Original Investigation

Familial Risk and Heritability of Cancer Among Twins in Nordic Countries

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IMPOR TANCE Estimates of familial cancer risk from population-based studies are essential components of cancer risk prediction.

OBJECTIVE To estimate familial risk and heritability of cancer types in a large twin cohort.

DESIGN, SETTING, AND PARTICIPANTS Prospective study of 80,309 monozygotic and 123,382 same-sex dizygotic twin individuals (N = 203,691) within the population-based registers of Denmark, Finland, Norway, and Sweden. Twins were followed up a median of 32 years between 1943 and 2010. There were 50,990 individuals who died of any cause, and 3,804 who emigrated and were lost to follow-up.

EXPOSURES Shared environmental and heritable risk factors among pairs of twins.

MAIN OUTCOMES AND MEASURES The main outcome was incident cancer. Time-to-event analyses were used to estimate familial risk (risk of cancer in an individual given a twin's development of cancer) and heritability (proportion of variance in cancer risk due to interindividual genetic differences) with follow-up via cancer registries. Statistical models adjusted for age and follow-up time, and accounted for censoring and competing risk of death.

RESULTS A total of 27,156 incident cancers were diagnosed in 23,980 individuals, translating to a cumulative incidence of 32%. Cancer was diagnosed in both twins among 1,383 monozygotic (2,766 individuals) and 1,933 dizygotic (2,866 individuals) pairs. Of these, 38% of monozygotic and 26% of dizygotic pairs were diagnosed with the same cancer type. There was an excess cancer risk in twins whose co-twin was diagnosed with cancer, with estimated cumulative risks that were an absolute 5% (95% CI, 4%-6%) higher in dizygotic (37%; 95% CI, 36%-38%) and an absolute 14% (95% CI, 12%-16%) higher in monozygotic twins (46%; 95% CI, 44%-48%) whose twin also developed cancer compared with the cumulative risk in the overall cohort (32%). For most cancer types, there were significant familial risks and the cumulative risks were higher in monozygotic than dizygotic twins. Heritability of cancer overall was 33% (95% CI, 30%-37%). Significant heritability was observed for the cancer types of skin melanoma (58%; 95% CI, 43%-73%), prostate (57%; 95% CI, 51%-63%), nonmelanoma skin (43%; 95% CI, 26%-59%), ovary (39%; 95% CI, 23%-55%), kidney (38%; 95% CI, 21%-55%), breast (31%; 95% CI, 11%-51%), and corpus uteri (27%; 95% CI, 11%-43%).

CONCLUSIONS AND RELEVANCE In this long-term follow-up study among Nordic twins, there was significant excess familial risk for cancer overall and for specific types of cancer, including prostate, melanoma, breast, ovary, and uterus. This information about hereditary risks of cancers may be helpful in patient education and cancer risk counseling.

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The global burden of cancer is considerable, with an estimated 12 million new cases and 8 million cancer deaths each year. In 2015 in the United States, 1.7 million individuals will be diagnosed with cancer and 590,000 will die of cancer, accounting for 1 in 4 deaths. In the Nordic countries, cancer is the leading cause of mortality, accounting for 30% of all deaths. Refinement of primary and secondary prevention strategies (ie, factors that would have the greatest influence on reducing cancer incidence and mortality) requires a detailed understanding of the contribution of genetic and environmental factors to disease pathogenesis.

Family-based studies have been helpful in describing familial aggregation of cancer. Many inherited risk loci have been identified by genome-wide association studies; however, these known loci explain only a small proportion of the variability in cancer incidence. Large twin studies of cancer can provide further insight into the relative contribution of inherited factors and characterize familial cancer risk by leveraging the genetic relatedness of monozygotic and dizygotic pairs of twins.

A study in 2000 found significant estimates of heritability (ie, the proportion of variability in disease risk in a population due to genetic factors) of 42% for prostate cancer, 35% for colorectal cancer, and 27% for breast cancer among twins from Sweden, Denmark, and Finland. The confidence intervals around these heritability estimates were wide, and the estimates were not interpretable for other common cancers.

