonella Anatum* for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

• **Type:** Microorganisms; **Name:** *Salmonella Blockley*; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Blockley (*Salmonella Blockley* for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

• **Type:** Microorganisms; **Name:** *Salmonella Braenderup*; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Braenderup (*Salmonella Braenderup* for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

• **Type:** Microorganisms; **Name:** *Salmonella Brandenburg*; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Brandenburg (*Salmonella Brandenburg* for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

• **Type:** Microorganisms; **Name:** *Salmonella Coeln*; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Coeln (*Salmonella Coeln* for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

• **Type:** Microorganisms; **Name:** *Salmonella Derby*; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Derby (*Salmonella Derby* for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

• **Type:** Microorganisms; **Name:** *Salmonella Eboko*; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Eboko
(Salmonella Eboko for short); unit: %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type:** Microorganisms; **Name:** *Salmonella* Give; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Give (*Salmonella* Give for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type:** Microorganisms; **Name:** *Salmonella* Heidelberg; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Heidelberg (*Salmonella* Heidelberg for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type:** Microorganisms; **Name:** *Salmonella* Hessarek; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Hessarek (*Salmonella* Hessarek for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type:** Microorganisms; **Name:** *Salmonella* Indiana; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Indiana (*Salmonella* Indiana for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type:** Microorganisms; **Name:** *Salmonella* Infantis; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Infantis (*Salmonella* Infantis for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type:** Microorganisms; **Name:** *Salmonella* Kedougou; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Kedougou (*Salmonella* Kedougou for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type:** Microorganisms; **Name:** *Salmonella* Kottbus; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Kottbus (*Salmonella* Kottbus for short); **Unit:** %; **Adverse Effect:** The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type:** Microorganisms; **Name:** *Salmonella* Lexington; **Description:** The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Lexington (*Salmonella* Lexington for short); **Unit:** %; **Adverse Effect:** The most
common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Liverpool; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Liverpool (*Salmonella Liverpool for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Livingstone; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Livingstone (*Salmonella Livingstone for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* London; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. London (*Salmonella London for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Mbandaka; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Mbandaka (*Salmonella Mbandaka for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Montevideo; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Montevideo (*Salmonella Montevideo for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Newport; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Newport (*Salmonella Newport for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Ohio; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Ohio (*Salmonella Ohio for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Rissen; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Rissen
(Salmonella Rissen for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Saintpaul; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Saintpaul (*Salmonella Saintpaul for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Senftenberg; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Senftenberg (*Salmonella Senftenberg for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Stanley; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Stanley (*Salmonella Stanley for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Tennessee; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Tennessee (*Salmonella Tennessee for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* Virchow; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* ser. Virchow (*Salmonella Virchow for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: rough *Salmonella*; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which contain rough strains of *Salmonella enterica* subsp. *enterica* with unspecified serovar (rough *Salmonella* for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.

- **Type**: Microorganisms; **Name**: *Salmonella* - other serotypes; **Description**: The model considers the prevalence among isolates from animal samples (pig, broiler, laying hen, turkey) which are positive for *Salmonella enterica* subsp. *enterica* with unspecified serotype (*Salmonella* - other serotypes for short); **Unit**: %; **Adverse Effect**: The most common symptoms of salmonellosis are diarrhea, fever, abdominal cramps, and vomiting.
Population

Name: People in Germany

Target Population: People in Germany that were identified by medical professionals to be suffering from salmonellosis

Scope

The model attributes human cases of the zoonotic disease salmonellosis to a certain source (namely, broilers, laying hens, pigs, turkeys, and unknown). It is based on a Bayesian microbial subtyping approach described by Hald et al. (2004) and subsequently modified by David et al. (2013). Data from two datasets using information from various years are used to parameterize the model. Both datasets are from Germany.

Temporal Information: In Jabin et al. 2019, data in Table 2 in Jabin et al. (2019) consider human salmonellosis in the years 2004-2007 and data in Table 3 in Jabin et al. (2019) consider the years 2010/2011. See Table 4 in Jabin et al. (2019) for details about the data sources for Salmonella in humans.

