A unifying biology of sex steroid-induced apoptosis in prostate and breast cancers

Philipp Y Maximov, Balkees Abderrahman, Ramona F Curpan, Yousef M Hawsawi, Ping Fan and V Craig Jordan

Abstract

Prostate and breast cancer are the two cancers with the highest incidence in men and women, respectively. Here, we focus on the known biology of acquired resistance to antihormone therapy of prostate and breast cancer and compare laboratory and clinical similarities in the evolution of the disease. Laboratory studies and clinical observations in prostate and breast cancer demonstrate that cell selection pathways occur during acquired resistance to antihormonal therapy. Following sex steroid deprivation, both prostate and breast cancer models show an initial increased acquired sensitivity to the growth potential of sex steroids. Subsequently, prostate and breast cancer cells either become dependent upon the antihormone treatment or grow spontaneously in the absence of hormones. Paradoxically, the physiologic sex steroids now kill a proportion of selected, but vulnerable, resistant tumor cells. The sex steroid receptor complex triggers apoptosis. We draw parallels between acquired resistance in prostate and breast cancer to sex steroid deprivation. Clinical observations and patient trials confirm the veracity of the laboratory studies. We consider therapeutic strategies to increase response rates in clinical trials of metastatic disease that can subsequently be applied as a preemptive salvage adjuvant therapy. The goal of future advances is to enhance response rates and deploy a safe strategy earlier in the treatment plan to save lives. The introduction of a simple evidence-based enhanced adjuvant therapy as a global healthcare strategy has the potential to control recurrence, reduce hospitalization, reduce healthcare costs and maintain a healthier population that contributes to society.

Introduction

Despite advances in understanding the molecular biology of prostate and breast cancers, they are still the most frequently diagnosed cancers in men and women, in the United States. There is no completely effective preventative for either prostate or breast cancer. Advances in the chemoprevention of prostate cancer remain controversial (Bosland 2016) and none are approved by the Food and Drug Administration (FDA). As a result, there were 220,800 new cases of prostate cancer reported with 27,540 deaths (Siegel et al. 2015) in men. Advances in chemoprevention have been made in breast cancer (Jordan 2014b, 2016, 2017a, Cuzick 2015, Cuzick et al. 2016), but the task of implementation is not trivial (Kaplan et al. 2005, Owens et al. 2011, Smith et al. 2016). There were 231,840 new breast cancer cases reported in 2015, accounting for almost 29% of the total estimated
female cancers (Siegel et al. 2015). Approximately, 40,290 deaths from breast cancer occurred in 2015 accounting for 14% of total deaths from cancers in women (Siegel et al. 2015). These figures present a major challenge in clinical research and for healthcare systems worldwide. Indeed, it is estimated that the incidence of breast cancer will increase by 50% from the level in 2011 for the combination of Indolent Lesion of Epithelial Origin (IDLE) and invasive disease by 2050 (Anderson et al. 2011). The increased survival of an aging population is the cause of the relentless rise in cancer. The goal of a cure remains. However, in practical terms, new affordable strategies are required for individuals affected by prostate or breast cancers to remain productive members of their families and society.

The sex steroid hormones i.e. androgens in men and estrogens in women play critical roles in the development and progression of prostate and breast cancers. Prostate cancer development relies on the androgen receptor (AR), whereas breast cancer development primarily relies on the estrogen receptor (ER). The majority of prostate and breast cancers are hormone dependent (Fig. 1). Antihormone therapies have had a profound impact in reducing the burden from breast cancer, worldwide (Jordan 2003, Santen et al. 2009a, Sledge et al. 2014). Here, we will address whether the lessons learned in breast cancer can

---

Figure 1
A schematic representation of the androgen and estrogen deprivation therapy in prostate cancer and pre- and postmenopausal women with breast cancer. (A) The hypothalamic–pituitary–gonadal axis in prostate cancer with their therapeutic targets. The hypothalamus produces gonadotropin-releasing hormone (GnRH), which stimulates the adenohypophysis of the pituitary to produce adrenocorticotropic hormone (ACTH). This in turn, stimulates the adrenal gland to produce androgens: dehydroepiandrosterone sulfate (DHEA-S) predominately, DHEA and androstenedione (AD) into the circulation. These androgens (A), alongside testosterone (T) from the testes, are converted in the prostate to their potent form, dihydrotestosterone (DHT). Dihydrotestosterone stimulates the growth of prostate cancer cells and exerts a negative feedback loop onwards to the hypothalamus and pituitary. Both, GnRH agonists/antagonists suppress LH production and cause a subsequent decline in serum testosterone to castrate levels. However, GnRH agonists (with chronic use) lead to the downregulation of GnRH receptors, whereas, GnRH antagonists usually cause an immediate blockade to the receptor. At the adrenal level, abiraterone inhibits adrenal androgen de novo steroidogenesis. At the prostate level, androgen receptor (AR) inhibitors are used and they have different mechanisms of action. For example, enzalutamide competitively inhibits the AR binding to DHT, inhibits nuclear translocation, and DNA and cofactor binding. Whereas, Bicalutamide is a highly selective, competitive and silent antagonist to the AR, which was also found to accelerate AR degradation. (B) The hypothalamic–pituitary–gonadal axis in premenopausal women with breast cancer and their therapeutic targets. The hypothalamus produces gonadotropin-releasing hormone (GnRH), which stimulates the adenohypophysis of the pituitary to produce luteinizing hormone (LH) and follicle-stimulating hormone (FSH). This in turn, stimulates the granulosa cells in the ovarian follicles to produce estrogen. However, FSH in particular stimulates the granulosa cells to produce inhibin, which suppresses FSH in a feedback loop and activin, a peripherally produced hormone that stimulates GnRH cells. Estrogen stimulates the growth of breast cancer cells, and exerts a negative feedback loop onwards to the hypothalamus and pituitary. Ovarian suppression can be achieved with LHRH superagonists such as goserelin, which is an analogue of LHRH, and a GnRH or LHRH agonist. Goserelin initiates a flare of LH production and ultimately leads to receptor downregulation. Antiestrogens can be estrogen receptor (ER) competitive blockers such as the Selective ER Modulators (SERMs, i.e. tamoxifen), or pure antiestrogens or what is known as a Selective ER Downregulators (SERDs, i.e. fulvestrant). Third-generation aromatase inhibitors (i.e. anastrozole, letrozole, exemestane) selectively block the aromatase enzyme system at the breast cancer level and therefore suppress estrogen synthesis. (C) The hypothalamic–pituitary–gonadal axis in postmenopausal women with breast cancer and their therapeutic targets. The differences from premenopausal women is that the ovarian follicles are depleted, therefore there is no active production of estrogen and progesterone. This leads to a dramatic increase in GnRH, an increase in FSH serum level relatively to that of LH through the feedback loops. Ovarian suppression is not used as a treatment option.

http://erc.endocrinology-journals.org
https://doi.org/10.1530/ERC-17-0416
© 2018 The authors
Published by Bioscientifica Ltd.
Printed in Great Britain

This work is licensed under a Creative Commons Attribution 4.0 International License.
be applied to prostate cancer therapy. Whether treatment strategies are the same or not for both diseases, resistance to antihormone treatments occurs in both prostate and breast cancers.

Currently, resistance to antihormone therapies in prostate and breast cancers are categorized as acquired resistance and de novo (intrinsic) resistance. It is considered that de novo resistance has the same mechanisms as the acquired resistance (Hoimes & Kelly 2010, Miller 2013), for the exception that these mechanisms are in place before the antihormone therapy is applied. We will focus on acquired resistance. In this review, we summarize the development of treatment approaches, the antihormonal agents used for the control of both diseases and the current understanding of the evolution of resistance to antihormonal therapies. We bring together these two major sex steroid-related diseases to define similarities and differences and compare and contrast treatments based on acquired antihormone resistance. We discuss the similarities of the phenomenon of sex steroid-induced apoptosis in both types of cancers after acquisition of antihormone resistance and explore the possibility that this new knowledge will have clinical applications. An innovative treatment approach that delivers affordable healthcare will save lives globally.

Hormonal therapies for prostate and breast cancer

A diagnosis of advanced prostate cancer or breast cancer was a death sentence before 1940s, with patients dying within 1–2 years after diagnosis. Today, these same patients will have an earlier diagnosis, better care, but will still die within 3 years of diagnosis of stage IV disease. The number of patients with advanced prostate cancer has declined in the past 70 years, as early detection and diagnosis with proper treatment and monitoring has increased the 5-year survival rate up to 80–90% (Kirby et al. 2011). The change in the approach to treatment started when Professor Charles Huggins reported the response of metastatic prostate cancer (MPC) to androgen deprivation therapy (ADT), using surgical castration or high-dose synthetic estrogen therapy (Huggins & Hodges 1941). Diethylstilbestrol (DES) became a standard of care. Huggins won the Noble Prize in 1966 for developing a logical treatment strategy for prostate cancer with the ADT. Since then, ADT has been used as the gold standard for the treatment of MPC.

Earlier, but parallel, advances were reported for the treatment of advanced breast cancer in women. The initial experiment of oophorectomy (Beatson 1896) was proven to be effective in 30% of premenopausal breast cancer patients with metastatic breast cancer (MBC) (Boyd 1900). This was followed by a number of surgical ablation strategies and additive hormonal therapies for MBC (Kennedy 1965).

In the mid-1940s, Alexander Haddow (Haddow et al. 1944) was the first to discover that high doses of synthetic estrogens, including DES, could be used to treat postmenopausal women with MBC with a 30% response rate. Haddow’s (Haddow et al. 1944) clinical trial showed that only breast and prostate cancers were responsive, whereas all other types were not. Nevertheless, at that time, the mechanism of action was not understood (Haddow 1970). However, one important clinical fact did emerge. High-dose estrogen was only effective as an antitumor agent in MBC if used 5 years or more after menopause. High-dose estrogen therapy became the gold standard for the treatment of women with MBC until the introduction of tamoxifen 30 years later (Jordan 2003). The biologic mechanisms and therapeutic significance of estrogen therapy was, at that time, obscure. However, the development of models to discover mechanisms of what became the new biology of estrogen-induced apoptosis (Jordan 2008, 2015a) is now the central theme of this position paper.

