Internal validity of the Swedish Maternal Health Care Register

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Abstract

Background: The Swedish Maternal Health Care Register (MHCR) is a national quality register that has been collecting pregnancy, delivery, and postpartum data since 1999. A substantial revision of the MHCR resulted in a Web-based version of the register in 2010. Although MHCR provides data for health care services and research, the validity of the MHCR data has not been evaluated. This study investigated degree of coverage and internal validity of specific variables in the MHCR and identified possible systematic errors.

Methods: This cross-sectional observational study compared pregnancy and delivery data in medical records with corresponding data in the MHCR. The medical record was considered the gold standard. The medical records from nine Swedish hospitals were selected for data extraction. This study compared data from 878 women registered in both medical records and in the MHCR. To evaluate the quality of the initial data extraction, a second data extraction of 150 medical records was performed. Statistical analyses were performed for degree of coverage, agreement and correlation of data, and sensitivity and specificity.

Results: Degree of coverage of specified variables in the MHCR varied from 90.0% to 100%. Identical information in both medical records and the MHCR ranged from 71.4% to 99.7%. For more than half of the investigated variables, 95% or more of the information was identical. Sensitivity and specificity were analysed for binary variables. Probable systematic errors were identified for two variables.

Conclusions: When comparing data from medical records and data registered in the MHCR, most variables in the MHCR demonstrated good to very good degree of coverage, agreement, and internal validity. Hence, data from the MHCR may be regarded as reliable for research as well as for evaluating, planning, and decision-making with respect to Swedish maternal health care services.

Keywords: Validity, Degree of coverage, National quality register, Medical records, Pregnancy outcomes, Antenatal care

Background

Health data registers and quality registers

Nordic countries have a long tradition of using population-based health data registers to monitor the general population. These health data registers include the Swedish Cause of Death Cause Register (1952), the Swedish Cancer Register (1958), the Norwegian Medical Birth Register (1967), and the Swedish Medical Birth Register (1973) [1]. Swedish health data registers are regulated by the Health Data Law in the Swedish code of statues (1998:543) and it is compulsory for patients, as well as for the health care services, to participate in these registers [2]. The health data registers use standardized data collection procedures, enabling surveillance of the health status of the population [3]. In addition, these registers are available to researchers [3-6]. Over the last several decades, a growing number of national quality registers surveying specific medical areas have been established in Sweden. Quality registers have been initiated and are administered by professional associations from different medical areas. In contrast to health data registers, participation in quality registers is voluntary for both patients and health care providers. That is, patients can choose not to contribute their individual data to a quality register. Quality registers are regulated by the
Swedish code of statutes (2008:355) [2]. The quality registers provide a unique possibility to survey different aspects of health care and health care outcomes. In addition, quality registers can be used to conduct research, to improve quality of health care, and to manage health care services [7]. Clearly, it is important that data in the registers are valid and representative [8]. The major part of quality registers are financed by the Swedish government and the Swedish Association of Local Authorities and Regions, government entities that have deemed it a national priority that quality registers should cover at least 80% of the population [7].

In Sweden, management of national quality registers is regulated by Swedish legislation and the National Board of Health and Welfare [7]. Collection and management of patient data in quality registers are regulated by the Swedish Patient Data Law, which charges health providers the responsibility of informing patients on the existence of a specific health register, the purpose of the register, and the type of data that are reported to the register. The patients are informed that their participation in the health register is voluntary and that removal of data is automatically granted if the patient desires [9].

The Swedish Maternal Health Care Register
The Swedish Maternal Health Care Register (MHCR) is a national quality register established in 1999. In 2007, a substantial revision was performed of its variables, Web application, and technical solutions. The revised version of MHCR was launched on January 1, 2010. The MHCR collects pregnancy, delivery, and postpartum data, including individual data on the pregnant women, foetuses, and infants. In 2010 and 2011, 81% and 85%, respectively, of the pregnant population were registered in the MHCR (personal communication). The main bulk of data registered in MHCR is related to pregnancy and delivery, but data on lifestyle, education, and socioeconomic factors are also reported. In accordance with the Swedish Patient Data Law [9], all antenatal care centres (ANC) are charged with informing each pregnant woman on the existence of the MHCR, its purpose, its content, and the fact that providing data is voluntary.

Data in the MHCR are entered on two different occasions by attending ANC midwives. Entering data into the MHCR is performed using a Web-based application specifically created for this purpose. To protect the integrity of the data, each midwife is provided with an individual user identity and a secure login procedure. The first dataset is entered when a pregnant woman registers in ANC. This dataset mainly includes information about background characteristics, such as educational level, weight, height, and smoking habits. On the first visit, Body Mass Index (BMI) is calculated using a software program built into the MHCR.

According to national recommendations for health care during pregnancy and after delivery, all women should be offered a postpartum meeting with a midwife in the ANC four to 16 weeks after delivery [10]. The second data entry takes place soon after the postpartum visit and includes items related to pregnancy, delivery, and the postpartum period. If a woman does not attend the voluntary postpartum visit, the midwife enters the second set of data at around 16 weeks postpartum using information from the medical records. The items in this second data entry address the outcomes of pregnancy and delivery.

Most of the registered items entered in the MHCR are data obtained from medical records manually registered by a midwife. The MHCR database is administrated by the Uppsala Clinical Research Centre (UCR), which specifically supports the maintenance of national quality registers and assists researchers using these registers.

No previous study has evaluated the validity of data included in the MHCR. As national quality registers are used for quality improvement and management within regional and local health care services as well as for research, it is important that the quality of data in the registers is high.

Aims
This study investigate the validity of data entered in the MHCR. The study has three specific aims: i) to explore degree of coverage of specified variables; ii) to investigate internal validity of data, including sensitivity and specificity of binary variables; and iii) to identify potential systematic errors.