Data background

Study Title: The role of parameterization in comparing source attribution models based on microbial subtyping for salmonellosis

Study Description: Two datasets from active monitoring in Germany were available. The data comprise four potential animal sources: broilers, laying hens, pigs, and turkeys. For each considered salmonellosis case, the serotype was determined. For cases caused by Salmonella Enteritidis or Salmonella Typhimurium additionally the phage type was determined (see Tables 2 and 3 in Jabin et al. 2019). Both datasets cover multiple years (for details see "Temporal Information" in the Subsection "Scope" of the Section "Model metadata" and Jabin et al. 2019). The data are used to analyze the source attribution model see Section "Material and methods" and Jabin et al. (2019) for details.

Material and methods

Data

Datasets covering studies on Salmonella in different sources for two time periods were compiled and used for this analysis. For both time periods, reliable data from active monitoring on four potential animal sources were available: broilers, laying hens, pigs, and turkeys. Cattle were not included in any study or program and were therefore not included in this analysis. The datasets, which cover the years 2004–2007 and 2010/2011, are called baseline data and monitoring data, respectively. In addition, data on human salmonellosis cases were considered.
Baseline data

The first dataset on *Salmonella* in sources was generated by four baseline studies conducted during 2004 and 2007 in Germany (EFSA (2007b), EFSA (2007a), EFSA (2008b), EFSA (2008a)). These data describe the prevalence $p_{ij}$ (in %) of each *Salmonella* subtype $i$ in each source $j$. The prevalence for all strains of *Salmonella* serotypes Enteritidis and Typhimurium as well as the relevant phage types are considered (see Jabin et al. (2019) for details).

Monitoring data

The second dataset on *Salmonella* in sources was compiled from monitoring programs during 2010 and 2011 in Germany (Käsbohrer et al. 2012, Käsbohrer et al. 2013). Data for broilers, laying hens, and turkeys were obtained from the *Salmonella* control programs in poultry in 2010. Since national monitoring of *Salmonella* prevalence in pigs was conducted only in 2011, the data on pigs from 2011 were combined with the poultry data from 2010, assuming that the serotype distributions in the animals were equal in both years (see Jabin et al. (2019) for details).

To summarize, the baseline and the monitoring data are comparable, i.e., the data were compiled in a similar way and the intention measures in the years were the same, thus, no significant difference in the data is expected.

Human data

Data on human *Salmonella* cases came from the Robert Koch Institute (RKI). The serotype distribution was obtained via their online database SurvStat@RKI (https://survstat.rki.de/, data access: 07.02.2012). In addition, phage type information for S. Enteritidis and S. Typhimurium strains were provided via personal communication by Wolfgang Rabsch (RKI). Since only a subset of all strains isolated from humans had been phage typed, we assumed that the phage type distribution among the typed strains is also representative for the untyped strains. To account for the four year time period of the baseline studies (from 2004 to 2007), we summed up all the corresponding sero- and phage types associated with human salmonellosis cases over that time period (see Jabin et al. (2019) for details).

Mathematical model

The presented Bayes data-based (DB) model is a source attribution model that is based on microbial subtyping (Jabin et al. 2019). It is derived from the model developed by Hald et al. 2004, the so-called Hald model. This model has been adopted widely (David et al. 2013, Pires et al. 2011, Ranta et al. 2011). The variation of the Hald model developed by David et al. 2013 reparameterizes the Hald model which leads to an improved robustness of source attribution estimates.

A note about terminology: the terms "subtype" and "type" are used interchangeably.
Hald model

The so-called Hald model (Hald et al. 2004) is a Bayesian modelling approach that uses microbial subtyping data to infer the sources for observed food-borne illnesses like salmonellosis. This model approach is based on inferring a posterior estimate of the considered outcome using prior assumptions and the use of data. In the Hald model, one assumption is that the number of human cases of salmonellosis is Poisson distributed:

$$o_i \sim \text{Poisson} \left( \sum_{j=1}^{J} \lambda_{ij} \right)$$  \hspace{1cm} (1)

where $o_i$ is the number of observed cases for *Salmonella* of subtype $i$. The number of subtypes run from $i = 1,2,\ldots, I$, where $I$ is the total number of *Salmonella* subtypes present in the data. The number of sources in the data and thus considered in the model is $J$. Here, $\lambda_{ij}$ is the number of expected cases caused by *Salmonella* subtype $i$ in source $j$ (with $j$ running from $j = 1,2,\ldots,J$). The Hald model defines $\lambda_{ij}$ as follows:

$$\lambda_{ij} = M_j \cdot p_{ij} \cdot q_i \cdot a_j$$  \hspace{1cm} (2)

where $M_j$ is the amount of source $j$ consumed (in tons, except for laying hens where it is number of eggs). The values $p_{ij}$ (in %) for the prevalence $p_i$ of *Salmonella* subtype $i$ in the source $j$. The parameter $q_i$ is a subtype-dependent factor which describes the ability of the *Salmonella* subtype $i$ to cause illness. The parameter $a_j$ is a source-dependent factor describing the ability of source $j$ to serve as a vector for *Salmonella*. Equation 2 represents the multiparameter prior of the model with the two parameters $a_j$ and $q_i$ of unknown value. For the parameter $a_j$ and $q_i$, uniform distributions where defined as prior distributions.

David model—a variation of the Hald model

Some authors describe difficulties with the convergence of the Hald model (Guo et al. 2011, Mullner et al. 2009). To address this issue, David et al. (2013) proposed a reparameterization of the Hald model based on unique types (or “specific types” as called in David et al. (2013)). A unique type is a subtype that is specific to a food-source and consequently is not found in another source under consideration. If there are one or multiple unique types for a source $j$ in the considered data, then the corresponding unique subtype-dependent parameters $q_{\text{ut},j}$ for these unique types are parameterized according to Equation 3 instead of Expression 6. The subscript, “ut” stands for unique type.

$$q_{\text{ut},j} = \frac{O_{\text{ut}}}{\sum_i O_i} \cdot \frac{1}{p_{\text{ut},j}}$$  \hspace{1cm} (3)

This reparameterization can only be done if all serotypes are phage typed. As not all the data of serotypes Enteritidis and Typhimurium considered by David et al. (2013) were phage typed, both serotypes were excluded from the reparameterization (see Section 2.2.5 in David et al. 2013).
Bayes data-based (DB) model—a variation of the David model

Following the idea of David et al. (2013) to use unique types for parameterizing the Hald model, the Bayes DB model uses the following parameterization setup:

1. Parameterization of the subtype-dependent parameter $q_i$
   - For each source $j$, choose freely one unique type and call the corresponding subtype-dependent parameter $q_{ut,j}$ (if available)
   - Parameterize $q_{ut,j}$ for the chosen unique type according to Equation 3
   - If there are no unique types, parameterize all $q_i$ according to Expression 6

2. Parameterization of the source-dependent parameter $a_j$
   - For each source $j$ where no unique type is available, the corresponding parameter $a_{nut}$ is defined as ("nut" in subscripts stands for "no unique type"):
     
     \[ a_{nut} = \frac{\sum_i o_i}{M_{nut}} \]  

   - This also applies to the case that no source has a unique type.

3. Parameterization of the consumption data $M_j$
   - $M_j$ is set according to consumption data (this is the case for the presented Bayes DB model).
   - If no consumption data are available, all $M_j$ are set to appropriate constant values. The values need to be large enough to assure consistent model results. These constant values are found through trial and error and depend on the considered dataset (for details see Section "The effect of consumption data on the consistency of source attribution estimates").

To estimate unknown parameters, uniform distributions are assumed as prior distributions for $a_j$ and $q_i$ (see Expressions 5 and 6, respectively). Unknown parameters are: 1) all $q_i$ which belong to non-unique types, 2) unique $q_i$ which have not been chosen according to the first step of the parameterization setup, 3) all $a_j$ which correspond to sources $j$ which have at least one unique type. Note that $M_j$ is always set to a fixed value. Consequently, if there are no unique types, all $a_j$ are parameterized according to Equation 4 and all $q_i$ according to Expression 6.

In the model presented in this paper the following prior distributions were assumed:

\[ a_j \sim uniform(0, 0.2) \quad (j = 1, 2, ..., J) \]  
\[ q_i \sim uniform(0, 1) \quad (i = 1, 2, ..., I). \]

The the limits of the prior distributions were chosen such that they produce complete posterior distributions for both datasets (baseline and monitoring data). Depending on the
data, one might have to adjust the limits of the distribution (see Section "The effect of prior distributions on completeness of posterior distributions" for details).

In the next section, we describe how to parameterize the model and run model simulations using FSKX format.

