Hospital factors and metastatic surgery in colorectal cancer patients, a population-based cohort study

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Abstract

Background: Only a limited proportion of patients with metastatic colorectal cancer (mCRC) receives metastatic surgery (including local ablative therapy). The aim was to investigate whether hospital volume and hospital level were associated with the chance of metastatic surgery.

Methods: This national cohort retrieved from the CRCBaSe linkage included all Swedish adult patients diagnosed with synchronous mCRC in 2009–2016. The association between annual hospital volume of incident mCRC patients and the chance of metastatic surgery, and survival, were assessed using logistic regression and Cox regression models, respectively. Hospital level (university/non-university) was evaluated as a secondary exposure in a similar manner. Both uni- and multivariable (adjusted for sex, age, Charlson comorbidity index, year of diagnosis, cancer characteristics and socioeconomic factors) models were fitted.

Results: A total of 1,674 (17%) out of 9,968 mCRC patients had metastatic surgery. High hospital volume was not associated with increased odds of metastatic surgery after including hospital level in the model, whereas hospital level was (odds ratio (OR) (95% confidence interval (CI)): 1.94 (1.68–2.24)). All-cause mortality was lower in university versus non-university hospitals (hazard ratio (95% CI): 0.83 (0.78–0.88)).

Conclusions: Patients with mCRC initially cared for by a university hospital experienced a greater chance to receive metastatic surgery and had superior overall survival. High hospital volume in itself was not associated with a greater chance to receive metastatic surgery nor a greater survival probability. Additional efforts should be imposed to provide more equal care for mCRC patients across Swedish hospitals.

Keywords: Colorectal neoplasms, High-volume hospitals, Low-volume hospitals, Neoplasm metastasis, Secondary, Surgery, Survival, University hospitals

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compared to systemic treatment only, is associated with superior OS for patients with unresectable liver metastases [4]. Metastatic surgery (including non-surgical local ablative treatments) in combination with primary tumour resection can offer selected patients the chance of cure.

In a universal healthcare system, such as that in Sweden, one of the most important principles is equal access to care [5]. The healthcare system in Sweden is decentralised into six healthcare regions and 21 regions, which are responsible for managing and prioritising their own healthcare resources. All six healthcare regions have one university hospital, except one that has two. A few university hospitals have multiple physical locations. Surgery of metastases for mCRC is limited to a few hospitals, mainly the university hospitals. Cytoreductive surgery with hyperthermic intraperitoneal chemotherapy (HIPEC) is further centralised to four university hospitals. Previous studies on mCRC have found that hospital factors, such as hospital level (university hospital/non-university hospital) and geographic location, are associated with the chance of receiving surgery for liver metastases [6–13], peritoneal metastases [14], pulmonary metastases [15] and mCRC overall [16, 17].

An increased chance of metastatic surgery for mCRC has also been associated with being discussed at a multidisciplinary team (MDT) conference [18, 19]. This has contributed to the recommendation that as of 2016, all CRC patients in Sweden should be discussed at MDT conferences, and in the presence of liver metastases, with a liver surgeon present [20]. In the recommendations from 2008 CRC patients with limited metastases were to be considered potentially curable and therefore further assessed in a multidisciplinary setting [21].

Increased hospital and surgeon volume are associated with improved outcomes for CRC patients undergoing surgery [22–25]. In the presence of metastases an equivalently important quality of care measure is related to the patients who do not undergo surgery. To explore whether differences in provided care exist, all patients with mCRC need to be identified and included. Studies of pancreatic cancer patients have shown that hospital volume based on all incident cases, both surgically and non-surgically treated, is associated to a higher frequency of surgery and/or chemoradiation [26].

Sweden has several high-quality registers including the Swedish colorectal cancer register (SCRCR), which had a coverage of 98.5% of colon adenocarcinomas and 98.8% of rectal adenocarcinomas in 2008–2016 [27]. All Swedish citizens and residents are registered in the Swedish Population Register and have unique personal identity numbers, which make linkage between registries possible. This provides an excellent opportunity to study all patients diagnosed with mCRC, regardless of what treatment they have later received. To our knowledge, the potential association between hospital volume of incident mCRC patients and the chance of having metastatic surgery has not been reported previously.

In this large nationwide Swedish study, variations in the proportion of mCRC patients receiving metastatic surgery between hospitals of different volume and level was evaluated. Being discussed at an MDT conference was explored as a mediator. It was hypothesised that hospital volume and level of hospital were associated with the chance of metastatic surgery and with OS.

