Hiding in plain view: the potential for commonly used drugs to reduce breast cancer mortality

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Hiding in plain view: the potential for commonly used drugs to reduce breast cancer mortality

Michelle D Holmes*1,2 and Wendy Y Chen1,3

Abstract
Many medications have been developed for one purpose but then are found to have other clinical activities. There is tremendous interest in whether non-cancer medications may potentially have effects on breast cancer survival. In this review article, we have presented and evaluated the evidence for several commonly used over-the-counter and prescription medications – including aspirin (and other non-steroidal anti-inflammatory drugs), beta-blockers, angiotensin-converting enzyme inhibitors, statins, digoxin, and metformin – that have been evaluated among breast cancer survivors in prospective studies. Substantial scientific evidence supports the hypothesis that some of these common and relatively safe drugs may reduce breast cancer mortality among those with the disease by an amount that rivals the mortality reduction gained by currently used therapies. In particular, the evidence is strongest for aspirin (approximately 50% reduction), statins (approximately 25% reduction), and metformin (approximately 50% reduction). As these drugs are generic and inexpensive, there is little incentive for the pharmaceutical industry to fund the randomized trials that would show their effectiveness definitively. We advocate that confirmation of these findings in randomized trials be considered a high research priority, as the potential impact on human lives saved could be immense.

Introduction
Many medications have been developed for one purpose but then are found to have other clinical activities. For example, minoxidil was originally developed as an anti-hypertensive but then was found to cause excessive hair growth. Because of the multiple potential pathways that can be involved with cancer growth and metastases, tremendous interest remains in whether currently used non-cancer medications may potentially have anti-cancer effects. In this review article, we will present and evaluate the evidence for several commonly used over-the-counter and prescription medications that have been evaluated among breast cancer survivors in prospective studies. Please note that we have not included a discussion of selective serotonin reuptake inhibitors and tamoxifen, since this appears to be more of a pharmacologic interaction rather than a true anti-cancer effect. We have focused our discussion on drugs that may influence cancer recurrence rather than primary incidence.

Methods
For this review article, we will focus on cohort studies, prospective nested case control studies, and randomized controlled trials that presented breast cancer-specific survival or recurrence data. We have omitted case control studies because these can be subject to bias. For our search strategy, we searched PubMed through July 2012 for relevant English language studies. The major search terms used were breast neoplasms and (mortality or survival or survival analysis or survivors or recurrence). For the individual drug search terms, we used (aspirin or anti-inflammatory agents, non-steroidal), adrenergic beta-agonists, (angiotensin-converting enzyme inhibitors), statins, digoxin, and metformin – that have been evaluated among breast cancer survivors in prospective studies. We also reviewed the references lists of all relevant papers for any additional studies. We did not include studies that were presented only in abstract form at a meeting or were published only as editorial letters.

Aspirin and other non-steroidal anti-inflammatory drugs

Biosynthesis rationale/preclinical data
Aspirin and non-steroidal anti-inflammatory drugs (NSAIDs) may influence breast cancer recurrence through a number of mechanisms. They inhibit production of
prostaglandins and cyclo-oxygenase (COX), which comes in two isoforms: COX-1 and COX-2 [1]. It has been known for over 20 years that elevated tissue levels of prostaglandins have been seen in breast tissue, especially hormone receptor-negative tumors [2]. Prostaglandins can stimulate angiogenesis [3] and inhibit apoptosis [4]. In addition, prostaglandins stimulate aromatase activity and thus may affect estrogen production [5]. Aromatase is an enzyme that catalyzes the conversion of androgen precursors to estrogen, the main source of estrogen production in post-menopausal women. Aromatase inhibitors are widely used for breast cancer treatment and lower estrogen levels. Aspirin and NSAIDs could improve survival if they acted as aromatase inhibitors. Cross-sectional studies provide suggestive evidence that aspirin can influence estrogen levels, since estrogen levels are lower among women using aspirin [6]. However, prostaglandin effects may not be limited to hormone receptor-positive tumors.

There is also strong evidence that aspirin and NSAIDs may prevent early metastasis but not advanced disease. COX-2 overexpression has been associated with human breast cancer that has metastasized [7]. This may explain why early trials of NSAIDs to treat advanced or metastatic breast cancer showed little effect [8]. A recent publication reviewed the extensive experimental evidence showing that platelets promote adhesion of circulating tumor cells to the endothelium and protect them from immune elimination within the circulatory system, thus enabling future establishment of metastases. Aspirin, but not NSAIDs, inhibits platelet function [9].