To address these limitations, we undertook an analysis within the Nordic Twin Study of Cancer (NorTwinCan), including twins from nationwide registers in Denmark, Finland, Norway, and Sweden followed up for an average of 32 years for cancer incidence and mortality. Statistical methods that take into account the potential statistical bias due to censoring and competing risk of death were used to estimate heritability and familial risk for specific types of cancer.

Methods

Study Population

The NorTwinCan study is an international, multidisciplinary collaboration of researchers working to investigate the genetic and environmental underpinnings of cancer. The cohort includes 357,377 individual twins from the population-based twin registers of Denmark, Finland, Norway, and Sweden and is composed of both monozygotic and same-sex dizygotic and opposite-sex dizygotic twins. In each country, the twin registries were assembled through the nationwide identification of twins across several birth cohorts.

Twins were identified through a range of methods, including review of national birth registries, church parish records, and civil registration systems. For example, the twin registry in Denmark was assembled for 4 birth cohorts during the study period of 1870 through 2004. The first cohort from 1870 through 1930 was assembled retrospectively in the 1950s through review of birth register records in each of the local parishes for which births were recorded.

Subsequent birth cohorts in Denmark were identified through review of the civil registration system and national birth registry records. Nationwide coverage of twin registries ranges between 30% for the oldest birth cohort (from 1870-1930) and 70% for the birth cohort from 1931 through 1952. Coverage was approximately 100% for later cohorts. The lower coverage in the earlier cohorts is due in large part to the fact that pairs of twins had to survive to the age of 6 years to be included in the registry.

Twins zygoity for same-sex pairs of twins is determined primarily by validated questionnaire methods that show a high degree of accuracy (>95% agreement with genetic markers; questionnaire given either to the twin member or a relative if twin deceased). We excluded data from 5376 twins with missing or inconsistent zygoity data. The ethical committees at each of the twin registries’ host institutions approved this project.

Cohort Follow-up

Residents in Nordic countries each have a unique national registration number that allows for linkage of data from the Nordic twin register to national cancer registers for each country, mortality registers, and registers of the total population to glean outcome and vital status information (date of death or emigration) for each individual. For cancer diagnoses, we obtained data by linkage to the national cancer register in each country.

Physicians and pathologists are mandated by law to report every newly diagnosed malignant tumor in each of the 4 countries. In addition, the nationwide death registries send information to the cancer registry for individuals when the death certificate mentions cancer. Case reporting to each registry is close to 100% complete. The quality of the data are assessed through careful review, and all reports of cancer diagnosis are verified at national registries.

Cancer register data include diagnoses of cancer type classified according to the International Statistical Classification of Diseases and Related Health Problems, Tenth Revision (ICD-10). For this study, we grouped the ICD-10 codes to categories defined in comparable ways across the cancer registries using the NORDCAN (a system designed for the standardization of cancer codes across the Nordic registries) classification of cancers.

Cohort follow-up is initiated at the start of cancer registration in each country or at a later time for birth cohorts born after the start. For example, the cancer register in Denmark was initiated in January 1943. For twins born in the cohorts to 1930, follow-up began in 1943. For Danish twins born after 1943, follow-up began in 1968 at the time when the national registration number system was started. We have follow-up through the end of 2008 in Norway, 2009 in Denmark and Sweden, and 2010 in Finland. Details of the country-specific dates of start and end of follow-up are summarized in Table 1 and additional details are provided in Hjelmborg et al.
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Statistical Methods

After combining cancers of the head and neck (ICD-10 codes OC 00-14), 36 cancer types remained. We present results for familial risk for 23 cancer types with more than 1 concordant monogygotic and dizygotic pair, and heritability estimates for malignancies with at least 5 concordant pairs of twins. In a twin study, familial or concordance risk is defined as the risk of a specific cancer type in a twin, given that the co-twin was diagnosed with the same cancer. Comparing the conditional risk with the cumulative incidence in the population provides an estimate of the excess familial risk of a cancer.