**Methods**

**Register data**

This study is based on the Colorectal Cancer Database (CRCBaSe), a mega-linkage originating from the SCRCR, which was further linked to national registers at the National Board of Welfare and Statistics Sweden. The SCRCR was used to identify patients, to get information on the registering hospital and on surgery of metastases and primary tumour, and to retrieve data on potential confounders and descriptive variables. When a patient is diagnosed with CRC, the hospital is responsible for registering the patient in the SCRCR and the managing hospital is automatically recorded when starting a registration in the SCRCR.

The Swedish Cancer Register (SCR), which dates back to 1958, was used to retrieve data on other cancer diagnoses and include data on previous CRC diagnoses not captured by the SCRCR, to ensure that only the first CRC diagnosis was included. Information from the Cause of Death Register was used to retrieve information on cause and date of death. The In- and Out-Patient Register was used to obtain information on comorbidities and to define presence of metastatic disease, receipt of primary tumour resection and surgery of lung, liver and peritoneal metastases (Supplementary Table 1). From the Register of the Total Population, data on civil status and migration was extracted. From the Longitudinal Integrated Database for Health Insurance and Labour Market Studies data on disposable income per family unit and educational level were extracted. Civil status and highest achieved level of education were defined at the time of CRC diagnosis, while for income, information on the year prior to CRC diagnosis was used. If data was unavailable for the mentioned year, data from 0–3 years before year of diagnosis was used instead.

**Inclusion and exclusion criteria**

All Swedish adult patients (≥ 18 years old) with a first-time CRC diagnosis in the SCRCR during 2009–2016, with no previous diagnosis of CRC in SCR, were included (n = 46,160). The following exclusions criteria were
applied: no presence of synchronous metastatic disease according to a clinical or pathological assessment \((n=35,737)\), no specified metastatic location \((n=451)\), and missing data on registering hospital \((n=4)\). The final study population consisted of 9,968 patients (Fig. 1).

**Exposures and outcomes**

Our primary exposure was annual hospital volume of incident mCRC patients, i.e., the total number of patients diagnosed with mCRC at a specific hospital during a given year. The annual hospital volume was divided into quartiles \((\leq 25^\text{th}, > 25^\text{th} \text{ to} \leq 50^\text{th}, > 50^\text{th} \text{ to} \leq 75^\text{th}, \text{ and} > 75^\text{th})\). As such, the same hospital could be exposed as a 1st quartile hospital one year and then fall into another quartile of hospital volume another year. Additionally, level of hospital (university or non-university) and its association with odds of metastatic surgery was investigated. Our primary outcome was receiving metastatic surgery \((\text{yes/no})\), which was defined as receiving surgical resection including a non-surgical locally ablative procedure of liver, and/or peritoneal, and/or lung metastases within 90 days prior to and 270 days after the date of primary CRC diagnosis date (Supplementary Table 2). This time window was chosen to encompass the majority of treatments for synchronous metastases, but not include treatment for metachronous metastases. Furthermore, it was required that the patient underwent primary tumour resection, either before, simultaneously or after the surgery of the metastases, to be defined as having received the primary outcome. Our secondary outcome was OS from date of diagnosis.

**Statistical methods**

Frequencies and proportions of patient and tumour characteristics were calculated separately within hospital volume quartiles. The proportion of patients receiving metastatic surgery by year of diagnosis, hospital volume quartiles, hospital level, and healthcare region, were visualised using bar charts. Chi-squared tests were used for formal testing of differences in distributions, and logistic regression was applied to evaluate trends.

Crude and adjusted logistic regression models were fitted to estimate odds ratios \((\text{ORs})\) with 95% confidence intervals \((\text{CIs})\) of receiving metastatic surgery. The first multivariable model included one exposure (hospital volume or level) and adjusted for sex (female, male), age \((18–64 \text{ years}, 65–79 \text{ years and} \geq 80 \text{ years old})\), Charlson comorbidity index \((\text{CCI}) [28] \text{ (0, 1 and} \geq 2)\), year of diagnosis \((\text{continuous} 2009–2016)\), location of the primary tumour \((\text{right colon, transverse colon, left colon, and} \text{unknown location in colon or rectum})\), clinical tumour \((\text{cT})\) stage \((1–3, 4, X)\), clinical nodal \((\text{cN})\) stage \((0, 1–2, X)\), extent of metastatic disease \((\text{single location/multiple locations})\), civil status \((\text{married/registered partner, or not married})\), highest commenced educational level \((\text{primary school, secondary school, or higher education})\) and disposable income per family unit \((\text{divided into four quartiles per year of diagnosis})\). A second multivariable model was constructed including both exposures (hospital volume and level) and adjusted for the variables mentioned above. An additional multivariable model also included the potential mediator MDT \((\text{yes/no})\). Tests for interactions between hospital volume and hospital level, and hospital level and MDT conference, were performed using a Wald test. Missing data were included in the adjusted models by use of the missing-indicator method [29].