Epidemiologic data
Three out of four large prospective observational studies have shown a potential survival benefit among women with breast cancer who use aspirin or NSAIDs. Kwan and colleagues [10] reported from the Life After Cancer Epidemiology (LACE) study, a prospective cohort of 2,292 survivors whose stage I to III breast cancer was diagnosed between 1997 and 2000 and who were drawn primarily from Kaiser Permanente Northern California. The authors found a reduced risk of recurrence for current regular (>3 days per week) use of ibuprofen (relative risk (RR) = 0.56, 95% confidence interval (CI) = 0.32 to 0.98) but not aspirin (RR = 1.09, 95% CI = 0.74 to 1.61). However, short follow-up (mean of 2.5 years) may have precluded the detection of an association. Blair and colleagues [11] reported a borderline reduced risk of breast cancer death (RR = 0.64, 95% CI = 0.39 to 1.05) for any use of NSAIDs (aspirin and non-aspirin NSAIDs combined) after diagnosis among 591 post-menopausal women with breast cancer and a reduced risk of breast cancer death for aspirin use alone (RR = 0.53, 95% CI = 0.30 to 0.93). In the combined group of any NSAID use, use of aspirin only (43%) was considerably more common than use of non-aspirin NSAIDs only (10%) or use of both (27%). In the Nurses’ Health Study [12], we reported on 4,164 women with early-stage breast cancer and found a reduced risk of breast cancer death for aspirin use after diagnosis (RR = 0.51, 95% CI = 0.41 to 0.65). The survival benefit was similar for estrogen receptor (ER)-positive and -negative tumors. There was a suggestion of a protective association with NSAID intake as well, but power was limited. Intriguingly, among a subset of 2,001 subjects for whom we had tumor samples to perform COX-2 immunohistochemistry, we found a similar association for aspirin use among those with COX-2-positive tumors (RR = 0.64, 95% CI = 0.43 to 0.96) and COX-2-negative tumors (RR = 0.57, 95% CI = 0.44 to 0.74), suggesting that the aspirin mechanism for breast cancer may be independent of COX-2 [7]. Aspirin binds covalently to and inhibits both COX-1 and COX-2. In breast carcinogenesis, in contrast to colon cancer, COX-1 activity may be relatively more important [13].

Among 1,024 breast cancer cases from a population-based case control study followed as a cohort for an average of 7 years, Li and colleagues [14] reported a non-statistically significant reduced risk of overall mortality among those using aspirin (RR = 0.82, 95% CI = 0.54 to 1.24) and a similar risk for breast cancer mortality (RR = 0.89, 95% CI = 0.53 to 1.52).

In addition to the prospective studies, randomized trial data have demonstrated an effect of aspirin on cancer recurrence. In the UK, Rothwell and colleagues [15] pooled data from five large randomized trials of aspirin to prevent vascular disease. The purpose of the pooled analysis was to examine the effect of aspirin on cancer metastases presenting during or after the trials’ follow-up. In the pooled data, those subjects allocated to aspirin had a reduced risk of cancer with distant metastasis, mainly due to a reduced risk of metastatic adenocarcinoma (RR = 0.52, 95% CI = 0.35 to 0.75). In addition, patients with adenocarcinoma who did not have metastasis at initial diagnosis and who remained on aspirin up to or after diagnosis had a markedly reduced risk of metastasis during follow-up (RR = 0.31, 95% CI = 0.15 to 0.76). Examination of case fatality by individual cancers was hampered by small numbers, but there was a suggestion of reduced case fatality for breast cancer (RR = 0.16, 95% CI = 0.02 to 1.19). Because these dramatic pooled findings were similar in the one trial which used a low-dose (75 mg) slow-release formulation of aspirin designed to inhibit platelet function only in the portal circulation and not to have systemic effects, the authors speculate that aspirin’s effect on platelet-mediated formation of metastases is the likely mechanism [15].

Corroboration was provided by a linked meta-analysis comparing data from observational studies with those
from the randomized trials. The risk of breast cancer with distant metastases pooled from observational studies (RR = 0.58, 95% CI = 0.20 to 1.71) was similar to that found in randomized trials but, owing to small numbers, did not reach statistical significance [16].