The discovery of the AR in the late 1960s by three independent groups of Liao (Anderson & Liao 1968), Bruchovsky (Bruchovsky & Wilson 1968) and Mainwaring (1969), was an important breakthrough, as it triggered the search for androgen antagonists. Similar advances were made with the discovery of the ER in the early 1960s. Jensen first described the binding of radiolabeled estradiol in rat estrogen target tissues (Jensen & Jacobson 1962), and three years, later in 1966 Toft and Gorski identified the actual ER protein (Toft & Gorski 1966). Nevertheless, the therapeutic breakthrough of non-steroidal antiestrogens was focused on the modulation of fertility in rodents and women during the 1960s before the discovery of the ER (Jordan 1984, Lerner & Jordan 1990).

In the early 1970s, the first non-steroidal antiandrogen flutamide was discovered (Neri et al. 1972) and was approved in 1989 by the FDA for the treatment of prostate cancer. This discovery was followed by other non-steroidal antiandrogens including nilutamide (Raynaud et al. 1979) and bicalutamide (Furr et al. 1987), which were compared to castration in MPC patients in randomized trials. Results showed that antiandrogen drugs were better tolerated than castration (Chodak et al. 1995, Seidenfeld et al. 2000). However, they are inferior therapies in regard
to overall survival (OS) and progression-free survival (PFS) (Chodak et al. 1995, Seidenfeld et al. 2000).

In 1971, an advance in physiology was made when Schally discovered the structure of the hypothalamic hormone known as the luteinizing hormone (LH)-releasing hormone (LHRH; called the gonadotropin-releasing hormone GnRH) (Schally et al. 1971). This led to an understanding of the sex steroid feedback control mechanisms orchestrated by the hypothalamo-pituitary axis (Fig. 1). Advanced prostate cancer patients who were treated with daily doses of the LHRH agonists had a 75% decrease in serum testosterone levels, a decrease or normalization of plasma acid phosphatase levels, and a significant decrease in cancer-associated bone pain (Tolis et al. 1982). In 1977, Schally received the Nobel Prize in Physiology and Medicine for discovering peptide hormone production in the brain. Many synthetic LHRH superagonists were subsequently developed for clinical use (Schally et al. 2000), such as buserelin, goserelin, leuprolide and nafarelin. Additionally, many LHRH antagonists have been developed and tested for the treatment of men with advanced prostate cancer such as orgalutran, cetrorelix and abarelix (Schally et al. 2000). An antagonist was considered to be necessary as the superagonists first stimulate gonadotropin release (which causes an androgen burst) before a desensitized and refractory state occurs. Estrogen has been used to treat prostate cancer by lowering gonadotropin levels and as a result androgen levels. Estrogen was evaluated successfully to block the stimulatory rise in gonadotropin caused by LHRH superagonists (Ahmann et al. 1987).

Fernand Labrie (Labrie et al. 1986) was one of the pioneers who developed the idea of a complete androgen blockade using combination antiandrogen therapy with flutamide and LHRH agonists or surgical castration in patients with MPC increasing the PFS and OS. Crawford and colleagues (Crawford et al. 1989) demonstrated that the combination of flutamide and leuprolide resulted in a slightly longer PFS. As a result, many physicians in the United States shifted toward combined androgen blockade as initial therapy for advanced prostate cancer. The signaling pathways of the AR and mechanism of action of different antiandrogens is depicted in Fig. 2.

In contrast, breast cancer treatment strategies followed a separate path with an early move from the treatment of MBC to adjuvant therapy following breast surgery. A key factor in the differences in the treatment strategies of prostate and breast cancer is the fact that the majority of breast cancer occurs after menopause whereas there is no hypothalamos–pituitary–ovarian communication to alter estrogen levels (Fig. 1). By contrast, a recognized menopause does not occur in men and, as a result, hormonal communication from the pituitary to the testicular target remains. Currently clinical strategies are being defined and refined to address breast cancer treatment in the premenopausal patients (Abderrahman & Jordan 2016, Rossi & Pagani 2017).

The ER became the target for tamoxifen to treat breast cancer (Jordan & Koerner 1975) based on the National Cancer Institute consensus conference in Bethesda in 1974 on ERs in human breast cancer (McGuire et al. 1975). Treatment strategies in the 1970s for breast cancer proposed, long-term adjuvant antihormone therapy (Jordan 1978, 2014b) and the possibility of chemoprevention (Jordan 1976). These treatment strategies were proposed before tamoxifen was approved for the treatment of MBC in the United States (December 29th, 1977).

The actual development of tamoxifen was not initially a major priority by the pharmaceutical industry, but dependent upon chance and the investment in young scientists (Jordan 2006, 2015b). Tamoxifen’s withdrawal from clinical development and resurrection in the 1970s with a clear strategic plan for the development of the medicine was the key to success (Jordan 2006, 2014b). Tamoxifen became the standard for antihormonal therapy of ER+ MBC (Furr & Jordan 1984, Jordan 2003, 2006). Five years of adjuvant treatment with tamoxifen improved clinical outcome compared to shorter adjuvant therapy (EBCTCG 1998), and for more than a decade, 5 years of adjuvant tamoxifen (Davies et al. 2011) (or aromatase inhibitors, AIs) was the standard of care for ER+ breast cancer. Tamoxifen was the first medicine, in a new group of medicines called the selective estrogen receptor modulators (SERMs) (Maximov et al. 2013). Ultimately, tamoxifen was the first antiestrogen to be approved by the FDA for the prevention of breast cancer in women (Jordan 2003).

Another approach to treat breast cancer inhibits the aromatase enzyme system (CYP19) that catalyzes estrogen biosynthesis in postmenopausal women. This group of medicines is called the AIs, and these are currently used for the treatment of postmenopausal breast cancer patients. Aminoglutethimide was the first AI introduced, which has an efficacy in MBC patients (Lipton & Santen 1974). Nevertheless, all the AIs used in the early 1970s were not specific for CYP19 and showed side effects with depression of adrenal function. Glucocorticoids needed to be used to compensate (Santen et al. 1981). As a result, the first-generation AIs (aminoglutethimide and testololactone) were not suitable for adjuvant treatment.
Figure 2
A schematic representation of the signal transduction pathways in ER-positive breast cancer cells and prostate cancer cells. (A) At the adrenal level, adrenal androgen de novo steroidogenesis occurs. Cholesterol is produced and converted to Pregnenolone with the aid of CYP11A1 enzyme. Pregnenolone is converted to dehydroepiandrosterone (DHEA) with the aid of CYP17A1. Finally, DHEA is converted to androstenedione (AD) with the aid of 3β-hydroxysteroid dehydrogenase enzyme. Then, AD is converted to testosterone via 17β-hydroxysteroid dehydrogenase. At the adipose tissue level, both androstenedione and testosterone are converted with the aid of the aromatase enzyme system to estrone (predominant in postmenopausal women), and estradiol (predominant in premenopausal women), sequentially. Estrogen normally binds to the ER in the cytoplasm, the estrogen:ER complex translocates to the nucleus, gets phosphorylated, and binds to estrogen responsive elements (EREs) with the recruitment of coactivators. This creates a transcription complex (TC). This in turn, will initiate a cascade of protein synthesis and subsequent tumor proliferation through the activation of estrogen-sensitive genes. Whereas, SERMs:ER follow a similar pattern but recruits corepressors and inhibits protein synthesis; causing tumor regression. For SERDs, they bind to the ER causing an alien conformation. This leads to the destruction of the ER through the ubiquitin proteasome system; subsequently tumor regression. (B) At the adrenal level, adrenal androgen de novo steroidogenesis occurs. Cholesterol is produced and converted to Pregnenolone with the aid of CYP11A1 enzyme. Pregnenolone is converted to dehydroepiandrosterone with the aid of CYP17A1. Finally, DHEA is converted to DHEA-S with the aid of steryl-sulfatase (STS) and bile salt sulfotransferase. At the prostate level, DHEA-S in Leydig cells is converted back to DHEA via STS and then DHEA is converted to AD via enzyme 3β-HSD. Then, AD is converted to testosterone via enzyme AKR1C3, and finally to DHT via steroid 5α-reductase. Dihydrotestosterone normally binds to the AR in the cytoplasm, the DHT:ER complex translocates to the nucleus, gets phosphorylated, binds to androgen responsive elements (AREs) with the recruitment of coactivators. This creates a transcription complex (TC). This in turn, will initiate a cascade of protein synthesis and subsequent tumor proliferation through the activation of androgen-sensitive genes. Whereas, AR inhibitors:AR complex follows a similar pattern but recruits corepressors and inhibits protein synthesis; causing tumor regression. For SARDs, they bind to the AR causing the degradation of the receptor; subsequently tumor regression. Androgen receptor inhibitors vary in their mechanisms of action. For example, enzalutamide competitively inhibits the AR binding to DHT, inhibits nuclear translocation of AR, and DNA and cofactor binding. Whereas, bicalutamide is a highly selective, competitive and silent antagonist to the AR, which was also found to accelerate AR degradation. Abiraterone inhibits CYP17A1 and subsequently adrenal androgen de novo steroidogenesis. Dutasteride is a 5α-reductase inhibitor that blocks testosterone conversion into DHT.
Steroid-induced apoptosis in endocrine cancers

By P Y Maximov et al.

