Evaluation of using ICD-10 code data for respiratory syncytial virus surveillance

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Abstract
Background: Respiratory syncytial virus (RSV) is the most common cause of acute lower respiratory tract infection (ALRI) in young children. ICD-10-based syndromic surveillance can transmit data rapidly in a standardized way.

Objectives: We investigated the use of RSV-specific ICD-10 codes for RSV surveillance.

Methods: We performed a retrospective descriptive data analysis based on existing ICD-10-based surveillance systems for ALRI in primary and secondary care and a linked virological surveillance in Germany. We described RSV epidemiology and compared the epidemiological findings based on ICD-10 and virological data. We calculated sensitivity and specificity of RSV-specific ICD-10 codes and in combination with ICD-10 codes for acute respiratory infections (ARI) for the identification of laboratory-confirmed RSV infections.

Results: Based on the ICD-10 and virological data, epidemiology of RSV was described, and common findings were found. The RSV-specific ICD-10 codes had poor sensitivity 6% (95%-CI: 3%-12%) and high specificity 99.8% (95%-CI: 99.6%-99.9%). In children <5 years and in RSV seasons, the sensitivities of RSV-specific ICD-10 codes combined with general ALRI ICD-10 codes J18.-, J20.- and with J12.-, J18.-, J20.-, J21.-, J22 were moderate (44%, 95%-CI: 30%-59%). The specificities of both combinations remained high (91%, 95%-CI: 86%-94%; 90%, 95%-CI: 85%-94%).

Conclusions: The use of RSV-specific ICD-10 codes may be a useful indicator to describe RSV epidemiology. However, RSV-specific ICD-10 codes underestimate the number of actual RSV infections. This can be overcome by combining RSV-specific and general ALRI ICD-10 codes. Further investigations are required to validate this approach in other settings.

KEYWORDS
epidemiology, ICD-10 code, respiratory syncytial virus, sensitivity, specificity, surveillance
Respiratory syncytial virus (RSV) is a worldwide distributed pathogen of acute respiratory infection (ARI) of all ages. In infants and young children, RSV is the most common cause of acute lower respiratory tract infection (ALRI) and a major cause of hospital admission for ALRI. Worldwide in 2015, 21.6–50.3 million RSV-associated ALRI episodes occurred in children younger than 5 years, with about 2.7–3.8 million hospital admissions.1,2 Currently, only passive immunization with palivizumab against RSV is available for children at high risk.3 In 2015, the World Health Organization (WHO) Product Development for Vaccines Advisory Committee highlighted the development of safe and efficacious RSV vaccines for global use. Several novel RSV vaccines have shown promising results in clinical trials and are expected to enter the market by 2025.4,5 The planning of future RSV vaccination strategies and the evaluation of RSV vaccination impact rely on timely RSV epidemiological data and long-term observation of RSV seasonality through RSV surveillance systems.

International Statistical Classification of Diseases and Related Health Problems (ICD) diagnosis codes have been used to describe the burden of respiratory diseases and the impact of vaccination.6–9 ICD-based digital syndromic surveillance is a relatively novel surveillance practice, compared to the traditional surveillance. It can not only describe epidemiology of disease, but also capture and transmit data rapidly in a standardized and sustainable way at lower costs, and provide very early warning of potential public health threats.10–12 ICD-based digital syndromic surveillance systems have been linked with a virological surveillance and a sentinel electronic data collection system based on ICD-10 codes (SEEDARE).13 SEEDARE was evaluated as a valid system for syndromic influenza surveillance.14 In secondary care, an ICD-10 code-based surveillance system for sentinel practices report influenza and other ARI voluntarily through a syndromic influenza surveillance system. This system has been linked with a virological surveillance and a sentinel electronic data collection system based on ICD-10 codes (SEEDARE).13 SEEDARE was evaluated as a valid system for syndromic influenza surveillance.14

The Robert Koch Institute (RKI) established the 10th revision of ICD (ICD-10)-based digital syndromic surveillance systems for influenza and other ARI in primary and secondary care in Germany (Appendix S1). In primary care, general practitioners, internists, and pediatricians of sentinel practices report influenza and other ARI data voluntarily through a syndromic influenza surveillance system. This system has been linked with a virological surveillance and a sentinel electronic data collection system based on ICD-10 codes (SEEDARE).13 SEEDARE was evaluated as a valid system for syndromic influenza surveillance.14

