Positive predictive value of the ICD-10 hospital diagnosis of pleural empyema in the Danish National Registry of Patients

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Objective: Health care databases are a valuable source for epidemiological research in respiratory diseases if diagnoses are valid. We validated the International Classification of Diseases, 10th revision (ICD-10) diagnosis of pleural empyema in the Danish National Registry of Patients (DNRP).

Methods: We randomly selected hospitalized patients registered in the DNRP with a discharge diagnosis of pleural empyema between 1995 and 2009 in the North Denmark Region. We retrieved and reviewed medical records and estimated the positive predictive value (PPV) of the empyema diagnosis. Analyses were stratified by study period, hospital type (referral versus district), department type (pulmonary medicine or thoracic surgery versus other), cause of empyema (medical, surgical, or traumatic), and age group. To assess changes over time, we included chi-square tests for linear trend.

Results: We retrieved the medical records of 224/225 sampled patients with empyema (99.6%). Of those, 182 were classified as being definite cases, and 21 were probable cases, yielding a PPV of 90.6% (95% confidence interval [CI]: 86.0–94.1). The PPV decreased from 95.7% in patients aged 15–39 years to 87.5% in patients aged 80 years and over but was uniformly high regardless of study period, hospital or department type, or cause of empyema.

Conclusion: Our finding of a high overall PPV indicated good agreement between ICD-10 codes for pleural empyema and medical records. Registry-based discharge codes may be a suitable source of data on pleural empyema for epidemiological research.

Keywords: pneumonia, positive predictive value, validation, International Classification of Disease Codes, empyema

Introduction
Pleural empyema is a serious infection characterized by the accumulation of pus in the pleural space.1 In the United States, 65,000 patients suffer from empyema each year, with 30-day mortality up to 15% and an estimated hospital cost of $500 million due to prolonged hospitalizations and multiple invasive procedures.2–5 The empyema usually develops as a complication of pneumonia (40%–60% of all empyemas), thoracic surgery (20%), or trauma (4%–10%).6 Less commonly, empyema can develop as a result of esophageal rupture and subdiaphragmatic spread.

Many of the studies addressing risk factors and outcome of empyema rely on hospital discharge diagnosis codes recorded in administrative health care databases.7–9 Use of these databases for epidemiological research can be a cost-effective way of collecting data on empyema.10–12 The large size of many databases offers the potential for precise estimates of effect and for studying rare exposures or outcomes.
At the same time, the risk of some biases, such as recall bias and nonresponse, is reduced. On the other hand, the researchers have no control of data selection and quality, and the usability of these databases is therefore highly reliant on the validity of the registered data. 10-12 Thus, to draw valid inferences from database studies of pleural empyema, the validity of the diagnosis needs to be quantified. 11,13

To our knowledge, no study has validated pleural empyema diagnosis in hospital registries. We therefore examined the positive predictive value (PPV) of empyema discharge diagnosis in the population-based Danish National Patient Registry (DNRP) using data from medical record reviews as the reference. In addition, we determined whether the validity of the empyema diagnosis varied by calendar period, hospital or department type, cause of empyema, or patient age.

Materials and methods
Denmark has 5.3 million inhabitants, and the National Health Service provides universal tax-supported health care. Since 1968, a unique 10-digit civil registry number (CPR number) has been assigned to all Danish residents at birth or emigration by the Danish Civil Registry System. 14 In all Danish registries, patients are identified by their CPR-number. We therefore linked national data on hospital diagnoses and medical records using the CPR number and identified patients diagnosed with pleural empyema between 1995 and 2009 residing in the North Denmark Region (with a population of 579,741 inhabitants or 10% of the population of Denmark in 200915).

The DNRP
The DNRP tracks each hospital admission in Denmark since 1977, recording dates of admission and discharge, and up to 20 discharge diagnoses. The registry covers 99.4% of all discharge records from Danish hospitals. 16 Diagnoses are coded by physicians using the Danish version of the International Classification of Diseases, 8th revision (ICD-8) (1977–1993) and 10th revision (ICD-10) (1994 onward). Since 1995, outpatient contacts and visits to the emergency room have been recorded in addition to the hospital stays.

Identification of coded pleural empyema cases
Using the DNRP, we identified all discharges between 1995 and 2009 associated with a primary or secondary diagnosis of empyema (ICD-10 codes J86.0 Pyothorax with fistula and J86.9 Pyothorax without fistula). We did not evaluate the coding of tuberculous empyema (ICD-10 code A15.6) because we consider these cases to constitute a distinct disease group that is categorized separately in the ICD-10 system; that is, under A15 “Respiratory tuberculosis, bacteriologically and histologically confirmed”. 17 Because we were interested in first incident episodes of empyema, we removed all hospitalizations for any recurrent episode of empyema during 1995–2009, and patients who had been hospitalized for empyema between 1977 and 1994 (ICD-8 code 510 Empyema pleurae). We also removed pediatric patients <15 years of age.