Methods
Study design and study sample
This cross-sectional observational study compared data on pregnancy and delivery using medical records and the MHCR. The Regional Ethical Board at Umeå University (Umeå, Sweden) approved the national study (Dno 2012-44-31 M).

A power estimation was performed to determine the sample size; to obtain kappa values of 0.6 (considering the null value of kappa to be 0.4) and to achieve 90% power, a sample size of 540 was required if the prevalence was 0.1 (or 0.9) and 220 if the prevalence was 0.5. However, kappa is very sensitive to prevalence and as the categorical variables vary considerably with respect to prevalence, a sample of 900 medical records was judged to be a sufficient sample size to respond to the research questions under study. This study uses a national sample comprising nine Swedish hospitals, 100 medical records from each hospital. The hospitals were selected because they provided a variation in geographic and demographic characteristics. In Sweden, there were...
109,752 deliveries in 2011. The data collection was performed at hospitals representing delivery units ranging from 1,298 to 10,363 births in 2011 [11] and covered the northern and southern regions of Sweden. To some extent, the selection of hospitals was influenced by convenience, as most of the authors of this study constituted a subset of the board of the MHCR and are affiliated with five of the selected hospitals included in the study. These circumstances provided a better opportunity to supply instructions and support to the local administrators who were extracting the data from the medical records.

Inclusion criteria for the study were medical records of women with data on pregnancy and delivery both in the medical records and in the MHCR. Exclusion criteria were data lacking in either of these two data sources.

Medical records of pregnancy, delivery, and the postpartum period
The software program Obstetrix™ is widely used in Sweden and contains pregnancy, delivery, and postpartum data, accounting for approximately 90% of medical records on pregnant women in Sweden. Other software programs used in clinical practice are Partus™ and Cosmic Birth™. A few clinics still document medical data using pen and paper.

Data collection procedures
Before the start of the study, the heads of all participating clinics provided verbal consent to participate. After the consent was secured from the heads of the clinics, local administrators, one administrator at each hospital, were contracted to supply the data registration. Most of the local administrators were medical secretaries, but in a few hospitals midwives or other staff were contracted.

In 2011, data on 85% of all pregnant women were included in the MHCR. Therefore, the personal identity numbers of 120 consecutively delivered women were extracted from the birth log at each clinic to ensure that 100 women were identified from each clinic with data both in the medical records and in the MHCR. From the nine clinics, we selected 100 women per clinic who gave birth from March 1st to March 9th. Extracted data from the medical records for the 900 women were transferred in encrypted form to the UCR. The UCR combined the extracted data in the medical records with the corresponding data in the MHCR. The goal was to collect data on 100 women from each hospital; i.e., we wanted to have data from 900 medical records. For seven hospitals, data on 100 women were incomplete. Despite repeated reminders by e-mail and by telephone, no further data were delivered, resulting in a final dataset of extracted data from 878 medical records.

Study protocol
An Excel™-protocol was developed by the authors to register categorical and numeric variables extracted from the medical records and to secure that data were extracted in a similar manner at all hospitals. In general, registration of data from the medical records into the Excel™-protocol was done manually by the local administrator. However, in one hospital data were electronically collected from the medical records and imported into the Excel™-protocol. Then the content of each Excel™-protocol was encrypted and sent to the UCR.

Data from Excel™-protocols and data from the MHCR were merged by the UCR using the personal identity number for each woman. To ensure that individuals could not be identified, the merged dataset was delivered to the authors with each individual given a unique serial number.

Presentation of included variables
All variables included in the MHCR and the selected variables for this study are presented in Table 1. Some variables available in the MHCR were excluded for the validity control, such as variables regarding the postpartum period and variables with no corresponding data in the medical records (e.g., the variables of self-reported health before, during, and after pregnancy).

Most of the categorical variables in the MHCR had the response options of “yes”, “no”, or “don’t know”. However, two variables had other response options: “mode of delivery” (“caesarean section”, “instrumental vaginal delivery”, or “non-instrumental vaginal delivery”) and the variable “gender” (“girl”, “boy”, or “unknown gender”). Three of the categorical variables with response options “yes”, “no”, or “don’t know” had an additional question if the response “yes” was noted. These variables had the following additional options: i) Alcohol Use Disorder Identification Test (AUDIT) scores; ii) the options elective caesarean section (CS) or emergency CS, if mode of delivery was registered as CS; and iii) the two-hour plasma glucose value was requested if an oral glucose tolerance test (OGTT) had been performed.

Quantitative variables were registered as continuous numeric values. Birth weight was registered in grams. Maternal body weight was recorded in whole kilograms and maternal height in centimetres. AUDIT-scores ranged from 0 to 40. Variables addressing dates were registered in a pre-set calendar format. Some deliveries were multiple births. Data on first twin, such as mean birth weight and mode of delivery, were included in the presentation of singleton pregnancies. Mean birth weight for second twin was also calculated.
To investigate to what extent data from the medical records had been correctly registered in the Excel®-protocol, a second data extraction was performed (i.e., re-collection of data). Three of the participating hospitals – Östersund Hospital (Östersund), Södersjukhuset (Stockholm), and Umeå University Hospital (Umeå) – were selected for this control procedure. Two of the authors (KP and IH, both midwives with extensive experience with ANC) performed this re-collection of data. An identical Excel®-protocol as used for the first data collection from medical records was used for this second data collection procedure. The goal was to include every second woman from the primary dataset from each of the three selected hospitals in this second validation procedure of data (i.e., data were collected from medical records on 50 women from each hospital, resulting in data from 150 medical records).

