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New use of prescription drugs prior to a cancer diagnosis

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ABSTRACT

Purpose Cancers often have considerable induction periods. This confers a risk of reverse causation bias in studies of cancer risk associated with drug use, as early symptoms of a yet undiagnosed cancer might lead to drug treatment in the period leading up to the diagnosis. This bias can be alleviated by disregarding exposure for some time before the cancer diagnosis (lag time). We aimed at assessing the duration of lag time needed to avoid reverse causation bias.

Methods We identified all Danish patients with incident cancer between 2000 and 2012 (n = 353,087). Incident use of prescription drugs was assessed prior to their cancer diagnosis as well as among population controls (n = 1,402,400). Analyses were conducted for all cancers and for breast, lung, colon and prostate cancer individually. Further, analyses were performed for a composite measure of all incident drug use as well as for nine pre-specified individual drug classes, representing drug treatment likely to be prescribed for symptoms of the given cancers.

Results The incidence rate for new drug treatment among cancer cases was stable around 130 per 1000 persons per month until 6 months prior to cancer diagnosis where it increased gradually and peaked at 434 in the month immediately preceding the cancer diagnosis. Considerable variation was observed among cancers, for example, breast cancer showed almost no such effect. The pre-selected drug classes showed a stronger increase prior to cancer diagnoses than drugs overall.

Conclusions Incident use of drugs increases in the months prior to a cancer diagnosis. To avoid reverse causation, 6 months' lag time would be sufficient for most drug-cancer associations. © 2016 The Authors. Pharmacoepidemiology and Drug Safety published by John Wiley & Sons Ltd.

INTRODUCTION

Many cancers have considerable induction periods, and patients will therefore often have a lengthy pre-diagnostic period.1 A yet undiagnosed cancer might cause symptoms that are misinterpreted as benign illness. For example, gastrointestinal cancers might give rise to abdominal pain, which is treated with, for example, proton pump inhibitors or laxatives, before any attempts of diagnostic work-up are made. Further, the more frequent health care contact for the individual patient leading up to the diagnosis might in itself lead to the initiation of drug treatment, for example, the clinical work-up might reveal a previously undiagnosed diabetic condition or hypertension, which then leads to initiation of new drug therapy. Both of these factors might be observed as an increase in new drug treatments given to patients before their cancer diagnosis.

Such a drug use pattern would have implications for pharmacoepidemiological studies of cancer risk associated with use of drugs. In these studies, onset of new drug treatment prior to diagnosis raises the possibility for reverse causation bias.2 For example, if we were to study the association between proton pump inhibitors and gastric cancer, we might find an association solely attributable to the fact that early cancer symptoms (before the diagnosis is made) are misinterpreted as acid related disorders and treated with proton-pump inhibitors. The conventional approach to this reverse causation problem is to disregard a certain period (lag time) before the cancer diagnosis when accounting for drug exposure.3,4 This solution is applicable both in cohort and case-control designs. To our knowledge, however, there has been no systematic appraisal of this phenomenon.

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In this study, we aimed to inform the choice of lag-time period in studies on drug–cancer associations to avoid reverse causation bias. We did so using the Danish nationwide health registries on cancer\(^5\) and prescription drugs\(^6\) to investigate the onset of new drug treatments in the period leading up to a cancer diagnosis.

**METHODS**

Sampling all Danish patients with incident cancer between 2000 and 2012, we assessed drug use in a 24-month window prior to the date of cancer diagnosis. Drug use patterns among cancer cases were compared with the drug use among population controls, matched to cases by sex, age and calendar time.

**Study population**

Using the Danish Cancer Registry\(^5\), we identified all incident cancer cases in Denmark between 2000 and 2012. For each cancer case, we identified four population controls, matched on sex, age and calendar time (date of the case diagnosis). For both cancer cases and the population controls, we excluded children, individuals with a prior cancer diagnosis (except non-melanoma skin cancer) and individuals that had migrated in or out of Denmark within the last 10 years.

**Drug use**

Incident drug use was defined as an individual’s first-ever filling of a prescription drug according to the Danish National Prescription Registry\(^6\), defining drugs at the level of the single substance. As the registry holds data from 1995, this ensured a minimum of 5 years prescription data when identifying new users.

For cancer cases and population controls, we assessed incident use of prescription drugs in monthly intervals relative to the date of the cancer diagnosis (or day of sampling for controls), going back 24 months. In each month, we estimated the incidence rate (IR) of drug use by—within that month—counting the number of new drug treatments and dividing by follow-up among cancer cases (or population controls), thus calculating the IR with a unit of new treatments per person-month.