In particular, dizygotic twins are as genetically alike as full siblings and thus familial risk among dizygotic twins can be generalized to siblings. Heritability is defined as the proportion of variance in cancer risk on the liability scale due to interindividual genetic differences in the population. For both familial risk and heritability, estimates are a function of follow-up time and the age of the cohort.

Individuals were followed up prospectively through the registries until cancer diagnosis, death or emigration during follow-up, or the end of the study. We defined the dates of entry and follow-up separately for each cohort depending on ascertainment procedures and data availability in each country (Table 1).

In statistical models, we accounted for left truncation from differing start dates of follow-up and right censoring for those censored at the end of follow-up, those censored when lost to follow-up due to emigration, or at competing risk of death. Cumulative risk of cancer was calculated using the nonparametric Aalen-Johansen estimator.13 We modeled potential competing deaths as described in Scheike et al,8 which accounted for competing causes in both the twin and co-twin.

We used quantitative genetic models to estimate the relative contribution of genetic and environmental factors with the variation in cancer risk. This approach assumes that there is a normally distributed liability to develop a genetically complex disease such as cancer. The probability that an individual will express the disease is modeled as a function of the latent unmeasured liability, and disease occurs only when an individual surpasses the threshold. This approach analyzes the disease covariance within monozygotic and dizygotic pairs and decomposes the variance into a sum of components: additive genetic effects (A), common environmental effects shared among twins (C), and individually unique environmental effects (E).

Within-pair covariance is expressed as $\kappa \times \text{var}(A) + \text{var}(C)$, where $\kappa = 1$ for monogygotic pairs because they share 100% of their genomes and $\kappa = \frac{1}{2}$ for dizygotic pairs because they share on average half of their segregating genes. To test whether there is evidence of a genetic component for each of the cancer types, we compared the tetrachoric correlation for monozygotic and dizygotic twins in a model in which the marginal estimates were the same.

The biometric modeling approach is comparable with that of Lichtenstein et al,7 except that we adjusted for censoring by weighting individuals by the inverse probability of being censored at the time of follow-up using the same weights within pairs of twins.9 The probabilities of being censored were estimated using the Kaplan-Meier method stratified by zygosity and country.

We similarly used the inverse probability of weighting to estimate the median difference in age at diagnosis for the pairs of twins concordant for cancer. All analyses were performed using the R mets package version 1.1.1 (R Foundation for Statistical Computing). Two-sided P values were used with an alpha level of less than .05.

Results

This analysis comprised 203 691 individual twins in the cohort, 80 309 monozygotic and 123 382 same-sex dizygotic twins, of whom 104 251 were women (Table 1). During an average of 32 years of follow-up, we identified 27 156 incident cancers among 23 980 individuals. In addition, 50 990 individu-
Cancer was diagnosed in both twins among 1383 monozygotic (2766 individuals) and 1933 dizygotic (2866 individuals) pairs. Of these, 38% of monzygotic pairs (n = 522) and 26% of dizygotic pairs (n = 496) were diagnosed with the same type of cancer.

The estimated cumulative incidence of cancer in the overall cohort and the familial risk among monzygotic and dizygotic twins appear in the Figure. The estimated cumulative incidence of cancer, accounting for competing causes of death, was 8% by the age of 65 years, 25% by the age of 80 years, and 32% by the age of 100 years using twins as individuals in a standard cohort analysis. These risks are similar to the nationwide rates in the Nordic populations, showing representativeness of the twins. The lifetime familial risk of any cancer for those whose co-twins were also diagnosed with cancer was 37% (95% CI, 36%-38%) among dizygotic pairs and 46% (95% CI, 44%-48%) among monzygotic pairs by the age of 100 years.