### Table 1 Presentation of all variables registered at first and second data entry in the Swedish Maternal Health Care Register (MHCR)

| First data entry                                                                 | Second data entry                                                                 |
|---------------------------------------------------------------------------------|----------------------------------------------------------------------------------|
| Data collected at first visit in antenatal care (ANC)                           | Data collected at postpartum visit in antenatal care (ANC) 4 to 16 weeks after delivery |
| **Variables**                                                                   | **Variables**                                                                    |
| Date of first visit in ANC<sup>a</sup>                                          | Live born child                                                                   |
| Country of birth                                                                 | Still birth/termination of pregnancy                                              |
| No. of previous deliveries                                                       | Date of delivery estimated by ultrasound                                           |
| Maternal weight (kilograms)<sup>b</sup>                                         | Estimated date of delivery (ultrasound)                                           |
| Maternal height (centimetres)<sup>b</sup>                                       | Estimated date of delivery (last menstruation)                                    |
| Smoking three months prior to pregnancy                                         | Ultrasound examination at gestational age 16-21 week                               |
| No. of cigarettes/day three months prior to pregnancy                           | Combined Ultrasound and Biochemical screening (CUB)                                |
| Smocking at first ANC visit                                                      | Second trimester serum screening                                                  |
| No. of cigarettes/day at first ANC visit                                         | Chorionic villus sampling (CVS)                                                   |
| Use of snuff three months prior to pregnancy                                     | Amniocentesis (AC)                                                                |
| Use of snuff at first ANC visit                                                  | Number of antenatal visits until estimated date of delivery (determined by ultrasound) |
| Assessment of use of alcohol prior to pregnancy with screening instrument Alcohol Use Disorder Identification Test (AUDIT) | Number of midwives surveying the pregnant woman in ANC                            |
| AUDIT-score                                                                      | Use of authorized interpreter                                                     |
| Education level                                                                  | Smoking at 32 weeks of gestation                                                  |
| Employment status                                                                | No. of cigarettes/day at 32 gestational weeks                                     |
| Self-rated health prior to pregnancy                                             | Use of snuff at 32 weeks of gestation                                             |
| Maternal weight (in kilograms), last data entry after 35 gestational weeks      | Maternal weight (kilograms), last data entry after 35 gestational weeks           |
| Participated in prenatal education group (pregnant woman)                       | Participation in prenatal education group (partner)                                |
| Counselling due to fear of childbirth                                           | Breast feeding at 4 weeks postpartum                                              |

**Variables presented in bold text were selected for the comparison of data in medical records and in MHCR.**

<sup>a</sup>Gestational age at registration in ANC is calculated by the software program.

<sup>b</sup>Body Mass Index (BMI) at registration in ANC is calculated by the software program.

<sup>c</sup>Foetal growth proportionality – i.e., appropriate for gestational age (AGA), large for gestational age (LGA), and small for gestational age (SGA) – is calculated by the software program.

<sup>d</sup>In cases of multiple births, birth weight and gender are also registered for second twin.
Statistical analysis

Data from the medical records were considered the gold standard. The proportions of available data in the medical records and in the MHCR and the proportions of data available in both data sources were calculated for each variable. In addition, the proportion of cases where the medical records and the MHCR presented identical information was calculated for each variable. For the subset of data (re-collected dataset) where the categorical variables with a subsequent explorative question in the case of a “yes” response, the number of “yes” responses constituted the denominator in the calculations. Degree of agreement was estimated using Cohen’s kappa for categorical data and Pearson’s correlation coefficient was used for normally distributed, continuous data. Spearman’s correlation coefficient was used to evaluate dates. Sensitivity and specificity were analysed for binary variables. Sensitivity was defined as the proportion of actual positives, that were correctly identified as such. Specificity was defined as the proportion of negatives that were correctly identified as such. Sensitivity and specificity were analysed for binary variables. SPSS version 19 was used for all calculations. The level of significance was set at 0.05.

Results

Background presentation

Corresponding data on pregnancy and delivery from medical records and the MHCR were collected from 878 medical records at nine hospitals. These hospitals and their characteristics are presented in Table 2. The number of deliveries at the included hospitals corresponds to 28.0% of the total number of deliveries in Sweden in 2011. The data collected from medical records included mean age (30.7 years, SD ±5.0), mean BMI (24.6, SD ±4.6), and mean birthweight of infant (3515 g, SD ±568). Eleven pregnancies were multiple births. The mean gestational age was 278.2 days (SD ±12.5) or 39.7 weeks (SD ±1.8) for singleton births and 241.6 days (SD ±36.2) or 34.5 weeks (SD ±5.2) for multiple births. Mean birth weight of second twin was 1810 g (SD ±1003).

Degree of coverage of data in medical records and in the MHCR

The degree of coverage of all investigated variables is presented in Table 3. The degree of coverage of variables included in medical records varied from 48% to 100% and most variables presented high degree of coverage in medical records. There was a high degree of coverage for the categorical variable OGTT (98.9%) in medical records. However, there was a lower degree of coverage for the associated variable “OGTT two-hour value of plasma glucose” (48.0%) in medical records.

Degree of coverage of data registered in the MHCR varied between 90.0% and 100%. The variables with a relatively lower degree of coverage in the MHCR, although in fact a high degree of coverage, addressed various forms of prenatal diagnostics with a degree of coverage of approximately 90%.

Data available in both data sources (medical records and MHCR) ranged from 46.0% to 100%. Variables with complete data in both data sources were variables addressing date of birth and whether the child was born alive or stillborn. Other variables with a high level of data available in both data sources included “singleton birth/multiple births” (99.8%), “mode of delivery” (99.5%), and “gender of child” (99.5%).

