Peripartum Blood Transfusions are Associated with Increased Risk of Cancer: A National Retrospective Cohort Study

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Background: The effect of blood transfusions on the risk of developing primary cancer remains unclear, especially when administered in the peripartum period.

Materials and Methods: We conducted a retrospective cohort study of 270,529 pregnant women who delivered between January 1, 2007 and December 31, 2009, with data obtained from three national databases in South Korea. From this cohort, we identified 4569 patients who received peripartum blood transfusions. We calculated hazard ratios (HRs) for new diagnoses of cancer and adjusted them for relevant clinical factors using a Cox proportional hazards model.

Results: During follow-up, patients who received peripartum transfusions had an increased risk of developing cancer, with an adjusted HR of 1.16 (95% confidence interval [CI], 1.01–1.34). In a subgroup analysis, this risk was significant only among patients who received 3 or more units of blood, with an adjusted HR of 1.40 (95% CI, 1.10–1.79). Increased risk after transfusions were seen with brain, lung, ovarian, and gallbladder cancers. The difference in cancer risk between the transfusion and no-transfusion groups remained significant during both the first (1.29% vs 1.07%, p < 0.01) and second year (0.74% vs 0.56%, p < 0.01) after delivery.

Conclusion: Receipt of 3 or more blood transfusions in the peripartum period was associated with a significantly increased risk of developing cancer. Prospective studies should be pursued to further understand the link between blood transfusions and long-term oncologic risks.

Keywords: blood transfusion, pregnancy, cancer, immunomodulation

Introduction
Blood transfusions are an increasingly common indicator of peripartum morbidity.¹,² Multiple concerns have been raised regarding the potential long-term risks of blood transfusions, including the possibility of transmitting infectious agents.³,⁴ Transfusions may also modify the immune system via a phenomenon termed transfusion-related immunomodulation (TRIM).⁵,⁶ According to the TRIM hypothesis, the influx of new antigens and immunologically active mediators during transfusions can contribute to immune tolerance and suppression.⁷ These post-transfusion changes include decreased cytotoxic T cell and natural killer cell activity and increased secretion of anti-inflammatory cytokines.⁸,⁹ The importance of preserving a robust immunological response to tumors has been highlighted by recent advances in immune checkpoint inhibitors in cancer treatment.¹¹ Transfusions may also promote cancer
development through the transmission of oncogenic viruses, including hepatitis B and C viruses, human immunodeficiency virus, and Epstein-Barr virus; however, this risk has markedly diminished in the modern era with screening efforts.12,13

Despite these concerns, multiple clinical studies over the past 30 years have yielded contradictory results. Several analyses have demonstrated an association between perioperative blood transfusions and cancer recurrence,10 especially in colorectal cancer,14 although no causal relationship has been definitively established.14,15 There is greater uncertainty in regards to the risk of developing primary malignancies. Most studies have been performed in older patients, who are more likely to have occult cancers that can predispose them to requiring transfusions,16,17 or did not control for risk factors such as alcohol and tobacco use.18

Pregnancy uniquely represents a situation during which transfusions are commonly indicated in a generally healthy cohort of patients. However, limited data exist on the effect of peripartum transfusions on subsequent cancer risk. We thus tested the hypothesis that blood transfusions in peripartum women would lead to an increased incidence of primary cancers.

Materials and Methods

Data Sources

This study was conducted by merging the databases of the Korea National Health Insurance (KNHI) claims, National Health Screening Examination (NHSE), and National Health Screening Program for Infants and Children (NHSP-IC).

In Korea, 97% of the population is enrolled in the KNHI program, and the KNHI claims database contains all claims information for these individuals. Therefore, nearly all information on patients’ diseases and their treatments can be obtained from this centralized database, with the exception of procedures that are not covered by insurance, such as cosmetic surgery. The KNHI database has been extensively validated as a comprehensive population-level dataset, including for cancer-related information.19,20

Using the KNHI claims database, we identified all pregnant women who delivered between January 1, 2007 and December 31, 2009, and obtained information on blood transfusions in the peripartum period. Subsequent development of cancer was tracked until December 31, 2015.