In secondary care, an ICD-10 code-based surveillance system for severe acute respiratory infections (ICOSARI) has been implemented in cooperation with a private hospital network in Germany.15

Studies estimating validity of ICD diagnosis codes for the identification of laboratory-confirmed influenza have shown mixed results.14,16–18 So far, few studies have looked at accuracy of RSV-specific ICD-10 diagnosis codes for the identification of true RSV infections. To our knowledge, only Pisesky et al19 reported high sensitivity (97.9%, 95%‐CI: 95.5%‐99.2%) and specificity (99.6%, 95%‐CI: 98.2%‐99.8%) of RSV-specific ICD-10 codes for the identification of hospitalized RSV among children.

The aim of this study was to evaluate the use of RSV-specific ICD-10 diagnosis codes for RSV surveillance.

We performed a retrospective descriptive data analysis based on the data derived from ICD-10-based influenza and other ARI surveillance systems SEEDARE and ICOSARI, and from the virological surveillance at the RKI. The SEEDARE system has functioned since 2007, the virological surveillance since 2010, and ICOSARI since 2015. The datasets of ICOSARI for the years 2009 to 2014 were collected retrospectively. The Appendix S1 provides details on the surveillance participants, data collection methods, collected data, total number of collected data, and study period (13–15, Appendix S1).

The SEEDARE system was approved by the German Federal Commissioner for Data Protection and Freedom of Information, and the ICOSARI system by the RKI and HELIOS Kliniken GmbH data protection authority. As SEEDARE and ICOSARI involved no interventions and the analysis was based on anonymized data only, no ethical clearance was required for them.14,15 The virological surveillance activities were approved by the German Federal Commissioner for Data Protection and Freedom of Information and the Ethical Committee of the Charité, Universitätsmedizin, Berlin.

We defined a RSV-ICD-case based on SEEDARE data as a medical consultation with any of the three RSV-specific ICD-10 code diagnoses (J12.1 RSV pneumonia, J20.5 acute bronchitis due to RSV, and J21.0 acute bronchiolitis due to RSV). We defined a RSV-ICD-case based on ICOSARI data as a hospitalization with any of the three RSV-specific ICD-10 code diagnoses as primary discharge diagnosis. In the virological surveillance, we defined a confirmed-RSV-case as a by realtime reverse transcriptase polymerase chain reaction (rtRT-PCR) confirmed RSV sample. In each data source, a RSV season was defined as the weeks when cumulative number of RSV-ICD-cases or confirmed-RSV-cases exceeded 1.2% of total RSV-ICD-cases or confirmed-RSV-cases. One gap week below the threshold was allowed.20,21

We estimated number of RSV-ICD-cases and confirmed-RSV-cases by gender, age group (0-1, 2-4, 5-14, 15-34, 35-49, 50-59, ≥60 years), and calendar week based on each data source, respectively.

We identified the sentinel practices that participated in both SEEDARE and the virological surveillance concurrently by practice-ID. We matched the medical consultations of SEEDARE with virological samples by practice-ID, age, gender, consultation date, and sampling date. Only one-to-one matches were included for the further data evaluation. We calculated sensitivity of RSV-specific ICD-10 code diagnosis as proportion of RSV-ICD-cases among confirmed-RSV-cases, and specificity as proportion of non-RSV-ICD-cases among non-confirmed-RSV-cases of the identified practices. We calculated sensitivity and specificity of RSV-specific ICD-10 code diagnosis among young children, in RSV seasons, and combined with different general ARI ICD-10 codes J06.- acute upper respiratory infections of multiple and unspecified sites (J06, J06.0, J06.8, J06.9), J11.- influenza, virus not identified (J11, J11.0, J11.1, J11.8), J12.- viral pneumonia, not elsewhere classified (J12, J12.8, J12.9), J18.- pneumonia, organism unspecified (J18, J18.0, J18.8, J18.9), J20.- acute bronchitis (J20, J20.8, J20.9), J21.- acute bronchiolitis (J21, J21.8, J21.9), J22 unspecified ALRI, and B34.9 unspecified viral infection, respectively.2 The sensitivities and specificities were calculated with 95% confidence interval (95%-CI). Additionally, we compared RSV-ICD-cases with confirmed-RSV-cases of the identified practices by calendar week.
We used Stata (version 15) and Microsoft Excel 2010 for the data analysis.