Simulations

All model parameters and their descriptions are presented in Table 1. Two simulation scenarios are provided in the fskx-model (see Table 2 for the parameter values of both scenarios and Suppl. material 1 for the fskx-model). The default simulation considers the data of the baseline study (see Section "Baseline data" for details). The second simulation setting is based on the monitoring data (see Section "Monitoring data" for details).

| Table 1. Description of the model parameters of the source attribution model. In the row that specifies the source, article always refers to the reference description of Jabin et al. (2019). |
|---|---|
| Id | list_sources |
| Classification | INPUT |
| Name | list_sources |
| Description | List all possible sources |
| Unit | [] |
| Data Type | INTEGER |
| Source | Article |
| Value | c('Broilers', 'Laying hens', 'Pigs', 'Turkeys') |

| Id | qfix_ind |
| Classification | INPUT |
| Name | qfix_ind |
| Description | Indices of subtype-dependent factor for subtype i (qi), which will be set to fixed values. These are the four values for the human cases concerning the “unique types”: S. Virchow, S.E. PT 1, S.T. DT 193, and S. Saintpaul |
| Unit | [] |
| Data Type | VECTOROFNUMBERS |
| Source | Data |
| Value | c(63,64,65,66) |
| Min Value | 1 |
| Id                | Description                  |
|-------------------|------------------------------|
| input_FileName    | Name of the file that contains the analysed data |
| OpenBUGS_parameter | The values that should be logged while running the OpenBUGS-model |

### Max Value
Number of considered subtypes

### Id
input_FileName

### Classification
INPUT

### Name
input_FileName

### Description
Name of the file that contains the analysed data

### Unit
[]

### Data Type
STRING

### Source
Article

### Value
"Table2.csv"

### Id
OpenBUGS_parameter

### Classification
INPUT

### Name
OpenBUGS_parameter

### Description
The values that should be logged while running the OpenBUGS-model

### Unit
[]

### Data Type
STRING

### Source
Article

### Value
c("source", "unknown", "a", "q", "lambdaexp")

### Id
OpenBUGS_niter

### Classification
INPUT

### Name
OpenBUGS_niter

### Description
Number of total iterations per chain used in the OpenBUGS-model

### Unit
[]

### Data Type
INTEGER

### Source
Article

### Value
30000

### Min Value
OpenBUGS_nburnin+1

### Id
OpenBUGS_nburnin

### Classification
INPUT

### Name
OpenBUGS_nburnin

### Description
Length of burn in, i.e. number of iterations to discard at the beginning.
| Id          | aValue          |
|-------------|-----------------|
| Classification | INPUT |
| Name        | aValue          |
| Description | Values for the source-dependent factors \(a_i\) that are used to determine initial values for the OpenBUGS model |
| Unit        | []              |
| Data Type   | INTEGER         |
| Source      | Article         |
| Value       | 10000           |
| Min Value   | 1               |

| Id          | qValue          |
|-------------|-----------------|
| Classification | INPUT |
| Name        | qValue          |
| Description | Values for the subtype-dependent factors \(q_i\) that are used to determine initial values for the OpenBUGS model |
| Unit        | []              |
| Data Type   | VECTOROFNUMBERS |
| Source      | Data            |
| Value       | c(0.002, 0.001, 0.19, 0.18, 0.178) |
| Min Value   | 0               |

| Id          | OpenBUGS_model |
|-------------|----------------|
| Classification | INPUT |
| Name        | OpenBUGS_model |
| Description | The filename of the txt-file that contains the OpenBUGS-model |
| Unit        | []              |
| Data Type   | STRING          |
| Source | The filename is freely chosen. The BUGS-model is described in the reference article. |
|--------|-----------------------------------------------------------------------------------|
| Value  | "BugsModel.txt"                                                                    |
| Id     | mean_res                                                                         |
| Classification | OUTPUT                                                                 |
| Name   | mean_res                                                                         |
| Description | Mean number of estimated human salmonellosiscases attribute to potential sources |
| Unit   | Cases                                                                            |
| Data Type | VECTOROFNUMBERS                                                                 |
| Min Value | 0                                                                                 |
| Max Value | 1                                                                                 |
| Id     | quantil_95                                                                       |
| Classification | OUTPUT                                                                 |
| Name   | quantil_95                                                                       |
| Description | 95%-quantile of estimated human salmonellosiscases attributed to the potential sources |
| Unit   | Cases                                                                            |
| Data Type | VECTOROFNUMBERS                                                                 |
| Min Value | 0                                                                                 |
| Max Value | 1                                                                                 |
| Id     | quantil_05                                                                       |
| Classification | OUTPUT                                                                 |
| Name   | quantil_05                                                                       |
| Description | 5%-quantile of estimated human salmonellosiscases attributed to the potential sources |
| Unit   | Cases                                                                            |
| Data Type | VECTOROFNUMBERS                                                                 |
| Min Value | 0                                                                                 |
| Max Value | 1                                                                                 |
The Bayes DB model is implemented in the programming language R (R Core Team 2019). In addition to R, the open source software OpenBUGS is required to successfully execute the model (Neal 2009). The linkage between both software tools is done by the R package "R2OpenBUGS" (Sturtz et al. 2005; see file "packages.json" in the fskx-model).