Enrollees in the KNHI system are invited to undergo a standardized NHSE. The NHSE involves a free, voluntary biennial cardiovascular and age-appropriate cancer screening and is offered to any patient who is employed or above the age of 40.20 The pre-pregnancy characteristics of our study population were evaluated via review of the NHSE data. The KNHI system also provides an NHSP-IC for all neonates that includes a health interview with the parents. Information on gestational age at delivery and birth weight was obtained using the NHSP-IC health interview.

This study was approved by the Institutional Review Boards of the Korea University Medical Center.

Study Population

A flowchart of patient enrollment is shown in Figure 1. Based on the KNHI claims dataset, we identified 666,554 women who had a first delivery between January 1, 2007 and December 31, 2009. To facilitate the evaluation of the pre-pregnancy characteristics, women were included in the analysis only if they underwent an NHSE within one year prior to their pregnancy. During our study period, 45.6% of pregnant women participated in the NHSE. In the general population, 66% of eligible patients participated in the NHSE in 2009.21 Among this group, women whose offspring did not participate in the NHSP-IC (10.3% of NHSE participants and 5.6% of full cohort) were excluded from the analysis. Finally, 35,003 women with a prior history of cancer were also excluded.

Outcomes

Using the KNHI claims dataset, a new diagnosis of cancer was identified by principal or secondary diagnoses based on International Classification of Diseases-10th Revision (ICD-10) codes. Women were classified as having cancer if they were newly diagnosed with cancer (ICD-10 code Cxx.x) between delivery and December 31, 2015. The timing of the initial diagnosis was confirmed by the lack of a medical claim for cancer as a primary or secondary diagnosis before pregnancy.

Pregnancy factors were identified using the KNHI claims dataset, and post-partum hemorrhage (PPH) and preeclampsia were identified according to ICD-10 codes. Based on the KNHI claims dataset, presence of multiple pregnancy, the delivery mode, the use of blood transfusions, and the number of transfusion units administered were confirmed. The peripartum period was defined as time encompassed by the hospitalization during which delivery occurred. All categories of blood product transfusion were included, such as packed red blood cells, pooled platelets, and fresh frozen plasma. Using data on gestational age at delivery from the NHSP-IC, preterm birth was also identified. Preterm birth was defined as a gestational age of <37 weeks.
Pre-pregnancy factors were evaluated using the NHSE data. Smoking status was identified using health questionnaires. The health examination included the calculation of body mass index (BMI, in kg/m²). Obesity was defined as BMI ≥ 25 kg/m² according to the cutoffs established by the Korean Society for the Study of Obesity. Hypertension (HTN) was defined as blood pressure ≥ 140/90 mmHg or the current use of antihypertensive medication. Blood samples for laboratory values were obtained after a fast of at least 8 hours. Diabetes mellitus was defined as fasting glucose ≥ 126 mg/dL or the current use of antidiabetic medication. Abnormal liver function tests were defined as aspartate aminotransferase (AST) ≥ 31 or alanine aminotransferase (ALT) ≥ 31 mg/dL. High total cholesterol level was defined as levels ≥ 200 mg/dL.

Statistical Analysis
Continuous and categorical variables are expressed as mean ± standard deviation (SD) and percentages, respectively. Clinical and biochemical characteristics were compared among groups using the Student’s t-test for continuous variables and the χ² test for categorical variables. The cumulative incidence of cancer was estimated using the Kaplan–Meier method and compared using the Log rank test. Cox proportional hazards models were used to estimate the adjusted hazard ratios (HRs) and 95% confidence intervals (CIs) for the development of cancer. Clinical variables were pre-defined and chosen to reflect both basic health indicators and cancer risk factors. Participants were censored if they developed cancer before or on December 31, 2015 in those without cancer. All tests were two sided, and p-values < 0.05 were considered statistically significant. Statistical analyses were performed using SAS for Windows, version 9.4 (SAS Inc., Cary, NC, USA).