3 | RESULTS

3.1 | Primary care

3.1.1 | Descriptive analysis of RSV-ICD-cases based on SEEDARE data

A total of 1165 RSV-ICD-cases were identified from the SEEDARE database from week 40/2007-13/2017. Among those, 338 (29%) were diagnosed with J12.1, 432 (37%) with J20.5, and 395 (34%) with J21.0. The proportion of RSV-ICD-cases among all ARI-ICD-cases was 0.1%.

About two-thirds (765; 66%) of RSV-ICD-cases were children aged <2 years. The number of RSV-ICD-cases declined rapidly from 2 years of age and remained at a constantly low level from 5 years of age onwards. Under 2 years of age, the number of RSV-ICD-cases was higher in boys (423) than in girls (339; Figure 1).

The cumulative number (88) of the RSV-ICD-cases within the observed 10-year period peaked in the 8th calendar week, and the proportion (0.3%) of RSV-ICD-cases among all ARI-ICD-cases peaked in the 2nd calendar week. The RSV season on average was from 41st to 16th calendar week with the season length of 28 weeks. Within the RSV seasons, 92% (1075) RSV-ICD-cases were captured.

3.1.2 | Descriptive analysis of confirmed-RSV-cases based on virological surveillance data

From week 40/2010-18/2017, 1785 (8%) respiratory specimens of ARI or influenza-like illness (ILI) patients were RSV positive.

The highest RSV positive rate (25%; 659) was among children aged <2 years. The RSV positive rate decreased from 2 years of age, reached the lowest level in the age group 15-34 years (2%; 98), then increased slightly again, and reached 6% (145) at the age of 60 years and older. Under 2 years of age, the RSV-positive rate was higher among boys (25%; 378) than girls (24%; 270; Figure 2).

The cumulative number (143) of confirmed-RSV-cases peaked in the 6th calendar week, and the RSV positive rate (18%) peaked in the 52nd calendar week. The RSV season on average was from 48th to 15th calendar week with the season length of 20 weeks. Within the RSV seasons, 94% (1671) confirmed-RSV-cases were captured.
3.2 | Integration of RSV data of practices participated in SEED\textsuperscript{ARE} and virological surveillance

Forty-eight sentinel practices participated in both SEED\textsuperscript{ARE} and the virological surveillance from week 40/2010-13/2017. In total, 5589 respiratory specimens of the 48 practices were tested for RSV. Of those, 400 (7%) were RSV positive, and 2624 (47%) could be matched with the medical consultations based on SEED\textsuperscript{ARE} one to one (Figure 3).

Overall, the sensitivity of RSV-specific ICD-10 code diagnosis was 6% (95%-CI: 3%-12%), and the specificity was 99.8% (95%-CI: 99.6%-99.9%). The sensitivity (16%, 95%-CI: 7%-29%) increased among children aged <5 years and during the RSV seasons based on the virological data (48th-15th calendar week), and the specificity (99.5%, 95%-CI: 97.5%-99.9%) remained high. In children aged <5 years and in RSV seasons, the sensitivities of RSV-specific ICD-10 codes combined with general ALRI ICD-10 codes J18.-, J20.-, and with J12.-, J18.-, J20.-, J21.-, J22 both reached 44% (95%-CI: 30%-59%), and the specificities of the two combinations were still at a high level (91%, 95%-CI: 86%-94%; 90%, 95%-CI: 85%-94%). The sensitivity of RSV-specific ICD-10 codes combined with all general ARI ICD-10 codes was 90% (95%-CI: 78%-97%), whereas the specificity was 16% (95%-CI: 11%-21%; Table 1).

Figures 4 and 5 indicate number and proportion of RSV-ICD-cases based on the SEED\textsuperscript{ARE} and confirmed-RSV-cases based on the virological surveillance in the 48 practices by calendar week, respectively. The trends of the curves were similar.

3.3 | Secondary care

3.3.1 | Descriptive analysis of RSV-ICD-cases based on ICOSARI data

Among 1 417 700 respiratory disease, hospitalizations from week 01/2009-15/2017, 7345 (0.5%) were hospitalizations with any of the RSV-specific ICD-10 codes as primary or secondary discharge diagnosis, and 3154 (0.2%) as admission diagnosis. Of the 7345 RSV hospitalizations, 6918 (94%) were with RSV-specific ICD-10 codes as primary discharge diagnosis. Of the three RSV-specific ICD-10 codes, J21.0 was most frequently diagnosed as primary discharge (2705; 39%) and also admission diagnosis (1679; 53%).