For the survival analyses, the patients were followed from date of diagnosis until date of death, migration, or 31st December 2017, whichever occurred first. The underlying time scale was time since diagnosis. The
Kaplan–Meier method was used to calculate OS proportions, and the log rank test to test for survival differences, by hospital volume and hospital level. Cox regression models were used to estimate hazard ratios (HRs) with 95% CIs of the all-cause mortality rate for the four hospital volume quartiles and hospital level. Both univariable and multivariable models were fitted, where the latter adjusted for the same variables as described in the multivariable logistic regression models above. The proportional hazard assumption was formally tested using the Schoenfeld residuals.

A sensitivity analysis was performed using a delayed entry model, in which follow-up started at 270 days after diagnosis. Patients who died (n = 3,964) or were administratively censored (n = 11) during these 270 days were not included in this analysis (n_included = 5,993). The Kaplan–Meier method was used to calculate OS proportions, and the log rank test to test for survival differences, by whether the patients received metastatic surgery or not.

The significance level was set to 5%. Statistical analyses were done using Stata (StataCorp. 2017. Stata Statistical Software: Release 16. College Station, TX: StataCorp, LLC).

Ethical approval
Ethical approval was given by the Regional Board of the Ethical Committee in Stockholm, Sweden (DNR: 2014/71–31, 2018/328–32, 2021–00,342) and the study was conducted in accordance with the ethical standards described.

Results
A total of 9,968 adult patients diagnosed with mCRC in Sweden 2009–2016 met the study criteria (Fig. 1). Of these, 1,674 (17%) had surgery of their metastases during the pre-defined time window around diagnosis and primary tumour resection (procedures performed are shown in Supplementary Table 2). The proportion of patients with mCRC who had metastatic surgery increased from 12% in 2009 to 20% in 2016 (Fig. 2A, p for trend < 0.001).

The annual volumes of incident mCRC patients at hospitals in the 1st, 2nd, 3rd, and 4th quartiles were 1–20, 21–31, 32–46, and 47–103 patients per year, respectively. The proportion of patients receiving surgery for mCRC increased from 13% in the 1st quartile to 21% in the 4th quartile (Fig. 2B, p for trend < 0.001). The proportion receiving metastatic surgery was 23% of patients initially cared for by a university hospital compared to 14% at non-university hospitals (Fig. 2C, p < 0.001). During the period, 20% of patients diagnosed with mCRC received surgery in healthcare region 1 and 3 compared with...
14% and 15% in healthcare region 5 and 2, respectively (Fig. 2D, \( p < 0.001 \)).

**Patient, hospital and tumour characteristics**

Patients managed at high-volume hospitals (4th quartile) were younger at diagnosis, had reached a higher level of education and a higher disposable income (Table 1). There were no differences in CCI (Table 1). Approximately half of the patients underwent resection of the primary tumour, a third had liver metastases only, and about half of the patients had metastatic disease at multiple locations (Table 2). Around 24% (\( n = 327/1,359 \)) of the patients who underwent primary tumour resection and liver surgery had synchronous primary tumour and liver surgery. The proportion of patients initially cared for at a university hospital ranged between 3% in the 1st quartile and 77% in the 4th quartile of hospital volume.

Data on race/ethnicity of study participants was not available.

**Metastatic surgery**

The chance of receiving metastatic surgery increased with hospital volume, compared to the 1st quartile; 2nd quartile OR (95% CI): 1.28 (1.08–1.52), 3rd quartile OR (95% CI): 1.33 (1.12–1.58), 4th quartile OR (95% CI): 1.68 (1.42–1.98), from adjusted models (Fig. 3 and Supplementary Table 3). The chance of metastatic surgery was higher in university hospitals than non-university hospitals (OR (95% CI): 1.89 (1.68–2.12). After including both exposures, hospital volume and hospital level, there was no association between hospital volume and outcome. The same analysis revealed that patients managed at a university hospital had increased odds of having metastatic surgery (OR (95% CI): 1.94 (1.68–2.24)) also after adjustment for hospital volume (Fig. 3 and Table 1)