In conclusion, abundant preclinical and epidemiologic data support a protective role for aspirin and NSAIDs in breast cancer survival [10-12,16]. In addition, pooled data from five large randomized trials of aspirin used to prevent vascular disease have demonstrated a reduced risk of metastatic and fatal adenocarcinoma, including breast cancer, among those allocated to aspirin. Results from the randomized trials hint that low-dose (75 mg) aspirin may be effective, suggesting that inhibition of platelet function may be the key mechanism in preventing metastases [15]. However, as we reported in a previous review, several other mechanisms may be involved for breast cancer [6]. Although COX-2 expression is strongly linked to the cancer process, for breast cancer these beneficial effects are not solely or primarily caused by inhibiting COX-2 [7]. Although non-aspirin NSAIDs may also improve breast cancer survival, evidence is currently strongest for aspirin [13,15].

**Beta-blockers**

**Biological rationale/preclinical data**

Patients and clinicians have shown great interest in the possible link between stress and cancer initiation and progression. In a 2006 review, Antoni and colleagues [17] elucidated how bio-behavioral influences (for example, life stress, psychological processes, and health behaviors) could plausibly affect cancer processes through neuroendocrine pathways. In fact, evidence is stronger for an effect on cancer progression than on cancer initiation [17].

The major neuroendocrine transmitters of the stress response are catecholamines, and beta-adrenergic receptors mediate most of the effects of catecholamines. Preclinical studies in several types of cancer (ovarian, nasopharyngeal, prostate, and pancreatic) have shown catecholamine stimulation to increase angiogenesis, tumor invasion, metastasis, and inhibit apoptosis; many of these effects could be inhibited by the use of beta-adrenergic blocker drugs such as propranolol [18].

In a recently published study, mice with mammary cancer subjected to chronic stress had neuroendocrine activation that did not affect growth of the primary tumors but increased distant metastases 30-fold. This tumor spread could be inhibited by treatment with propranolol [19]. Beta-adrenergic receptors have been found in human breast cancer cells [20].

**Epidemiologic data**

Four observational studies among women with breast cancer (three cohorts and one prospective nested case control) have examined the association between intake of beta-blockers and risk of either breast cancer mortality or recurrence. In 2010, Powe and colleagues [21] reported on 466 stage I to III UK breast cancer patients with more than 10 years of follow-up; 92 (20%) had pre-existing hypertension and 43 of these (9%) were treated with beta-blockers. In multivariate models controlling for age and tumor characteristics, women using beta-blockers had marked decreases in breast cancer mortality (RR = 0.29, 95% CI = 0.12 to 0.72) and distant recurrence (RR = 0.43, 95% CI = 0.20 to 0.93) [21]. This initial small study was rapidly followed by three larger ones published in 2011. Melhem-Bertrandt and colleagues [22] reported on 1,413 patients with stage I to III breast cancer at the MD Anderson Cancer Center; 102 (7%) used beta-blockers. The authors hypothesized that the higher prevalence of abdominal obesity and metabolic syndrome among women with triple-negative breast cancer and its link to adrenergic dysregulation and also high expression of beta-adrenergic receptors in triple-negative breast cancer cell lines could make these patients particularly sensitive to beta-blocker treatment. The authors reported a decreased risk of relapse (RR = 0.52, 95% CI = 0.31 to 0.88) for users of beta-blockers among all patients. This was most pronounced among the 377 patients with triple-negative breast cancer (RR = 0.30, 95% CI = 0.10 to 0.87) [22]. Ganz and colleagues [23] reported on 1,779 women with stage I to IIIA breast cancer from the LACE cohort, all of whom had linked pharmacy records. Mean follow-up was 8.2 years, and 270 of the women (15%) used beta-blockers. The authors found non-statistically significant decreased risks of breast cancer death (RR = 0.76, 95% CI = 0.44 to 1.33) and recurrence (distant, loco-regional, or contralateral, RR = 0.85, 95% CI = 0.57 to 1.32) among users of beta-blockers [23]. The fourth study is a nested case control reported by Barron and colleagues [24] linking the Irish national cancer registry and pharmacy registries. They studied 5,333 women with stage I to IV breast cancer. Five hundred ninety-five beta-blocker users (70 using propranolol and 525 using atenolol) were matched 1:2 on factors associated with breast cancer screening and other healthy behaviors (including socioeconomic status, smoking, aspirin, and statin use) to controls not using a beta-blocker. The 70 propranolol users had a markedly decreased risk of breast cancer mortality (RR = 0.19, 95% CI = 0.06 to 0.60) compared with non-beta-blocker users, but this was based on only four breast cancer deaths among propranolol users. Propranolol users also were less likely to present with locally advanced or metastatic tumors. No such association was seen for atenolol [24]. Propranolol is non-selective and blocks both beta-1 and beta-2 adrenergic receptors, whereas atenolol blocks only beta-1. Historically, over time, patterns of use have
moved from the non-selective to the cardioselective (beta-1) blockers. The authors present preclinical evidence that beta-2 signaling may be more important for cancer metastasis, and their results would seem to bolster this hypothesis [24].