Of the 6918 RSV-ICD-cases, 93% (6415) were children aged <2 years. The number of RSV-ICD-cases declined rapidly from 2 years of age. From 5 years of age, only a few RSV-ICD-cases were identified in each age group. In the age group 60 years and older, the number (32) of RSV-ICD-cases rose slightly. Under 2 years of age, number of RSV-ICD-cases was higher among boys (3623) than girls (2792; Figure 6).

The cumulative number (535) of RSV-ICD-cases peaked in the 5th calendar week, and the proportion (2.1%) peaked in the 52nd calendar week. The RSV season on average was from 48th to 16th calendar week with the season length of 21 weeks. Within the RSV seasons, 93% (6444) RSV-ICD-cases were captured.

4 | DISCUSSION

Using ICD-10-based surveillance, we identified age groups under high risk of RSV, and successfully described general trends and seasonality of RSV in primary and secondary care in Germany, as confirmed by data from the virological surveillance system. In primary care, RSV-specific ICD-10 codes had poor sensitivity and high specificity for the identification of laboratory-confirmed RSV infections. In young children, two combinations of RSV-specific ICD-10 codes with general ALRI ICD-10 codes increased the sensitivity without decreasing the specificity much.

The described RSV epidemiology based on ICD-10 code and virological data showed many common findings. Especially, high number of RSV cases among young children, and higher number of RSV cases among young boys than young girls were found in ICD-10 and...
Bronchiolitis is a very severe manifestation of RSV disease in young children and is a major cause of hospital admission in this group. It is often diagnosed in secondary care based on ICOSARI and J20.5 (acute bronchitis due to RSV) in primary care based on SEEDARE.

TABLE 1 Sensitivities and specificities of RSV-specific ICD-10 code diagnosis combined with different general ARI ICD-10 codes of the practices participated in both SEEDARE and virological surveillance, week 40/2010-13/2017

| RSV codes | Sensitivity % | 95%-CI | Specificity % | 95%-CI |
|-----------|---------------|--------|---------------|--------|
| <2 y of age RSV codes | 6 | 3.1 | 99.8 | 99.6-99.9 |
| <5 y of age RSV codes | 8 | 2.2 | 99.4 | 95.6-99.9 |
| In RSV seasons | 14 | 6.2 | 99.6 | 98.9-99.9 |

In RSV seasons, the proportion of young children among all RSV codes + general ARI codes was higher in secondary care based on ICOSARI than in primary care based on SEEDARE. This is in agreement with the explanations above (unpublished data).

In children aged <5 years and in RSV seasons, the sensitivity of RSV-specific ICD-10 codes grew more than twofold, and the specificity remained high. Physicians were probably more likely to encode RSV-specific ICD-10 codes than general ARI ICD-10 codes when testing is performed, an ICD-10 code diagnosis will probably no longer be recoded when laboratory findings are only available in the practice a few days later after the medical consultation. Therefore, suspected and also laboratory-confirmed RSV infections may be encoded with general ARI ICD-10 codes. These could be the reasons why most of the laboratory-confirmed RSV cases were not encoded with RSV-specific ICD-10 codes in the sentinel practices which participated in both SEEDARE and virological surveillance in the present study. In preparation for the present study, the RKI performed a survey to explore RSV coding behavior in primary care in Germany. The results of the survey are in line with the explanations above (unpublished data).

In children aged <5 years and in RSV seasons, the sensitivity of RSV-specific ICD-10 codes grew more than twofold, and the specificity remained high. Physicians were probably more likely to encode with RSV-specific ICD-10 codes for young children and in RSV seasons since RSV is more common in this group and during this time period. In the present study, we tried estimating the sensitivities and specificities of RSV-specific ICD-10 codes combined with different general ARI ICD-10 codes. RSV-specific ICD-10 codes combined with two groups of general ALRI ICD-10 codes achieved moderate sensitives and high specificities. The high sensitivity of RSV-specific
FIGURE 4 Cumulative number of RSV-ICD-cases based on SEED\textsuperscript{ARE} and cumulative number of confirmed RSV-cases based on virological surveillance by calendar week in the practices participated in both SEED\textsuperscript{ARE} and virological surveillance, week 40/2010-18/2017