Table 2 Characteristics of the nine participating hospitals and number of medical records extracted at each hospital

| City       | Participating hospital            | Level of health care | Inhabitants/km² 2011a | No. of births 2011b (%) | No. of medical records (%)b |
|------------|-----------------------------------|----------------------|------------------------|--------------------------|-----------------------------|
| Göteborg   | Sahlgrenska University Hospital   | University           | 66.8                   | 10363 (9.4)              | 91 (10.4)                   |
| Halmstad   | Halmstad Hospital                | County               | 55.6                   | 1799 (1.6)               | 96 (10.9)                   |
| Jönköping  | Ryhov Hospital                   | County               | 32.4                   | 2075 (1.9)               | 99 (11.3)                   |
| Stockholm  | Karolinska University Hospital   | University           | 320.5                  | 4462 (4.2)               | 96 (10.9)                   |
| Stockholm  | Södersjukhuset                   | University           | 320.5                  | 7331 (6.7)               | 98 (11.2)                   |
| Sundsvall  | Sundsvall Hospital               | Regional             | 11.2                   | 1536 (1.4)               | 100 (11.4)                  |
| Umeå       | Umeå University Hospital         | University           | 4.7                    | 1817 (1.6)               | 100 (11.4)                  |
| Örebro     | Örebro University Hospital       | University           | 3.3                    | 2867 (2.6)               | 99 (11.3)                   |
| Östersund  | Östersund Hospital               | Regional             | 2.6                    | 1298 (1.2)               | 99 (11.3)                   |

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aPopulation density in catchment area. Data from "Inhabitants per kilometer²" [Internet] Statistics Sweden; 2011 (cited 2013, June 6) http://www.scb.se/Pages/SSD/SSD_SelectVariables340487.aspx?px_tableid=ssd_extern%3aBefArealTathetKon&rid=ca8cabdd-0d60-488b-b047-4b5c5a89dc55.
bData from National Board of Health and Welfare/Graviditeter, förlossningar och nyfödda barn. Medicinska Födelseregistret 1973-2011. Assisterad befruktning 1991 – 2010' [in Swedish] http://www.socialstyrelsen.se/publikationer2013-2013-3-27.
cProportions are calculated by using the total no of births in Sweden 2011 (N = 109 752) as denominator.
dProportions are calculated by using the total no of medical records as denominator.
Table 3 Data in medical records and the Sweedish Maternal Health Care Register (MHCR); comparison between the two data-sets using correlation analysis, and analysis of sensitivity and specificity for binary variables

| Variable                                      | Data source: Medical records | Data source: MHCR | Data available in both data sources | Identical information in both data sources | Correlationa | Sensitivity | Specificity |
|-----------------------------------------------|------------------------------|-------------------|-------------------------------------|--------------------------------------------|---------------|-------------|-------------|
|                                              | n   | %   | n   | %   | n   | %   | n   | %   |                             |
| Date of first visit in ANC (numerical)       | 877 | 99.9 | 868 | 98.9 | 867 | 98.7 | 685 | 79.0 | 0.878 (S) |
| No of previous deliveries (numerical)        | 878 | 100 | 867 | 98.7 | 867 | 98.7 | 840 | 96.8 | 0.971 (P) |
| Maternal weight at first ANC visit (numerical) | 862 | 98.1 | 855 | 97.4 | 847 | 96.4 | 798 | 94.2 | 0.990 (P) |
| Maternal height (numerical)                  | 872 | 99.3 | 862 | 98.2 | 860 | 97.9 | 834 | 97.0 | 0.982 (P) |
| Smoking at first ANC visit (Yes/No)          | 875 | 99.7 | 872 | 99.2 | 868 | 98.9 | 843 | 97.1 | 0.742 (C) |
| Use of Snuff at first ANC visit (Yes/No)     | 878 | 100 | 871 | 99.2 | 871 | 99.2 | 861 | 98.9 | 0.540 (C) |
| Assessment of alcohol screening prior to pregnancy (AUDIT) (Yes/No) | 802 | 91.3 | 859 | 97.8 | 788 | 89.7 | 691 | 87.7 | 0.480 (C) |
| If Yes, AUDIT score (numerical)b             | 650/643 | 98.9 | 777/771 | 99.2 | 621 | 95.5 | 600 | 96.6 | 0.989 (P) |
| Prenatal diagnostics                         |     |     |     |     |     |     |     |     |                             |
| Amniocentesis (AC) (Yes/No)                  | 875 | 99.7 | 791 | 90.1 | 788 | 89.7 | 772 | 98.0 | 0.754 (C) |
| Chorion Villus Sampling (CVS) (Yes/No)       | 875 | 99.7 | 790 | 90.0 | 787 | 89.6 | 778 | 98.9 | 0.176 (C) |
| Combined Ultrasound and Biochemical screening (CUB) (Yes/No) | 780 | 88.8 | 791 | 90.1 | 700 | 89.7 | 665 | 95.1 | 0.888 (C) |
| Second trimester Serum Screening (Yes/No)    | 849 | 96.7 | 790 | 90.0 | 767 | 87.4 | 671 | 87.4 | 0.002 (C) |
| Ultrasound examination at 16 – 21 gestational weeks (Yes/No) | 862 | 98.2 | 791 | 90.1 | 779 | 88.6 | 755 | 96.9 | 0.064 (C) |
| Estimated date of delivery (ultrasound) (numerical)c | 871 | 99.2 | 874 | 99.5 | 868 | 98.9 | 781 | 90.0 | 0.946 (S) |
| Oral Glucose Tolerance Test (OGTT) performed (Yes/No) | 869 | 98.9 | 877 | 99.9 | 868 | 98.9 | 842 | 97.0 | 0.854 (C) |
| If Yes, 2-hour value of plasma glucose at OGTT (numerical)d | 100/48 | 48.0 | 104/97 | 93.3 | 46 | 46.0 | 34 | 73.9 | 0.902 (P) |
| Smoking at 32 gestational weeks (Yes/No)     | 858 | 97.7 | 876 | 99.8 | 856 | 97.5 | 849 | 99.1 | 0.864 (C) |
| Use of Snuff at 32 gestational weeks (Yes/No) | 832 | 94.8 | 876 | 99.8 | 830 | 94.5 | 826 | 99.5 | 0.712 (C) |
| Maternal weight, last data entry after 35 gestational weeks (numerical) | 777 | 88.5 | 843 | 96.0 | 760 | 86.6 | 706 | 92.9 | 0.989 (P) |
| No. of ANC visits until estimated date of delivery (numerical) | 877 | 99.9 | 868 | 98.9 | 867 | 98.7 | 627 | 72.3 | 0.915 (P) |
| Date of delivery (numerical)                 | 878 | 100 | 878 | 100 | 878 | 100 | 842 | 95.9 | 0.989 (S) |
| Live born child (Yes/No)                    | 878 | 100 | 878 | 100 | 878 | 100 | 874 | 99.5 | 0.598 (C) |
| Birth weight (numerical)                     | 876 | 99.8 | 869 | 99.0 | 868 | 98.9 | 813 | 93.7 | 0.989 (P) |
Table 3 Data in medical records and the Swedish Maternal Health Care Register (MHCR); comparison between the two data-sets using correlation analysis, and analysis of sensitivity and specificity for binary variables (Continued)

