



Methodological paper

Hospital Readmission Ratio

Methodological report of the 2017 model

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1. Introduction

1.1 Indicators of quality of hospital care

Overall quality of hospital care can be estimated using several types of quality indicators based on hospital admission data. Such indicators for identifying potentially suboptimal quality of hospital care might focus for example on unexpected in-hospital or post-discharge mortality, potentially preventable hospital readmissions or unexpected long duration of admissions. In the Netherlands, hospital admission and discharge data is registered in the LBZ, a national hospital discharge register covering all general, university and three specialised hospitals. Other specialised clinics, independent treatment centres and private clinics are not included. Inpatients as well as day cases and prolonged observations without overnight stay are registered. For each hospital discharge administrative data of the admission are registered, as well as diagnoses and procedures.

In the Netherlands, hospitals participating in the LBZ registration are annually provided by Dutch Hospital Data (DHD) with a set of indicators based on their performance in the previous year. Up to 2016 this set included the (unadjusted) hospital readmission rate, which is the ratio of the number of observed readmissions to the total number of hospital admissions. However, since this ratio does not correct for case-mix differences, it might be less indicative of differences in the true number of potentially preventable readmissions. Therefore, in 2017 DHD has asked Statistics Netherlands to develop a model to estimate the expected readmission risks adjusted for relevant covariates, in a fashion similar to the estimation of the hospital standardized mortality rates (HSMR). From 2017 onwards, Statistics Netherlands produces Hospital Readmission ratio models on a yearly basis.

1.2 Predictive value of the hospital readmission model

Internationally, models for estimating hospital readmission rates are used for the purpose of risk stratification but also as a quality indicator. From previous studies it is known that several patient characteristics can contribute to the risk to be readmitted to the hospital. In a systematic review by Kansagara *et al.* (2011), an overview is presented of the various validated models that have been used internationally, the covariates included in those models and their overall predictive value. Common covariates include comorbidity indexes, age, sex and/or prior use of medical services (hospitalizations). Regardless of the number of included covariates, the results of only a small fraction of the models are moderately discriminative (AUC/C-statistic > 0.70). The model developed by Statistics Netherlands includes additional covariates such as severity of the main diagnosis, urgency of the admission and socio-economic status. However, the overall predictive value of the model did not exceed previously published values (AUC = 0.69). It was demonstrated though, that the level of case mix correction applied by the model significantly improved comparability of the outcomes of the individual hospitals. So, although the case mix correction is probably incomplete, it does, to some extent, reduce effects due to differences in patient populations. As such, applying the model to calculate adjusted readmission ratios for individual hospitals is an improvement over calculating crude rates.

1.3 Aim of the current project

In 2018 Statistics Netherlands has published two models (referred to as the '2016 models'; Van der Laan *et al.* 2018a). The initial model, developed in 2017, was based on the linkage of

admissions and readmissions that occurred within the same hospital (intra-hospital readmissions). To improve that model, planned transfers to and from neighbouring or specialized hospitals were identified and excluded from the intra-hospital readmissions model. It is common practice for hospitals to refer inpatients to other hospitals for specific procedures, such as coronary interventions. Such planned transfers should not be labelled as readmissions. The results of this improved intra-hospital model were compared to that of a newly developed inter-hospital model, that also took into account readmissions in other hospitals, while excluding planned transfers. Since readmissions can also take place in other hospitals, including inter-hospital readmissions in the model might improve its predictive value.

The predictive value of both 2016 models was however largely comparable, and it was concluded that apart from views regarding the relevance of inter-hospital readmissions for measuring quality of care, practical considerations might determine which of both models will be used for calculating the readmission ratios of the individual hospitals. A practical disadvantage of the inter-hospital ratio is that hospitals need patient information from other hospitals to calculate the ratios and to study the files of the patients with readmissions. For this reason DHD decided to use the intra-hospital model in their regular hospital indicators reports.

In the current project we produced an updated version of the intra-hospital model ('2017 model'), excluding planned transfers, based on LBZ data of 2016 and 2017. The outcome is described in chapter 3.

1.4 Output

Statistics Netherlands has only calculated the model for the hospital readmission risks on the basis of LBZ data of 2016-2017, not the outcomes for the individual hospitals. For their regular hospital indicators reports, DHD will use this model to estimate the expected readmission risk, adjusted for relevant covariates, for each individual primary (index) hospital admission in 2018. For each hospital the standardized (adjusted) readmission ratio can be calculated as the observed number of readmissions (x 100) divided by the sum of the expected readmission risks of the index admissions of that hospital.

2. Methods

2.1 Changes compared to the previous intra-hospital model

In the current project we used the methods of the 2016 intra-hospital model which excludes transfers (admissions starting on the same day as the discharge date of the preceding admission) as readmissions, developed in 2018 (Van der Laan *et al.* 2018a). These methods were, without changes, applied for the model of 2017, using LBZ data of 2016-2017. The methods used in the model are described in more detail in the next paragraphs.

2.2 Readmission ratio

The (hospital) readmission ratio is calculated using the expected (hospital) readmission risk as the denominator and observed readmission as the numerator. The expected readmission risk is predicted for each individual admission within a given period, adjusted for patient and admission characteristics of that admission as covariates. Readmission risk was predicted for all (index) admissions that potentially could be followed by a readmission, excluding admissions for diagnoses with complex care paths where planned readmissions are often involved.

Readmissions are defined as those admissions that occurred within 30 days of the discharge date of the preceding index admission. Detailed information on the characteristics and criteria of index admissions and readmissions is given in paragraphs 0 and 2.3.6 respectively.

Expected readmission risk is determined for each of the included diagnosis groups, which are based on the CCS (*Clinical Classifications Software*), which clusters ICD codes of the main diagnoses of the admissions into 259 clinically meaningful categories¹. In accordance with the HSMR, we further clustered these groups into 157 diagnosis groups, which are partly the same clusters used for the SHMI (Summary Hospital-level Mortality Indicator) in the UK (HSCIC, 2016). To determine readmission risk we used logistic regression models, with an observed readmission as the target (dependent) variable and various variables available in the LBZ as covariates.

The methodology for estimating the expected readmission risk is very similar to that used for estimating expected mortality rates applied for calculating the HSMR rates, described in detail elsewhere (Van der Laan *et al.* 2018b). In the following section we therefore briefly describe the applied methods, while deviations from the HSMR methodology or other methods specific to the current project are described in more detail.

2.3 Target population and data set

2.3.1 Patient identifier

Statistics Netherlands has linked the LBZ data to the Dutch national population register, using a pseudonym of the national personal identification number, and the combination of date of birth, sex and postal code as linkage keys. Through this linkage a unique pseudonymised person ID could be added to the LBZ dataset. With this linkage >99% of all admissions could be uniquely linked to a person in the population register; thus the loss of data was minimal (<1%). Using this identifier not only allows identification of transfers to other hospitals, it also eliminates bias due to administrative errors in hospital-specific patient numbers.

¹ See <https://www.cbs.nl/-/media/excel/2018/40/classification%20of%20variables%20hsmr%202017.xlsx>

2.3.2 Admissions – general criteria

We consider both the population of hospitals and the population of admissions. Our population of (re)admissions consists of “all hospital stays (inpatient admissions) of Dutch residents in Dutch short-stay hospitals within the study period”. Only completely registered admissions with a registered main diagnosis were included. In the LBZ, the date of discharge, and not the day of admission, determines the LBZ year a record is assigned to. Therefore, the registered hospital stays of year t comprise all inpatient admissions that ended in year t . Day cases and prolonged observations were excluded, since subsequent readmissions might be elective, for example, for prolonged treatment.