### Table 1: Patient characteristics according to annual hospital volume of patients with metastatic colorectal cancer

|                                | First quartile | Second quartile | Third quartile | Fourth quartile |
|--------------------------------|---------------|-----------------|---------------|----------------|
| **Sex**                        |               |                 |               |                |
| Male                           | 1,325 (57%)   | 1,455 (56%)     | 1,386 (55%)   | 1,400 (56%)    |
| Female                         | 1,003 (43%)   | 1,136 (44%)     | 1,145 (45%)   | 1,118 (44%)    |
| **Age**                        |               |                 |               |                |
| Median (years)                 | 71            | 70              | 71            | 69             |
| 18–64 years                    | 631 (27%)     | 784 (30%)       | 757 (30%)     | 889 (35%)      |
| 65–79 years                    | 1,182 (51%)   | 1,265 (49%)     | 1,221 (48%)   | 1,169 (46%)    |
| ≥ 80 years                     | 515 (22%)     | 542 (21%)       | 553 (22%)     | 460 (18%)      |
| **Charlson comorbidity index** |               |                 |               |                |
| 0                              | 1,469 (63%)   | 1,600 (62%)     | 1,523 (60%)   | 1,543 (61%)    |
| 1                              | 229 (10%)     | 287 (11%)       | 290 (11%)     | 264 (10%)      |
| ≥ 2                            | 630 (27%)     | 704 (27%)       | 718 (28%)     | 711 (28%)      |
| **Year of diagnosis**          |               |                 |               |                |
| 2009–2012                      | 1,228 (55%)   | 1,359 (52%)     | 1,198 (47%)   | 980 (39%)      |
| 2013–2016                      | 1,040 (45%)   | 1,232 (48%)     | 1,333 (53%)   | 1,538 (61%)    |
| **Civil status**               |               |                 |               |                |
| Married/registered partner     | 1,238 (54%)   | 1,416 (55%)     | 1,329 (53%)   | 1,309 (52%)    |
| Not married: never married, divorced, widowed | 1,070 (46%) | 1,175 (45%) | 1,201 (47%) | 1,208 (48%) |
| Missing                        | 0 (0%)        | 0 (0%)          | 1 (0%)        | 1 (0%)         |
| **Highest level of commenced education** |               |                 |               |                |
| Primary school (maximum 9 years) | 909 (39%) | 1,011 (39%)     | 965 (38%)     | 775 (31%)      |
| Secondary school (3–4 additional years) | 945 (41%) | 1,011 (39%) | 993 (39%) | 1,039 (41%) |
| Higher education               | 459 (20%)     | 531 (20%)       | 550 (22%)     | 680 (27%)      |
| Missing                        | 14 (1%)       | 38 (1%)         | 24 (1%)       | 24 (1%)        |
| **Disposable income per family unit divided into annual quartiles.*** |               |                 |               |                |
| 1st quartile                   | 676 (29%)     | 653 (25%)       | 624 (25%)     | 543 (22%)      |
| 2nd quartile                   | 609 (26%)     | 646 (25%)       | 648 (26%)     | 584 (23%)      |
| 3rd quartile                   | 569 (24%)     | 685 (26%)       | 633 (25%)     | 605 (24%)      |
| 4th quartile                   | 474 (20%)     | 607 (23%)       | 625 (25%)     | 785 (31%)      |
| Missing                        | 0 (0%)        | 0 (0%)          | 1 (0%)        | 1 (0%)         |

Includes 9,968 patients diagnosed with metastatic colorectal cancer in 2009–2016 in Sweden

* From chi square test
** Year of/up to 3 years before if missing
*** Year before diagnosis or up to 3 years before or year of if missing
There was a significant interaction between hospital volume and hospital level (p from Wald test < 0.001). This can be interpreted as the effect of hospital volume was significantly different for university hospitals compared with non-university hospitals.

**MDT and temporal trends**

The proportion discussed at a pretherapeutic MDT conference increased from 60% in 2009 to 86% in 2016 (p-value from trend test < 0.001). In 2009, 73% of mCRC patients at university hospitals were discussed at MDT conferences and 54% at non-university hospitals (p < 0.001). The proportion being discussed at MDT conferences were similar in 2016, 86% vs 85% at university and non-university hospitals, respectively (p = 0.777).

In light of the diminishing differences in the proportion being discussed at an MDT by hospital level at the end of the study period, we investigated if the same was evident for differences in metastatic surgery. The proportion receiving metastatic surgery at a university hospital compared to a non-university hospital was 18% vs. 10% in 2009 (p < 0.001) and 28% vs 16% in 2016 (p < 0.001).