| Gender of infant (Boy/Girl/Sex unknown) | 878 | 100 | 874 | 99.5 | 874 | 99.5 | 862 | 99.2 | 0.973 (C) |
| Singleton birth/multiple births | 877 | 99.9 | 878 | 100 | 877 | 99.8 | 875 | 99.7 | 0.908 (C) |
| Mode of delivery (vaginal/instrumental vaginal/caesarean section) | 876 | 99.8 | 876 | 99.8 | 874 | 99.5 | 857 | 98.0 | 0.946 (C) |
| If caesarean section, elective CS/emergency CS* | 130/115 | 88.5 | 129/128 | 99.2 | 110 | 84.6 | 102 | 92.7 | 0.841 (C) |

Comparison between the two data-sets using correlation analysis, and analysis of sensitivity and specificity for binary variables.

*Correlation analysis: C = Cohen’s kappa, P = Pearson’s correlation coefficient, S = Spearman’s correlation coefficient;

*Measures are calculated for those who have undergone alcohol screening (n = 650). The denominator is the total no of “Yes” responses. Denominator in the Medical records =650. Denominator in the MHCR = 771.

*Measures are calculated for those who have undergone ultrasound.

*Measures are calculated for those who have undergone OGGT. The denominator is the total no of “Yes” responses. The denominator for the medical records = 100. The denominator for the MHCR = 104.

*Measures are calculated for those who have undergone caesarean section. The denominator is the total no of “Yes” responses. The denominator for the medical records = 130. The denominator for the MHCR = 129.

Agreement of data in medical records and in the MHCR

Identical data in both data sources ranged from 73.9% to 99.7%. For more than half of the investigated variables (17 of 27 variables), agreement of data in both data sources reached 95% or more. Five variables reached an agreement of data in both data sources of less than 90% (Table 3). Variables with the highest frequencies of identical information in the MHCR and in the medical records were mainly data related to delivery, such as “singleton birth/multiple births”, “live born child”, and “gender of child”. For the eleven multiple births, the agreement of birth weights of second twin was identical in both data sources (100%).

Table 4 presents the comparison between the primary data collection from the medical records and the re-collection of variables from 150 reinvestigated medical records. The degree of coverage of data in the reinvestigated medical records ranged from 86.7% to 100%; frequencies of available data in medical records were similar or improved at the re-collection with one exception. The re-collection contributed to an improvement of the number of variables with 100% available data in both data sources. In addition, the number of variables with identical data increased in comparison to the first data collection. Identical data in both data sources ranged from 64.0% to 100%. Twenty-two of the 27 variables reached agreement between data sources for 95.0% or more in the reinvestigated data collection. Furthermore, the re-collection of data improved the agreement of data, resulting in only two of the 27 variables showing an agreement in both data sources to less than 90% in the reinvestigated material.

Sensitivity and specificity

Analyses of sensitivity and specificity were performed on binary variables (Table 3). The medical record was considered to represent the true value. Sensitivity varied from 0.043 (second trimester screening) to 0.999 (live born child), and specificity ranged from 0.393 (assessment of alcohol screening prior to pregnancy) to 0.999 (use of snuff at 32 gestational weeks). For nine out of the 12 binary variables, specificity was 0.900 or higher, whereas only four out of 12 binary variables had a sensitivity of 0.900 or higher.

Systematic errors

Possible systematic errors were identified for two variables: “second trimester serum screening” and “number of ANC visits”. The variable “second trimester serum screening” demonstrated identical information in both data sources for 87.4%. One of the hospitals reported an unexpected large number of performed second trimester screenings in both data sources. The reported number of “second trimester serum screening” was not consistent with the clinical practice, so we discussed this issue with the midwives working in the catchment area of this hospital. These discussions revealed that the variable “second trimester serum screening” probably had been misunderstood, resulting in incorrect reporting of data.