In conclusion, tantalizing evidence from preclinical studies supports a role for beta-blockers to inhibit breast cancer metastasis and potentially improve survival. Observational studies are hampered by a relatively low prevalence (approximately 10% to 15%) of beta-blocker use and the fact that only a subset of non-selective beta-blockers may be effective, and these have been used less over time. Additionally, the beta-blocker effect may differ by tumor subtype, with a stronger effect seen among triple-negative tumors.

**Angiotensin-converting enzyme inhibitors and angiotension type I receptor blockers**

**Biological rationale/preclinical data**

The renin-angiotensin-aldosterone system (RAAS) has a potential role in breast cancer control. Angiotensin I is cleaved into angiotensin II by angiotensin-converting enzyme (ACE). Angiotensin II interacts with angiotensin type I receptors to promote aldosterone secretion and vasoconstriction. ACE inhibitors (ACEIs) and angiotensin type I receptor blockers (ARBs) are commonly used to treat hypertension, congestive heart failure, and chronic kidney disease [18].

Polymorphisms of the RAAS genes leading to increased activity of the system have been associated with increased risk of breast cancer [25,26]. Breast cancer cells have been found to express components of the RAAS [27]. RAAS stimulation of breast cancer cells can increase cell proliferation via protein kinase C activation and epidermal growth factor receptor transactivation as well as activating the P13K-kinase B (AKT) pathway [28,29]. RAAS stimulation of hormone receptor-negative breast cancer cells has been shown to increase expression of angiogenesis-related genes [27].

**Epidemiologic data**

Two observational studies that previously reported on use of beta-blockers and breast cancer survival also reported on ACEI/ARB use. Contrary to the hypotheses generated by the preclinical evidence, neither the MD Anderson cohort of 1,413 patients reported by Melhem-Bertrandt and colleagues [22] nor the LACE cohort (n = 1,779) reported by Ganz and colleagues [23] found any evidence of decreased recurrence, breast cancer mortality, or total mortality among women with breast cancer using ACEIs or ARBs (Table 1) [22,23]. In fact, an elevated risk of recurrence was found among the LACE cohort (RR = 1.56, 95% CI = 1.02 to 2.39) [23]. In a smaller cohort of 703 stage II/III breast cancer patients from Albert Einstein Medical Center, Chae and colleagues [30] reported a reduced risk of breast cancer recurrence among those using ACEI/ARB (RR = 0.49, 95% CI = 0.31 to 0.76), but total mortality was not reduced. Therefore, despite promising preclinical evidence for ACEIs/ARBs, substantial evidence for a protective effect among women with breast cancer is currently lacking.

**Statins**

**Biological rationale/preclinical data**

Statins – HMG-CoA (3-hydroxy-3-methyl-glutaryl-CoA) reductase inhibitors – are widely used lipid-lowering drugs. Interestingly, lipophilic statins (for example, simvastatin, lovastatin, and fluvastatin) have been shown in vitro to inhibit breast cancer cell growth and proliferation with a variety of hypothesized mechanisms. In multiple cell lines, statins can inhibit prenylation (post-translational modification) of multiple proteins, including those in the Ras family, which is involved in signal transduction and presumed to be important in carcinogenesis [31]. Statins may also inhibit histone deacetylase activity [32]. Drugs targeting histone deacetylation are already approved for lymphoma and have activity in other cancers as well. Several clinical trials in cancers other than breast cancer have suggested that statins used in conjunction with chemotherapy may improve efficacy [18]. In terms of breast cancer incidence, studies on the effects of statins are mixed. However, the only published, cohort studies on the association between statins and breast cancer recurrence have consistently shown a decreased risk of recurrence.

**Epidemiologic data**

The first published study used the LACE population. Statin use was confirmed via pharmacy records, and health outcomes were verified by questionnaire and medical record review. Use of statins for more than 100 days after diagnosis compared with shorter-term use was associated with a non-significant decreased risk of cancer recurrence (RR = 0.67, 95% CI = 0.39 to 1.33) after adjustment for age at diagnosis, race, body mass index (BMI), cancer stage, and tamoxifen use. Breast cancer recurrence risk decreased with increasing duration of post-diagnosis statin use (P for trend = 0.02). However, power was limited, as there were only 16 recurrences among survivors who used statins more than 100 days after diagnosis. The primary statin used in this cohort was the lipophilic lovastatin, which accounted for 84% of statin use among regular statin users [33]. The study by Chae and colleagues [30], which was previously cited on ACEI/ARB, also evaluated the association with statin use and reported a decrease risk of recurrence (multivariate hazard ratio (HR) = 0.40, 95% CI = 0.24 to 0.67) and no
Table 1. Cohort studies of aspirin, beta-blockers, angiotensin-converting enzyme inhibitors, statins, and metformin and breast cancer survival