When investigating how the OR for the association between hospital level and receiving metastatic surgery

### Table 2  Hospital and tumour characteristics according to annual hospital volume of patients with metastatic colorectal cancer

|                      | First quartile n = 2,328 | Second quartile n = 2,591 | Third quartile n = 2,531 | Fourth quartile n = 2,518 | All n = 9,968 | P-value* |
|----------------------|---------------------------|---------------------------|---------------------------|---------------------------|----------------|----------|
| Hospital level       |                           |                           |                           |                           |                |          |
| University           | 79 (3%)                   | 492 (19%)                 | 669 (26%)                 | 1,928 (77%)               | 3,168 (32%)    | <0.001   |
| Non-university       | 2,249 (97%)               | 2,099 (81%)               | 1,862 (74%)               | 590 (23%)                 | 6,800 (68%)    |          |
| Hospital region      |                           |                           |                           |                           |                |          |
| 1                    | 221 (9%)                  | 370 (14%)                 | 635 (25%)                 | 741 (29%)                 | 1,967 (20%)    | <0.001   |
| 2                    | 505 (22%)                 | 608 (23%)                 | 869 (34%)                 | 273 (11%)                 | 2,255 (23%)    |          |
| 3                    | 370 (16%)                 | 449 (17%)                 | 231 (9%)                  | 0 (0%)                    | 1,050 (11%)    |          |
| 4                    | 390 (17%)                 | 459 (18%)                 | 407 (16%)                 | 763 (30%)                 | 2,019 (20%)    |          |
| 5                    | 390 (17%)                 | 401 (15%)                 | 288 (11%)                 | 741 (29%)                 | 1,820 (18%)    |          |
| 6                    | 452 (19%)                 | 304 (12%)                 | 101 (4%)                  | 0 (0%)                    | 857 (9%)       |          |
| Primary tumour location |                        |                           |                           |                           |                |          |
| Right colon          | 615 (26%)                 | 637 (25%)                 | 643 (25%)                 | 676 (27%)                 | 2,568 (26%)    | <0.001   |
| Transverse colon     | 314 (13%)                 | 292 (11%)                 | 316 (12%)                 | 306 (12%)                 | 1,227 (12%)    |          |
| Left colon           | 743 (32%)                 | 761 (29%)                 | 732 (29%)                 | 673 (27%)                 | 2,912 (29%)    |          |
| Unknown location in colon |                 |                           |                           |                           |                |          |
| Rectum               | 625 (27%)                 | 882 (34%)                 | 831 (33%)                 | 844 (34%)                 | 3,193 (32%)    |          |
| Pretherapeutic multidisciplinary team conference | | | | | | |
| Yes                  | 1,507 (65%)               | 1,967 (76%)               | 1,950 (77%)               | 2,011 (80%)               | 7,435 (75%)    | <0.001   |
| No                   | 812 (35%)                 | 621 (24%)                 | 570 (23%)                 | 498 (20%)                 | 2,510 (25%)    |          |
| Missing              | 8 (0%)                    | 3 (0%)                    | 3 (0%)                    | 9 (0%)                    | 23 (0%)        |          |
| Resection of primary tumour | | | | | | |
| No                   | 1,128 (48%)               | 1,349 (52%)               | 1,397 (55%)               | 1,322 (53%)               | 5,196 (52%)    | <0.001   |
| Yes                  | 1,199 (52%)               | 1,242 (48%)               | 1,135 (45%)               | 1,196 (47%)               | 4,772 (48%)    |          |
| cT                   |                           |                           |                           |                           |                |          |
| 1–3                  | 659 (28%)                 | 959 (37%)                 | 824 (33%)                 | 824 (34%)                 | 3,264 (33%)    | <0.001   |
| 4                    | 564 (24%)                 | 757 (29%)                 | 735 (29%)                 | 795 (32%)                 | 2,849 (29%)    |          |
| X                    | 619 (27%)                 | 485 (19%)                 | 500 (20%)                 | 611 (24%)                 | 2,218 (22%)    |          |
| Missing              | 486 (21%)                 | 290 (15%)                 | 472 (19%)                 | 288 (11%)                 | 1,637 (16%)    |          |
| cN                   |                           |                           |                           |                           |                |          |
| 0                    | 450 (19%)                 | 531 (20%)                 | 413 (16%)                 | 401 (16%)                 | 1,795 (18%)    | <0.001   |
| 1–2                  | 1,099 (47%)               | 1,434 (55%)               | 1,521 (60%)               | 1,362 (54%)               | 5,414 (54%)    |          |
| X                    | 660 (28%)                 | 543 (21%)                 | 540 (21%)                 | 702 (28%)                 | 2,447 (25%)    |          |
| Missing              | 119 (5%)                  | 83 (3%)                   | 57 (2%)                   | 53 (2%)                   | 312 (3%)       |          |
| Metastatic location  |                           |                           |                           |                           |                |          |
| Liver                | 744 (32%)                 | 913 (35%)                 | 908 (36%)                 | 805 (32%)                 | 3,370 (34%)    | <0.001   |
| Lung                 | 205 (9%)                  | 176 (7%)                  | 170 (7%)                  | 128 (5%)                  | 679 (7%)       |          |
| Peritoneum           | 62 (3%)                   | 70 (3%)                   | 85 (3%)                   | 111 (4%)                  | 328 (3%)       |          |
| Other                | 73 (3%)                   | 93 (4%)                   | 83 (3%)                   | 105 (4%)                  | 354 (4%)       |          |
| Multiple locations   | 1,244 (53%)               | 1,339 (52%)               | 1,285 (51%)               | 1,369 (54%)               | 5,237 (53%)    |          |