The variable “number of ANC visits” showed an agreement of data in both data sources for 72.3% of the cases. The information addressing this variable in the Web application was defined as the number of visits to see a midwife at an ANC (noted on the ANC registration) until estimated date of delivery as established by ultrasound (not the actual date of birth). As pregnant women may meet other health care providers during pregnancy, such visits may have been included in the figure entered in the MHCR. A misfit of ± 1 visit was seen in 19.3% of the cases. The variation of misfitting values ranged from -7 visits to + 8 visits.
Table 4 Comparison between primary collection and re-collection of data from medical records using correlation analysis, and analysis of sensitivity and specificity for binary variables

| Variable                                                                 | Medical records<sup>a</sup> | Medical records re-collection<sup>b</sup> | Data available in both data sources | Identical information in both data sources | Correlation<sup>c</sup> | Sensitivity | Specificity |
|--------------------------------------------------------------------------|-------------------------------|-------------------------------------------|-------------------------------------|---------------------------------------------|------------------------|-------------|-------------|
| Variables collected at first antenatal care (ANC) visit                  |                               |                                           |                                     |                              |                        |             |             |
| Date of first visit in ANC (numerical)                                   | 150 100                       | 150 100                                   | 150 100                             | 116 77.3                                   | 0.773 (S)              |             |             |
| No of previous deliveries (numerical)                                    | 150 100                       | 150 100                                   | 150 100                             | 149 99.3                                   | 0.988 (P)              |             |             |
| Maternal weight at first ANC visit (numerical)                          | 147 98.0                      | 148 98.7                                  | 147 99.3                            | 146 99.3                                   | 0.959 (P)              |             |             |
| Maternal height (numerical)                                             | 149 99.3                      | 149 99.3                                  | 149 100                             | 148 99.3                                   | 1.000 (P)              |             |             |
| Smoking at first ANC visit (Yes/No)                                     | 149 99.3                      | 149 99.3                                  | 149 100                             | 148 99.3                                   | 0.794 (C)              |             |             |
| Use of Snuff at first ANC visit (Yes/No)                                | 150 100                       | 150 100                                   | 150 100                             | 146 97.3                                   | 0.793 (C)              | 0.667       | 1.000       |
| Assessment of alcohol screening prior to pregnancy (AUDIT) (Yes/No)     | 130 86.7                      | 130 86.7                                  | 130 86.7                            | 121 93.1                                   | 0.729 (C)              |             |             |
| If Yes, AUDIT score (numerical)<sup>h</sup>                            | 113 96.5                      | 108 98.1                                  | 102 90.3                            | 100 98.0                                   | 0.987 (P)              |             |             |
| Variables collected at 4 to 16 weeks postpartum                         |                               |                                           |                                     |                              |                        |             |             |
| Prenatal diagnostics                                                    |                               |                                           |                                     |                              |                        |             |             |
| Amniocentesis (AC) (Yes/No)                                             | 150 100                       | 150 100                                   | 150 100                             | 150 100                                   | 1.000 (C)              | 1.000       | 0.983       |
| Chorion Villus Sampling (CVS) (Yes/No)                                  | 150 100                       | 150 100                                   | 150 100                             | 150 100                                   | 1.000 (C)              |             | 0.992       |
| Combined Ultrasound and Biochemical screening (CUB) (Yes/No)            | 147 98.0                      | 149 99.3                                  | 147 98.6                            | 142 96.6                                   | 0.912 (C)              | 0.919       | 0.941       |
| Second trimester Serum Screening (Yes/No)                               | 148 98.7                      | 150 100                                   | 148 98.7                            | 148 100                                   | 0.246 (C)              |             |             |
| Ultrasound examination at 16 – 21 gestational weeks (Yes/No)            | 147 98.0                      | 147 98.0                                  | 147 98.0                            | 145 99.0                                   | 0.955 (S)              |             |             |
| Estimated date of delivery (ultrasound) (numerical)<sup>i</sup>         | 147 100                       | 147 100                                   | 147 100                             | 145 98.7                                   | 0.246 (C)              |             |             |
| Oral Glucose Tolerance Test (OGTT) performed (Yes/No)                   | 149 99.3                      | 149 99.3                                  | 148 98.0                            | 144 98.0                                   | 0.819 (C)              | 1.000       | 0.986       |
| If Yes, 2-hour value of plasma glucose at OGTT (numerical)<sup>j</sup>  | 13/10 77.0                    | 10/9 90.0                                 | 9 69.2                              | 9 100                                      | 1.000 (P)              |             |             |
| Smoking at 32 gestational weeks (Yes/No)                                | 145 96.7                      | 145 96.7                                  | 145 100                             | 145 100                                   | 1.000 (C)              | 1.000       | 1.000       |
| Use of Snuff at 32 gestational weeks (Yes/No)                           | 145 96.7                      | 145 96.7                                  | 145 100                             | 144 99.3                                   | 0.797 (C)              | 1.000       | 1.000       |
| Maternal weight, last data entry after 35 gestational weeks (numerical) | 142 94.7                      | 141 94.0                                  | 141 99.3                            | 137 97.2                                   | 1.000 (P)              |             |             |
| No. of ANC visits until estimated date of delivery (numerical)          | 150 100                       | 150 100                                   | 150 100                             | 96 64.0                                    | 0.890 (P)              |             |             |
| Date of delivery (numerical)                                            | 150 100                       | 150 100                                   | 150 100                             | 149 99.3                                   | 0.975 (S)              |             |             |
| Live born child (Yes/No)                                                | 150 100                       | 150 100                                   | 150 100                             | 150 100                                   | 0.975 (S)              |             |             |
Discussion

This is the first time that the validity of data entered in the MHCR has been investigated. Data from 878 medical records were compared with corresponding data registered in the MHCR. The information registered in the medical records was regarded as the gold standard. Data entered in the MHCR presented a strong correlation to corresponding data in the medical records. More than half of the variables under study demonstrated identical information in both data sources to a level of 95% or more. Five of the 27 studied variables showed an agreement of less than 90% in both data sources. A second re-collection of the same variables of a subset of 150 medical records of the original sample, performed to further validate the primary data collection in this study, increased the number of variables with identical information in both data sources. Possible sources of systematic errors in the MHCR were identified for two variables.