Results
Our study cohort included 270,529 women, with 4569 (1.7%) of them receiving transfusions. Patients who required transfusions were more likely to be older or to have hypertension, a history of diabetes mellitus, or abnormal liver function tests at baseline. They were also more likely to undergo caesarean section, experience postpartum hemorrhage, develop pre-eclampsia, or have a preterm birth or multiple pregnancy (Table 1).

Women in this cohort were followed for an average of 7.14 ± 1.59 years to assess for the development of a cancer diagnosis, with the transfusion group followed for 6.98 ± 1.75 years and the no-transfusion group followed for 7.14 ± 1.58 years. Receipt of perioperative transfusions was associated with a greater likelihood of developing cancer during the study period (HR 1.25, 95% CI 1.08–1.44). This association remained significant even after adjusting for multiple clinical variables (HR 1.16, 95% CI 1.01–1.34). These variables included age, co-morbidities (diabetes, hypertension, obesity, abnormal...
liver function tests, or hypercholesterolemia), smoking status, and obstetric factors (history of multiple pregnancies, caesarean section, preterm birth, or pre-eclampsia).

The relation between transfusions and cancer occurrence was strongly influenced by the number of units of blood product that the patients received (Table 2). There was not a significantly increased risk of developing cancer among patients who received only 1 unit (HR 1.23, 95% CI 0.91–1.65) or 2 units (HR 1.11, 95% CI 0.89–1.38) of blood. However, patients who received 3 or more units of blood had a significantly higher occurrence of cancer (HR 1.40, 95% CI 1.10–1.79).

Further analysis showed that transfusions did not increase the risk for all cancer types (Table 3). Significantly increased risk of occurrence was specifically seen with brain (HR 4.90, 95% CI 1.93–12.43), lung (HR 4.57, 95% CI 1.76–11.86), ovarian (HR 2.79, 95% CI 1.41–5.54), and gallbladder (HR 2.30, 95% CI 1.01–5.26) cancers.

The increased risk of cancer after transfusions was persistent over two years. During the first year after delivery, 1.29% of patients who received transfusions were diagnosed with cancer, as opposed to 1.07% of patients who did not receive transfusions (p < 0.01). This difference remained significant during the second year, with an incidence of cancer of 0.74% in the transfusion group and 0.56% in the no-transfusion group (p < 0.01) (Figure 2).

### Discussion

Our data showed that 1.7% of women required blood transfusions in the peripartum period, which is similar to rates observed in other modern cohorts. Those patients who received transfusions had a greater incidence of primary cancers in the follow-up period. This effect was only significant among patients who were transfused 3 or more units of blood products.

Prior studies have demonstrated an increased incidence of cancer among patients who received transfusions, but it has been unclear if the transfusions themselves drove this effect. In a large Scandinavian cohort study, the increased rate of new cancers among transfusion recipients was most pronounced in the first 6 months after the transfusion. The authors thus inferred a reverse causation effect in which patients with subclinical cancer were more likely to require transfusions. Similarly, a case-control study of US patients utilizing Medicare claims data showed that transfusions were only associated with elevated risk of cancer within the first 12 months after transfusion.