*From chi square test

Abbreviations: cN (clinical nodal status), CT (clinical tumour status)

Includes 9,968 patients diagnosed with metastatic colorectal cancer in 2009–2016 in Sweden

Supplementary Table 3). There was a significant interaction between hospital volume and hospital level (p from Wald test < 0.001). This can be interpreted as the effect of hospital volume was significantly different for university hospitals compared with non-university hospitals.
was altered by including the potential mediator MDT, the results were mainly unchanged and university hospital remained significantly associated with the chance of metastatic surgery (OR (95% CI): 1.90 (1.64–2.19)), whereas hospital volume remained not associated (Supplementary Table 3). This can be interpreted as the full effect of hospital level is not mediated by MDT, but is a direct effect or mediated by some other factor. Furthermore, MDT was an independent predictor of receiving metastatic surgery in the model (OR (95% CI): 1.54 (1.30–1.82)). The effect of being discussed at an MDT was different between different hospital levels (p from Wald test < 0.001), but MDT was positively associated with receiving metastatic surgery at both university hospitals and non-university hospitals.

Overall survival
The 1- and 5-year OS were slightly higher for patients managed at hospitals in the 4th volume quartile (56% and 14%, respectively) compared with patients managed at hospitals in the 1st volume quartile (50% and 11%, respectively, Fig. 4A-C). High hospital volume was associated with a decreased all-cause mortality rate (4th versus 1st quartile of hospital volume, adjusted HR (95% CI): 0.92 (0.86–0.98), Table 3 and Supplementary Table 3). When further adjustment was made for hospital level the association between hospital volume and all-cause mortality disappeared (Table 3 and Supplementary Table 3).

Similarly, there were differences in OS based on whether the managing hospital was a university hospital or not. The 1- and 5-year OS were 59% vs 50% and 15% vs 12% for patients managed at a university hospital vs. a non-university hospital, respectively (HR (95% CI): 0.83 (0.78–0.88), results from second model including both exposures). Neither the effect of hospital volume nor hospital level violated the proportionality assumption (p-value from test of proportional hazards > 0.05).

In the sensitivity analysis with delayed entry, the 1- and 5-year OS estimates were 97% and 48%, respectively for patients who received metastatic surgery, and 84% and 12% respectively in the group who did not receive metastatic surgery (Fig. 4C).
Figs 4  A-C Cumulative overall survival (OS) estimates by A) hospital volume, B) hospital level, C) metastatic surgery. Kaplan–Meier estimates of OS including 9,968 patients followed from diagnosis of synchronous metastatic colorectal cancer. The stratification by metastatic surgery was based on a delayed entry of patients in model to 270 days post diagnosis (n = 5,993). P-value from log-rank test was $p = 0.001$ for A and $p < 0.001$ for B and C.
Discussion

In this large nationwide cohort study of almost 10,000 patients with synchronous mCRC 17% of patients had surgery of their primary tumour and metastases. The most important hospital characteristic was management at a university hospital, where patients experienced almost a doubled chance of receiving metastatic surgery and an improved survival. The proportion receiving metastatic surgery varied significantly by healthcare region.