In addition, admissions of foreigners were excluded from the model, since readmissions might have also taken place in a hospital in their residential country. Furthermore, foreigners cannot be linked to the Dutch population register. The number of admissions of foreigners is relatively small.

Lastly, duplicate admissions were removed. This included admissions with identical values for date and time of admission and of discharge in combination with identical values for either (1) hospital ID and hospital-specific patient ID or (2) the pseudonymised person ID. In case of duplicate admissions, the admission with the lowest LBZ registration number was removed and the one with the highest LBZ registration number was kept, since we assumed that the latter admission might have been registered as a corrected version of the first. Duplicate admissions rarely occur in the LBZ.

2.3.3 Hospitals

Hospitals report admission data (hospital stay data) in the LBZ. However, not all hospitals participate in the LBZ. In principle, the hospital readmission risk model includes all general hospitals, all university hospitals and short-stay specialised hospitals with inpatient admissions participating in the LBZ in the study period.

One of the short-stay specialised hospitals was excluded since it mostly treats patients with oncological diseases, which are excluded from the data (see paragraph 0). In addition, one of the general hospitals stopped operating on October 1st 2017. Therefore, admissions with a discharge date in August and September in that hospital were only considered as potential readmissions and not as index admissions, since those admissions could not logically be followed by readmissions in the same hospital after October 1st.

The readmission ratio is calculated using LBZ data on admissions, using the pseudonymised personal ID as the unique key for identifying (re)admissions. The combination of the person ID (for identifying patients) and the hospital ID number (for identifying the same hospital) was used for linking admissions. In case of merging hospitals, the hospital ID number the hospital used in the LBZ registration year (based on the discharge dates of the admissions), was used for the associated study period in the models. For example, two hospitals that had merged in 2017 were analysed as separate units for study period 2016 and as a single unit for study period 2017. If the hospitals would have been analysed as a single merged hospital for both years, transfers between both hospitals could not be identified and excluded in 2016 and would be labelled as readmissions, thus negatively influencing their readmission rates.

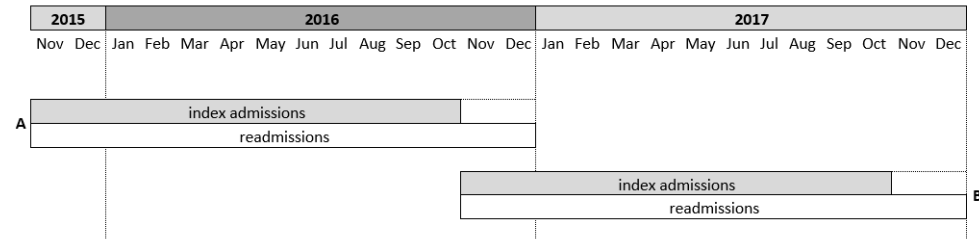
2.3.4 Study periods

For the calculation of the current model, LBZ data of 2016 and 2017 was used. Previously we showed that to identify the highest percentage of readmissions ending in year t, using index admissions with a discharge date from November 1st of year t-1 up to October 31st of year t (study period) is optimal (Van der Laan *et al.* 2017a). Thus, for study period 2016 ('year'=2016 in the model) we selected index admissions with a discharge date from November 1st 2015 up to October 31st 2016 and for study period 2017 we selected index admissions with a discharge date from November 1st 2016 up to October 31st 2017 ('year'= 2017 in the model). The occurrence of readmissions was analysed in the period between November 1st 2015 up to December 31st 2017.

Using these study periods, admissions with a discharge date between November 1st 2016 and December 31st 2016 can potentially be linked to index admissions from either 2016 or 2017. However, hospital ID numbers could have changed between 2016 and 2017 due to merging and previously we had decided to use the hospital ID number the hospitals had used in the year of the actual registration of data. As a result, admissions from the above-mentioned two-month period should potentially be analysed using old and new hospital IDs and should be added twice to the dataset with different hospital IDs.

To avoid this issue, data from both study periods were processed separately for the identification of index admissions, transfers and readmissions (see figure 2.3.4.1). After this processing, all index admissions of both study periods were combined into a single dataset that was entered into the model.

2.3.4.1 Study periods used for identifying index admissions and readmissions.



A=study period 2016 using hospital IDs from 2016 for all admissions; B=study period 2017 using hospital IDs from 2017 for all admissions

2.3.5 Criteria for index admissions

Expected readmission risk was only calculated for those inpatient admissions (meeting the general criteria for admissions, see 2.3.2) for which readmission was possible (i.e. patient did not die during the index admission), and excluding some specific diagnosis groups. These admissions are referred to as index admissions. Thus, in summary, the index admissions had to meet the following criteria:

- The patient did not die during the admission.
- The main diagnosis of the admission was not related to oncology (CCS groups 11-45) or psychiatry (CCS groups 65-75) since hospital care for these diagnoses is usually complex and follow-up care might be required. In addition the main diagnosis was not related to obstetrics (CCS groups 176-196; 218), since most deliveries do not take place during inpatient admissions, so it cannot be determined whether an admission for this purpose is the 'true' index admission.

- The date of discharge was from November 1st 2015 up to October 31st 2016 ('year'= 2016) or from November 1st 2016 up to October 31st 2017 ('year'= 2017).

2.3.6 Criteria for potential readmissions

Inpatient admissions only qualified as potential readmissions (meeting the general criteria for admissions, see 2.3.2) if the following criteria were matched:

- The main diagnosis of the admission was not related to oncology (CCS groups 11-45) or psychiatry (CCS groups 65-75) since hospital care for these diagnoses is usually complex and follow-up care might be required. In addition the main diagnosis was not related to obstetrics (CCS groups 176-196; 218), since most deliveries do not take place during inpatient admissions, so it cannot be determined whether an admission for this purpose is a "true" readmission.
- The main diagnosis of the admission was not related to social, socio-economic or psychosocial circumstances or administrative purposes (ICD10: Z55-Z65), other circumstances (ICD10: Z70-Z76) or screening, follow-up care or rehabilitation (CCS groups 254-258), since admissions for these purposes are usually planned.
- The discharge date of the admission was before or on December 31st of year t .
- The maximal time lapse between the admission date of the readmission and the discharge date of the index admission is 30 days (29 days interval at maximum). For example, when an index admission has a discharge date of January 1st, a subsequent admission on January 30th is classified as a readmission, while a subsequent admission on January 31st is not.
- If a readmission in the same hospital started on the same day as the discharge date of the index admission, the minimal time lapse between both admissions is one hour. If the hour of discharge of the index admission or the hour of admission of the readmission is unknown in this specific situation, the subsequent admission is not identified as a readmission.

Note that the main diagnosis of the readmission does not have to be related to the main diagnosis of the index admission.

2.3.7 Transfers

Transfers were not labelled as readmissions. Transfers are defined as admissions with a date of admission that was identical to the date of discharge of the previous admission in another hospital. In case of 'overlapping admissions' in two different hospitals (i.e. the start date of the second admission preceded the date of discharge in the first hospital) the second admission was also labelled as a transfer. Transfers affect the identification of readmissions in two ways:

First, when index admissions are followed by a transfer, these index admissions (by definition) cannot have a readmission. Although index admissions that are followed by a transfer cannot have readmissions, these index admissions were not removed from the model.