<table>
<thead>
<tr>
<th>Authors (year)</th>
<th>Study population</th>
<th>Number taking drug (percentage)</th>
<th>Number</th>
<th>Years of follow-up</th>
<th>Recurrence RR (95% CI)</th>
<th>Breast cancer mortality RR (95% CI)</th>
<th>Total mortality RR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aspirin</td>
<td>Kwan et al. [10] (2007) LACE cohort, stages I-IIIA, with linked pharmacy records</td>
<td>-</td>
<td>2,292</td>
<td>Mean 2.5</td>
<td>1.09*</td>
<td>-</td>
<td>-</td>
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<tr>
<td></td>
<td>Blair et al. [11] (2007) Iowa Women’s Health Study, post-menopausal</td>
<td>-</td>
<td>591</td>
<td>Maximum 9.5</td>
<td>0.53 (0.30-0.93)</td>
<td>0.53 (0.36-0.79)</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Holmes et al. [12] (2010) Nurses’ Health Study, stages I-III</td>
<td>-</td>
<td>4,164</td>
<td>Maximum 30</td>
<td>0.51 (0.41-0.65)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Li et al. [14] (2012) Cases from a population-based case control study followed as</td>
<td>-</td>
<td>1,024</td>
<td>Mean 7.3</td>
<td>0.89 (0.52-1.52)</td>
<td>0.82 (0.54-1.24)</td>
<td>-</td>
</tr>
<tr>
<td>Beta-blockers</td>
<td>Powe et al. [21] (2010) UK, stages I-III</td>
<td>-</td>
<td>466</td>
<td>Mean 10.3</td>
<td>0.43* (0.20-0.93)</td>
<td>0.29 (0.12-0.71)</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Melhem-Bertrandt et al. [22] (2011) MD Anderson triple-negative stages I-III</td>
<td>-</td>
<td>1,413</td>
<td>Median 4.6</td>
<td>0.52* (0.31-0.88)</td>
<td>-</td>
<td>0.64 (0.38-1.07)</td>
</tr>
<tr>
<td></td>
<td>Ganz et al. [23] (2011) LACE cohort, stages I-IIIA, with linked pharmacy records</td>
<td>-</td>
<td>1,779</td>
<td>Mean 8.2</td>
<td>0.86* (0.57-1.32)</td>
<td>-</td>
<td>0.76 (0.44-1.33)</td>
</tr>
<tr>
<td></td>
<td>Barron et al. [24] (2011) Nested case control linked Irish cancer and pharmacy registries, stages I-IV</td>
<td>-</td>
<td>5,333</td>
<td>Median 3.5</td>
<td>-</td>
<td>0.19 (0.06-0.60)</td>
<td>-</td>
</tr>
<tr>
<td>Angiotensin-converting enzyme inhibitors (ACEIs) and angiotensin type I receptor blockers (ARBs)</td>
<td>Melhem-Bertrandt et al. [22] (2011) MD Anderson triple-negative stages I-III</td>
<td>-</td>
<td>1,413</td>
<td>Median 4.6</td>
<td>0.82* (0.54-1.26)</td>
<td>-</td>
<td>0.99 (0.65-1.51)</td>
</tr>
<tr>
<td></td>
<td>Ganz et al. [23] (2011) LACE cohort, stages I-IIIA, with linked pharmacy records</td>
<td>-</td>
<td>1,779</td>
<td>Mean 8.2</td>
<td>1.56* (1.02-2.39)</td>
<td>1.27 (0.74-2.19)</td>
<td>1.04 (0.72-1.51)</td>
</tr>
<tr>
<td></td>
<td>Chae et al. [30] (2011) Stage II/III hospital patients</td>
<td>-</td>
<td>703</td>
<td>Median 4.6</td>
<td>0.49* (0.31-0.76)</td>
<td>-</td>
<td>NS</td>
</tr>
<tr>
<td>Statins</td>
<td>Kwan et al. [33] (2008) LACE cohort, stages I-IIIA, with linked pharmacy records</td>
<td>-</td>
<td>1,811</td>
<td>Mean 5.0</td>
<td>0.67* (0.39-1.13)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Ahern et al. [34] (2011) Danish registry cohort with linked pharmacy records, stages I-III</td>
<td>-</td>
<td>18,769</td>
<td>Mean 6.8</td>
<td>0.83* (0.70-0.98)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Chae et al. [30] (2011) Stage II/III hospital patients</td>
<td>-</td>
<td>703</td>
<td>Median 4.6</td>
<td>0.40* (0.24-0.67)</td>
<td>-</td>
<td>NS</td>
</tr>
<tr>
<td>Metformin</td>
<td>He et al. [42] (2012) MD Anderson HER2-positive stages II-IV</td>
<td>-</td>
<td>1,983</td>
<td>Median 4.0</td>
<td>-</td>
<td>0.47 (0.24-0.90)</td>
<td>0.52 (0.28-0.97)</td>
</tr>
<tr>
<td></td>
<td>Bayraktar et al. [46] (2012) MD Anderson, triple-negative on adjuvant chemotherapy, stages I-III</td>
<td>-</td>
<td>1,448</td>
<td>Median 5.2</td>
<td>1.63* (0.87-3.06)</td>
<td>for diabetics not on metformin, compared with diabetics on metformin</td>
<td>1.22 (0.66-2.28) for diabetics not on metformin, compared with diabetics on metformin</td>
</tr>
<tr>
<td></td>
<td>Currie et al. [44] (2012) UK, patients with solid tumors, including breast cancer stages I-IV</td>
<td>-</td>
<td>112,408 (25,575 breast cancer)</td>
<td>Mean 9.3, median 6.8 (overall survival)</td>
<td>-</td>
<td>0.96* (0.64-1.43)</td>
<td></td>
</tr>
</tbody>
</table>