To the best of our knowledge, annual hospital volume of incident patients with mCRC has not previously been associated with whether or not metastatic surgery is performed. However, patients managed at a university hospital are known to experience an increased likelihood of having surgery for colorectal liver metastases [7, 8, 13] and peritoneal metastases [14]. Our results clearly show that the associations between hospital volume and metastatic surgery were essentially explained by hospital level. In Sweden, centralisation of most metastatic surgery analysed in this study has been to the university hospitals. In Sweden, specific cancer centres/hospitals are not present, whereas this is rather common in other healthcare systems, where they may or may not be connected to university hospitals. Further, metastatic surgeries may also be performed in non-university hospitals. Therefore, the results regarding hospital level probably holds the highest external validity in comparison with countries with a similar healthcare organisation, chiefly the other Nordic countries but also countries like the Netherlands. Hospital factors e.g. presence of a liver-centre on site [6, 30], or a HIPEC-centre on site [14], hospitals volume of liver resections [9], and management at a high-volume hospital of treatments for mCRC [16], have been associated with an increased likelihood of receiving metastatic site surgery. The finding that patients diagnosed at university hospitals in Sweden are more likely to receive metastatic surgery may be mediated through many factors which are more common in the university hospitals. Even if our finding of hospital level associated with metastatic surgery may not be immediately be applicable to other healthcare environments, we can more confidently conclude that hospital volume on its own is not associated with more metastatic surgery and better outcomes for mCRC patients.

The reasons why patients at non-university hospitals are less likely to be referred to metastatic surgery are probably multifactorial, reflecting differences in opinion or even knowledge of the treating physicians or the patient’s wish not to be referred. One explanation for the association seen is that interventional clinical studies are common at university hospitals, a factor that has been associated with improved outcome for CRC patients [31]. Patients with mCRC have improved OS when treated at academic programmes/hospitals compared with community programmes/district hospitals [7, 8, 14, 32]. Another important factor is whether the metastases are deemed resectable, upfront or after conversion therapy, or not. Although all regions in Sweden but a few have a dedicated clinical oncology department with specialists in both radiation and medical oncology, the university hospitals have larger oncology departments with more specialists also devoted to research. It is possible that patients at university hospitals receive more active conversion chemotherapy resulting in greater possibilities for, e.g., liver surgery. Previous studies have also illustrated that the assessment of resectability of liver metastasis is highly dependent on the observer, where referring physicians can incorrectly consider some metastatic lesions as non-resectable [33] and liver specialists are more prone to deem metastases as resectable [11,
During the 8-year period analysed here, a marked change in what is considered resectable has occurred. This is probably a major reason for the increasing temporal trend in the proportion of metastatic surgery noted, but could also be responsible for the differences seen between healthcare regions. Such differences tend to decrease with time, although we could not detect any indications of this. Individual surgeons, medical oncologists and interventional radiologists at the different university hospitals may also have affected the results. In summary, the importance of avoiding individual decisions and instead making use of MDT conferences, with specialised oncologists and subspecialised surgeons present, must be stressed.

MDTs are positively associated with the chance of metastatic surgery for mCRC [18, 19]. To explore whether presence of MDTs could be a mediator for the observed differences in metastatic surgery by hospital level we performed an explanatory model adjusting for MDT. Our results showed that the elevated chance of metastatic surgery for patients managed at a university hospital remained, indicating that there are other reasons for the dissimilarities observed. It is possible that the quality of the discussion varied at different hospital levels, but this information is not possible to obtain. An alternative to local or even regional MDTs could be centralised repeated assessments of resectability, which increased the proportion eligible for surgery of CRC liver metastases in a Finnish material [34]. Future studies are needed to evaluate if this is sufficient to diminish the demonstrated variability in provided care by hospital level, or between different healthcare regions, for mCRC patients.

**Strengths and weaknesses**

The greatest strength of this national cohort study is its size and quality of data. Thanks to the recently established CRCBaSe, we had access to high-quality data from SCRCR and several other large national registries. The previous Swedish study on selection for metastatic surgery for mCRC in 2007–2011 was based on the SCRCR and restricted to data on liver resections only [8]. In the current study we could complement the SCRCR data with data from several other registers e.g., treatment codes from the patient register and therefore also study metastatic surgery of the lungs and peritoneum. Another strength is that we could ascertain that the patients did not have a previous CRC diagnosis as far back as 1958. Moreover, by linkage to national patient registries, we could limit our study population to patients with at least one defined metastatic location, yielding high internal validity. In terms of external validity, the nationwide approach increased generalisability. We believe the results to be applicable to countries with similar hospital systems, especially those that have a similar centralisation of metastatic surgeries to university hospitals. In countries with specialised cancer centres outside of the university organisation, the conclusions regarding the benefits of being managed at university hospitals may be more difficult to transfer.