There was an excess cancer risk in twins whose co-twin was diagnosed with cancer, with estimated cumulative risks that were an absolute 5% (95% CI, 4%-6%) higher in dizygotic (37%; 95% CI, 36%-38%) and an absolute 14% (95% CI, 12%-16%) higher in monzygotic twins (46%; 95% CI, 44%-48%) whose twin also developed cancer compared with the cumulative risk in the overall cohort (32%).

The types of cancers with the highest estimated cumulative incidence in the cohort were prostate (10.5%), breast (9.4%), lung (3.2%), nonmelanoma skin (1.9%), and colon (2.9%) (Table 2). There were elevated familial risks among monzygotic and dizygotic pairs of twins for most cancer types as indicated by the familial risk estimates compared with the cumulative incidence. These risks were substantially higher for monzygotic than dizygotic pairs for cancers of the prostate and breast.

Some of the strongest familial associations were observed for somewhat less common cancers. For testicular cancer, for which the cumulative risk in the cohort was 0.5%, the risk was substantially higher when his co-twin was also diagnosed with testicular cancer; for dizygotic twins, the familial risk estimate was 6% (95% CI, 2%-17%) and for monzygotic twins it was 14% (95% CI, 6%-30%) if his co-twin had previously been diagnosed with testicular cancer.

Familial cancer risk was also high for melanoma of the skin, with familial risks of 6% (95% CI, 3%-13%) for dizygotic twins and almost 20% (95% CI, 12%-31%) for monzygotic twins compared with a cumulative risk of 1.2% for the overall cohort. Familial risk of nonmelanoma skin cancer was also evident, although less than for melanoma. Familial risk of ovarian cancer for women whose twins also had ovarian cancer was similarly greater in monzygotic (9%; 95% CI, 4%-18%) than dizygotic (3%; 95% CI, 1%-7%) twins.

Results from the quantitative genetic modeling used to decompose the familial associations into the genetic and shared environmental components that affect twins the same way appear in Table 3 and in the eTable in the Supplement. The heritability for cancer overall was 33% (95% CI, 30%-37%), with no evidence of a shared environmental component.

A high estimate of heritability of 57% (95% CI, 51%-63%) was found for prostate cancer. For breast cancer, 31% of variability may be associated with genetic factors and 16% with shared environmental factors. The strong familial effect noted for testicular cancer among monzygotic and dizygotic twins was associated with both significant genetic (37%) and shared environmental (24%) factors. Moreover, we found significant heritability estimates for cancer of the kidney (38%), skin melanoma (58%), and skin nonmelanoma (43%).
Lung cancer had one of the highest estimates for shared environmental factors (24%). Heritability estimates for gastrointestinal cancers of the colon (15%; 95% CI, 0%-45%), rectum (14%; 95% CI, 0%-50%), and stomach (22%; 0%-55%) were smaller relative to other malignancies.

Data on the median difference in age at diagnosis among concordant pairs of twins for selected cancer types appear in Table 4. For all cancer cases combined, the median age difference was slightly shorter in monozygotic pairs (8.0 years) than in dizygotic pairs (9.3 years). With the
exception of prostate cancer, none of these differences in median age differed between monozygotic and dizygotic pairs for 11 cancer types.

Discussion

This prospective Nordic twin cohort study provides familial risk estimates for cancer overall, for 23 types of cancer, and for relatively rare cancer types. Overall, there was a significantly excess familial risk of developing any cancer, 37% in dizygotic pairs and 46% in monozygotic pairs compared with 32% in the whole twin cohort. The data provide strong evidence of an excess familial risk for 20 of the 23 cancer types, as shown by the comparison of familial risks for those cancers with the cumulative risk in the twin cohort overall.

Testicular cancer and nonmelanoma and melanoma skin cancers showed substantial excess familial risks, particularly among monozygotic pairs. Although the excess familial risk for breast, prostate, and other cancer types were more modest, the absolute differences in risk were considerable.