By utilizing a dataset limited to peripartum transfusions, we potentially reduced the risk of a reverse causation effect, as the prevalence of predisposing co-

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### Table 1 Clinical Characteristics of Participants Stratified by Transfusion Status

<table>
<thead>
<tr>
<th></th>
<th>No Transfusion (n = 265,960)</th>
<th>Transfusion (n = 4569)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>29.15 ± 3.04</td>
<td>30.01 ± 3.54</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>20.59 ± 4.21</td>
<td>20.58 ± 2.62</td>
<td>0.89</td>
</tr>
<tr>
<td>Obesity (%)</td>
<td>5.83 ± (15,493)</td>
<td>6.26 ± (286)</td>
<td>0.21</td>
</tr>
<tr>
<td>Systolic BP (mmHg)</td>
<td>111.0 ± 11.11</td>
<td>111.4 ± 11.35</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Diastolic BP (mmHg)</td>
<td>69.99 ± 8.33</td>
<td>70.45 ± 8.61</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>HTN (%)</td>
<td>4.14 ± (11,021)</td>
<td>5.41 ± (247)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Fasting glucose (mg/dL)</td>
<td>86.15 ± 13.58</td>
<td>86.62 ± 20.26</td>
<td>0.12</td>
</tr>
<tr>
<td>DM (%)</td>
<td>3.21 ± (8,544)</td>
<td>4.73 ± (216)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>TC (mg/dL)</td>
<td>173.5 ± 31.83</td>
<td>173.3 ± 30.87</td>
<td>0.58</td>
</tr>
<tr>
<td>High TC (%)</td>
<td>17.03 ± (45,293)</td>
<td>17.33 ± (792)</td>
<td>0.59</td>
</tr>
<tr>
<td>AST (mg/dL)</td>
<td>19.47 ± 10.78</td>
<td>19.79 ± 10.21</td>
<td>0.03</td>
</tr>
<tr>
<td>ALT (mg/dL)</td>
<td>15.21 ± 16.67</td>
<td>15.63 ± 14.99</td>
<td>0.06</td>
</tr>
<tr>
<td>Abnormal LFT (%)</td>
<td>5.76 ± (15,316)</td>
<td>6.94 ± (317)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Smoking (%)</td>
<td>3.55 ± (9444)</td>
<td>4.03 ± (184)</td>
<td>0.01</td>
</tr>
<tr>
<td>Multiple pregnancy (%)</td>
<td>1.60 ± (4246)</td>
<td>9.81 ± (448)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Cesarean section (%)</td>
<td>32.87 ± (74,417)</td>
<td>52.40 ± (2394)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Preterm birth (%)</td>
<td>2.96 ± (7862)</td>
<td>12.41 ± (567)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Preeclampsia (%)</td>
<td>2.19 ± (5821)</td>
<td>9.46 ± (432)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Postpartum hemorrhage (%)</td>
<td>5.51 ± (14,655)</td>
<td>41.48 ± (1895)</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

**Notes:** Values are expressed as mean (SD) or %). Obesity was defined by BMI ≥25 kg/m². Abnormal liver function test (LFT) was defined as AST ≥31 mg/dL or ALT ≥31 mg/dL.

**Abbreviations:** BMI, body mass index; BP, blood pressure; HTN, hypertension; DM, diabetes mellitus; TC, total cholesterol; AST, aspartate aminotransferase; ALT, alanine aminotransferase; LFT, liver function tests.

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### Table 2 Risk of the Development of Cancer According to Transfusion Status

<table>
<thead>
<tr>
<th></th>
<th>Unadjusted HR (95% CI)</th>
<th>Adjusted HR* (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any transfusion</td>
<td>1.25 (1.08, 1.44)</td>
<td>1.16 (1.01, 1.34)</td>
</tr>
<tr>
<td>1 unit (N = 1,072)</td>
<td>1.23 (0.91, 1.65)</td>
<td>1.13 (0.84, 1.52)</td>
</tr>
<tr>
<td>2 units (N = 2196)</td>
<td>1.11 (0.89, 1.38)</td>
<td>1.04 (0.84, 1.30)</td>
</tr>
<tr>
<td>≥3 units (N = 1,301)</td>
<td>1.51 (1.18, 1.93)</td>
<td>1.40 (1.10, 1.79)</td>
</tr>
</tbody>
</table>