Second, transfers cannot be readmissions. In case of 'to and fro' transfers from hospital A to hospital B and back to hospital A, the latter admission in hospital A is *not* a readmission of the first admission in hospital A. In fact, an admission in hospital A that is a transfer from hospital B can (by definition) never be a readmission of any other previous admission.

The general criteria for admissions, the additional criteria for index admissions and readmissions and the role of transfers are summarised in table 2.3.7.1.

2.3.7.1 General criteria, additional criteria for index admissions and readmissions and the influence of transfers.

	Criteria for index admissions	Criteria for potential readmissions
General	<ul style="list-style-type: none"> - Inpatient admissions registered in the LBZ - Completely registered admissions with a registered main diagnosis - Admissions of Dutch residents 	<ul style="list-style-type: none"> - Inpatient admissions registered in the LBZ - Completely registered admissions with a registered main diagnosis - Admissions of Dutch residents
Follow-up	The patient did not die during the admission.	
Diagnosis	The main diagnosis of the admission was not related to oncology (CCS groups 11-45), psychiatry (CCS groups 65-75) or obstetrics (CCS groups 176-196; 218).	The main diagnosis of the admission was not related to oncology (CCS groups 11-45), psychiatry (CCS groups 65-75), obstetrics (CCS groups 176-196; 218), social, socio-economic or psychosocial circumstances or administrative purposes (ICD10: Z55-Z65), other circumstances (ICD10: Z70-Z76) or screening, follow-up care or rehabilitation (CCS groups 254-258).
Period	For year t in the model the date of discharge was from November 1 st year $t-1$ up to October 31 st year t ('year' = t).	The discharge date of the admission was before or on December 31 st of year t .
Maximal time lapse		The maximal time lapse between the admission date of the readmission and the discharge date of the index admission is 30 days (29 days interval at maximum)
Minimal time lapse		If the readmission started on the same day as the discharge date of the index admission, the minimal time lapse between both admissions is one hour
Influence of transfers¹	Index admissions followed by a transfer cannot have a readmission.	Transfers cannot be readmissions.

¹ A transfer is an admission in hospital B with a date of admission that is identical to the date of discharge of a previous admission in hospital A.

2.4 Target variable

The target variable for the regression analysis of the model is the occurrence of a readmission within 30 days of the discharge date of the preceding index admission.

The pseudonymised person ID (resulting after linkage of the LBZ to the national population register) was used as the unique key for identifying admissions of the same patient in a single hospital and for the identification of transfers to other hospitals.

The dataset was composed based on the criteria presented in section 2.3. According to the criteria for index admissions and readmissions, two variables were added to the dataset to mark both types of admissions. Readmissions can also count as index admissions in case they are followed by another readmission.

After that, the dataset was processed to allocate readmissions to index admissions: index admissions and potential readmissions of the same patient (person ID) are identified within the same hospital only. As was explained in section 2.3.4, this allocation is done for each year separately.

Within the set of admissions per patient, for each index admission the presence of a readmission within 30 days is determined. Each index admission can only be followed by a single subsequent readmission, and a single readmission can also be only allocated to a single index admission. If an index admission is followed by multiple potential readmissions within 30 days, only the first occurring readmission is marked as such. Based on this algorithm, for each index admission the presence of a readmission is marked.

Transfers are identified according to the method presented in section 2.3.7. After that, the previously described rules are applied ('an admission followed by a transfer cannot have a readmission' and 'a transfer cannot be a readmission'), with the result that some of the admissions are no longer regarded as readmissions. The index admissions associated with those readmissions were initially marked as having a readmission, but since these readmissions are no longer categorized as such after applying the transfer rules, the presence of a readmission is cleared from the respective index admissions.

Subsequently, all index admissions and the corresponding covariates are selected, plus the target variable (whether the primary admission was followed by a readmission or not) and these were entered into the model.

To illustrate the implementation of excluding transfers from the model, an example is given in table 2.4.1.

2.4.1 Example of the identification of readmissions after excluding transfers.

Admission	Hospital	Step 1	Step 2		Step 3
		Is the admission followed by a readmission?	Is the admission followed by a transfer?	Is the admission a transfer?	Is the admission followed by a readmission (after correction for transfers)?
A1	A	Yes (A2)	No	No	No, A2 is a transfer
<i><patient is home></i>					
B1	B	Yes (B2)	Yes (A2)	No	No, B2 is a transfer
A2	A	Yes (A3)	Yes (B2)	Yes (of B1)	No, A2 is followed by a transfer (B2)
B2	B	No	No	Yes (of A2)	No
<i><patient is home></i>					
A3	A	No	No	No	No

In this example a patient is admitted five times to two different hospitals within a period of 30 days. All admissions are index admissions, and admissions B1, A2 and B2 are consecutive admissions (date of admission of A2 is equal to date of discharge of B1; and date of admission of B2 is equal to date of discharge of A2). According to the criteria for readmissions, in step 1 the presence of readmissions is determined. After that, the presence of transfers is determined in step 2. Finally, the information of steps 1 and 2 is combined into step 3: the presence of readmissions corrected for transfers, where we apply the rules ‘an index admission followed by a transfer cannot have a readmission’ and ‘a transfer cannot be a readmission’.

For example, A2 is a possible readmission to A1, but since A2 is a transfer, it cannot be a readmission. As a result, A1 is not followed by a readmission. In addition, A3 is not a readmission of A2, since A2 is followed by a transfer (B2) and thus cannot have a readmission.

2.5 Stratification

Instead of performing one logistic regression for all admissions, we performed a separate logistic regression for each main diagnosis group. These sub-populations of index admissions are more homogeneous than the entire population. Hence, this stratification may improve the precision of the estimated readmission probabilities. As a result of the stratification, covariates are allowed to have different regression coefficients across diagnosis groups. Due to the exclusions of specific CCS groups for the index admissions, 35 of the 157 diagnosis groups (as used for the HSMR) are fully excluded. Therefore, the model included 122 separate logistic regressions, one for each diagnosis group selected (see Appendix II for the diagnosis groups included).

2.6 Covariates (explanatory variables or predictors of readmission risk)

By including covariates of patient and admission characteristics of the index admissions in the model, the hospital readmission risk is adjusted for these characteristics. For this purpose we selected the same covariates that are also regularly used in the (H)SMR model estimations, which are variables (available in the LBZ) known to be associated with in-hospital mortality. During the development of the readmission model, it was demonstrated that these covariates indeed contributed to the predictive value of the model (Van der Laan *et al.* 2017a).

The LBZ variables that are included in the model as covariates are age, sex, socio-economic status, severity of main diagnosis (based on mortality risk categories), urgency of admission, Charlson comorbidities, source of admission, month of admission and year. These variables are described below. Detailed information on these variables and their content is available in the HSMR methodology report (Van der Laan *et al.* 2018b). For the variables socio-economic status, severity of main diagnosis and source of admission the detailed classifications are presented in the file 'Classification of variables', published together with the methodology report of the HSMR (Van der Laan *et al.* 2018b). The variable 'year' is different from the variable used for the HSMR model, since it reflects the study period the index admission belongs to, rather than year of discharge. The specific (modified) definitions of 'year' for the readmission model are described in 2.3.4.