Medical record validation
To validate the diagnoses of empyema identified in the DNRP, we randomly selected 15 patients in each calendar year using the SAS RANUNI function (SAS Institute, Cary, NC). We then used CPR numbers to identify the corresponding medical records. We confined the review to the North Denmark Region, as data quality is considered uniform in the hospital discharge registries. 13 The records were reviewed by one of two reviewers (MS or JBK), who assessed all the available information, including written radiology reports and results from laboratory tests. The actual imaging films were not reinterpreted, and the available information was in most cases identical with the information available to the physician at the time of discharge. All cases with an uncertain diagnosis based on the available information were discussed, and diagnoses were made according to consensus.

Based on the available information, we classified the empyema cases as either definite, probable, or no evidence of empyema. Definite empyema was defined by frank pus aspirated from the pleural space, a positive Gram stain/subculture for pathogenic microorganisms in the pleural fluid, and/or an autopsy diagnosis of empyema. 1,6,18,19 Patients with clinical symptoms suggestive of empyema in association with compatible radiographic features (eg, pleural thickening, loculated and/or septated pleural effusions 6,20) who did not meet the criteria for definite empyema were classified as having probable empyema. We ruled out empyema when results of diagnostic tests were negative, or if no tests were performed because symptoms had disappeared, or were explained by another disease. We further categorized the most probable cause of the patients’ empyemas as: previous surgery (patients who had an operation involving the thoracic cavity before their empyema), trauma, esophageal rupture, or any medical cause (patients who did not have thoracic surgery, trauma, or esophageal rupture and were presumed to have a medical cause for their empyema, most frequently pneumonia).

Statistical analysis
Our study outcome was the PPV of an empyema diagnosis, defined as the proportion of patients with an empyema
diagnosis in the DNRP who had definite or probable empyema according to their medical record. We calculated the PPV with 95% confidence intervals (CIs). The PPV was calculated for the whole study population and stratified by calendar period (1995–1999, 2000–2004, and 2005–2009). This stratification was done to examine whether the validity differed over time and included a chi-square test for linear trend. In addition, PPVs were calculated according to hospital type (referral versus district), department type (pulmonary medicine or thoracic surgery versus other departments), cause of empyema, and age group (15–39, 40–64, 65–79, 80 years and older).

We analyzed the data with Stata Software (v. 11.1; Stata Corp. College Station, TX). The study was approved by the Danish Data Protection Agency (Journal no. 2010-41-4969).

**Results**

**Description of study population**

From 509 adult patients in Northern Denmark registered with empyema in the DNRP between 1995 and 2009, we randomly selected 225 cases for this validation study. We excluded one patient who was transferred to a hospital outside the region with a suspected empyema. Of the remaining 224 patients with available medical records, we classified 182 (81.3%) as being definite empyema cases. Twenty-one patients did not meet the criteria for definite empyema and were classified as probable cases. Among the remaining 21 patients whose empyema diagnosis was rejected, 19 patients had other pulmonary disorders; one had a pulmonary abscess, one had pulmonary tuberculosis, one had sarcoidosis, two had emphysema, eight had pneumonia, one had chylothorax, one had multiple rib fractures, one had chronic sequelae after previous pneumonia, and three had pulmonary cancer. The two remaining patients had severe muscle dystrophia and surgical site infection, respectively. Table 1 shows demographic and diagnostic characteristics of patients registered in the DNRP according to whether they had definite, probable, or no empyema.

![Table 1](image)

**Validation of empyema diagnosis**

The overall PPV including definite and probable empyema hospitalizations was 90.6% (95% CI: 86.0–94.1) and 81.3% (95% CI: 81.3–86.1) when focusing only on definite cases. PPVs of empyema diagnosed in 1995–1999, 2000–2004, and 2005–2009 were 90.7% (95% CI: 81.7–96.2), 94.6% (95% CI: 86.7–98.5), and 86.7% (95% CI: 76.8–93.4), respectively, indicating that no major changes in empyema coding validity occurred over the 15-year study period (P-value for trend = 0.40). The PPVs in different strata of calendar period, hospital type, department type, and cause of empyema showed only slight differences from the overall PPV. However, the PPV appeared to decrease with increasing age; that is, from 95.7% (95% CI: 78.1–99.9) in patients aged 15–39 years to 87.5% (95% CI: 67.6–97.3) in patients aged 80 years and over (P-value for trend < 0.001) (Table 2).
Discussion

Our finding of a high overall PPV indicated good agreement between ICD-10 codes for pleural empyema and medical records. In addition, we detected little variation in the PPV by calendar period, hospital type, department type, or cause. However, PPV was slightly higher in younger age groups than in elderly patients.