Published by Bioscientifica Ltd.
Printed in Great Britain

Current treatment strategies for prostate and breast cancers

Various parameters, such as the tumor volume and the pathological grade, have been correlated with prostate cancer malignancy (Bostwick et al. 2000). A strong correlation with an excellent prognosis was evident in prostate cancer presenting with a high percentage of AR-positive cells (Barboro et al. 2014). Prostate cancer that is AR negative is very rare; therefore, little attention has been given to this subtype. The aggressiveness of prostate cancer is based on the Gleason score, a system based on pathological grade. In prostate cancer, the lowest Gleason score sum found in a tumor biopsy is 6, which are low grade or well differentiated, less aggressive with slow growth and limited invasion and metastasis. The Gleason score sum of 8–10 are found in high-grade tumors, poorly differentiated, tending to be aggressive and quickly grow and spread. Gleason score sum of 7 is called intermediate grade and is found in moderately differentiated tumors.

Since prostate cancer is an indolent disease, the majority of men diagnosed will not be treated with any type of therapy. It was found that the majority of men with prostate cancer have lower prostate cancer-specific mortality rates and are more likely to die from age-related comorbidities (Lu-Yao et al. 2009, Albertsen et al. 2011). However, if prostate cancer is graded as aggressive, then surgery and sometimes adjuvant radiotherapy are the therapies of choice. Radical prostatectomy alone in men with localized prostate cancer has a 7-year recurrence-free survival (RFS) of approximately 70% (Kattan et al. 1999) and the biochemical PFS of approximately 50% (Bolla et al. 2005). However, application of immediate adjuvant radiotherapy can further significantly increase clinical PFS (Bolla et al. 2005). If the disease has progressed in spite of primary therapies, has metastasized or is an advanced poor prognosis or/and high-grade tumor only then is hormonal therapy applied. Recurrent tumors that are nonmetastatic or for locally advanced tumors (tumors that have spread to nearby tissue or local lymph nodes) are sometimes treated with adjuvant hormonal therapy concomitantly with adjuvant radiotherapy. This can further increase PFS and OS, especially if applied at earlier time points (Fleschner et al. 2008, Payne & Mason 2011, Omrcen et al. 2015, Shipley et al. 2017). Current treatments strategies for prostate cancer are summarized in Fig. 2.

By contrast, breast cancer is a highly heterogeneous tumor with different malignant subtypes. Prat and Perou (2011) used gene expression profiling to classify breast cancer into subtypes based on the expression of the main receptors, ER, progesterone receptor (PR) the erythroblastosis oncogene (ErBB2, HER2/neu) and the AR: Luminal A (ER+, PR+, HER2– and low Ki-67, low grade), Luminal B (ER+, PR+, HER2+/−, high Ki-67 and high grade), human epidermal growth factor receptor 2 (ER–, PR– and HER2+), basal-like or triple-negative (TNBC) (ER–, PR– and HER2–), claudin-low (often TNBC with low expression of cell-to-cell contact proteins and E-cadherin, in particular, with infiltration of lymphocytes), Luminal ER–/AR+ (AR+ and respond to antihormonal therapy with antiandrogens (Gucalp & Traina 2016)) and normal-like (ER+, PR+, HER–, low Ki-67 and normal like) breast cancers. Patients with ER+ early-stage breast cancer account for about 75% of breast cancer cases (Harvey et al. 1999). Though the primary therapies for early-stage breast cancers, regardless of their subtype are surgery and radiotherapy, long-term hormonal adjuvant therapy is used in most cases with ER+ breast cancers. The first FDA-approved antiestrogen tamoxifen is usually prescribed to premenopausal patients as they have a very low risk of developing endometrial cancer as a side effect from long-term tamoxifen treatment. The Early Breast Cancer Trialists’ Collaborative Group in 2011 confirmed that five years of using tamoxifen as adjuvant treatment reduced the risk of death and reduced the 15-year recurrence risk by 40% (Davies et al. 2011). The benefits of the 10-year extended therapy with tamoxifen were presented in two studies in 2013 with the 15-year follow-up of the Adjuvant Tamoxifen: Longer Against Shorter (ATLAS) trial (Davies et al. 2013) and the Adjuvant Tamoxifen To Offer More (aTTom) (Gray et al. 2013) trial. The outcome of the ATLAS trial is that, ER-positive patients with extended tamoxifen therapy reduced the risk of breast cancer recurrence,
mortality and reduced overall mortality (Davies et al. 2013). The outcome of the aTTom trial was similar to the ATLAS trial confirming that, continuing tamoxifen treatment in ER-positive breast cancer patients for 10 years rather than just 5 years leads to further reductions in recurrence and subsequent decrease in mortality. The documentation for the clinical characteristics of high-risk patients eligible for extended tamoxifen therapy (>5 years) have recently been published (Pan et al. 2016).

For postmenopausal women with thromboembolic or no osteoporotic comorbidities, AIs are usually prescribed. Anastrozole was the first of the third-generation AIs used in a clinical trial called the Tamoxifen Alone or Combination (ATAC) trial (Baum et al. 2002). Anastrozole has some advantages over tamoxifen as a first-line adjuvant treatment for early breast cancer in postmenopausal patients. Results for the combination treatment are the same as tamoxifen alone (Baum et al. 2002, Cuzick et al. 2010). This was to be expected. A rule of pharmacology is that a partial agonist (tamoxifen) that binds to a receptor when combined with a therapy that removes the ligand from the body produces a response of the partial agonist alone. In 2010, a meta-analysis (Dowsett et al. 2010) was performed and demonstrated the superiority of 2–3 years of tamoxifen followed by an AI for 2–3 years over 5 years of tamoxifen alone. Other clinical trials, referred to as the Breast International Group (BIG 1–98) (Breast International Group 2005) and the adjuvant tamoxifen and exemestane in early breast cancer (TEAM-trial) (van de Velde et al. 2011), were designed to address the question whether AIs would be superior to tamoxifen or not after 2–3 years of tamoxifen followed by switching to an AI for five years, showed no significant decrease in disease-free survival (DFS) or the RFS. A meta-analysis of individual data from postmenopausal patients with early-stage ER-positive breast cancers comparing 5 years of AIs against 5 years of tamoxifen or switching to an AI up to year 5 after 2–3 years of tamoxifen compared to 5 years of tamoxifen or an AI alone has shown that AIs have a significantly more favorable recurrence rates (RR) than tamoxifen by 30% (Early Breast Cancer Trialists’ Collaborative Group 2015). AIs also caused more bone fractures, but fewer cases of endometrial cancers than tamoxifen (Early Breast Cancer Trialists' Collaborative Group 2015).

Currently, the period of adjuvant tamoxifen/AI treatment is extended up to 10 years based on the National Cancer Institute of Canada Clinical Trials Group (NCIC
CTG MA17), which showed the superiority of 5 years of tamoxifen followed by five years of letrozole compared to 5 years of tamoxifen alone (Goss et al. 2003). Recently, Goss and colleagues found in the MA17 extension using an additional 5 years of AI for a 10 years total significantly increases the rates of DFS and decreases the incidence of contralateral breast cancer but the rate of OS was not increased (Goss et al. 2016). It should be noted, however, that in the ATLAS trial (Davies et al. 2013) and the combined analysis of ATLAS and aTTom trials mortality did not decrease significantly during extended adjuvant therapy but only a decade after extended therapy. As tamoxifen is a competitive inhibitor of estrogen action at the ER (Jordan 1984), and is not cytotoxic, it is suggested that decreases in mortality occur by cell selection and subsequent estrogen-induced apoptosis from the woman’s own estrogen (Jordan 2014a, 2015a). Breast cancer remains the only cancer with an option of long-term adjuvant antihormone therapy proven to save lives. Current treatment strategies for breast cancer are summarized in Fig. 3.

The application of antihormone therapy is crucial in ER+ breast cancer as it is able to reduce the recurrence of breast cancer at least by half, and, unlike prostate cancer, which is an indolent disease, breast cancer will progress faster and recur without treatment. All prostate cancer patients and half of breast cancers develop acquired resistance to antiestrogen therapy.

Understanding acquired resistance to hormonal therapies in prostate cancer

ADT is the primary therapy for prostate cancers that are classified as an aggressive type (high Gleason score sum), advanced or locally advanced. The average duration of clinical responses to antiandrogen therapies in advanced prostate cancer is 12–18 months after which practically all patients evolve to castration-resistant prostate cancer (CRPC) tumor phenotype. CRPC is characterized by consistent elevation of prostate-specific antigen (PSA) despite ADT and/or metastases. It is estimated that 10–20% of all non-advanced prostate cancer patients will progress to CRPC after surgery or radiotherapy (Kirby et al. 2011).

Currently multiple examples exist for the molecular mechanisms of antihormone resistance in prostate cancer with an analogous classification for breast cancer. Each mechanism or their combinations may have clinical applications in individual cases. The mechanisms of acquired resistance to antihormone therapies for prostate cancer can be categorized based either on the dependence on the AR or dependence on the ligand (Table 1).

| Category                                                                 | Mechanisms                                                                 |
|--------------------------------------------------------------------------|-----------------------------------------------------------------------------|
| Ligand-dependent, receptor-dependent                                      | • hypersensitivity of the receptor to the ligand due to point mutations     |
|                                                                         | • increased receptor expression                                              |
|                                                                         | • increased transcriptional activity of the receptor due to changes in coregulators and corepressors levels |
| Ligand-independent, receptor-dependent                                   | • increased levels of endogenous or circulating ligand                      |
| Bypass pathway (ligand-independent, receptor-independent)               | • gain-of-function mutations in the receptor                                |
|                                                                         | • cross-talk mechanisms with other growth factor pathways                   |
| Hormone receptor negative, ligand-independent                           | • deactivation of tumor suppressor pathways                                  |
|                                                                         | • high expression of anti-apoptotic and low expression of pro-apoptotic molecules |
|                                                                         | • activation of cell proliferation survival pathways                         |
|                                                                         | • activation of growth factor receptor pathways                              |
|                                                                         | • employment of other types of hormone receptors                            |

They have been categorized by their dependence on the hormone receptors or hormones themselves.