The fskx-model can be executed, developed further, and easily adapted to new data on the local computer, e.g., using the KNIME extension FSK-Lab (see https://foodrisklabs.bfr.bund.de/fsk-lab/ and de Alba Aparicio et al. (2018)).
Executable model

In order to execute the model, please register at the virtual research environment "FMJ_Lab".

Execute with default simulation parameters: execute

The default simulation runs for 2 minutes 11 seconds on the virtual research environment.

Execute another simulation scenario or create a personalized scenario: execute

Results

The main result is that the existing source attribution model previously published in Jabin et al. (2019) is available in the ready-to-use FSKX-format. The format is an open information exchange format and uses model metadata and controlled vocabulary to harmonise annotations. The transformation into an FSKX compliant model took about one day of work for a person already familiar with the format. In the FSKX compliant format the model predicts the same source attribution as in the original version as the R and the OpenBUGS code is nearly identical to the code used in Jabin et al. (2019).

To be able to successfully use the model, it is important to know how to set up and run the model as well as assess the appropriateness of the results. We present these practical issues since this is a purely technical paper it seems appropriate to provide this level of technicality here.

Successfully executing a Bayesian model using Markov chain Monte Carlo simulations

When running our Bayesian model using Markov Chain Monte Carlo (MCMC) methods, we studied three important aspects of model diagnostics. To ensure a high quality estimation of unknown parameters, we check the following aspects of a MCMC method: the convergence behaviour of the Markov chains, the completeness of posterior distributions, and the consistency of results.

The effect of prior distributions on completeness of posterior distributions

The limits for the uniform distribution have a strong influence on the completeness of the posterior distributions. The limits are incorporated into the OpenBUGS code of the model (see file "BugsModel.txt" in the fskx-model). In the Bayes DB model, the lower limit is 0 and the upper limits are 0.2 for $a_j$ and 1 for $q_i$ for both datasets (see Section "Bayes data-based (DB) model—a variation of the David model" and Expressions 5 and 6). The upper limits were chosen such that the model provides complete posterior distributions. This was assured by examining visually the plots of the posterior distributions of $a_j$ and $q_i$ (Hald et al. 2004; and Fig. 1 in this paper). If one changes the upper limit of the prior distribution of $q_i$ to 0.2, incomplete posteriors were obtained. In Fig. 1, there is an example for a complete
and an incomplete posterior distribution (Subfigures A) and B), respectively). The incomplete posterior distribution is a trimmed version of the complete one. Please note that issues in the completeness of posterior distributions might only occur in some but not all of the model parameters.

The effect of initial values on convergence and uncertainty estimates

The choice of the starting values of the Markov chains (also known as initial values) has an impact on the convergence and uncertainty estimates of the model calculation. The model runs with five Markov chains. The default starting values for the five chains are listed in Table 1; the parameter names for $a_j$ and $q_i$ are "aValue" and "qValue", respectively. This means that the Markov chains for each unknown parameters start with five different, but predefined, starting points. The effect of initial values on the convergence and uncertainty estimates will be exemplified using the baseline data in three parameter scenarios. The parameter scenarios differ in their corresponding set of five starting points for their five Markov chains. The parameter scenarios exemplify the following effects: successful convergence (Parameter scenario 1), slow convergence (Parameter scenario 2), and no convergence (Parameter scenario 3). The parameter scenarios are represented graphically in Figs 2, 3, 4 which show the starting points for the Markov chains in scatter plots, convergence behaviour of the Markov chains in trace plots, and the result of the source...
Attribution estimates in bar plots. The bar plots include error bars which correspond to a 90% equal-tailed interval (i.e. the interval between the 0.05-quantile and the 0.95-quantile of the posterior distribution of the number of human cases). The error bars represent the uncertainty in the model estimation; the bigger the bars the higher the uncertainty.