For the regressions, all categorical covariates are transformed into dummy variables (indicator variables), having scores 0 and 1. A patient scores 1 on a dummy variable if he/she belongs to the corresponding category, and 0 otherwise. As the dummy variables for a covariate are linearly dependent, one dummy variable is left out for each categorical covariate. The corresponding category is the so-called reference category. We used the first category of each covariate as the reference category.

Covariates:

- **Age at admission** (in years): 0, 1-4, 5-9, 10-14, ..., 90-94, 95+.
- **Sex** of the patient: *male, female*.
- **SES (socio-economic status)** of the postal area of patient's address: *lowest, below average, average, above average, highest, unknown*.
- **Severity of main diagnosis** groups: [0-0.01], [0.01-0.02], [0.02-0.05], [0.05-0.1], [0.1-0.2], [0.2-0.3], [0.3-0.4], [0.4-1], *Other*.
- **Urgency** of the admission: *elective, acute*.

- **Comorbidity_1 – Comorbidity_17.** All 17 covariates are dummy variables, having categories: *0 (no)* and *1 (yes)*.
- **Source of admission:** *home, nursing home or other institution, hospital.*
- **Month of admission.** Six 2-month periods: *January/February, ..., November/December.*
- **Year.** Year of the study period (generally for index admissions year *t* is defined by a discharge date from November 1st of year *t-1* up to October 31st): *2016, 2017.*

2.7 Estimation of the model

Logistic regression models were estimated for each of the 122 diagnosis groups using the variables of the index admissions mentioned in the previous paragraph and the dichotomous variable indicating whether an admission was followed by a readmission as the target variable. Computations were performed using the `glm` function in R (R Core Team, 2015). Categories, including the reference category, are collapsed if the number of index admissions is smaller than 50 or when there are no readmissions in the category. For more information on this see the aforementioned methodology report for the HSMR.

The results of the model are described in chapter 3.

3. Outcome of the 2017 model

3.1 Dataset

Table 3.1.1 shows the number of hospitals that were included in the model. All general and university hospitals could be included in both study periods (2016 and 2017). One short stay specialised hospital was excluded since its patients are mostly treated for oncological disease, which is excluded from the model. Another short stay hospital (clinic for lung diseases) stopped operating as an independently registered clinic in 2017 and merged with a university hospital, and was therefore not included as a separate hospital the 2017 dataset. Due to mergers, the number of general hospitals was lower in study period 2017 compared to 2016.

3.1.1 Number of hospitals in the 2017 model.

Study period		General hospitals ^{a)}	University hospitals	Selected specialised hospitals ^{b)}	Total hospitals
2016	Total number	71	8	3	82
	Used in model	71	8	2	81
2017	Total number	67	8	2	77
	Used in model	67	8	1	76

a) Excluding military hospital

b) One clinic for lung diseases (up to 2016), one cancer hospital and one eye hospital

The number of index admissions included in the model, the total number of identified readmissions and the unadjusted readmission rate for both study periods are listed in Table 3.1.2.

3.1.2 Admissions and readmissions in 2017 model.

	2016	2017
Total number of index admissions included in model	1 334 251	1 283 437
Number of identified readmissions	122 078	115 773
Unadjusted readmission rate	9.1%	9.0%

3.2 Impact of the covariates on readmission rate

Appendix I shows which covariates have a statistically significant (95 percent confidence) impact on readmission rate for each of the 122 regression models (one for each diagnosis group). Tables 3.2.1 and 3.2.2 show the total number of significant covariates and the total Wald statistics for the 122 regression models. The tables are sorted in descending order (most important variables at the top). The first table shows in how many diagnosis groups a variable is significant in the model. The effect of variables on the predicted probabilities, and, therefore, the importance of the variables for the case mix correction performed by the models, is better measured with the Wald-statistics (shown in the second table).

The order of the variables differs somewhat in both tables, but in both tables age, urgency, sex and severity are in the top 5 of the most important variables for model estimation. For the HSMR 2017 model (Van der Laan *et al.* 2018b) this is also the case for age, urgency and severity, indicating that these variables are relevant for both predicting readmissions and in-hospital mortality. For the 2017 readmission model, sex is more important than for the 2017 HSMR

model, while for the 2017 HSMR model the Charlson comorbidities 9 ((severe) liver disease), 17 (severe liver disease) and 16 (metastatic cancer) are more important than for the 2017 readmission model. Apparently severe liver disease has a higher influence on estimating in-hospital mortality than on estimating readmissions. The difference in importance of Charlson group 16 in both models can be explained by the fact that cancer-related main diagnoses are excluded from the readmission model, since planned readmissions for those diagnoses are frequent.

3.2.1 Statistical significance of the covariates for the 122 logistic regressions (summary), model 2017.

Covariate	No. of significant results	Covariate	No. of significant results
Age	106	Charlson_9	36
Charlson_13	84	Charlson_5	30
Urgency	77	Charlson_7	26
Severity	71	SES	20
Sex	68	Charlson_16	18
Charlson_3	65	Charlson_4	17
Charlson_1	62	Year	16
Charlson_6	62	Month of admission	16
Charlson_2	60	Charlson_17	13
Charlson_10	54	Charlson_12	12
Charlson_14	54	Charlson_15	1
Charlson_11	45	Charlson_8	1
Source of admission	42		

3.2.2 Wald chi-square statistics for the 122 logistic regressions, model 2017.

Covariate	Sum of Wald statistics	Sum of df	Covariate	Sum of Wald statistics	Sum of df
Age	15 997	2 013	Month of admission	800	610
Urgency	9 993	121	SES	785	550
Severity	4 997	296	Charlson_11	664	85
Source of admission	2 219	216	Charlson_9	651	96
Sex	2 172	120	Charlson_5	374	91
Charlson_13	2 099	112	Charlson_7	284	90
Charlson_6	1 199	117	Charlson_16	236	96
Charlson_1	1 177	113	Year	225	122
Charlson_3	1 136	109	Charlson_12	150	63
Charlson_2	1 102	96	Charlson_17	108	27
Charlson_14	1 023	106	Charlson_8	18	9
Charlson_10	937	118	Charlson_15	5	2
Charlson_4	865	89			

Compared to the 2016 intra-hospital readmission model (Van der Laan *et al.* 2018a), the current model was based on more data (2 years of LBZ data versus 1.5 years in the 2016 model). As a result, the number of significant covariates and the Wald statistics were higher in the current model compared to the 2016 model. Nevertheless, the order of the variables remained largely comparable.