Table 1 Mechanisms of resistance to antihormone treatments are similar in both prostate and breast cancer cancers.
The modeling results show that, when compared with the wtAR:DHT complex (Fig. 4A), the helix 12 of the mutantAR:bicalutamide complex closes over the LBD of the receptor, which provides agonist conformation of the AR and its subsequent activation (Fig. 4C). It should be noted that the precise mechanism of antiandrogen action at the LBD of the AR remains unclear (Bisson et al. 2008, Duke et al. 2011, Tan et al. 2015). Activating mutations at the ER can explain the phenomenon of antiandrogen withdrawal syndrome, when the termination of therapy with antiandrogens is followed by regression of tumors (Hara et al. 2003).

Besides various mutations of the AR that contribute to the endocrine resistance in CRPC, a new role of membrane-associated AR isoforms in CRPC is emerging. Membrane-bound ARs have been identified in LNCaP cells and in hormone-insensitive DU145 cells and are associated with rapid non-genomic hormone responses in cells (Papakonstanti et al. 2003, Papadopoulou et al. 2008a,b). However, very little is known about the significance of the membrane-bound ARs in CRPC, but recent report identifies an AR splice variant called the AR8 that is shown to be associated with castration resistance in prostate cancer (Yang et al. 2011). Overexpression of the AR8 isoform increases the association of the receptor with the EGFR in CRPC cells and promotes cell proliferation and survival (Yang et al. 2011).

The AR in CRPC cells can become hypersensitive to low doses of androgens. This hypersensitivity is associated with mutations in the AR itself, leading to an increased sensitivity of the receptor to low concentrations of circulating androgens (Gregory et al. 2001b). Additionally, overexpression of the AR can be another AR-dependent mechanism that creates hypersensitivity. Indeed, it was shown that 30% of CRPC tumors overexpress the AR at high levels in the cells and 80% of patients show an elevated gene copy number (Feldman & Feldman 2001, Waltering et al. 2012), which may be a result of selection of cell populations with high levels of the AR under androgen deprivation pressure (Rau et al. 2005).

### Figure 4
Molecular modeling of the wild-type and mutant ER and AR bound with agonists and antagonists. (A) wtAR:DHT LBD complex (PDB ID: 3L3X); (B) the best docking pose of the wtAR:bicalutamide complex (PDB ID: 3RLJ), obtained via flexible docking (the experimental structure used for docking was selected based on the 3D similarity between bicalutamide and the available ligands co-crystalized with AR WT, thus the experimental structure 3RLJ was selected due high similarity between the native ligand, S-22 and bicalutamide). The major interactions are shown in dashed lines and colored as follow: hydrophobic interactions in lavender, pi-pi interactions in purple, water-mediated H-bonds are shown in blue, and classical H-bonds are depicted in green.; (C) T741L AR mutant:bicalutamide LDB complex (PDB ID: 1Z95), helix 12 is colored in green and the major interactions are shown in dashed lines and colored as follow: hydrophobic interactions in lavender, pi-pi interactions in purple, water-mediated H-bonds are shown in blue, and classical H-bonds are depicted in green.; (D) wtER:E2 LBD complex (PDB ID: 1GWR); (E) wtER:endoxifen LBD complex; (F) Superposition of E2 D538G mutant with ERα D358G apo LBD structures (helix 12 is shown in red for apo conformation and pink in the E2 bound mutant structure). The major interactions are shown in dashed lines and colored as follow: hydrophobic interactions in lavender, pi-pi interactions in purple, water-mediated H-bonds are shown in blue, and classical H-bonds are depicted in green.
An increase in the AR expression is associated with the amplification of the AR gene and in some cases attributed to polysomy of the X chromosome (Ropke et al. 2004). However, androgen action is not dependent upon the AR alone but is modulated by coregulators. The levels of expression of these coactivators, particularly SRC1 and SRC2, are higher in poorly differentiated prostate tumors or in recurrent prostate cancers and provide cells with higher AR activity in a low-dose androgen environment (Fujimoto et al. 2001, Gregory et al. 2001a). Additional AR-specific coactivators of note have been identified: ARA70 increases the AR activity and even facilitates the binding of estradiol to the AR (Yeh et al. 1998), FKBP51 stabilizes the AR with HSP90 heat-shock protein complex and facilitates the binding of androgens (Ni et al. 2010), and TRIM24 increases AR transcriptional activity (Groner et al. 2016).

Long-term antiandrogen therapy can affect the hypothalamo–pituitary axis as a negative feedback loop in men, leading to a compensatory increase of circulating androgens (Rau et al. 2005). This, by itself, can activate the AR by the law of mass action, but testosterone is not the physiologic ligand as it is required to be converted to dihydroxytestosterone (DHT) in the prostate cancer cell. Increased 5α-reductase enzyme activity contributes to the increase of endogenous androgens in the tumor. This results in the selection of CRPC cells able to convert androgen to endogenous DHT to produce a growth advantage (Navarro et al. 2002, Titus et al. 2005, Chang et al. 2014). A polymorphism in the 5α-reductase gene is noted in men of African-American descent, which is responsible for higher enzymatic activity in prostate cancer cells, as well as in prostate cancer cases with bad prognosis (Ruijter et al. 1999). In fact, the intratumoral levels of androgens can be as high as 40% above the baseline levels before ADT (Nishiyama et al. 2004). Recently another polymorphism in HSD3B1, which encodes 3β-hydroxysteroid-dehydrogenase-1 has been identified in a retrospective study of CRPC as a factor that correlates with an increased DHT synthesis (Hearn et al. 2016).

### Ligand-independent and receptor-dependent mechanisms of resistance

Most mutations are point gain-of-function mutations and are mostly located in the LBD of the AR and allow other sex steroids, such as glucocorticoids to bind to the AR and activate it (Zhao et al. 2000). Resistance to abiraterone was demonstrated in some mCRPC tumors with mutated AR (Cai et al. 2011, Chen et al. 2015). Additionally, alternative AR mRNA splice variants occur that generate constitutively active AR proteins (Dehm et al. 2008, Watson et al. 2010, Bubley & Balk 2017).

Increased expression of certain growth factors are associated with increased activity of the AR in mCRPC as well. The subversion of the AR transcriptional activity via growth factor receptor-mediated growth is called the cross-talk. Epidermal growth factor (EGF), keratocyte growth factor (KGF) and insulin-like growth factor 1 (IGF-I) can activate the AR and can be reversed by antiandrogens (Culig et al. 1994). Tyrosine kinase receptors, such as HER2, which is highly expressed in CRPC cells, can also activate the AR via phosphorylation, through activation of the MAPK and the Akt pathways (Lin et al. 2001). The growth factor IL-6 is responsible for the progression of CRPC. This occurs by increasing AR activity by 50% more than observed with DHT alone (Culig et al. 2002). Resistance to growth inhibition occurs through the MAPK and STAT3 pathways, induces autophosphorylation of HER2 to activate the AR-mediated cascade independent of the ligand (Chen et al. 2000, Culig et al. 2002). High expression levels of the AR mRNA are maintained by NF-κB in CRPC cells, which sustains high AR protein levels (Zhang et al. 2009).

### Bypass pathway

One of the mechanisms of resistance involves recruitment of cellular survival pathways in CRPC, but the tumors still express the AR. However, the tumor cells do not require active AR to proliferate and survive. This mechanism is called the bypass pathway. Over expression of the antiapoptotic genes like Bcl-2, Bcl-xl and NF-κB (Gleave et al. 1999) are characteristic of the bypass pathway. Activation of other cell survival signaling pathways like PI3K/Akt has also been examined and linked with CRPC progression (Taylor et al. 2010). Mutations in tumor suppressor genes, like PTEN, also play a role in hormone resistance and allow the cells to rapidly progress though the cell cycles (Li et al. 1997, Mulholland et al. 2011, Edlind & Hsieh 2014). Mutated BRCA1 and BRCA2 tumor suppressor genes that are strongly associated with breast cancer incidence and progression have also been shown to be present in CRPC cells and associated with progression of prostate cancer cells to a CRPC phenotype (Rosen et al. 2001, Kote-Jarai et al. 2011, Leongamornlert et al. 2012, Robinson et al. 2015). Several other proteins have been identified that are associated with AR bypassing.
and progression and survival of therapy-resistant CRPC, such as TWIST1, DKK3 and VAV3 (Marques et al. 2010).

Recently, other possible contributing factors to the progression of CRPC disease were identified. Some estrogens are synthesized in males and the ERα is expressed in CRPC cells. The activation of the ERα stimulates proliferation and migration of the tumor cells (Attia & Ederveen 2012, Mishra et al. 2015). The estrogen-related receptor (ERR) induces bone metastases and activation of VEGF-A, WNTSA and TGFβ1 in mCRPC cells (Fradet et al. 2016). The glucocorticoid (GR) and the mineralocorticoid receptors (MR) have been associated with the progression of CRPC and being able to substitute and bypass the blocked AR (Arora et al. 2013). It appears that the GR and the AR have an array of common response genes due to homologous DNA-binding domains of the receptors (Sahu et al. 2013, Grindstad et al. 2015).

**Ligand-independent mechanisms of resistance with the loss of the AR**

Antiandrogen resistance in prostate cancer can also be ligand independent and AR negative. Loss of expression of the AR in CRPC has been recorded in 30% of cases (Suzuki et al. 2003) due to hypermethylation of the AR gene (Jarrard et al. 1998). This epigenetic deregulation of the AR expression is much more common for the CRPC compared to only 10% of de novo hormone-resistant prostate cancers (Suzuki et al. 2003). Selected cells, with loss of the AR expression after antihormonal therapy, have adapted and ‘hijacked’ pathways enabling them to grow using other growth stimulatory pathways and even employ other hormone receptor pathways.

**Understanding acquired resistance to hormonal therapies in breast cancer**

The mechanisms of antihormone resistance in breast cancer cells are very similar to the mechanisms in CRPC (Table 1) (Rau et al. 2005, Risbridger et al. 2010).