*aAny recurrence; *b* distant recurrence; *cACEI or ARB use; *dACEI only; *e* breast cancer survivors only; see [44] for all cancer survivor numbers. CI, confidence interval; LACE, Life After Cancer Epidemiology; NS, not significant; RR, relative risk.
impact on overall survival. Power was also limited in this study, as there were only 19 recurrences among those who used statins at least 6 months [30].

The largest study to date was a population-based Danish cohort of 18,769 survivors of stage I to III breast cancer diagnosed between 1996 and 2003. Statin use was assessed via linkage to the National Registry of Medicinal Products, which tracks all prescriptions since 1995. Breast cancer recurrence was confirmed via cancer registry data. Women who used primarily lipophilic statins had a lower risk of recurrence compared with non-users (10-year HR = 0.73, 95% CI = 0.60 to 0.89) after adjustment for age at diagnosis, menopausal status, cancer stage, ER status, cancer treatment, and use of other relevant non-prescription medications. In contrast, women who used primarily hydrophilic statins (for example, atorvastatin, pravastatin, or rosuvastatin) had the same risk of breast cancer recurrence as non-users (10-year adjusted HR = 1.2, 95% CI = 0.79 to 1.70). It should be noted that the analyses with hydrophilic statins were limited for power; there were only 39 recurrences at 10 years compared with 182 recurrences among users of lipophilic statins. The primary statin used in the Danish cohort was simvastatin (accounting for 72% of prescriptions among statin users), which is the most lipophilic statin. Stratified models showed no difference by grade, ER status, or type of treatment [34]. In the US, prior to the introduction of the generic statins in 2006, Lipitor (atorvastatin; Pfizer Inc, New York, NY, USA) had the largest market share [35].

Although the pharmacologic differences between lipophilic and hydrophilic statins in terms of their cholesterol-lowering effects have been well characterized, less is known about statins' pleiotropic effects. Hydrophilic, but not lipophilic, statins may increase mevalonate synthesis in extra-hepatic tissues and this may result in differential effects on cancer development. Lipophilic statins also tend to accumulate more in fat and have higher plasma protein binding than hydrophilic statins, and this could result in more extra-hepatic activity and systemic effects [36].

In summary, the observational data on statins influencing breast cancer recurrence risk are compelling and provide a strong justification for a randomized trial. Furthermore, similar to the aspirin data, data from the multiple randomized trials of statins for cardiovascular disease prevention/treatment should be pooled to evaluate for possible effects on cancer recurrence and mortality.

**Digoxin**

**Biological rationale/preclinical data**

Cardiac glycosides (the most widely used of which is digoxin) have also demonstrated anti-tumor effects in vitro, presumably through inhibition of Na’K’-ATPase. A variety of anti-tumor effects, including induction of apoptosis and inhibition of DNA topoisomerase II, have been observed in cell lines [18,37,38].