Degree of coverage of data

The findings of this study presented a sufficient degree of coverage of data in the medical records under study. Data from the medical records have been transferred to the Swedish Medical Birth Register (MBR) since 1973. Previous studies have shown that most variables in the MBR demonstrate sufficient degree of coverage of data [12,13].

The estimated proportion of registered pregnancies in MHCR during 2010 and 2011 were 81% and 85%, respectively (personal communication). Missing MHCR data could be the result of midwives failing to enter data for all pregnant women as this work task is not compulsory and the fact that providing data is voluntarily (i.e., pregnant women can choose to opt out). However, missing data related to opting out is considered a minor issue (personal communication).

The degree of coverage of data entered in the MHCR was high for most variables in our study. The data in the MHCR were entered by the midwife working in the ANC; some information was available in the medical records and some information was provided orally by the pregnant woman. The variables regarding prenatal diagnostics in the MHCR demonstrated a relatively lower degree of coverage than other included variables, although it was still high. A possible explanation for this relatively lower degree of coverage may be the design of this question in the MHCR Web application. Only after the midwife registered “yes” for the question “Have any foetal diagnostics been performed?” is the second option displayed. In the Swedish MBR, an improvement of data quality regarding amniocentesis and chorionic villus sampling was found when the location of these variables in the medical records was changed [11]. Hence, rephrasing and redesigning these questions in the Web application may further improve the degree of coverage of data for variables related to prenatal diagnostics.

To our knowledge, no previous studies have monitored how primary data are registered in the medical records or have investigated the validity of primary data in relation to data included in the medical records. Our study shows that some variables demonstrated a higher degree of coverage in the MHCR than in the medical records. Some studies that use vital statistics databases...

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Table 4 Comparison between primary collection and re-collection of data from medical records using correlation analysis, and analysis of sensitivity and specificity for binary variables (Continued)

| Birth weight (numerical) | 150 100 150 100 150 100 140 93.3 0.997 (P) |
|--------------------------|-----------------------------------------------|
| Gender of infant         | 150 100 150 100 150 100 149 99.3 0.987 (C) |
| Singleton birth/multiple births | 150 100 150 100 150 100 100 1.000 (C) |
| Mode of delivery         | 150 100 150 100 150 100 149 99.3 0.983 (C) |
| If caesarean section     | 23/22 95.7 23/23 100 22 95.7 22 100 1.000 (C) |

*Primary collection of data from medical records.
*Re-collection of data from medical records.
Correlation analysis: C = Cohen’s kappa, P = Pearson’s correlation coefficient, S = Spearman’s correlation coefficient.
Sensitivity or specificity not possible to calculate since one or more of the cells in the calculation includes zero.
*Ma...
for perinatal epidemiology have a major limitation: the
data these studies use, although considered the gold
standard, have not been evaluated for their reliability
and validity [14].

**Agreement between data sources**
The agreement of data in both data sources was high for
most variables (Tables 3 and 4). To analyse correlation
of categorical data, Cohen's kappa was used. Cohen's
kappa is defined only for a square table and is strongly
influenced by prevalence (e.g., number of “yes” responses).
When there is a high level of correlation between two
variables and when one of four cells is empty, the per-
formance of Cohen's kappa declines. This decline was the
case for the variable “use of snuff”, where Cohen's kappa
was calculated to 0.540, although data were identical for
98.9% of cases in the medical records and in the MHCR.
Another example was the variable “chorionic villus sam-
pling”, where Cohen's kappa was calculated to be 0.176,
although the proportion of identical data in medical
records and in the MHCR reached as high as 98.9%. In
these cases, the proportion of identical information in
both data sources provided more valuable information
than Cohen's kappa provided.

Our findings of agreement between the data sources
were similar to the findings reported in a pilot study that
assessed data quality in the Uniform Data Set (UDS)
used by the American Association of Birth Centers [15].
In this pilot study, a care provider entered data online
on four occasions; the data addressed demographic charac-
teristics, risk factors, process of care, and maternal
and infant outcomes. The agreement of variables from
medical records and the UDS varies from 87.5% to
100%.

In an American evaluation of the use of electronic
health records in emergency medical services, electronic
data processing was compared to manual data processing.
The results show good to excellent agreement between
the two approaches [16]. In the Swedish setting, there is a
disadvantage when data are entered in the MHCR, as data
from the medical records currently cannot be automa-
tically exported to the MHCR. All registrations in the
MHCR are made manually by midwives in an ANC.
Despite these potential sources of manual mistakes when
registering data, the findings in our study indicate that
the accuracy of data registered in the MHCR reaches a level
of good to very good.

**Sensitivity and Specificity**
Variables characterized by one of the binary response
options (“yes” and “no”) demonstrated either a high
specificity and a low sensitivity or a low specificity
and a high sensitivity. Binary variables demonstrating a
high specificity and a low sensitivity were “use of snuff”,
“smoking”, “chorion villus sampling”, and “second trimes-
ter screening”. In contrast, variables characterized by a
majority of “yes” responses demonstrated high sensitivity
and low specificity (i.e., “assessment of alcohol screening
prior to pregnancy”, “ultrasound examination at 16-21
gestational weeks”, and “live born child”). These results
indicate that midwives performing data entry are more
prone to enter results that are expected than unexpected.
Similarly, an American study investigating the correctness
of data in a computerized perinatal database found that
there is greater likelihood to overlook a given diagnosis
than to score positive a disease that does not occur [17]. A
review on quality of data in perinatal health databases,
including 43 validation studies, shows that most con-
ditions and procedures demonstrate high specificities,
indicating few false positives [18]. Most of the binary
variables in our study demonstrated a low prevalence
of the investigated outcome. This finding may explain
why only four of 12 variables showed a sensitivity ex-
ceeding 0.900.