These estimates of familial risk are in line with other family- and twin-based cohort studies. Precise estimates of familial cancer risk from population-based studies are essential components of accurate cancer risk prediction, and could be used in clinical practice to guide genetic counseling. Dizygotic pairs of twins are as genetically similar as siblings.

As such, familial risk estimates among dizygotic pairs are relevant for siblings who are born at separate times. Risk estimates among monozygotic pairs (who share nearly 100% of their inherited genomes) can be used to derive an upper bound on the ability of genetic studies to discriminate individuals who will experience different disease outcomes.

Shared environmental factors can include parental factors, such as socioeconomic status, lifestyle, and occupation and experiences and exposures shared by siblings during childhood and adolescence, and screening patterns in adult life. For many cancer types, we did not observe evidence of shared environmental associations, even though our models specifically allowed for estimation of these environmental factors that members of a family share.

Lung cancer had one of the highest shared environmental components, likely due to shared smoking habits of pairs of twins. Testicular and breast cancer also had significant estimates of shared environment, which may reflect in part the hypothesized in utero origins of these cancer types. It is possible that the shared environment components may be

Table 3. Estimates of Heritability and Shared Environment for Specific Types of Cancer in the NorTwinCan Cohort

<table>
<thead>
<tr>
<th>Familial Risk, % (95% CI)</th>
<th>Heritability</th>
<th>Shared Environment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall cancer</td>
<td>33 (30-37)</td>
<td>0</td>
</tr>
<tr>
<td>Head and neck</td>
<td>9 (0-60)</td>
<td>26 (0-65)</td>
</tr>
<tr>
<td>Stomach</td>
<td>22 (0-55)</td>
<td>6 (0-31)</td>
</tr>
<tr>
<td>Colon</td>
<td>15 (0-45)</td>
<td>16 (0-38)</td>
</tr>
<tr>
<td>Rectum and anus</td>
<td>14 (0-50)</td>
<td>10 (0-38)</td>
</tr>
<tr>
<td>Lung</td>
<td>18 (0-42)</td>
<td>24 (7-40)</td>
</tr>
<tr>
<td>Skin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Melanoma</td>
<td>58 (43-73)</td>
<td>0</td>
</tr>
<tr>
<td>Nonmelanoma</td>
<td>43 (26-59)</td>
<td>0</td>
</tr>
<tr>
<td>Breast</td>
<td>31 (11-51)</td>
<td>16 (0-31)</td>
</tr>
<tr>
<td>Corpus uteri</td>
<td>27 (11-43)</td>
<td>0</td>
</tr>
<tr>
<td>Ovary</td>
<td>39 (23-55)</td>
<td>0</td>
</tr>
<tr>
<td>Prostate</td>
<td>57 (51-63)</td>
<td>0</td>
</tr>
<tr>
<td>Testis</td>
<td>37 (0-93)</td>
<td>24 (0-70)</td>
</tr>
<tr>
<td>Kidney</td>
<td>38 (21-55)</td>
<td>0</td>
</tr>
<tr>
<td>Bladder, other urinary organs</td>
<td>30 (0-67)</td>
<td>0</td>
</tr>
<tr>
<td>Leukemia, other</td>
<td>57 (0-100)</td>
<td>0</td>
</tr>
</tbody>
</table>

Abbreviation: NorTwinCan, Nordic Twin Study of Cancer.

a Not calculated for cancer types with less than 5 concordant pairs.