**Epidemiologic data**

Only one study (which has been published several times at varying times of follow-up) has evaluated the association between breast cancer recurrence and digoxin use. With 22 years of follow-up, breast cancer survivors who used digoxin had a lower rate of death (6%, n = 32) from breast cancer than non-users (34%, n = 143) [39]. To date, no other study on this topic has been published.

**Anti-diabetic medications**

**Biological rationale/preclinical data**

Although insulin, sulfonylureas, thiazolidinediones, and metformin all lower glucose levels and have been used successfully for diabetes treatment, they have different mechanisms of action and different associations with breast cancer survival. In terms of its anti-diabetic effect, metformin inhibits hepatic gluconeogenesis and improves insulin sensitivity in peripheral tissue [40]. The mechanism of action for thiazolidinediones (for example, rosiglitazone) is not completely understood, but they increase glucose utilization in adipose, muscle, and hepatic tissue, most likely by activating peroxisome proliferator-activated receptors (PPARs). Both metformin and thiazolidinediones are associated with lower fasting insulin and C-peptide levels. Importantly, insulin is a known mitogen and can activate insulin-like growth factor 1 (IGF-1) receptors. Sulfonylureas stimulate the release of insulin from pancreatic beta cells. Higher circulating insulin has been associated with worse breast cancer mortality [41]. Since most of the studies that have evaluated the effect of anti-diabetic medications on cancer survival have compared recurrence risks across categories of drugs, all classes will be discussed concurrently [42].

**Epidemiologic data**

Because of the important biologic differences among the anti-diabetic medications, in addition to the overall search criteria, we limited our review to studies that reported results separately for types of anti-diabetic medication. In addition, we reviewed an intriguing study looking at pathologic complete response rates by anti-diabetic medication use. Interestingly, the studies have been surprisingly consistent, showing a decreased risk of all-cancer mortality among diabetics who use metformin compared with those who use sulfonylureas or insulin. For a variety of reasons, these observational studies can be challenging to interpret. Diabetics have greater comorbidity and shorter life expectancy and so may get
less aggressive cancer screening or treatment and may have a higher risk of adverse events from treatments [43]. In addition, the use of certain anti-diabetic drugs may be associated with certain prognostic factors. For example, metformin users tend to be younger than subjects who use other anti-diabetic medications. Analyses are further complicated by the fact that diabetic patients switch back and forth over time between mono-therapy and combined therapy.

The study with the largest number of breast cancer cases was a retrospective study of 112,408 subjects from the UK with a diagnosed solid tumor; 8,392 of the subjects had type II diabetes. Medication use was confirmed with pharmacy records. Among 25,575 breast cancer survivors (1,182 with type II diabetes), there was an increased risk of breast cancer mortality associated with having type II diabetes (unadjusted HR = 1.32, 95% CI = 1.17 to 1.49). However, metformin use among cancer survivors was associated with a decreased risk of overall mortality (adjusted HR = 0.85, 95% CI = 0.78 to 0.93) compared with non-diabetics. In contrast, diabetics who used sulfonylureas (HR = 1.13, 95% CI = 1.05 to 1.21) or insulin (HR = 1.13, 95% CI = 1.01 to 1.27) had an increased risk of mortality. These differences were not significant in analyses limited to breast cancer survivors [44]. In two companion studies, investigators from MD Anderson focused on outcomes among specific breast cancer subgroups. In the first study, they retrospectively reviewed 1,983 consecutive patients with stage II to IV HER2-positive breast cancer (154 diabetics) and found again that diabetes was associated with worse overall survival (adjusted HR = 1.42, 95% CI = 1.04 to 1.94). However, in multivariate analyses, survival differed by anti-diabetic therapy, and insulin users had a shorter survival than diabetics who did not use insulin and non-diabetics. In contrast, diabetics who used metformin had significantly longer survival compared with diabetics who did not use metformin or non-diabetics. In multivariate analyses, both metformin use and thiazolidinedione use were associated with improved survival after adjustment for age, BMI, ER status, and use of insulin or insulin secretagogue therapy. In an analysis of competing risks among diabetic patients, metformin and thiazolidinediones were associated with decreased breast cancer-specific mortality [42]. The second study focused on triple-negative breast cancer and included 1,448 women (including 63 diabetics on metformin and 67 diabetics not on metformin) with stage I to III triple-negative breast cancer treated with adjuvant chemotherapy between 1995 and 2007. In multivariate analyses, both diabetics not on metformin and non-diabetics had a non-significant increased risk of distant metastases (HR = 1.63, 95% CI = 0.87 to 3.06 and HR = 1.62, 95% CI = 0.97 to 2.71, respectively) compared with diabetics on metformin. The study was limited by the small number of distant recurrences among the diabetics (18 among metformin users and 26 among non-metformin users) and limited data on metformin use, which were available for the adjuvant chemotherapy period only [46].