FIGURE 5 Proportion of RSV-ICD-cases based on SEED\textsuperscript{ARE} and RSV positive rate based on virological surveillance by calendar week in the practices participated in both SEED\textsuperscript{ARE} and virological surveillance, week 40/2010-18/2017

FIGURE 6 Number of RSV-ICD-cases by age group and gender based on ICOSARI, week 01/2009-15/2017
ICD-10 codes combined with all general ARI ICD-10 codes suggests that in addition to RSV-specific ICD-10 codes, most laboratory-confirmed RSV infections were diagnosed with general ARI ICD-10 codes. Thus, the misclassification related to inaccurate labeling of RSV infections with other disease-specific or pathogen-specific ICD-10 codes was uncommon in the present study.

The present study has some limitations. The sensitivity and specificity of RSV-specific ICD-10 code diagnoses in secondary care could not be evaluated on a case by case basis since virological data of the ICOSARI network were not available for the present study. However, in the ICOSARI network, suspected RSV cases in young children were tested by rapid antigen detection tests and rtRT-PCR, and laboratory-confirmed RSV infections were encoded with RSV-specific ICD-10 codes. Although whether the testing and coding took place in a 100% frequency is not verified, these have been as a standard procedure in the pediatric units and the coding quality could have increased in recent years (personal communication). In addition, high validity has been reported in the literature for RSV-specific ICD-10 codes for the identification of hospitalized RSV among children.²⁹

The RSV coding behavior of physicians in primary care may vary during and out of RSV season, based on use of laboratory diagnostics, age of patient, and level of coding awareness. The differences in coding behavior may lead to information bias. The number of confirmed-RSV-cases and RSV-ICD-cases increased slightly among older adults based on virological as well as ICOSARI data, and it remained at a low level based on SEEDARE. The RSV infection normally goes unrecognized with milder symptoms among adults; however, it is a common pathogen of ARI in older adults and can lead to severe disease.²⁹,³⁰ Therefore, the RSV infections were probably underestimated among older adults in SEEDARE. This could be another limitation. However, the evaluation of the accuracy of ICD-10 codes was exactly the objective of the present study due to the potential information bias.

The present study was based on anonymized data. According to practice-ID, age, gender, consultation date, and sampling date alone, more than half of the virological samples could not be matched to medical consultations one to one and were excluded for the evaluation of sensitivity and specificity of RSV-specific ICD-10 codes which might lead to selection bias. However, the probability of the selection bias was low since no conspicuous deviations were found between the matched and the excluded virological data (data not shown).

5 | CONCLUSIONS

The use of RSV-specific ICD-10 code data may be a useful indicator to identify age groups under high risk of RSV, to monitor general trends, and to observe seasonality of RSV. The RSV epidemiology based on ICD-10 code data from different data sources and virological data showed similar age and sex distribution, percent positivity, and seasonality patterns. Therefore, RSV-specific ICD-10 codes are appropriate for RSV surveillance. However, in primary care, RSV-specific ICD-10 code diagnosis was less sensitive, and relying on RSV-specific ICD-10 codes alone will underestimate the actual number of RSV infections. RSV-specific ICD-10 codes combined with the general ALRI ICD-10 codes J18.−, J20.−, and with J12.−, J18.−, J20.−, J21.−, J22 achieved moderate sensitivities and high specificities, respectively. Thus, when establishing an ICD-10-based digital RSV surveillance system in young children, an extended ICD-10-based RSV case definition using the two combinations of ICD-10 codes seems to better capture the true RSV disease burden. Further investigations are required to validate the use of the two combinations of ICD-10 codes in RSV surveillance systems in other countries as the RSV coding behavior may differ in different countries, to find out an even better combination of ICD-10 codes for the identification of RSV infections in primary care, and to evaluate the use of RSV-specific ICD-10 codes in secondary care.

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CONFLICT OF INTEREST

None.

AUTHOR CONTRIBUTIONS

Wei Cai, Silke Buda, Walter Haas, Siddhivinayak Hirve, and Wenqing Zhang were involved in designing the study. Kristin Tolkdorf, Silke Buda, and Ekkehard Schuler participated in the collection of ICOSARI data. Wei Cai analyzed the data, and Kristin Tolkdorf helped analyze the data. Wei Cai drafted the manuscript. All authors reviewed and approved the final manuscript.

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SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section at the end of the article.

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