In Parameter Scenario 1, the starting points are evenly spaced in the lower fifth of the plane of possible starting values (see Fig. 2). The error bars in the source attribution results are small (see bar plot in Fig. 2).

In Parameter Scenario 2, the starting points are concentrated near two points: one point is (0, 0) the other (0.18, 0.18) (see Fig. 3). The Markov chains converge slowly for some parameters, e.g., \( a_2 \) and \( q_3 \) in the trace plot of Fig. 3. The error bars of the model results are large (much larger than in the previous parameter scenario) (see bar plot in Fig. 3).

Finally, the starting points cluster near two points: one point is (0, 0.18) the other (0.18, 0.18) (see Fig. 4). In this parameter scenario, the Markov chains do not converge at all within the 30,000 iterations for some parameters, e.g., for \( a_2 \) and \( q_3 \) (the trace plot of Fig. 4). Consequently, the error bars of the model results are larger than in the previous two parameter scenarios (see bar plot in Fig. 4).
The effect of consumption data on the consistency of source attribution estimates

Some authors pointed out that the parameter for consumption data, \( M \), are not essential for the approach (Mughini-Gras and van Pelt 2014, Mullner et al. 2009, Wahlstrom et al. 2011). According to them, \( M \) serves as a scaling factor for \( a \) and could be omitted (as done in Mullner et al. (2009), Wahlstrom et al. (2011)). The approach of setting \( M \) to 1, caused problems for the Bayes DB model. Problems are either numerical issues or inconsistent results. Inconsistency means that the predicted number of human cases for a certain subtype is not in the same order of magnitude of the number of human cases found in the data (in such cases we found that the number of observed cases could be up to a factor of 2000 larger than the model estimates).

Simplifying the Bayes DB model for the baseline data by setting all \( M \) to 1 and keeping the prior distributions as they were defined in Expression 5 and Expression 6 caused problems. OpenBUGS was not able to successfully execute the model, due to numerical problems (OpenBUGS reports an “conjugate gamma updater error” for one of the \( q_i \)). This problem disappeared when the prior distributions for \( q_i \) were changed to

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**Figure 3.**

Starting points for the Markov chains of Parameter scenario 2 and their effects on the convergence behaviour and the model predictions. The starting points are concentrated near the points (0, 0) and (0.18, 0.18) (see the points in the scatter plot in the upper right corner). With this set of starting points, Markov chains converge slowly as can be seen in the four trace plots on the left hand side which show how the parameter values that the model estimates change through the iteration steps of the model calculations. Each of the four trace plots correspond to one model parameter (\( a_1, a_2, q_2 \) and \( q_3 \), where types 1, 2 and 3 correspond to *S. enterica* serotype Enteritidis PT 11, PT 14b, and PT 19, respectively). In each trace plot there are five traces, one trace for each Markov chain. Each Markov chain has its own colour. The predicted source attribution shows small error bars (see the bar plot).
For the monitoring data, setting all $M_j$ to 1 and using prior distributions as defined in Expression 5 and 6 led to inconsistent results. The model worked properly when priors distributions were set to $a_j \sim \text{uniform}(0,30000)$ while the priors for $q_i$ remained the same as in Expression 6 (cf. Fig. 5).

One way to interpret the need for enlarging the priors for $a_j$ and $q_i$ is that parameters $a_j$ and $q_i$ must compensate for the restrictions applied to $M_j$. One may consider $a_j$ and $q_i$ as complex priors distributions that combine estimates of the potential of the Salmonella of type $i$ and the source $j$ to cause salmonellosis. In summary, if $M_j$ is simplified, the prior distributions may need to be adjusted.
Source attribution determination

Source attribution methods aim to identify and quantify the contribution of different sources to disease burdens like salmonellosis (Jabin et al. 2019). The human salmonellosis cases are attributed to different sources (namely broilers, laying hens, pigs, turkeys, and unknown). In Fig. 6A, the number of human cases of *Salmonella* subtypes in animal sources from the baseline studies 2004–2007 are presented. The source that causes the majority of salmonellosis cases is laying hens while turkeys cause the smallest burden of the considered sources. The results for the monitoring data (2010/2011) are presented in Fig. 6B. The majority of cases here results from unknown sources. Closely followed by laying hens and pigs. A relatively low burden results from turkeys.