**Note:** Adjusted for age, DM, HTN, obesity, abnormal LFT, high total cholesterol, smoking status, multiple pregnancy, caesarean section, preterm birth, and preeclampsia.
morbidities and undiagnosed cancers was presumably decreased in this cohort of younger women. We also limited our study population to first deliveries to further limit potential confounders related to prior pregnancies and transfusions. We could find only one comparable study, which evaluated cancer risk after blood transfusion during obstetric delivery in 621 Swedish women.\textsuperscript{24} No association between transfusions and cancer incidence was noted during the 32-year follow-up period,\textsuperscript{24} though this cohort may have been too small to detect any existent effect. It is still possible that residual confounding remained a factor in our study, with subclinical or precancerous lesions leading to increased need for transfusions. However, the increased risk of cancer after blood transfusions remained persistent past the first year, arguing against the possibility that undiagnosed, subclinical cancers were responsible for our result.

A putative mechanism for our observed effect is not clear from our data. Transmission of unscreened carcinogenic agents has been proposed as a potential etiologic factor.\textsuperscript{17,25} Donor seeding of cancer has also been explored as a causative mechanism. This route has been documented most notably in

<table>
<thead>
<tr>
<th></th>
<th>No Transfusion</th>
<th>Transfusion</th>
<th>Adjusted HR\textsuperscript{*} (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liver</td>
<td>0.33 (882)</td>
<td>0.24 (11)</td>
<td>0.71 (0.39, 1.28)</td>
</tr>
<tr>
<td>Breast</td>
<td>0.33 (873)</td>
<td>0.22 (10)</td>
<td>0.57 (0.30, 0.96)</td>
</tr>
<tr>
<td>Thyroid</td>
<td>1.12 (2974)</td>
<td>1.44 (66)</td>
<td>1.22 (0.95, 1.56)</td>
</tr>
<tr>
<td>Gallbladder</td>
<td>0.05 (143)</td>
<td>0.13 (6)</td>
<td>2.30 (1.01, 5.26)</td>
</tr>
<tr>
<td>Brain</td>
<td>0.02 (62)</td>
<td>0.11 (5)</td>
<td>4.90 (1.93, 12.43)</td>
</tr>
<tr>
<td>Uterus</td>
<td>0.10 (271)</td>
<td>0.18 (8)</td>
<td>1.55 (0.76, 3.16)</td>
</tr>
<tr>
<td>Colon</td>
<td>0.80 (2133)</td>
<td>0.88 (40)</td>
<td>1.06 (0.77, 1.45)</td>
</tr>
<tr>
<td>Lung</td>
<td>0.02 (52)</td>
<td>0.11 (5)</td>
<td>4.57 (1.76, 11.86)</td>
</tr>
<tr>
<td>Stomach</td>
<td>0.10 (272)</td>
<td>0.15 (7)</td>
<td>1.45 (0.68, 3.10)</td>
</tr>
<tr>
<td>Ovary</td>
<td>0.06 (171)</td>
<td>0.20 (9)</td>
<td>2.79 (1.41, 5.54)</td>
</tr>
<tr>
<td>Others</td>
<td>0.25 (669)</td>
<td>0.35 (16)</td>
<td>1.22 (0.74, 2.01)</td>
</tr>
</tbody>
</table>

Note: \textsuperscript{*}Adjusted for age, DM, HTN, obesity, abnormal LFT, high total cholesterol, smoking status, multiple pregnancy, cesarean section, preterm birth, and preeclampsia.