**Ligand-dependent and receptor-dependent mechanisms of resistance**

The hypersensitivity of breast cancer cells to low doses of estrogens during estrogen ablation therapy has been associated with increased levels of ER expression. For instance, the ER protein levels were shown to be higher in long-term estrogen-deprived (LTED) MCF-7 cells by as high as 10-fold (Katzenellenbogen et al. 1987, Welshons & Jordan 1987). This can also happen in estrogen depletion with tamoxifen treatment (Berstein et al. 2004). One of the possible pathways of such hypersensitivity to estrogens was explained by a non-genomic activity of the ER, when it phosphorylates Shc, which in turn binds to signaling proteins Grb-2 and Son of Sevenless (SoS). As a result, this activates MAPK/ERK via Ras and Raf and promotes the phosphorylation of the ER at the AF-1 motif and activation of the receptor (Santen et al. 2003). The increased transcriptional activity of the ER can also be upregulated by overexpressed coactivators. Estrogen receptor coactivator SRC3 is the most important for breast cancer as its expression is restricted to only a few tissues, including the breast (Suen et al. 1998). Clinical studies (Osborne et al. 2003, Alkner et al. 2016) noted that high levels of SRC3 coactivator were associated with worse outcomes in tamoxifen-treated breast cancer patients. Low corepressor expression has been described in tamoxifen-resistant tumors and has been reviewed elsewhere (Legare & Basik 2016). Asides from the levels of the ER protein and its activity modulating cofactors, high levels of circulating and intratumoral hormones can also provide antihormone resistance. As tamoxifen binds and blocks the ER in breast tumor cells, it can also bind to the ER in pituitary gland and hypothalamus and disrupt the negative feedback loop. Tamoxifen induced an elevation of the circulating levels of estrogens secreted from the ovaries by increasing gonadotropin-releasing hormone production. This mechanism has been used to explain the elevated levels of estrogens in tamoxifen-treated premenopausal patients (Ravdin et al. 1988, Jordan et al. 1991). The aromatase enzyme, that converts androgens to estrogens, can also be elevated in estrogen-deprived cells adaptively in vitro (Yue et al. 2003) and can be stimulated through stromal cells that express prostaglandin E2, IL-6, 11 and tumor necrosis factor α (TNFα) (Schrey & Patel 1995). Indeed, it was shown that breast tumor tissues have higher levels of aromatase expression than peritumoral tissues (Bulun et al. 1996).

In recent years, antihormone resistance was also linked to the expression of membrane-associated ERs. The first membrane-associated ER that was identified was GPR30. It was demonstrated that the translocation of GPR30 to the cell surface significantly increased after estrogen treatment in tamoxifen-resistant breast cancer cells and its activity was mediated through the EGFR (Ignatov et al. 2010, Mo et al. 2013) and it is able to attenuate the inhibition of MAPK as well (Mo et al. 2013). It was also shown that GPR30 is able to upregulate aromatase expression in tamoxifen-resistant breast cancer.
cells, which can be linked to the sensitivity to AIs in breast cancer patients with acquired or de novo resistance to tamoxifen (Catalano et al. 2014). Another novel variant of ER that was recently identified is the membrane-bound ER-α36 that is associated with tamoxifen resistance in vitro (Wang & Yin 2015, Gu et al. 2017). However, the clinical roles these findings are yet to be determined.

Ligand-independent and receptor-dependent mechanisms of resistance

Just like in the case with CRPC, one of the ligand-independent receptor-positive mechanisms of resistance in breast cancer could be the activating mutations of the ER. Mutations of the ESR1 gene, that encodes the ER, have been identified in the LBD of the receptor in 14–54% of clinical samples from metastatic breast cancer patients and have also been linked to antihormone resistance (Robinson et al. 2013, Jeselsohn et al. 2014). These mutations are found most often in the metastases rather than in the primary tumors. Most mutations occur in positions Y537 and D538 and are described as gain-of-function mutations, which lead to constitutive ligand-independent activation of the ER (Robinson et al. 2013, Toy et al. 2013). Amino acid residues 537 and 538 are positioned in the AF-2 motif of the ER LBD so that mutations of these residues can induce a ligand-independent agonistic conformation of H12, closing the unoccupied LBD by interacting with residue at position 351 (Jordan et al. 2015). The ligand-free ER then recruits coactivators and activates the ER, even with the binding of tamoxifen (ligand dependent) (Nettles et al. 2008, Jordan et al. 2015, Fanning et al. 2016). We have also performed molecular dynamics modeling to demonstrate the conformational perturbation of the ER LBD with D538 mutation (Fanning et al. 2016) (Fig. 4).

ER-positive resistance in breast cancer is also attributed to the activation of growth factor pathways, such as HER2, IGF-1R and FGFR and stress-related kinases, such as AKT, JNK, MAPKs, c-SRC and others, that regulate posttranslational modifications of the ER and its coactivators that increase the receptor activity (Schiff et al. 2004, Shou et al. 2004, Santen et al. 2009b, Theoret et al. 2011). There is clinical evidence that proves that differential expression of various growth factor receptors in tamoxifen-resistant tumors are associated with resistance to tamoxifen and can play a role of a predictive clinical marker for therapy efficacy (Busch et al. 2015, Tomiguchi et al. 2016). Interestingly, these mechanisms can increase the membrane-associated ER activity with 17β-oestradiol (E2) or even tamoxifen, also contributing to resistance in breast cancer. Increased levels of NF-κB and AP-1 can tether more ER to certain gene promoters and promote hormonal resistance (Zhou et al. 2007).

Bypass pathway

Ligand and ER-independent mechanism depends upon MYC, Cyclin E1 and D1, p21 and p27 can promote progression through cell cycle despite tamoxifen therapy (Span et al. 2003, Butt et al. 2005, Perez-Tenorio et al. 2006, Chu et al. 2008). Antiapoptotic molecules, such as Bcl-xL, can be overexpressed to inhibit pro-apoptotic molecules and promote survival (Riggins et al. 2005).

Evolution of acquired resistance in prostate and breast cancers

Antihormonal therapy is standard for the treatment of recurrent and metastatic prostate cancer, however, up to 90% of these patients will ultimately fail and develop CRPC disease within 12–33 months after ADT. To understand this process of acquired resistance, numerous studies in vitro and in vivo were performed to simulate long-term ADT in prostate cancer cells to decipher the evolving mechanisms of acquired resistance (Kokontis et al. 1994, Umekita et al. 1996, Joly-Pharaboz et al. 2000) (Fig. 5). The studies to mimic long-term antiandrogen therapy in prostate cancer were performed using LNCaP cells, a popular AR-positive human cell line. Initial studies used various durations of steroid deprivation, with culture media containing charcoal-treated serum (Kokontis et al. 1994). Continuous passaging of cells in androgen-deprived conditions led to the selection of clones hypersensitive to androgens. However, longer androgen starvation (2 years) of these clones let to the isolation of cells that grow independently from androgen, with an unanticipated vulnerability (Kokontis et al. 1994). Low doses of androgen reduced the number of viable cells after 6 days of treatment (Kokontis et al. 1994). The authors (Kokontis et al. 1994) also noted a high level of AR protein and mRNA in these resistant cells compared to wild-type LNCaP cells. Expression of PSA protein and mRNA increased when treated with an androgen. Experiments in vivo using the same cells showed that the wild-type LNCaP tumors grew well in mice with androgen treatment; however, the derived resistant cell line grew only in castrated mice and treatment with DHT caused regression of the tumors (Umekita et al. 1996). The authors (Umekita et al. 1996) used further androgen deprivation of resistant cells in their experiments in vitro to derive a cell
line that grew in androgen-deprived conditions as well as the wild-type cell line under testosterone stimulation. Interestingly, these resistant tumors were stimulated to grow with E2 and medroxyprogesterone acetate (MPA) and 5α-reductase inhibitor finasteride was able to partially reverse the tumoricidal actions of testosterone (Umekita et al. 1996).

Another group has performed similar in vitro and in vivo studies with long-term androgen-deprived LNCaP cells (Joly-Pharaboz et al. 2000). Wild-type LNCaP cells were passaged in culture medium supplemented with charcoal-treated serum for 1 year (Joly-Pharaboz et al. 2000). The resulting cell line grew independently from androgen, however, treatment with various androgens, and even E2 resulted in retarded cell growth due to apoptosis (Joly-Pharaboz et al. 2000). Experiments in vivo showed that androgen induced apoptosis and tumor regression with this model (Joly-Pharaboz et al. 2000).

Liang and coworkers (Chuu et al. 2011b) used variants of LNCaP prostate cancer cell lines to demonstrate that antiandrogen-resistant LNCaP cell lines with an AR-rich phenotype have a G1 cell cycle blockade in the presence of androgens by regulating cMyc, Skp2 and p27kip via the AR. Additionally, they found that higher dosages of testosterone lead to more growth inhibition of relapsed tumors suggesting that the manipulation of androgen/AR signaling pathway may be a potential therapeutic target in AR-positive metastatic prostate cancer. Kawata and coworkers (Kawata et al. 2010) reported that prolonged treatment of a bicalutamide-resistant subline (LNCaP-BC2) with bicalutamide induces AR overexpression and androgen hypersensitivity to low levels of androgen. The authors identified the phosphorylated AR (pAR210) overexpression and a possible mechanism for androgen hypersensitivity. However, after long-term androgen deprivation, LNCaP prostate cancer cells evolve to be a cell population vulnerable to androgen-induced apoptosis (Chuu et al. 2011a). Nevertheless, continuous treatment with androgens eventually selects for cells that will be resistant to the apoptotic actions of androgens and grow. The authors speculated that it would be possible to use intermittent androgen deprivation (IAD) to slow the progression of resistance and use androgen therapy during the relapse after the ADT cycle to further control the tumor progression (Chuu et al. 2011a).