**Analysis**

In an overall analysis, the IR for any new drug treatment was calculated in the 2 years preceding any cancer diagnosis and compared with that of population controls. Similar analyses were carried out specifically for the four most common cancers: lung, breast, prostate and colon cancer. Lastly, we performed the analysis on these four types of cancer for six pre-specified drug classes, selected as drugs that could likely be prescribed for symptoms for a given cancer: drugs against overactive bladder and drugs against prostatic hyperplasia (possibly associated with a later diagnosis of prostate cancer), inhaled beta-agonists and cough suppressants (lung cancer), and drugs against constipation or diarrhoea and proton pump inhibitors (colon cancer). No relevant drug could be identified for early symptoms of breast cancer. Furthermore, we included use of opioids, which is possibly related to all cancer diagnoses, as well as oral antidiabetics and statins, as markers for drugs that might be initiated following health care contacts. For a full list of definitions for these drugs, see Appendix A. In these analyses, we only considered individuals at risk for incident drug use, that is we disregarded those with ever-use of the drugs in question prior to the time window of assessment.

**RESULTS**

We identified 353,087 eligible cancer cases that were matched to 1,402,400 population controls. The four most common types of cancer were breast \((n = 51,774)\), colon \((n = 29,505)\), lung \((n = 45,509)\) and prostate cancer \((n = 41,115)\).

In all analyses, the pattern of incident drug use was very similar between cancer cases and populations controls in the time period of 24 to 12 months prior to cancer diagnosis. Thus, only results from the 12-month window prior to diagnosis are presented throughout.

The overall IR for new drug treatment among all cancer cases was stable around 130 per 1000 persons per month until 6 months prior to cancer diagnosis where it increased and peaked at 434 new drug treatments per 1000 persons per month in the month immediately preceding the date of cancer diagnosis. This is illustrated in Figure 1.

Figure 1. Number of new drug treatments per 1000 persons per month in a 12-month window prior to cancer diagnosis among cancer cases and compared with the pattern among age- and sex-matched population controls.

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Figure 2. Number of new drug treatments per 1000 persons per month in a 12-month window prior to cancer diagnosis among breast, colon, prostate and lung cancer cases, compared with the pattern among age- and sex-matched population controls.

Figure 3. Number of new drug treatments per 1000 persons per month in a 12-month window prior to cancer diagnosis among cancer cases for pre-selected drug classes. Note the logarithmic Y-axis.
preceding the cancer diagnosis (Figure 1). This pattern varied considerably between the four most common cancers (Figure 2): Almost no increase was observed for breast cancer cases, while prostate, colon and lung cancer peaked at 397, 428 and 741 new drug treatments per 1000 persons per month in the month prior to cancer diagnosis, respectively. No noticeable changes were observed among the population controls.

The pre-selected site-specific drugs all showed a noticeable increase prior to cancer diagnosis (Figure 3). New use of the three non-specific drugs included in the analysis all showed an increase prior to cancer diagnosis, especially opioid analgesics, which increased prior to all cancers except breast cancer, most pronounced for lung cancer, increasing 15-fold and peaking at 70 new drug treatments per 1000 per month (Figure 3).

In a post hoc analysis, we explored the drug classes contributing to the increased overall IR observed among cases. For the 6-month time window prior to the cancer diagnosis, we calculated cumulative incidence proportions (risks) for incident use of individual drug classes (fourth ATC-level, e.g. A02BC proton pump inhibitors), while restricting to those at risk, that is, never users of these drugs by the start of the time window. We reported the 20 drug classes with the largest absolute risk difference, comparing cancer cases to population controls. The results for all cancers (Table 1), showed that the increase in drug use was driven by proton pump inhibitors, analgesics and antibiotics. Similar analyses for the four most common cancers (eTable1a-d), showed that the increased drug use was driven by therapy either specific to the individual cancer, such as laxatives prior to a colon cancer diagnosis, or related to cancer diagnoses in general (analgesics and antibiotics).

### DISCUSSION

Our study demonstrated a very clear increase in new drug use in the months preceding a cancer diagnosis. We have demonstrated both a specific and an unspecified component, the first apparently related to specific symptoms of the cancers, the second probably to unrelated conditions revealed during frequent physician contacts before the cancer diagnosis. The pattern differs between cancers. For most drug-cancer associations, six months’ lag time appears to be sufficient to avoid substantial reverse causation.

Our study has several strengths. The Danish cancer registry has virtually complete and valid registration of all cancers in a well-defined, stable population, and the prescription registry has offered complete coverage of the Danish population since 1995. Among the weaknesses is that we cannot account for the handling of the earliest cancer symptoms in primary care. However, we find it unlikely that GPs to any great extent would treat for example colon cancer-related abdominal pain with proton pump inhibitors if they suspected the pain to be cancer-related. Diagnostic process delay is unlikely to play a major role; according to Danish law, there is an expedited diagnostic work-up if a cancer suspicion is voiced.

In general, there are two reasons to apply lag time in studies of drug–cancer associations; the problem of reverse causation as demonstrated in this paper and the fact that exposure immediately before the cancer diagnosis usually cannot be considered as contributing to the development of the cancer. By including such etiologically irrelevant exposure, true associations would be attenuated. At least for colon and prostate cancer, there is good evidence of a long latency from cancer development to a clinically overt cancer.