Table 4. Difference in Age at Diagnosis Among Monozygotic and Dizygotic Pairs of Twins for Specific Types of Cancer

<table>
<thead>
<tr>
<th>No. Concordant Pairsa</th>
<th>Age Difference at Diagnosis, Median (95% CI), y</th>
<th>P Valueb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall cancer</td>
<td>1383</td>
<td>1933</td>
</tr>
<tr>
<td>Head and neck</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>Stomach</td>
<td>14</td>
<td>15</td>
</tr>
<tr>
<td>Colon</td>
<td>30</td>
<td>31</td>
</tr>
<tr>
<td>Rectum, anus</td>
<td>14</td>
<td>13</td>
</tr>
<tr>
<td>Lung</td>
<td>50</td>
<td>74</td>
</tr>
<tr>
<td>Skin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Melanoma</td>
<td>11</td>
<td>6</td>
</tr>
<tr>
<td>Nonmelanoma</td>
<td>16</td>
<td>10</td>
</tr>
<tr>
<td>Breast</td>
<td>124</td>
<td>141</td>
</tr>
<tr>
<td>Corpus uteri</td>
<td>9</td>
<td>6</td>
</tr>
<tr>
<td>Prostate</td>
<td>197</td>
<td>148</td>
</tr>
<tr>
<td>Bladder</td>
<td>18</td>
<td>13</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>Monzygotic</th>
<th>Dizygotic</th>
<th>Monzygotic</th>
<th>Dizygotic</th>
</tr>
</thead>
<tbody>
<tr>
<td>8.0 (0.4-22.8)</td>
<td>9.3 (0.3-33.9)</td>
<td>8.3 (3.3-12.3)</td>
<td>10.1 (3.5-17.8)</td>
</tr>
<tr>
<td>10.3 (5.2-15.4)</td>
<td>14.2 (10.0-18.4)</td>
<td>8.3 (5.8-11.1)</td>
<td>6.1 (3.8-11.7)</td>
</tr>
<tr>
<td>10.3 (6.0-15.3)</td>
<td>5.3 (2.7-11.6)</td>
<td>7.8 (6.1-9.5)</td>
<td>7.7 (5.4-9.4)</td>
</tr>
<tr>
<td>8.9 (7.5-21.8)</td>
<td>15.8 (3.4-23.6)</td>
<td>6.1 (3.5-11.9)</td>
<td>7.4 (2.4-7.7)</td>
</tr>
<tr>
<td>9.3 (6.9-12.0)</td>
<td>10.5 (9.1-12.2)</td>
<td>12.4 (6.7-17.2)</td>
<td>9.3 (3.6-10.5)</td>
</tr>
<tr>
<td>3.7 (4.4-6.1)</td>
<td>6.1 (4.6-7.9)</td>
<td>3.7 (4.4-6.1)</td>
<td>6.1 (4.6-7.9)</td>
</tr>
<tr>
<td>7.1 (4.1-9.9)</td>
<td>14.3 (7.2-20.5)</td>
<td>7.1 (4.1-9.9)</td>
<td>14.3 (7.2-20.5)</td>
</tr>
</tbody>
</table>

a The numbers refer to the number of twin pairs not the individual twins.
b For difference in median ages between twin types (monozygotic vs dizygotic).
somewhat higher in twins than the estimates expected for siblings because twins may share a more similar child-raising experience.

Most pairs of twins were discordant for a specific cancer. Indeed, among the pairs of twins in which both members developed cancer, more than two-thirds were diagnosed with a different malignancy. This finding of familial aggregation across different cancer types is in line with data from an Icelandic family-based study that showed excess familial aggregation among 17 different cancer types.26

There are novel insights from genetic epidemiology studies suggesting pleiotropy of genetic variants across multiple cancer types.27,28 A more detailed investigation of shared familial risk across cancer types may provide key insights into the underlying cancer susceptibility.