Clinically, there is evidence to support androgen-induced apoptosis in CRPC. Bruchovsky and coworkers (Akakura et al. 1993, Bruchovsky et al. 2000) used IAD to demonstrate that androgen action would inhibit growth of antiandrogen-resistant prostate cancer. There is evidence that IAD is able to prolong progression of resistant disease, and testosterone restoration between
ablation therapy cycles can induce tumor regression in the laboratory in vivo (Sato et al. 1996) and in the clinic (Pether et al. 2003, Mathew 2008). In a recent viewpoint by Klotz and Higano (Klotz & Higano 2016), IAD was described as a viable alternative to the continuous androgen deprivation (CAD) in men with no underlying cardiovascular diseases. The IAD strategy was preferable with improved quality of life, cheaper health care costs, despite no observed advantage over CAD in terms of OS. Recently, Schweizer and coworkers (Schweizer et al. 2015) found a 50% response rate to androgen therapy when monitoring either PSA levels or radiologically identified CRPC disease.

Similar advances were made in studies of the antiestrogen resistance in breast cancer, and the evolution of breast cancer cells in the estrogen-free environment (Fig. 5). The evolution of MCF-7 breast cancer cells in estrogen deprivation conditions is similar to the evolution of LNCaP cells in response to androgen deprivation (Jordan et al. 2016).

Tamoxifen is a competitive inhibitor of estrogen action (Jordan 1984) and long-term adjuvant tamoxifen therapy was predicted to be essential to suppress breast tumor cell growth (Jordan 2014b). Early studies using MCF-7 breast cancer cell line transplanted into oophorectomized athymic mice demonstrated that although tumors eventually developed despite tamoxifen therapy (Osborne et al. 1987), the tumors, in fact, grew because of tamoxifen therapy (Gottardis & Jordan 1988, Gottardis et al. 1989a,b). Tamoxifen-stimulated tumors were growth stimulated by either tamoxifen or physiologic estradiol. As a result, no estrogen treatment or treatment with a pure antiestrogen (Gottardis et al. 1989a,b) prevented tumor growth. Discovery of this biology of early acquired resistance to tamoxifen preceded the clinical finding that either an AI or the pure antiestrogen fulvestrant were appropriate second-line therapies after tamoxifen failure in MBC (Howell et al. 2002, Osborne et al. 2002). This unique form of acquired resistance has clinical relevance in SERM pharmacology with a withdrawal response in MBC to SERMs tamoxifen and raloxifene (Howell et al. 1992,Dosik & Kaufman 2004, Lemmo 2016). The recent development (Fan et al. 2014a,b,c) of an in vitro model of acquired resistance to SERMs has provided important insight into how either tamoxifen (SERMs) or estrogen can stimulate tumor cell growth. Estrogen-stimulated growth in early acquired resistance to tamoxifen in vivo is via a genomic pathway, but with estrogen action at genomic sites blocked by tamoxifen. By contrast, tamoxifen stimulates tumor cell growth non-genomically by enhancing the IGFR1β pathway.

It is important to reemphasize that high-dose synthetic estrogen therapy was the first chemical therapy to treat any cancer (Haddow et al. 1944). However, Haddow (1970) noted that high-dose synthetic estrogen therapy was only effective at producing a 30% response rate in MBC 5 years following menopause. If estrogen was administered therapeutically nearer to the menopause then MCB grew. The reasons for this clinical observation were unknown and mechanisms were not deciphered during the 1950s–1970s, when high-dose estrogen was the standard of care for postmenopausal MBC. In the 1970s, tamoxifen, a non-steroidal antiestrogen (Jordan 2003), became the standard of care for all stages of breast cancer until the introduction of AIs in the late 1990s. There was no interest in understanding how high-dose estrogen therapy killed breast cancer cells despite the fact that high-dose DES produced a survival advantage over tamoxifen in a small trial in MBC (Ingle et al. 1981, Peethambaram et al. 1999).

It is therefore ironic that the study of acquired resistance to tamoxifen treatment in breast cancer should expose a vulnerability of antihormone-resistant breast cancer i.e.: estrogen-induced apoptosis (Wolf & Jordan 1993, Yao et al. 2000). Most importantly, the MCF-7 breast tumors developed acquired resistance to tamoxifen by cell selection over a 5-year period. Within two years, acquired resistance is evidenced by tamoxifen-stimulated growth and estrogen-stimulated growth; the growth stimuli are interchangeable. However, between 3 and 5 years of tamoxifen exposure, tamoxifen stimulates tumor growth but physiologic estrogen causes complete regression of small (<0.3 cm) tumors. The MCF-7 tumors rapidly regressed in response to E2. It was proposed (Yao et al. 2000) that estrogen treatment of recurrent breast cancer following the failure of long-term tamoxifen treatment, will result in a tumor regression and breast cancer cells will regain their responsiveness to estrogen for growth. Tamoxifen causes a decrease in mortality and prevents disease recurrence after 5 years of stopping the therapy, i.e., does not cause a rebound effect anticipated for a competitive inhibitor of estrogen action. The reason suggested is that a woman’s own estrogen causes estrogen-induced apoptosis in populations of vulnerable micrometastases that has long-term acquired resistance (Yao et al. 2000). This hypothesis in now supported by considerable clinical evidence reviewed elsewhere (Jordan 2014a, 2015a).
Song and coworkers (Song et al. 2001) reported that long-term estrogen deprivation leads to estrogen-induced apoptosis in LTED breast cancer cell population in vitro. Estrogen deprivation for a short time causes an elevation in the ER protein levels (Katzenellenbogen et al. 1987, Welshons & Jordan 1987). After 8–11 months of estrogen deprivation, MCF-7 cells acquire adaptive hypersensitivity to estrogen (Masamura et al. 1995), which is similar to LNCaP cells and hypersensitivity to androgen (Feldman & Feldman 2001). This may explain the early mechanism of AI resistance in breast cancer. Various cell models were developed over the years to study long-term estrogen deprivation in estrogen-free environment and using dilution cloning selection (Jiang et al. 1992, Pink et al. 1995, Song et al. 2001, Lewis et al. 2005b). Two breast cancer cell lines were selected after long-term estrogen deprivation (2 years). MCF-7:5C and MCF-7:2A cell lines were at first characterized as ER positive and non-responsive to estrogens or antiestrogens (Jiang et al. 1992, Pink et al. 1995); however, optimization of culture conditions dramatically altered these characteristics (Lewis et al. 2005b). The MCF-7:5C cells were shown to undergo low-concentration estrogen-induced apoptosis within a week of treatment in a concentration-dependent manner (Lewis et al. 2005b), and the intrinsic mechanism of estrogen-induced apoptosis was described (Lewis et al. 2005a, Fan et al. 2012, 2015). The MCF-7:2A cells undergo slow apoptotic alterations that occur within two weeks of treatment with estrogen. Both of these cell lines were used to investigate genome-wide alterations in estrogen-regulated gene expression profile involved in apoptosis (Ariazi et al. 2011).

**Current therapies for hormone-resistant prostate and breast cancers**

Resistance to antihormonal therapy occurs in prostate and breast cancers, as new cell populations are selected after long-term sex steroid deprivation. These cells are characterized by sex hormone-independent growth.

It is believed that the AR in CRPC is still functional and can be abrogated to stop disease progression. Cytotoxic chemotherapy was routinely utilized to treat aggressive disease in the absence of targeted alternatives for CRPC prostate cancer. De Bono and coworkers (De Bono et al. 2010) compared cabazitaxel with the topoisomerase type II inhibitor mitoxantrone in mCRPC patients previously treated with docetaxel. Mortality was significantly decreased in the cabazitaxel group (De Bono et al. 2010). Smith and coworkers (Smith et al. 2013) evaluated cabozantinib (XL184), which is an orally bioavailable tyrosine kinase inhibitor that acts against MET and vascular endothelial growth factor receptor 2 (VEGFR2), in CRPC patients. They concluded that cabozantinib has clinical efficacy in CRPC improving PFS with a decrease of soft tissue lesions, resolution of bone scans, decline of bone turnover markers, pain and use of narcotic painkillers. However, the major strategic advance for the treatment of CRPC is the realization that the AR is still functional in CRPC and, like in breast cancer, remains a potential target.

New antihormonal agents are improving the prognosis of CRPC. Abiraterone acetate (Barrie et al. 1994) is an inhibitor of cytochrome P450 (CYP17) (Fig. 1), which plays an essential role in de novo intratumoral androgen production from cholesterol in CRPC tumors (Locke et al. 2008). This therapeutic approach to treat prostate cancer is analogous to the use of adjuvant therapy with AIs in postmenopausal breast cancer patients (Fig. 3). De Bono and coworkers (de Bono et al. 2011) evaluated abiraterone acetate in patients with mCRPC who have received chemotherapy and demonstrated that the inhibition of androgen biosynthesis by abiraterone prolonged the OS. Other approaches target the AR with new antiandrogens.

Scher and coworkers (Scher et al. 2010) evaluated the antitumor activity and safety of enzalutamide, which blocks AR activity in men with CRPC (Fig. 2). Increasing doses of enzalutamide reduced serum PSA and stabilized bone disease in 56% of patients (Scher et al. 2010).

Recently, Penson and coworkers (Penson et al. 2016) compared the efficacy of enzalutamide and bicalutamide in CRPC. Enzalutamide decreased the mortality of patients by 76% with a median PFS of 19.4 months compared to bicalutamide with a median PFS of 5.7 months. There was a significant increase in PFS with enzalutamide in the proportion of patients with a ≥50% PSA response, time to PSA progression and radiographic PFS in metastatic patients. Advantages of enzalutamide were observed in both metastatic and nonmetastatic subgroups. However, evidence is emerging on acquired resistance to abiraterone and enzalutamide (Attard & Antonarakis 2016, Bubley & Balk 2017, Gupta et al. 2017).