3.3 Model evaluation for the 122 regression analyses

Appendix II shows the Areas Under the Curve (AUCs; also known as C-statistics) for each of the 122 regression models. From these AUCs it can be concluded that most models have weak predictive power (this was also the case in the 2016 intra-hospital model). Of the 122 diagnosis groups, only 14 have an AUC of 0.70 or above:

- Joint disorders and dislocations; trauma-related; sprains and strains (diagnosis nr. 132): AUC = 0.75
- Tuberculosis (diagnosis nr. 1): AUC = 0.74
- Fracture of upper limb (diagnosis nr. 135): AUC = 0.74
- Other connective tissue disease (diagnosis nr. 126): AUC = 0.73
- Other non-traumatic joint disorders (diagnosis no. 123): AUC = 0.73
- Disorders of mouth, teeth, and jaw (diagnosis nr. 91): AUC = 0.73
- Superficial injury; contusion (diagnosis nr. 144): AUC = 0.73
- Other upper respiratory disease (diagnosis nr. 89): AUC = 0.72
- Intracranial injury (diagnosis nr. 138): AUC = 0.72
- Residual codes; unclassified (diagnosis nr. 157): AUC = 0.71
- Open wounds of extremities (diagnosis nr. 141): AUC = 0.70
- Nonmalignant breast conditions (diagnosis nr. 116): AUC = 0.70
- Other skin disorders, chronic ulcer of skin (diagnosis no. 120): AUC = 0.70
- Open wounds of head; neck; and trunk (diagnosis nr. 140): AUC = 0.70

Apparently the models that have better predictive power relatively often concern index admissions with a main diagnosis related to injuries.

3.4 Regression coefficients

The file “coefficients intra-hospital readmission index 2017.xlsx” contains the estimated regression coefficients (columns ‘Estimate’) for each of the 122 logistic regressions as well as their standard errors (columns ‘Std. Err.’). For the sake of clarity, the reference categories are given in the first row of the corresponding covariates, and by definition have zero coefficient for each regression. In many cases categories are collapsed. This results in equal coefficients for the collapsed categories. If all categories were collapsed into one category for a certain variable and for a certain diagnosis group (i.e. if there was only one category with ≥ 50 admissions and ≥ 1 readmission), the variable was dropped from the model and all associated coefficients were set to zero. The significance of each of the coefficients is shown in Appendix I.

3.5 Limitations

The readmission indicator has largely the same limitations as the HSMR. Below we will address some issues that are specific to the readmission indicator.

- In principle all readmissions are included in the model: planned and unplanned; related and not related to the index admission. Ideally only unplanned readmissions should be included. However, these are not registered as such in the LBZ. The LBZ contains the variable urgency (acute versus not acute). An admission is registered 'acute' if care is needed within 24 hours and therefore does not seem to reflect the difference between planned and unplanned readmissions. To avoid the inclusion of planned readmissions, some diagnosis groups where planned readmissions are likely (for example the various groups concerning cancer) are excluded as index and readmissions. Also diagnoses that are likely planned readmissions (for example follow-up care and rehabilitation) are excluded as potential readmissions. Furthermore, in the present model (planned) transfers are excluded as readmissions. However, there will still be planned readmissions remaining in the dataset.
- Unlike with the HSMR, Statistics Netherlands does not provide readmission ratios for 2017, based on the model of 2017. DHD will use the estimated models to calculate the ratios using hospital data from 2018. This means that the models are applied to a different year than that on which they were estimated. As was shown for the readmission model 2015 (Van der Laan *et al.* 2017a), this results in a bias and extra variance. Fortunately, the bias can be estimated and the overall average of the ratio can be presented to the hospitals.
- It is difficult to predict readmissions using the variables present in the models: the models explain only a small part of the observed variation. This makes it more likely that there are unobserved population differences that are not corrected for, that influence the readmission probability. This means that some of the differences in the current readmission ratio can be caused by unobserved population differences.
- The model described identifies intra-hospital readmissions only and readmissions that occur in another hospital are not identified. As a result, for hospitals where patients are often readmitted in another hospital, the indicator could underestimate the readmission ratio and vice versa.

4. Conclusion

The outcome of the 2017 readmission model is largely comparable to that of the 2016 intra-hospital model.

Like in the 2016 model, 'to and fro' transfers are excluded as readmissions. This removes some of the noise from the model, as these transfers can be considered as planned readmissions, which are not of interest when the readmission ratio is used as indicator of quality of care. Although several diagnosis groups consisting of diseases that require treatment during multiple, consecutive admissions have been excluded from the model, it is possible that the data still contains planned readmissions, resulting in less reliable outcome. Although the predictive power of the model is generally low, the case mix correction performed by the model does remove some of the differences between the hospitals caused by population differences. However, because of the weak predictive power of the models, it is likely that there are still population differences remaining for which the model does not correct. Nevertheless, applying the model for calculating readmission ratios for individual hospitals is preferable to calculating crude rates.

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Appendix I: Results of the logistic regressions

Statistical significance (95% confidence) of the covariates for the 122 logistic regressions (1=significant; 0=non-significant; "-"=variable dropped because all categories are collapsed, due to < 50 admissions or no readmissions in all but one category).

Diagnosis group	Sex	Age	Urgency	Severity	Comorbidity_1	Comorbidity_2	Comorbidity_3	Comorbidity_4	Comorbidity_5	Comorbidity_6	Comorbidity_7	Comorbidity_8	Comorbidity_9	Comorbidity_10	Comorbidity_11	Comorbidity_12	Comorbidity_13	Comorbidity_14	Comorbidity_15	Comorbidity_16	Comorbidity_17	SFS	Month admission	Source admission	Year
1	0	1	0	0	-	-	-	-	-	-	-	-	-	0	-	-	-	-	-	-	-	0	0	0	0
2	1	1	0	1	1	0	1	1	1	0	1	-	1	0	1	0	1	1	-	0	0	0	1	1	0
3	1	0	0	0	0	0	0	0	0	0	0	-	0	0	0	-	1	1	-	0	-	0	0	0	0
4	0	0	0	0	0	-	-	-	-	0	-	-	-	0	-	-	1	1	-	0	-	0	0	0	0
5	0	0	1	0	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1	-	-	0	0	0	0
6	0	1	1	1	0	0	0	0	1	0	0	-	1	0	0	0	0	1	-	0	1	0	0	0	0
37	1	1	1	1	1	1	1	1	0	1	0	-	0	1	0	0	1	1	-	0	-	0	0	0	0
38	1	1	1	1	1	1	0	0	0	1	0	-	0	0	0	0	1	0	-	0	-	0	1	0	1
39	0	1	1	1	1	1	1	0	0	1	0	-	0	0	1	-	1	0	-	0	-	0	0	0	0
40	1	1	0	1	1	1	1	0	0	0	0	-	0	1	1	-	1	0	-	0	-	0	0	0	0
41	1	1	1	1	1	1	0	1	1	1	0	-	0	0	1	0	1	0	-	0	-	0	0	0	1
42	1	1	1	0	1	1	1	0	1	0	0	-	1	0	1	0	1	1	-	1	1	0	0	1	0
43	0	1	0	0	-	-	-	-	-	0	-	-	0	0	-	-	-	-	-	-	-	0	0	0	0
44	0	1	1	1	0	0	1	0	1	0	0	-	0	0	-	-	1	1	-	0	-	1	0	0	0
45	1	1	1	1	1	1	1	0	1	1	0	1	1	1	1	0	1	0	-	0	1	0	0	0	0
46	0	1	0	-	0	-	0	-	-	0	0	-	0	0	-	-	0	0	-	1	-	0	0	0	0
51	0	1	1	1	0	-	0	0	-	0	-	-	1	0	-	0	0	0	-	-	-	0	0	1	0
52	0	0	0	0	1	-	-	-	0	0	-	-	-	0	-	-	1	-	-	-	-	0	0	0	0
53	1	1	0	1	0	-	0	0	1	0	-	-	-	0	-	1	0	0	-	0	-	0	0	1	0
54	1	0	0	0	0	-	0	0	-	1	-	-	-	0	-	0	0	0	-	0	-	0	0	0	0
55	0	1	0	1	0	1	1	0	0	0	0	-	0	1	0	0	1	1	-	0	-	0	0	1	0
56	0	0	0	0	0	-	0	0	0	0	-	-	1	0	-	-	1	0	-	0	-	0	0	0	0
57	1	1	1	1	1	1	1	1	0	1	1	-	0	1	0	0	1	1	-	0	-	1	0	0	0
58	1	1	1	1	1	1	1	0	0	1	0	-	1	1	1	0	1	0	-	0	0	1	1	0	0
59	1	1	1	0	0	1	0	0	0	1	0	-	0	0	1	0	1	1	-	1	-	0	0	1	0
60	0	1	1	1	1	1	0	0	-	0	0	-	0	1	0	-	1	1	-	0	-	0	0	1	1
61	0	1	0	1	1	0	1	0	0	0	1	-	0	0	1	-	1	0	-	-	-	0	0	0	0
62	1	1	1	1	1	1	1	0	1	1	1	-	0	1	1	0	1	1	-	0	-	0	1	1	0
63	1	1	1	0	0	1	1	1	0	1	1	-	1	1	1	0	1	0	-	0	-	1	0	1	1
64	1	1	0	-	1	1	1	1	0	1	0	-	1	1	1	0	1	1	-	1	-	1	0	0	0
65	0	1	1	1	1	1	0	0	0	1	1	-	0	1	0	0	1	1	-	1	-	0	1	1	0
66	0	0	1	0	1	0	0	-	-	0	-	-	-	0	-	-	1	-	-	-	-	0	0	1	0
67	0	1	1	0	1	0	1	0	0	1	0	-	0	0	0	-	1	1	-	0	-	0	0	0	1
68	1	1	1	1	1	1	1	0	0	1	1	-	0	1	1	0	1	1	-	0	1	1	0	0	1
69	1	1	1	1	0	1	1	0	-	1	-	-	0	0	-	-	1	1	-	-	-	0	0	0	0
70	1	1	1	1	1	0	1	0	1	1	1	-	0	1	1	0	1	0	-	1	0	1	0	1	0
71	1	1	0	1	1	1	1	1	0	1	1	-	1	1	1	0	1	1	-	1	-	0	0	1	0