The presented results allow to analyse the quantity of the burden assignable to each source and provide the basis to compare different datasets. Although the baseline and the monitoring data are comparable and no significant difference between the datasets is expected (see Section "Data" for details), it provides the basis for comparison. There is much more to say about the model and its results but we focus here on the technical aspects of making the model FSKX compliant and some of the model mechanics. For a more detailed discussion of the model and its results see Jabin et al. (2019).
Discussion

Modelling of source attribution is a powerful approach that can contribute to the reduction of human zoonotic cases, in particular salmonellosis. However, model results are highly sensitive to changes of multiple parameters that can differ for each model. In the presented model, these parameters include the initial values for observed Salmonella cases and the assumption about the consumption data. If someone aims to reproduce the model results, this is only possible if the parameter settings are identical to the original settings. In other words, slight changes in a model parameter might result in a big change in the model prediction and thus, the results presented in an article or report cannot be reproduced. The issue of reproducible results is a general challenge in science (Baker 2016, Goodman et al. 2016). It might seem that in computational work it is in principle easy to re-use and share the used data and the used code in order to reproduce results (except for variations when the model calculations include probabilistic elements). Nonetheless, there is a problem with reproducibility in this area as well (Waltemath and Wolkenhauer 2016, Stodden et al. 2018, Tiwari et al. 2021, Miłkowski et al. 2018). It can be hard to re-use one own modestly documented models; it can be particularly challenging to re-use models developed by other authors (see Topalidou et al. 2015 for an illustrative example). Problems arise from the limited amount of documentation and versioning of the code and/or insufficient information about the model scope (Stodden et al. 2018, Waltemath and Wolkenhauer 2016). The consequence of the reproducibility/sharing problem is that models are re-invented and re-implemented; a time-consuming and/or error-prone process. Several approaches have been discussed in the literature to remedy these problems (Grimm et al. 2014, Wilson et al.}
All approaches come down to a combination of standardized way to document the model and to choose ways to store and share computational resources like data and model code platform independently. FSKX format is an open information exchange format that provides a way to create well-documented and reproducible mathematical risk assessment models that are annotated in a harmonised way using model metadata and controlled vocabulary (de Alba Aparicio et al. 2018).

The implementation of a model in a standardized and annotated exchange format like FSKX-format is a way that focuses on long-term usability and understandability of the model. The community as well as the creators would benefit from such an approach. One example where a creator developed a model with an FSKX conform end-product in mind is the work of Plaza-Rodríguez et al. (2019).

Much time-consuming and/or error-prone work can be saved in the future if model development is done with a mind-set of long-term usability, reproducibility, and understandability. The FSKX format enables sharing model code reliably and reproducibly and thus paves the way for successful collaboration and further development of models.

Conclusion

In this work, we demonstrated that it is straight forward to take a Bayesian source attribution model running under R and OpenBUGS originally published in Jabin et al. (2019), and translate it into the Food Safety Knowledge Exchange (FSKX) format. This standardized format provides an annotated model together with relevant simulation settings. The ready-to-use model can be executed in this "executable paper" and on the local computer, e.g., using software like the KNIME extension FSK-Lab (de Alba Aparicio et al. 2018). In addition, it is easy to re-use the model code and interpret simulation results. In conclusion, we provide an annotated, ready-to-use source attribution model and the considered Salmonella datasets and by that facilitate model exchange, adjustment, and re-use by the international and multi-sectoral One Health community.

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Author contributions

Esther M. Sundermann: Conceptualization, Data Curation, Project administration, Software, Visualization, Writing - Original Draft, Writing - Review & Editing. Guido Correia Carreira: Conceptualization, Formal analysis, Writing - Original Draft, Visualization, Writing - Review & Editing. Annemarie Käsbohrer: Data Curation, Writing - Review & Editing. The author contributions are taken from https://www.elsevier.com/authors/policies-and-guidelines/credit-author-statement

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Supplementary material

Suppl. material 1: SourceAttributionModel.fskx  [doi]

Authors: Esther M. Sundermann
Data type: fskx-model
Download file (1.44 MB)