Several new antiandrogens are in early clinical development. The antiandrogen ARN-509 developed by Janssen Research & Development is an example of a potent competitive pure antiandrogen that has been evaluated in phase I/II trials in CRPC patients. In phase I trial, ARN-509 was well tolerated with fatigue being the most reported side effect (Rathkopf et al. 2013). In the phase II study, ARN-509 demonstrated an 80–90% efficacy in patients...
with naïve CRPC in both metastatic and nonmetastatic settings. There was a 29% response rate in mCRPC patients previously treated with abiraterone, reducing the PSA levels by more than 50% (Rathkopf et al. 2012). The novel small peptide EPI-001 targets the N-terminal domain of the AR containing the activating function-1 region (AF-1). This interrupts the AR’s interaction with other proteins and androgen response elements in the androgen-responsive genes promoters. As a result, transcriptional activity is disrupted (Andersen et al. 2010). This peptide has not entered clinical trial, but showed promising results in the CRPC xenograft models (Andersen et al. 2010). A novel selective AR downregulating drug (SARD) AZD3514 had limited tolerability in CRPC patients in a phase I trial with modest antitumor activity; however, it did show activity in 17–25% of patients reducing PSA by more than 50% (Omlin et al. 2015). The authors concluded that developing SARDs in the future for treatment of CRPC may hold merit (Omlin et al. 2015).

Despite the use of long-term antihormone adjuvant therapy for breast cancer, approximately 50% of patients have disease recurrence. The question we must ultimately address is how we improve response rates? Though tamoxifen was approved initially for treatment of MBC in both pre- and postmenopausal women, AIs became the first-line therapy for postmenopausal breast cancer patients who did not have any prior hormonal therapy or have recurred within 12 months after previous adjuvant AI therapy. However, if the tumors recur in less than 12 months after hormonal therapy with an AI, then tamoxifen is recommended or a pure antiestrogen fulvestrant as second-line therapies. Recently, Robertson and an international team of colleagues (Robertson et al. 2016) in a phase III clinical trial have demonstrated superiority of fulvestrant over anastrozole as first-line therapy in postmenopausal patients with metastatic of locally advanced breast cancer. For premenopausal women, tamoxifen can be prescribed as first-line adjuvant hormonal therapy and AIs or fulvestrant can be used as second- and third-line therapies in case of cancer recurrence, but only with ovarian function suppression (Abderrahman & Jordan 2016). Antihormone resistance eventually occurs after exhaustive antihormone therapy fails. However, depending on the size and location of the metastasis cytotoxic chemotherapy is more likely to be used after a failed AI therapy rather than second or third-line antihormone agents.

New strategies for the treatment of hormone-refractory breast cancer are evolving based on inhibition of aberrant pathways. Abnormalities in the CDK4/6 and the mTOR pathways play a crucial role in the pathogenesis of breast cancer. These pathways are therapeutic targets for the treatment of naïve MBC or antihormone-resistant breast cancer. In phase I/II clinical studies (Schwartz et al. 2011), palbociclib, which is a specific CDK4/6 inhibitor (O’Leary et al. 2016), demonstrated an excellent bioavailability, mild to moderate adverse effects, and a well-tolerated toxicity. In phase III clinical study called PALbociclib Ongoing trials in the Management of breast cAncer-3 (PALOMA-3) (Turner et al. 2015) the combination of palbociclib with endocrine therapy significantly improves PFS. All these data resulted in palbociclib receiving an FDA approval in 2015 as a first-line treatment for advanced postmenopausal ER-positive/HER2-negative breast cancer in combination with letrozole.

Numeral studies with mTOR inhibitors (i.e. everolimus, temsirolimus, deforolimus) show promise in the ER-positive and/or HER2-positive breast cancer (Fasolo & Sessa 2008, Vicier et al. 2014, Baselga et al. 2017). The combination of everolimus with either an AI (Beck et al. 2014, Finn et al. 2015) or fulvestrant (Beaver & Park 2012, Sun et al. 2016, Pritchard et al. 2017) demonstrated clinical efficacy. The Breast Cancer Trials of OraL EverOlimus-2 (BOLERO-2) (Baselga et al. 2012), combined everolimus and exemestane for women with advanced ER-positive/HER2-negative breast cancer who previously failed AI therapy. In BOLERO-2, everolimus improved PFS in trastuzumab-resistant patients. Interestingly, in an early study with an mTOR inhibitor (deGraffenried et al. 2004) rapamycin ester (CCI-779) treatment restored tamoxifen response in tamoxifen-resistant breast carcinoma (Yu et al. 2001).

Regrettably, combination therapies with CDK4/6 inhibitors or mTOR inhibitors with an antihormonal therapy do not result in lives saved, although life extension is a positive benefit. The question now becomes: how can adjuvant endocrine therapy be advanced based on what we now know from current clinical trials? There is a linear progression from therapeutic success in MBC to trials of adjuvant therapy, but we suggest this may not be that simple with CDK4/6 inhibitors and mTOR inhibitors.

The high monthly cost for both CDK4/6 inhibitors and mTOR inhibitors (Carey & Perou 2015), and the toxicity profile of grade 3/4 side effects with palbociclib (Finn et al. 2015) (i.e. neutropenia, leukopenia, and lymphopenia), and grade 1/2 side effects with everolimus (Baselga et al. 2012) (i.e. fatigue, stomatitis, anorexia, diarrhea, noninfectious pneumonitis, metabolic disorders with hyperglycemia and hematologic disorders) hinder their utilization as a useful long-term adjuvant treatment.
These systemic side effects and financial costs will reduce patient compliance and the value of antihormone therapy will be lost. It is difficult to maintain compliance for current antihormonal agents for 5 years, so an increase in side effects will result in the failure to control disease recurrence. We suggest another path in the final sections of this review.

Consideration for implementing a pathway forward that saves lives following a diagnosis of prostate and breast cancer

Enormous progress has occurred in the last 40 years in the approach to treating prostate and breast cancer. In the period 1967–1977 there were no proactive detection programs, diagnosis was usually late stage disease and the word cancer was not used. Quite rightly, cancer had the reputation as a death sentence. Radical surgery and radiotherapy were the major weapons in the physicians armamentarium and chemical therapy (chemotherapy) was primitive. Medical oncology was an emerging specialty. High-dose estrogen therapy was effective in 30% of both metastatic breast and prostate cancers, but this was a paradox as both breast and prostate cancers were known to be sex steroid dependent. Mechanisms were unknown.

A significant step forward occurred in breast cancer treatment with the publication of a symposium at King’s College, Cambridge (28–29th September, 1977) in the October Supplement of Reviews in Endocrine-Related Cancer (Jordan 1978), the fore-runner of the current Society for Endocrinology journal Endocrine-Related Cancer. The conclusions, which hold true today, were: (1) treating animals with a large tumor burden cannot affect a cure; (2) the tumor ER is important to predict a response to tamoxifen; (3) treating with tamoxifen early in tumorigenesis; i.e. low tumor burden, produces some protection for animals; (4) longer treatment with tamoxifen is superior to short treatment in animals with microscopic disease.

This and subsequent publications (Jordan 1978, Jordan et al. 1979, 1980, Jordan & Allen 1980) triggered the move to long-term adjuvant antiestrogen therapy proven to save lives (Goss et al. 2005, Davies et al. 2013). As illustrated in this current review of prostate and breast cancer treatments, the diseases run different courses. Adjuvant therapy in prostate cancer is not implemented in the same way as is routine for breast cancer. In breast cancer, antihormone therapy is used to benefit patients in all stages of breast cancer, but the same is not true for prostate cancer. ADT is only used in MPC, locally advanced or recurring cases. Nevertheless, our review illustrates that the evolution of acquired resistance for both breast and prostate cancer is similar. Mechanisms of acquired resistance are broadly the same or the adaptions of alternate growth stimulating pathways are similar. The major risk factor for both prostate and breast cancer, is age. A primary consideration is to seek effective therapeutic solutions for our aging population. Resources are scarce and our goal of achieving chemoprevention of breast and prostate cancers has fallen short. We still do not know precisely who will develop breast or prostate cancer, and why. Treating large population to benefit a few, who do not know their disease was prevented, was an ineffective approach. Side effects from any chemopreventive intervention are unacceptable to any but the most committed high-risk woman who wishes to prevent breast cancer. A strategy to prevent prostate cancer using an inhibitor of 5α-reductase was scientifically sound (Homma et al. 1997, Andriole et al. 2004, Thorpe et al. 2007) but outcomes were controversial due to potential risks of high-grade prostate cancers and this advance in health care was abandoned (FDA 2011, Theoret et al. 2011). The chemoprevention solution has overwhelmed healthcare systems. There is neither physician time to address individual needs for chemoprevention (Smith et al. 2017) nor, it seems, physician knowledge about options (Smith et al. 2017). We must, therefore, do what can be done to aid patients with breast and prostate cancers. This strategy must be inexpensive, globally applicable and aim to keep as many individuals well who can continue to contribute effectively to the welfare of the family. This essential goal will impact on the welfare of countries as each family unit can contribute to the economy of that country. In the final section we will address what can be done, how and why the approach is feasible.

An approach to global health care maintenance in prostate and breast cancers

Tamoxifen has taught us the fundamental laws of clinical therapeutics. To this day, antihormone therapy of MBC plus/minus chemotherapy or precision medicines (to block cell replication or the survival pathways that subvert antihormone action away from the ER growth pathway) can delay but not prevent death (Abderrahman & Jordan 2016). The same medicine tamoxifen or now an AI (letrozole) applied as a long-term adjuvant therapy, can delay recurrence and decrease mortality. Laboratory studies of acquired resistance to antihormone therapies
opened the door to understanding the ‘carry over’ effect of long-term adjuvant antiestrogen therapy that decreases mortality after adjuvant therapy is stopped (Fisher et al. 2005, Cuzick et al. 2007, Powles et al. 2007).