Year	Source admission	Month admission	SES	Comorbidity_17	Comorbidity_16	Comorbidity_15	Comorbidity_14	Comorbidity_13	Comorbidity_12	Comorbidity_11	Comorbidity_10	Comorbidity_9	Comorbidity_8	Comorbidity_7	Comorbidity_6	Comorbidity_5	Comorbidity_4	Comorbidity_3	Comorbidity_2	Comorbidity_1	Severity	Urgency	Age	Sex	Diagnosis group
72	1	0	0	1	0	1	1	1	0	1	1	0	-	0	1	0	1	1	1	1	1	1	0	1	
73	1	0	1	-	0	-	0	1	0	1	1	1	-	1	1	1	0	1	1	1	1	1	1	1	
74	1	1	0	-	0	-	0	1	0	0	0	0	-	0	0	0	0	0	0	1	1	1	0	1	
75	1	0	0	-	0	-	0	1	-	1	1	-	-	0	0	0	0	1	1	0	1	1	1	1	
76	1	1	0	-	0	-	0	1	0	1	1	0	-	0	1	1	0	1	1	1	1	1	0	1	
77	1	0	0	-	0	-	0	1	-	1	1	0	-	1	1	0	0	1	1	1	0	1	1	1	
78	1	1	0	-	0	-	0	1	1	1	1	1	-	0	1	1	1	1	1	1	1	1	1	1	
79	1	0	0	-	0	-	0	1	1	0	0	0	-	0	1	1	0	1	1	1	1	0	1	1	
80	1	1	0	-	0	-	0	1	1	0	0	0	-	1	1	1	0	1	1	1	1	0	1	1	
81	1	1	0	-	0	-	0	1	-	-	-	-	-	0	1	0	0	0	0	0	0	0	0	1	
82	1	1	1	-	0	-	0	1	1	1	1	1	-	0	1	0	0	1	1	1	1	1	1	1	
83	0	1	0	-	0	-	0	0	-	0	0	-	-	0	1	0	1	1	1	0	0	-	0	1	
84	1	1	1	-	0	-	0	0	0	0	0	0	-	-	0	0	1	0	1	0	0	-	0	1	
85	0	1	1	-	0	-	0	0	-	0	0	-	-	1	1	0	0	0	0	1	0	0	0	1	
86	0	1	0	-	0	-	0	0	-	0	1	1	-	0	1	0	0	0	0	1	0	0	0	1	
87	0	0	0	-	0	-	0	-	-	-	0	-	-	-	0	-	-	-	-	-	-	-	-	0	
88	1	1	0	-	0	-	0	1	0	0	1	0	-	0	1	0	0	1	1	1	0	1	1	1	
89	0	1	1	-	0	-	0	1	1	1	1	1	-	1	1	1	0	1	1	1	1	1	1	1	
90	0	1	1	-	0	-	0	1	1	1	1	1	-	1	0	0	0	0	1	1	1	0	1	0	
91	0	1	1	-	0	-	0	1	-	-	-	-	-	1	-	-	-	0	1	-	0	-	0	1	
92	0	1	1	-	0	-	0	1	-	1	1	-	-	1	0	0	1	0	0	0	0	0	0	0	
93	0	0	0	-	0	-	0	0	-	-	0	-	-	0	0	-	-	1	-	-	-	-	-	0	
94	1	1	0	-	0	-	0	1	-	1	0	-	-	0	0	0	0	0	1	0	0	0	0	0	
95	0	1	0	-	0	-	0	0	-	0	1	-	-	0	0	-	-	0	1	0	0	0	0	1	
96	0	1	1	-	0	-	0	0	-	-	0	-	-	0	0	-	-	0	0	0	0	0	0	0	
97	1	1	1	-	0	-	0	1	0	1	1	1	-	0	1	0	0	0	0	0	0	0	0	1	
98	1	0	1	-	0	-	0	-	-	0	0	-	-	0	0	-	-	0	0	0	0	0	0	0	
99	1	1	0	-	0	-	0	0	0	0	0	0	-	0	0	0	0	1	1	0	0	0	1	0	
100	0	1	1	-	0	-	0	1	1	1	1	-	1	1	0	0	0	0	0	0	0	0	0	0	
101	1	1	1	-	0	-	0	1	-	0	0	-	0	0	0	1	0	0	0	0	0	0	0	0	
102	1	1	1	-	0	-	0	1	1	1	1	1	-	0	1	1	0	1	1	1	1	0	0	0	
103	0	1	1	-	0	-	0	1	-	1	1	-	-	0	0	0	0	0	0	0	0	0	0	1	
104	1	1	1	-	0	-	0	1	-	1	0	-	-	1	1	0	0	0	0	0	0	0	0	1	
105	0	1	1	-	0	-	0	0	-	0	0	-	-	0	0	0	0	0	0	0	0	0	0	0	
106	1	1	0	-	0	-	0	1	1	1	1	0	-	0	1	1	0	1	1	1	0	0	0	0	
107	0	1	1	-	0	-	0	0	-	0	0	-	-	0	0	0	0	0	0	0	0	0	0	0	
108	1	1	1	-	0	-	0	1	0	0	1	1	-	0	1	0	0	1	1	1	0	0	0	0	
109	1	1	0	-	0	-	0	1	-	1	1	-	-	0	0	0	0	0	0	0	0	0	0	0	
110	0	1	0	-	0	-	0	0	-	0	1	-	-	0	0	0	1	0	0	0	0	0	1	1	
111	0	1	1	-	0	-	0	0	-	0	0	-	-	1	0	0	0	0	0	0	0	0	0	0	
112	1	1	1	-	0	-	0	1	1	1	0	1	-	0	1	1	0	1	1	1	1	0	1	1	
113	1	1	1	-	0	-	0	1	1	1	1	1	-	0	1	1	0	0	0	0	0	0	0	1	
114	1	1	1	-	0	-	0	1	0	0	0	-	-	0	0	0	0	0	0	0	0	0	0	0	
115	-	1	1	-	0	-	0	1	0	0	0	-	-	0	0	0	0	0	0	0	0	0	0	0	