Twin studies can provide context for genome-wide association studies, many of which have identified multiple risk loci for cancer incidence.29-32 Estimates of heritability in twin studies, as well as those derived from genome-wide association studies, allow for the calculation of the extent to which cancer variability is explained by established genetic risk loci. For prostate cancer, the 100 risk loci identified to date explain approximately one-third of the genetic contribution.33,34 For breast cancer, the estimated proportion of heritability explained by the known genetic risk loci may approach up to 30%.35

Few genome-wide association studies of more rare cancers have been undertaken, but our data suggest that such studies may be essential in elucidating the etiology of certain cancer types, such as melanoma, ovarian, kidney, and testicular. A recent analysis of genome-wide association studies for 13 cancer types suggests that the identified genetic variants for most cancer types do not explain the majority of heritability.36 Moreover, the heritability estimate of 33% for developing any cancer type suggests there are shared genomic regions associated with multiple cancer types. The results from 2 family-based studies are also in line with this observation.37,38 A systematic genome-wide association study of individuals with any cancer compared with controls free of any cancer may be a powerful and feasible approach to identify novel loci given that multiple cancer type-specific data sets already exist.

The concept of heritability has its limitations39 and is often misinterpreted as an estimate of population-attributable risk. Heritability can be thought of as the proportion of the variation in cancer risk in a population that can be accounted for by interindividual genetic differences.

A high estimate of heritability for cancer does not translate to a low population-attributable risk associated with lifestyle and environmental factors, nor does a high heritability exclude the possibility of an effective preventive action. The observed genetic estimates based on family relationships for specific cancer types are due to both cancerspecific genetics and genetic contributions to cancer risk factors, such as obesity and smoking, which have a genetic component.

The interval between diagnosis of cancer among concordant pairs of twins was fairly long. The median time between cancer diagnoses in concordant pairs of twins ranged from 4 to 15 years. The difference in pairs of twins with any cancer was similar to the specific numbers for cancer type.

These data suggest that there may be unique environmental factors that influence the timing of disease development or diagnosis. Moreover, cancer likely involves stochastic processes inherent in the carcinogenic process that may ultimately influence the timing of cancer initiation and diagnosis.

Our study has some strengths and limitations to consider in interpreting the study results. To our knowledge, this is the largest familial study of cancer to date and includes more than 3 decades of follow-up. In prior studies, the number of concordant pairs with cancer was small, leading to imprecise estimates of familial risk and heritability for common cancer types and uninterpretable estimates of heritability for less common cancer types. The large number of cancer cases and long follow-up in the cohort allowed us to provide more precise estimates of familial risk and heritability for several malignancies.

Notwithstanding the large size of the cohort and long-term follow-up, we were unable to provide estimates of familial risk or heritability for some of the more rare cancer types, including many of the hematological malignancies. The linkage with national population-based registers and the high quality of cancer case registration allowed for complete follow-up of the study population.

The study is based on twins from the Nordic countries, primarily a white population. For the majority of cancer types, the cumulative incidence of cancer in the twin cohort was similar to that of the entire country, suggesting these data can be generalized to the Nordic countries. However, it is unclear the extent to which these data can be generalized to multiethnic populations.

Given the older age at which many cancer types occur, long-term follow-up provides greater clarity on estimates of familial risk and heritability by allowing the cohort to attain sufficient age at which most cancer types occur. On a related note, some individuals at risk of developing cancer may also be at higher risk of dying from another chronic condition, thus influencing the estimates.

Our statistical approach addressed this challenge and accounted for differential follow-up time, censoring, and competing causes of death. The twin modeling estimates of heritability assume that there are similar shared environments between monozygotic and dizygotic pairs of twins, which cannot be formally tested in this setting.

**Conclusions**

In this long-term follow-up study among Nordic twins, there was significant excess familial risk for cancer overall and for specific types of cancer, including prostate, melanoma, breast, ovary, and uterus. This information about hereditary risks of cancers may be helpful in patient education and cancer risk counseling.
ARTICLE INFORMATION

Correction: This article was corrected online January 5, 2016, to fix incorrect author affiliations for 2 authors.

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Author Contributions: Drs Mucci and Hjelmborg had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. Drs Mucci and Hjelmborg contributed equally to this article and share first authorship. Drs Adami and Kaprio contributed equally to this article and share last authorship.

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REFERENCES