### Table 1

<table>
<thead>
<tr>
<th>ATC</th>
<th>Drug class</th>
<th>Cases (%)</th>
<th>Controls (%)</th>
<th>Absolute risk difference (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A02BC</td>
<td>Proton pump inhibitors</td>
<td>6.8</td>
<td>1.6</td>
<td>5.2</td>
</tr>
<tr>
<td>J01CA</td>
<td>Penicillins with extended spectrum</td>
<td>6.9</td>
<td>2.1</td>
<td>4.8</td>
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<tr>
<td>N02AX</td>
<td>Other opioids</td>
<td>6.0</td>
<td>1.5</td>
<td>4.4</td>
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<tr>
<td>J01CE</td>
<td>Beta-lactamase sensitive penicillins</td>
<td>7.6</td>
<td>4.0</td>
<td>3.6</td>
</tr>
<tr>
<td>J01FA</td>
<td>Macrolides</td>
<td>4.5</td>
<td>1.7</td>
<td>2.8</td>
</tr>
<tr>
<td>N02AA</td>
<td>Natural opioid alkaloids</td>
<td>3.5</td>
<td>0.9</td>
<td>2.6</td>
</tr>
<tr>
<td>M01AE</td>
<td>Propionic acid derivatives</td>
<td>5.1</td>
<td>2.5</td>
<td>2.6</td>
</tr>
<tr>
<td>N02BE</td>
<td>Anilides</td>
<td>4.0</td>
<td>1.7</td>
<td>2.3</td>
</tr>
<tr>
<td>A03FA</td>
<td>Propulsives</td>
<td>2.6</td>
<td>0.4</td>
<td>2.1</td>
</tr>
<tr>
<td>N05CF</td>
<td>Benzodiazepine related drugs</td>
<td>3.0</td>
<td>0.8</td>
<td>2.1</td>
</tr>
<tr>
<td>M01AB</td>
<td>Acetic acid derivatives and related substances</td>
<td>3.5</td>
<td>1.5</td>
<td>2.0</td>
</tr>
<tr>
<td>J01MA</td>
<td>Fluoroquinolones</td>
<td>2.4</td>
<td>0.7</td>
<td>1.7</td>
</tr>
<tr>
<td>J01EB</td>
<td>Short-acting sulphonamides</td>
<td>2.6</td>
<td>0.9</td>
<td>1.7</td>
</tr>
<tr>
<td>N05BA</td>
<td>Benzodiazepine derivatives</td>
<td>2.3</td>
<td>0.6</td>
<td>1.7</td>
</tr>
<tr>
<td>H02AB</td>
<td>Glucocorticoids</td>
<td>2.6</td>
<td>1.0</td>
<td>1.6</td>
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<tr>
<td>C03CA</td>
<td>Sulfonamides, plain</td>
<td>2.3</td>
<td>0.9</td>
<td>1.4</td>
</tr>
<tr>
<td>A12BA</td>
<td>Potassium</td>
<td>2.1</td>
<td>0.8</td>
<td>1.3</td>
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<td>A06AD</td>
<td>Osmotically acting laxatives</td>
<td>1.8</td>
<td>0.5</td>
<td>1.2</td>
</tr>
<tr>
<td>G04CA</td>
<td>Alpha-adrenoceptor antagonists</td>
<td>1.6</td>
<td>0.3</td>
<td>1.2</td>
</tr>
<tr>
<td>R05DA</td>
<td>Opium alkaloids and derivatives</td>
<td>2.0</td>
<td>0.8</td>
<td>1.2</td>
</tr>
</tbody>
</table>

Notes:
ATC = Anatomical-Therapeutical-Chemical

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late critical steps in carcinogenesis. Empirically, this may appear as a relatively short interval between exposure and clinically overt cancer. If we routinely apply lag time in all analyses, we would overlook such associations. Late-stage cancer promotion has been demonstrated for some immunomodulating drugs particularly in patients with organ transplant, where exposure as short as six months may trigger a clinical cancer diagnosis. Other good examples are, however, quite rare, especially associated with such shortness of exposure. Researchers are encouraged to explore the exposure pattern leading up to a cancer diagnosis in a given study, to inform the choice of lag-time.

We conclude that lag time should be considered in studies of drug–cancer associations and that 6 months is usually sufficient to avoid reverse causation bias. However, sound clinical reasoning should prevail concerning the manifestations of the specific cancers.

CONFLICTS OF INTEREST

The authors declare no conflict of interest.

KEY POINTS

- Cancers that are not yet diagnosed may cause symptoms that are confused with benign diseases and treated as such.
- This may result in reverse causation bias, whereby drug treatment that precedes the cancer diagnosis may be suspected of causing the cancer, while in reality it is used to treat early cancer symptoms. The bias can be alleviated by disregarding exposure for some time before the cancer diagnosis (lag time).
- Reverse causation has two components; a specific one that is related to the cancer’s early symptoms and an unspecific one that is related to frequent physician contact before the cancer diagnosis.
- Our study demonstrated both components, the specific being the dominant.
- Our data suggests that for most cancers, six months of lag time is sufficient to avoid reverse causation bias.

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REFERENCES


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