Year	Source admission	Month admission	SES	Comorbidity_17	Comorbidity_16	Comorbidity_15	Comorbidity_14	Comorbidity_13	Comorbidity_12	Comorbidity_11	Comorbidity_10	Comorbidity_9	Comorbidity_8	Comorbidity_7	Comorbidity_6	Comorbidity_5	Comorbidity_4	Comorbidity_3	Comorbidity_2	Comorbidity_1	Severity	Urgency	Age	Sex	Diagnosis group
116	0	0	0	-	-	-	0	-	-	0	-	-	-	0	0	-	-	-	-	-	-	1	1	0	116
117	0	0	0	0	0	-	1	1	-	1	0	-	-	0	1	0	-	1	0	1	1	1	1	-	117
119	0	0	1	0	0	-	0	1	1	1	1	-	-	1	1	0	0	1	1	1	1	0	0	0	119
120	0	0	0	0	0	-	0	0	0	1	1	-	-	0	1	0	0	1	1	1	1	1	1	1	120
121	0	0	0	0	0	-	0	0	0	1	0	-	-	0	0	0	0	1	0	0	0	0	0	0	121
122	0	0	1	0	0	-	0	1	1	1	0	-	-	1	1	0	0	1	1	0	1	1	1	1	122
123	0	0	0	0	0	-	0	0	0	0	1	-	-	0	1	0	0	0	0	0	0	1	1	0	123
124	0	0	0	0	0	-	0	1	1	0	1	-	-	0	1	0	0	0	0	1	1	1	1	0	124
125	0	0	0	0	0	-	0	0	0	1	-	-	-	0	1	0	-	1	1	0	0	0	0	1	125
126	0	0	1	0	0	-	0	1	0	0	1	-	-	0	1	0	0	1	1	1	1	1	1	1	126
127	0	1	0	0	0	-	0	-	-	-	0	-	-	-	0	0	1	0	1	0	-	0	0	0	127
128	0	0	1	0	0	-	0	0	0	-	0	-	-	1	0	0	-	-	-	-	-	-	-	0	128
129	0	0	0	0	0	-	0	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1	129
130	0	0	0	0	0	-	0	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1	130
131	0	1	1	0	0	-	0	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0	1	131
132	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	-	-	-	-	-	-	-	0	132
133	0	1	0	0	1	1	1	0	1	0	0	-	-	0	1	0	0	1	0	1	1	1	0	1	133
134	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	-	-	-	-	-	-	-	0	134
135	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	135
136	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	1	0	0	0	0	0	0	136
137	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	137
138	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	138
139	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	139
140	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	-	-	-	-	-	-	-	0	140
141	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	-	-	-	-	-	-	-	0	141
142	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	142
143	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	143
144	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	144
145	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	-	-	-	-	-	-	-	0	145
146	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	146
147	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	147
148	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	148
149	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	149
150	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	-	-	-	-	-	-	-	0	150
151	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	-	-	-	-	-	-	-	0	151
152	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	152
153	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	153
154	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	154
155	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	-	-	-	-	-	-	-	0	155
156	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	156
157	0	0	0	0	0	-	0	-	-	-	0	-	-	-	0	0	0	0	0	0	0	0	0	0	157
68	106	77	71	62	60	65	17	30	62	26	1	36	54	45	12	84	54	1	18	13	20	16	42	16	

The numbers of the comorbidity groups in the header of the table above are the following comorbidities:

- Comorbidity_1 - Acute myocardial infarction
- Comorbidity_2 - Congestive heart failure
- Comorbidity_3 - Peripheral vascular disease
- Comorbidity_4 - Cerebral vascular accident
- Comorbidity_5 - Dementia
- Comorbidity_6 - Pulmonary disease
- Comorbidity_7 - Connective tissue disorder
- Comorbidity_8 - Peptic ulcer
- Comorbidity_9 - Liver disease / Severe liver disease
- Comorbidity_10 - Diabetes / Diabetes complications
- Comorbidity_11 - Diabetes complications
- Comorbidity_12 - Paraplegia
- Comorbidity_13 - Renal disease
- Comorbidity_14 - Cancer
- Comorbidity_15 - HIV
- Comorbidity_16 - Metastatic cancer
- Comorbidity_17 - Severe liver disease

Appendix II: AUC

The area under the curve (AUC) or C-Statistic for the logistic regressions of the 122 main diagnosis groups.

Diagnosis group*)	Number of index admissions	Number of readmissions	AUC
1 Tuberculosis	802	76	0.74
2 Septicemia (except in labor)	7 630	1 066	0.62
3 Bacterial infection; unspecified site	3 696	582	0.62
4 Mycoses	1 034	193	0.65
5 HIV infection	500	77	0.68
6 Hepatitis, viral and other infections	13 599	1 097	0.62
37 Other and unspecified benign neoplasm	38 548	2 282	0.65
38 Thyroid and other endocrine disorders	12 577	989	0.69
39 Diabetes mellitus without complication	8 422	650	0.66
40 Diabetes mellitus with complications	12 193	2 074	0.66
41 Nutritional deficiencies and other nutritional, endocrine, and metabolic disorders	28 983	2 489	0.66
42 Fluid and electrolyte disorders	16 614	2 141	0.65
43 Cystic fibrosis	1 221	189	0.63
44 Immunity and coagulation disorders, hemorrhagic disorders	5 818	996	0.63
45 Deficiency and other anemia	22 573	3 536	0.63
46 Diseases of white blood cells	4 031	673	0.59
51 Meningitis, encephalitis, and other central nervous system infections	5 324	474	0.64
52 Parkinson`s disease	3 498	300	0.58
53 Multiple sclerosis and other degenerative nervous system conditions	7 436	672	0.64
54 Paralysis and late effects of cerebrovascular disease	2 345	177	0.66
55 Epilepsy and convulsions	22 324	1 896	0.59
56 Coma, stupor, and brain damage	1 417	147	0.67
57 Headache and other disorders of the sense organs	40 213	1 740	0.64
58 Other nervous system disorders	55 100	2 703	0.67
59 Heart valve disorders	19 901	1 955	0.62
60 Peri-, endo-, myocarditis, and cardiomyopathy	10 495	1 067	0.64
61 Essential hypertension, hypertension with compl., and secondary hypertension	6 600	544	0.66
62 Acute myocardial infarction	65 028	5 271	0.62
63 Coronary atherosclerosis and other heart disease	73 208	6 010	0.62
64 Nonspecific chest pain	44 929	2 837	0.63
65 Pulmonary heart disease	16 932	1 310	0.66
66 Other and ill-defined heart disease	1 205	108	0.69
67 Conduction disorders (heart disease)	12 055	901	0.63
68 Cardiac dysrhythmias	69 998	6 246	0.64
69 Cardiac arrest and ventricular fibrillation	4 676	289	0.66
70 Congestive heart failure, nonhypertensive	53 872	8 195	0.59
71 Acute cerebrovascular disease	66 012	4 645	0.62