Figure 2: Cumulative incidence of cancer diagnoses. The incidence of cumulative cancer incidence is illustrated for transfusion recipients and non-recipients over a 9-year span. The greater incidence in cancer diagnoses among transfusion recipients remained persistent during this time, suggesting that this difference was not due to occult malignancies that predisposed those patients to requiring transfusions.
the context of organ transplants, including a case report of two kidney transplant recipients who developed melanoma after receiving organs from a deceased donor who had been successfully treated for melanoma 16 years prior.26 However, a large retrospective cohort study did not demonstrate an elevated cancer risk after receiving transfusions from donors with subclinical cancer.27 Meanwhile, multiple immunomodulatory effects have been described after blood transfusions.28 Preclinical studies have suggested that these effects are mediated by allogeneic leukocytes, leukocyte-derived cytokines, and/or the presence of soluble HLA antigens.5,29 These findings demonstrate that transfusions can perturb the immune surveillance and editing process, which is essential to inhibiting carcinogenesis.30

Only certain cancers – namely, brain, lung, ovarian, and gallbladder cancers – were more common among patients with history of blood transfusions. The difference in lung cancer is notable particularly given the extremely low rates of smoking in this cohort and further supports the importance of immunologic factors in lung cancer.31 Alterations in immune function have not been widely discussed as contributing to ovarian carcinogenesis,32 which is typically protected against by pregnancy,33 but may be worth exploring in light of our results. Previous studies have observed an increased risk for liver cancer after blood transfusions.34,35 including one study that noted a persistently elevated risk after 5 years.17 Liver cancers were not more common among transfusion recipients in our cohort, which may in part be due to our ability to control for other relevant risk factors. We also note that the above cancers typically grow aggressively. Given our mean follow-up time of about 7 years, it is possible that malignancies known to develop over a longer time course, such as colon cancer, would require longer follow-up before exhibiting any impact from transfusions.

The strengths of our study include the use of a large population-based cohort from a national registry, which allowed for a complete and fully representative dataset. This database also allowed for long-term follow-up of patients and adjustment for many relevant confounding factors. The focus on post-partum patients, who generally represent a young and healthy population, may have further reduced the potential for confounders, such as the presence of occult malignancy. In addition, information on the number of transfusions was unique to our study compared to many similar studies. In studies that did incorporate this variable, several demonstrated a dose-response relationship between number of transfusions and degree of immunomodulation, possibly because a stronger antigen exposure is better able to trigger immunologic changes.36 Due to the recency of these data, our findings also reflect modern advances in blood product screening methods and transfusion strategies.

Notable limitations include the retrospective nature of our study and lack of detailed data regarding the transfusions. For example, characteristics of donors were unavailable, specifically in regards to presence of undiagnosed cancer or viral infections at the time of blood donation. Factors that may influence the immunomodulatory effect of transfusions, including storage time37 and use of leukocyte-reduced blood,38 were not obtained. We note that South Korea does not mandate leukocyte depletion,39 and in fact, one study found that only 10.3% of blood products in South Korea were leukocyte reduced in 2009.40 The recency of our data had the disadvantage of limiting the possible follow-up period to 8 years, and any longer-term effects of transfusions were unable to be evaluated. We also note that less than half of women who delivered were ultimately included in the analysis, raising the possibility that this is not a representative subset. The vast majority of patients were excluded due to lack of sufficient NHSE data; it is unclear if this would correlate at all with cancer risk factors.

Finally, our results may be influenced by physiologic factors unique to pregnant women. Pregnancy is known to cause complex immunologic changes that may alter the maternal response to blood transfusions.41,42 A recent study, for example, showed that pregnant women may be more likely to have adverse transfusion reactions than non-pregnant women,43 possibly due to higher rates of HLA alloimmunization.44 It is thus unclear whether our findings relate to pregnancy-specific factors, or if they are generalizable to other patients.

In conclusion, we found that transfusion of 3 or more units of blood in the peripartum period was associated with an increased risk of developing cancer in a comprehensive national cohort. The potential mechanisms for such an effect have not been fully elucidated, though evidence of an immunomodulatory effect has been previously reported. Our results suggest that restrictive strategies for blood transfusions in the obstetric setting should be considered to reduce long-term oncologic risks.

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