**Systematic errors**
This study revealed two potential systematic errors when
registering data in the MHCR. First, the analysis demon-
strated a misinterpretation at one of the participating
hospitals regarding the registration of “second trimester
serum screening” in the catchment area. An English
study reveals that some midwives (7.7%) believe that
they are not sufficiently prepared to inform patients
about available foetal screening methods. The majority
of midwives feel they are prepared to offer their patients
information about screening, but when testing the level
of knowledge of the conditions detectable by the avail-
able screening tests, the knowledge does not match the
preparedness [19]. The situation presented in the English
study might be applicable to the Swedish setting as well.
The available methods for prenatal screening and prenatal
diagnostics have rapidly increased over the last decade,
resulting in more complex information and counselling
needs [10], so some midwives working in an ANC might
not have had sufficient knowledge to correctly enter data
in the MHCR. The second possible systematic error found
was when addressing the number of ANC visits during
pregnancy. A quality study of the Swedish Medical Birth
Register found that information on the number of ANC
visits is missing in approximately 11% of the cases [10].
Our study found that the degree of coverage of this
variable was high for both data sources, but the agreement
between the data sources was not as high. A possible
source for the lower accuracy could be related to insuf-
ficient instructions in the MHCR manual. Most of the
incorrect values ranged ± one visit; a possible explanation
for this is that visits after 40 gestational weeks or visits to
the outpatient specialised clinic might have been included

http://www.biomedcentral.com/1472-6963/14/364
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in the MHCR data. Improvements in the MHCR user manual might increase the level of correct data in the MHCR.

Clinical importance
Quality register data are used for quality improvement and management within the health services as well as for research purposes. Therefore, it is of considerable importance that the improvements, decision-making, and results presented must be grounded in reliable and valid data. The benefit of the MHCR is the composition of the data, which include demographic, medical, and psychological aspects of the pregnancy, the delivery, and the postpartum period. Additionally, the data may be presented on a local, regional, and national level of the ANC, enabling comparisons of provided health care and outcomes of pregnancy and delivery. Despite manually registering data in the MHCR, the vast majority of variables included in the MHCR show very good agreement with corresponding information in the medical records. The findings in this study indicated that the data available from the MHCR are reliable enough to be used in clinical quality work and for research purposes.

Further studies
As the data are registered manually in the MHCR by midwives in an ANC, the experiences of midwives is important to address – How do midwives experience this work? Furthermore, it would be of interest to find out how data available in the MHCR are used for clinical improvements and quality aspects of health care at the local and regional levels of the ANC.

Methodological considerations
One of this study’s strengths is its design. Data were extracted and analysed in two steps: a primary data extraction from 878 medical records and a secondary data extraction of the same variables for a subset of 150 medical records from the primary sample. The re-collection of data was performed by two midwives (i.e., two of this paper’s authors) with extensive experience working in an ANC. This experience may have contributed to the improved quality of the data with increased statistical agreement between datasets. Data extracted by professionals other than midwives might be less accurate as these professionals may have much less experience evaluating and registering this type of data, a disadvantage that may have led to problems identifying the correct information.

Another strength of this study is the geographical variation of the included hospitals. The data extraction was performed at clinics in large cities as well as in small clinics located in more rural areas in Sweden. The selection of hospitals, in part, was determined by convenience as some of the authors were affiliated with five of these hospitals. Four other hospitals were selected with complementary characteristics in relation to the first five selected hospitals. The first author had close contact with the administrators at these hospitals in order to enhance the quality of the data collection. We believe that the selected hospitals sufficiently reflect the general characteristics of clinical settings in contemporary hospitals and ANC in Sweden.

The goal was to collect data for 900 medical records, 100 medical records from each hospital. In 2011, the degree of coverage of data was 85% in the MHCR (personal communication); that is, data were not available in the MHCR for 15% of pregnant women in Sweden for 2011. To identify 100 consecutive individuals with data in both medical records and the MHCR, we first collected the personal identity number of 120 individuals in the birth logs (from March 1), resulting in the identification of 100 women who had delivered at each hospital. Despite considerable efforts, this goal was not achieved as some administrators did not fully complete the Excel-protocols. Administrators of seven of the nine hospitals did not provide complete datasets. However, the number of missing cases (n = 22) corresponds to 2.4% of the goal, indicating that these missing cases could not have critically influenced the results of this study. Mean background characteristics on maternal age, height, weight, and BMI were 30.7 yrs, 166.2 cm, 67.9 kg, and 24.6 kg/m², respectively in our study. The corresponding results in the MHCR for 2011 (N = 89,313) were 30.7 yrs, 166.2 cm, 68.4 kg, and 24.7 kg/m², indicating that the study sample was representative for the year under study (personal communication).

Conclusions
Comparing data from medical records – the gold standard – with data registered in the MHCR, we found that most variables in the MHCR demonstrated good to very good degree of coverage of data, agreement, and internal validity. Hence, data from the MHCR may be regarded as reliable when used for evaluation, planning, and decision-making in Swedish maternal health care services as well as for research purposes.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
KP, IM, MP, and ML designed the study, organized the data collection, performed analyses of materials, and drafted the manuscript. ML contributed specifically with statistical competence. MH contributed to study design, interpretation of results, and drafting of the manuscript. CN contributed to study design, data collection, and drafting of manuscript. IH and YS contributed to data collection and drafting of the manuscript. All authors read and approved the final manuscript.
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