Diagnosis group*)	Number of index admissions	Number of readmissions	AUC
72 Transient cerebral ischemia, and other cerebrovascular disease	24 783	2 042	0.65
73 Peripheral and visceral atherosclerosis	19 755	3 497	0.64
74 Aortic and other artery aneurysms	13 390	1 562	0.58
75 Aortic and arterial embolism or thrombosis	7 937	1 332	0.62
76 Other circulatory disease	16 830	2 405	0.62
77 Phlebitis, varicose veins, and hemorrhoids	7 338	764	0.64
78 Pneumonia	68 296	7 298	0.62
79 Influenza	7 214	631	0.65
80 Tonsillitis and upper respiratory infections	45 150	2 686	0.66
81 Acute bronchitis	13 831	1 119	0.60
82 Chronic obstructive pulmonary disease and bronchiectasis	66 244	12 047	0.56
83 Asthma	16 765	1 613	0.61
84 Aspiration pneumonitis; food/vomitus	3 140	499	0.63
85 Pleurisy; pneumothorax; pulmonary collapse	12 441	1 776	0.60
86 Respiratory failure; insufficiency; arrest	2 965	411	0.65
87 Lung disease due to external agents	829	139	0.62
88 Other lower respiratory disease	13 877	1 709	0.60
89 Other upper respiratory disease	38 982	2 852	0.72
90 Intestinal infection	27 413	2 668	0.64
91 Disorders of mouth, teeth, and jaw	11 086	335	0.73
92 Esophageal disorders	7 804	865	0.63
93 Gastroduodenal ulcer	2 298	223	0.67
94 Gastritis, duodenitis, and other disorders of stomach and duodenum	4 226	578	0.65
95 Appendicitis and other appendiceal conditions	33 188	2 237	0.57
96 Peritonitis and intestinal abscess	2 228	436	0.64
97 Abdominal hernia	26 361	2 192	0.64
98 Regional enteritis and ulcerative colitis	9 738	1 420	0.57
99 Intestinal obstruction without hernia	15 710	2 294	0.57
100 Diverticulosis and diverticulitis	19 482	2 072	0.62
101 Anal and rectal conditions	12 011	1 113	0.61
102 Biliary tract disease	71 840	9 552	0.64
103 Liver disease; alcohol-related	3 132	798	0.64
104 Other liver diseases	8 876	2 237	0.66
105 Pancreatic disorders (not diabetes)	16 191	3 062	0.55
106 Gastrointestinal hemorrhage	18 307	2 623	0.62
107 Noninfectious gastroenteritis	7 735	888	0.61
108 Other gastrointestinal disorders	22 450	2 878	0.60
109 Nephritis; nephrosis; renal sclerosis	7 418	832	0.66
110 Acute and unspecified renal failure	8 315	1 401	0.59
111 Chronic kidney disease	7 827	1 541	0.58
112 Urinary tract infections	46 935	5 847	0.59
113 Calculus and other diseases of urinary tract	44 440	6 234	0.62
114 Genitourinary symptoms and ill-defined conditions	14 902	1 893	0.66
115 Hyperplasia of prostate and other male genital disorders	22 903	1 934	0.60
116 Nonmalignant breast conditions	9 309	268	0.70

Diagnosis group*)	Number of index admissions	Number of readmissions	AUC
117 Prolapse and other female genital disorders	36 233	1 665	0.67
119 Skin and subcutaneous tissue infections	28 987	2 412	0.66
120 Other skin disorders, chronic ulcer of skin	11 114	1 136	0.70
121 Infective arthritis and osteomyelitis	7 013	860	0.61
122 Osteoarthritis, rheumatoid arthritis, and other musculoskeletal deformities	130 410	5 678	0.65
123 Other non-traumatic joint disorders	8 688	491	0.73
124 Spondylosis, back problems, and osteoporosis	48 989	2 625	0.66
125 Pathological fracture	3 358	392	0.65
126 Other connective tissue disease	25 781	1 080	0.73
127 Cardiac and circulatory congenital anomalies	4 870	446	0.60
128 Noncardiac congenital anomalies	15 473	1 063	0.66
129 Short gestation; low birth weight; and fetal growth retardation	30 765	2 455	0.66
130 Intrauterine hypoxia, perinatal asphyxia, and jaundice	25 647	1 160	0.56
131 Other perinatal conditions	101 410	4 758	0.54
132 Joint disorders and dislocations; trauma-related; sprains and strains	18 646	450	0.75
133 Fracture of neck of femur (hip)	39 045	2 562	0.60
134 Skull and face fractures, spinal cord injury	6 083	266	0.66
135 Fracture of upper limb	24 527	1 323	0.74
136 Fracture of lower limb	27 236	2 311	0.68
137 Other fractures	23 450	1 281	0.62
138 Intracranial injury	20 302	851	0.72
139 Crushing injury or internal injury	11 325	593	0.68
140 Open wounds of head; neck; and trunk	3 271	146	0.70
141 Open wounds of extremities	2 903	229	0.70
142 Complication of device, implant or graft	49 071	6 997	0.64
143 Complications of surgical procedures or medical care	51 498	6 995	0.59
144 Superficial injury; contusion	27 251	1 187	0.73
145 Burns	2 029	111	0.69
146 Poisoning by psychotropic agents, drugs, or other medications	16 250	1 219	0.64
147 Other injuries and conditions due to external causes	5 625	414	0.68
148 Syncope	25 050	1 552	0.63
149 Fever of unknown origin	13 510	1 882	0.62
150 Lymphadenitis and gangrene	3 243	414	0.69
151 Shock	481	67	0.68
152 Nausea and vomiting	7 642	1 203	0.59
153 Abdominal pain	27 009	2 986	0.58
154 Malaise and fatigue	6 326	677	0.67
155 Allergic reactions	5 335	302	0.65
156 Rehabilitation and other aftercare, medical examination/evaluation/screening	63 225	4 249	0.61
157 Residual codes; unclassified	35 916	1 963	0.71

*) The diagnosis group numbers refer to the file 'Classification of variables' published together with the HSMR 2017 methodological report (see Van der Laan *et al.* 2018b). In this file, the CCS-groups and

corresponding ICD10-codes of the 157 diagnosis groups used for the HSMR are given. For the readmission ratio only 122 of these groups are used, but the numbering was kept the same.

Explanation of symbols

Empty cell	Figure not applicable
.	Figure is unknown, insufficiently reliable or confidential
*	Provisional figure
**	Revised provisional figure
2018–2019	2018 to 2019 inclusive
2018/2019	Average for 2018 to 2019 inclusive
2018/'19	Crop year, financial year, school year, etc., beginning in 2018 and ending in 2019
2016/'17–2018/'19	Crop year, financial year, etc., 2016/'17 to 2018/'19 inclusive

Due to rounding, some totals may not correspond to the sum of the separate figures.

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