Although we examined only a segment of the Danish national population, the results can be assumed to be generalizable, thanks to the uniform structure of record keeping and the universal nature of the health care system across Denmark. However, a number of limitations need to be considered in the interpretation of the results. First, we only examined one dimension of the validity: the predictive value of a positive registration of empyema. The optimal validation study would also examine sensitivity and specificity, but since no other data source with empyema exists, only the predictive value could be estimated. Second, most of the medical records were evaluated by one reviewer. However, any cases with an uncertain diagnosis were evaluated by both reviewers independently. Thereafter, final diagnoses were made according to consensus. The reviewers were not blinded to the registered discharge diagnosis codes, and blinding is usually recommended in diagnostic studies. However, since the result of the “test under study” (the register diagnosis) was by definition given for all the examined patients (as only patients who had a diagnosis were included), it is unlikely that lack of blinding of the reviewers would have influenced the PPV in general. Third, pleural effusions represent a spectrum of inflammatory fluid collections that ranges from transudative effusion to empyema, and the differentiation of this spectrum into patients with empyema and those without is somewhat arbitrary. Studies have shown that neither radiography, ultrasound, nor computed tomography can reliably identify the stage of pleural disease in adult patients with effusions and empyema. Although we carefully evaluated the reported clinical and radiographic findings, we may therefore have overestimated the PPV among patients with probable empyema. Finally, we do not know to what extent our findings can be extrapolated to other places, where the prevalence of empyema and the diagnostic strategy may be different.

Whether the data quality documented in our study is sufficient for registry-based studies depends on the proposed research questions and the study design used. If databases such as the DNRP are used to assess changes in incidence of empyema over time, the PPV must remain stable over time to obtain valid estimates. We found a slight variation in PPV by study period but there was not a significant linear trend over the 15-year period of our study. Thus, the increasing availability of advanced diagnostic tools, including computed tomography and magnetic resonance imaging, between 1995 and 2009 apparently did not improve the quality of diagnostic coding. We do not know whether there might have been any substantial time trends in the data quality before 1995. In analytical studies, inaccurate or missing data tend to bias the association toward the null hypothesis rather than to cause spurious associations.

Table 2 Validity of pleural empyema diagnosis

| Total No. of cases | No. definite cases | No. probable cases | PPV (%) | 95% CI |
|--------------------|-------------------|-------------------|---------|-------|
| All                | 224               | 182               | 21      | 90.6  | 86.0–94.1 |
| Period             |                   |                   |         |       |         |
| 1995–1999          | 75                | 63                | 5       | 90.7  | 81.7–96.2 |
| 2000–2004          | 74                | 65                | 5       | 94.6  | 86.7–98.5 |
| 2005–2009          | 75                | 54                | 11      | 86.7  | 76.8–93.4 |
| Hospital type      |                   |                   |         |       |         |
| Referral hospital  | 139               | 117               | 10      | 91.4  | 85.4–95.5 |
| District hospital  | 85                | 65                | 11      | 89.4  | 80.8–95.0 |
| Department type    |                   |                   |         |       |         |
| Pulmonary medicine or thoracic surgery | 101 | 85 | 7 | 91.1 | 83.8–95.8 |
| Other              | 123               | 97                | 14      | 90.2  | 83.6–94.9 |
| Cause of empyema   |                   |                   |         |       |         |
| Medical            | 179               | 144               | 19      | 91.1  | 85.9–94.8 |
| Surgical incl. esophageal rupture | 33 | 27 | 2 | 87.9 | 77.8–96.6 |
| Trauma             | 12                | 11                | 0       | 91.7  | 61.5–99.8 |
| Age group (years)  |                   |                   |         |       |         |
| 15–39              | 23                | 19                | 3       | 95.7  | 78.1–99.9 |
| 40–64              | 95                | 76                | 11      | 91.6  | 84.1–96.3 |
| 65–79              | 82                | 69                | 4       | 89.2  | 80.2–94.9 |
| 80 and older       | 24                | 18                | 3       | 87.5  | 67.6–97.3 |

Abbreviations: PPV, positive predictive value; CI, confidence interval.
as long as the sensitivity and specificity are similar in the compared groups. At the same time, if empyema is used as the outcome of interest, imperfect sensitivity, no matter how low, is not expected to bias relative measures of association as long as the specificity is 100%. We have shown that empyema cases that are registered largely represent true instances of this event. If the misclassification of empyema is associated with the examined exposure of interest, however, the risk estimates may be biased, both toward and away from the null, depending on the direction of the association.

Denmark and a few other places in the world are privileged by having a broad network of population-based registries, which enables use of routinely collected data to rapidly assemble large cohorts for epidemiological studies, including that of pleural empyema. The high PPV estimated in this study indicates a low false-positive number of empyema cases in the DNRP, suggesting that the DNRP is a valid source for identifying cases of empyema for epidemiological research.

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Disclosure
The authors report no conflicts of interest in this work.

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