The primary weakness of this study is its confinement to available register data. Detailed data on factors related to treatment strategy, such as tumour burden at each metastatic location, would have been preferable. Unfortunately, this has not been registered and is almost impossible to obtain retrospectively given the size of the study. Patients at high volume hospitals were younger and could therefore be deemed more suitable for the most demanding surgeries for mCRC. In addition, patients at high volume hospitals had higher socioeconomic status, a factor that has been associated with the chance of receiving surgery for mCRC [6, 7, 9, 10, 17, 30, 35]. However, the association between university hospital and the chance to receive metastatic surgery withheld after adjustment for mentioned confounders such as age, tumour characteristics, civil status, income and level of education. We acknowledge that there is a possibility of residual confounding. There may also be a risk of referral bias, if so, it is probably more pronounced in metropolitan areas where several hospitals are present within a short geographical distance. In the majority of cases, we believe the patients were referred to the closest hospital irrespective of tumour burden and comorbidities.

**Conclusions**

This study indicates that patients with mCRC receive different surgical care in Sweden. Initial management at a university hospital was associated with a greater chance of receiving metastatic surgery for patients with synchronous mCRC, and improved survival. High hospital volume in itself was not associated with a greater chance to receive metastatic surgery nor a greater survival probability. These results show that additional efforts should be imposed in order to provide more equal care for mCRC patients.

**Abbreviations**

CCI: Charlson comorbidity index; CI: Confidence interval; cN: Clinical nodal; CRC: Colorectal cancer; CRCBaSe: Colorectal cancer database; cT: Clinical tumour; HIPEC: Hyperthermic intraperitoneal chemotherapy; HR: Hazard ratio; mCRC: Metastatic colorectal cancer; MDT: Multidisciplinary team; OR: Odds ratio; OS: Overall survival; SCR: Swedish cancer register; SCRCR: Swedish colorectal cancer register.
**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1186/s12885-022-10005-8.

**Additional file 1: Supplementary table 1.** Treatment codes as described by the National Board of Health and Welfare in Sweden used to identify potentially curative treatments of metastases and primary tumour, translated from Swedish.*Defined as non-surgical treatment. Supplementary table 2.** Additional treatment-associated patient characteristics for 9,961 patients with synchronous metastatic colorectal cancer (mCRC) according to the managing hospital's annual incident number of patients with mCRC divided into 4 quartiles. *Data unspecifed on whether the neoadjuvant treatment was given for the primary tumour or the metastases or both. ** A total of 46 patients who underwent surgery for liver metastases (of which one had combination metastatic surgery) received solely non-surgical locally ablative therapy (percutaneous destruction or embolisation treatment) and four patients with lung metastases had solely non-surgical treatment (stereotactic body radiation, SBRRT). The solely non-surgical locally ablative treatments constituted 3% (n=50/1797) of the patients treated with metastatic surgery. Abbreviations: ASA (American Society of Anesthesiologists), pT (pathological tumour status). **Supplementary table 3.** Summary table of results. All multivariable models were adjusted for sex (male, female), age (18-64, 65-79, ≥80 years), Charlson comorbidity index (0,1,2), Primary tumour status (right colon, transverse colon, left colon, unknown unknown location in colon, rectum), cT-stage (1-3, 4, X), cN-stage (0, 1-2, X), metastases (single, multiple), civil status (married, not married), education (primary, secondary, higher education), income (4 annual quartiles), diagnosis year (continuous 2009-2016). Missing data handled using missing indicator method. Multivariable logistic regression models included 9,966 patients. Multivariable cox regression models included 9,968 patients. Abbreviations: CI: confidence interval, HR: hazard ratio, MDT: multidisciplinary team conference, OR: odds ratio, QT: quartile.

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**Authors’ contributions**

M.L. wrote the original draft and did the formal analysis. M.L., C.W. and C.N. were responsible for methodology and C.W. for data curation. A.M and C.N. provided the resources and funding acquisition. C.N. supervised the project. All authors participated in the conceptualisation and all authors reviewed the manuscript. All authors read and approved the final manuscript.

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**Availability of data and materials**

The study is based on depersonalized data which cannot be shared by the authors due to the instructions of the ethical permission. For further questions regarding the data availability we recommend contacting the board of CRCBaSe (email: ccrbase@mmk.ki.se).

**Declarations**

**Ethics approval and consent to participate**

The study was approved by the Regional Board of the Ethical Committee in Stockholm, Sweden (DNR: 2014/71–31, 2018/328–32, 2021–00342) and the study was conducted in accordance with the Declaration of Helsinki and the ethical standards described in the ethical permit. The requirement for informed consent was waived by the same ethics committee as mentioned above.

**Consent for publication**

Not applicable. No individual person's data in the manuscript.

**Competing interests**

Dr. Weibull is part of a research collaboration between Karolinska Institutet and Janssen Pharmaceutica NV for which Karolinska Institutet has received grant support. This was unrelated to this study. The other authors declare no competing interests.

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