Among 2,529 breast cancer patients who received neo-adjuvant chemotherapy for breast cancer at MD Anderson, the rate of pathologic complete response was higher in the metformin group (24%, n = 68) compared with diabetics who did not take metformin (8%, n = 87) and non-diabetics (16%, n = 2,374) (overall P for difference = 0.02). Metformin was also independently predictive of the chances of pathologic complete response (adjusted odds ratio = 2.95, 95% CI = 1.07 to 8.17) after adjustment for age, diabetes, BMI, stage, grade, ER status, and treatment [45].

The molecular basis for metformin’s inhibition of cancer cell growth is not known but is hypothesized to be its ability to inhibit P13-kinase/AKT/mammalian target to rapamycin (mTOR) signaling via activation of the LKB1/AMP-activated protein kinase (AMPK) pathway. Of all the medications presented in this review article, metformin is the only one that will have randomized trial data evaluating its effect on breast cancer recurrence within the near future. The National Cancer Institutes of Canada and US are enrolling subjects for a phase III study to evaluate the effect of metformin compared with placebo among women with higher-risk stage I and stage II or III breast cancer (NCIC MA32). The accrual of this study, which opened in April 2010 and is expected to close in 2016, is estimated to be 3,852, and the results are eagerly awaited. Because of their effects on the PPAR pathway, ongoing phase I clinical trials are using a variety of thiazolidinediones in combination with chemotherapy for advanced solid tumors.

Conclusions

Substantial scientific evidence supports the hypothesis that several common and relatively safe drugs may reduce breast cancer mortality among breast cancer survivors by an amount that rivals the benefit of currently used therapies. In particular, the evidence is strongest for aspirin (approximately 50% reduction), statins (approximately 25% reduction), and metformin (approximately 50% reduction).

We believe that randomized trials of aspirin, metformin, and statins are essential to move the field forward. Despite the compelling evidence presented in this review, it is based primarily on observational studies, which are subject to confounding. These drugs are generally safe, and their side effect profiles compare favorably with those of drugs used to treat cancer. However, we cannot estimate the overall risk-benefit ratio of these drugs without a randomized trial. For
example, aspirin has a measurable risk of gastrointestinal [47] and central nervous system [48] bleeding, and there is a suggestion of hepatotoxicity with metformin [49]. In addition, aspirin is not taken in a fixed dose; a randomized trial could help to establish the lowest effective dose.

If these findings are confirmed in randomized trials among breast cancer survivors, the public health impact would be immense. We estimate that, if aspirin is effective, using it to treat all patients with breast cancer in the US could potentially save 10,000 lives per year. In addition, if one considers the possible benefit in the developing world of an inexpensive, widely available medicine, the impact is truly staggering: an estimated 75,000 lives would potentially be saved each year.

In an era in which we struggle to contain health-care costs, the extra costs for patients with breast cancer in the US would be minimal. For developing countries, it could mean the difference between some adequate treatment and none. Whereas new cancer treatments typically benefit only patients in wealthy countries because of the costs, these drugs would be a breast cancer treatment available to every part of the world. The results of these trials could be truly transformative and change the treatment of breast cancer across the globe with what millions of people already have in their medicine cabinet.

Given the overwhelming weight of the biologic and observational data, randomized trials are the definitive way to assess the risk-benefit balance for breast cancer survivors. One such trial is under way for metformin. A similar trial for aspirin is definitely warranted, and possibly one for statins. We estimate that a trial of aspirin would require approximately 3,000 women with stage II or III breast cancer randomly assigned 1:1 and followed for 5 years and cost approximately $15 million USD. However, because these drugs are generic and widely available, there is little industry incentive to support such studies. We propose that the cost is small given the potential benefit. Who will fill this need?

**Abbreviations**

ACE, angiotensin-converting enzyme; ACEI, angiotensin-converting enzyme inhibitor; ARB, angiotensin type I receptor blocker; BMI, body mass index; CI, confidence interval; COX, cyclooxygenase; ER, estrogen receptor; HR, hazard ratio; LACE, Life After Cancer Epidemiology; NSAID, non-steroidal anti-inflammatory drug; PPAR, peroxisome proliferator-activated receptor; RAAS, renin-angiotensin-aldosterone system; RR, relative risk.

**Competing interests**

The authors declare that they have no competing interests.

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