The knowledge of mechanisms in adjuvant therapy in breast cancer can now be built upon to enhance survivorship and improve the quality of life during long-term adjuvant therapy. By contrast, the urologic community must decide whether select patients, destined to remain hormone responsive, could or should be treated with adjuvant ADT. An approach would be to correlate the genomics of indolent primary tumors with outcomes at recurrence that is antihormone responsive MPC. In this way, analysis of large data sets could save lives. The identification of those tumors that recur with MPC but subsequently respond to ADT would be candidates for adjuvant approaches in the future. Indeed, long drug holidays or androgen therapy may benefit patients with androgen-induced apoptosis of microscopic disease. Until that time, the strategy of long-term adjuvant control of prostate cancer cannot be considered.

For breast cancer, by contrast, the landscape holds numerous affordable possibilities. The AIs have reduced RR, with fewer serious side effects, but results of survivorship are less clear than with the SERM tamoxifen. However, the creation of the ‘estrogen-free’ woman for the remainder of her life, during adjuvant AI therapy, has concerns for general health. Osteoporosis is a concern, as is the less well-defined issues of coronary heart disease (CHD) and reduced mental capacity. This may include exacerbation of Alzheimer’s disease for our aging population. Clearly, large populations of patients with Alzheimer’s should be examined to determine whether breast cancer adjuvant treatment with either tamoxifen or AIs advance Alzheimer’s onset or exacerbates symptoms and severity.

The ‘SERMs Solution’ (Lerner & Jordan 1990) for the chemoprevention of breast cancer now has a role to improve long-term adjuvant therapy. The original proposal for SERM was:

‘Important clues have been garnered about the effects of tamoxifen on bone and lipids so it is possible that derivatives could find targeted applications to retard osteoporosis or atherosclerosis. The ubiquitous application of novel compounds to prevent diseases associated with the progressive changes after menopause may, as a side effect, significantly retard the development of breast cancer.’ (Lerner & Jordan 1990)

Following the success of the pioneering SERM tamoxifen, the medicinal chemistry community has advanced numerous safe and widely used new SERMs including raloxifene, bazedoxifene and ospemifene (Maximov et al. 2013). All are FDA approved for different indications in postmenopausal women’s health. Only raloxifene has a cancer indication; the chemoprevention of breast cancer in high-risk postmenopausal women. Lasofoxifene is not yet approved but promises not only to reduce fracture risk in osteoporosis, reduce breast cancer incidence, and reduce strokes, but is the only SERM proven to reduce CHD (Cummings et al. 2010). Turning around the ‘SERM solution’ for women’s health one more time, there is a strategic opportunity for medicinal chemists to solve one of the important molecular mechanisms of acquired AI resistance, i.e.: expansion and mutations of the ER. However, this must be achieved not with an orally active pure antiestrogen (Abderrahman & Jordan 2016), but a SERM that destroys the ER.

Orally active ‘pure antiestrogens’ are a current focus of medicinal chemistry with the goal of being effective therapies in MBC after the failure of AI therapy (Abderrahman & Jordan 2016). But this is not good enough. The oral pure antiestrogen solution as a future adjuvant therapy would still keep women estrogen free.

Medicinal chemists already know how to make a SERM that maintains bone density in ovarierectomized rats, but destroys the tumor cell ER (Willson et al. 1994, Bentrem et al. 2001). The compound GW-5638 (Etacstil), was reported 20 years ago! The acrylic ‘antiestrogenic’ side chain when attached to the triphenyethylene core, fits appropriately into the ER ligand-binding domain but causes perturbation of the ER complex, resulting in rapid destruction (Wu et al. 2005). This acrylic side chain is a recurrent feature of the ‘new pure antiestrogens’ under investigation (Abderrahman & Jordan 2016).

A new SERM that destroys tumor ER, used as an adjuvant therapy, would not only enhance survivorship by reducing recurrence noted with AIs, but also improve woman’s health. Current problems of compliance can be addressed and improved. Women struggle with poor quality of life with AIs. Even a 3 month trial of local estrogen (or testosterone) is currently being evaluated to eliminate vaginal atrophy (Melisko et al. 2017), but a SERM could also achieve the same result (Jordan 2017b). Quality of life and being well is an essential component of patient survival. Stopping long-term adjuvant therapy prematurely, because of a lack of compliance, is the same as deciding upon a couple of years of adjuvant therapy. To stop adjuvant therapy early is not recommended. Indeed, the value of more than 5 years of adjuvant therapy has been evaluated. Ten years of adjuvant tamoxifen is superior to eliminating vaginal atrophy (Song et al. 2001) opened the door to understanding the ‘carry over’ effect of long-term adjuvant antiestrogen therapy that decreases mortality after adjuvant therapy is stopped (Fisher et al. 2005, Cuzick et al. 2007, Powles et al. 2007).
to 5 years of adjuvant tamoxifen in lives saved, but only in the five years after completion of 10 years of adjuvant tamoxifen (Davies et al. 2013). This is the essential role of estrogen-induced apoptosis, but the value in lives saved with an adjuvant AI is less clear (Goss et al. 2016).

The Study of Letrozole Extension (SOLE) addressed the issue of 3 month drug holidays annually (Fig. 6) but now is an opportunity to advance a new adjuvant therapy strategy. The goal of the study was to establish that a woman’s own estrogen would benefit patients by triggering estrogen-induced apoptosis. Invoking a physiologic antitumor mechanism would reduce micrometastatic disease and decrease recurrence. The hypothesis was based on published laboratory evidence (Wolf & Jordan 1993, Yao et al. 2000, Song et al. 2001). Though recommended at the time, administration of low-dose estrogen was considered too dangerous for patients without clinical evidence of efficacy and safety. The clinical studies have now occurred (Ellis et al. 2009, Anderson et al. 2012) so the laboratory concept is sound. The SOLE study is now reported (Colleoni et al. 2017) but shows no benefit for intentional 3 month annual drug holidays for 4 consecutive years of letrozole adjuvant therapy. Nevertheless, the SOLE trial provides significant important new information for two further advances in women’s health.

Our goal here is to propose a long-term therapeutic strategy that not only builds on past clinical experience, but also introduces a new strategic concept for adjuvant therapy to improve patient care globally. The new information from the SOLE trial is a first step forward. Firstly, the fact that a patient can stop therapy for 3 months and then restart adjuvant therapy allows compliance issues, due to side effects, to be addressed. A vigilant breast team can now offer returning to continuous adjuvant AI therapy to patients in distress. Secondly, the 3 month adjuvant window can now be used to create an advance in adjuvant therapy to reduce the micrometastatic tumor burden for those patients known to be at high risk for recurrence and death if a second five years of adjuvant therapy is not enforced (Abderrahman & Jordan 2017). This is important, as a short-term intensive preemptive
salvage therapy (Fig. 7), because cost and toxic side effects for current precision medicines (palbociclib and everolimus) will make years of combination therapy with an innovative antiestrogen therapy impractical (Carey & Perou 2015). But how should the clinical community advance the new therapeutic innovations?

Firstly, there has to be clearly defined patient population that is at high risk of recurrence despite long-term adjuvant antihormone therapy. Recent data reported by the Early Breast Cancer Trialists Collaborative Group (EBCTCG) (Pan et al. 2017) define that high-risk population that recurs following 5 years of adjuvant tamoxifen. Follow-up is for 15 years and, not surprisingly, RR depend upon the size of the original primary tumor and the number of axillary lymph nodes. Secondly, there needs to be a defined combination of physiologic estrogen plus a cocktail of precision medicines to reduce the burden of micrometastatic disease and avoid recurrence (Jordan et al. 2016). The goal is to kill micrometastatic disease not just hold cancer cell growth. Understandable concerns are raised still about the safety of low-dose estrogen (Reeder-Hayes & Muss 2017) and this is appropriate, but medicinal chemists are already addressing the problem. Raloxifene derivatives that trigger apoptosis in LTED breast cancer,

Figure 7
A schematic representation of the proposed design (alongside a proposed optimized version) for the preemptive salvage therapy. (A) Breast cancer patients who are ER-positive after surgery and at high risk of recurrence (this includes large primary tumors and positive lymph nodes at diagnosis), can harness the benefits of long-term estrogen deprivation, with a preemptive salvage therapy, aiming at clearing occult micrometastases. After 5 years of adjuvant antihormonal therapy with either tamoxifen or AIs, breast cancer cell populations undergo selection pressure. The new long-term estrogen-deprived (LTED) breast cancer cell populations are now vulnerable to a woman’s own estrogen through apoptosis (aka estrogen-independent). Whereas, they would normally grow with estrogen within 5 years past menopause (aka estrogen-dependent). The clinically observed response rate to low-dose estrogen therapy was 30% in metastatic breast cancer. Estrogen can act in synergy with other FDA approved breast cancer cell survival inhibitors or apoptosis promoters. This synergy can potentially increase the response rate above 30%. (B) Panel A can be optimized. Estrogen deprivation can be achieved with a new SERM that degrades the ER, preventing future drug resistance and receptor mutations. As one example, there is an orally active SERD, GW5638, which is metabolically hydroxylated to GW7604, in the same way, tamoxifen is metabolically activated to 4-hydroxytamoxifen. Unlike tamoxifen, GW7608 triggers the destruction of ER in BC cells, while retaining an estrogenic tickle at ER elsewhere (i.e. bones and serum lipids). Although GW7608 is a SERD for degrading the ER, it is also a SERM due to its agonistic and antagonistic mechanism of action at different tissue levels. A similar mechanistic SERM/SERD compound can improve estrogen deprivation (with an AI) by destroying the ER, while maintaining women’s health. In addition, estrogen in the proposed 3-month drug holiday can be replaced with selective human estrogen receptor partial agonist (ShERPA). These compounds mimic estrogen without causing significant uterine growth and were found to inhibit the growth of endocrine-independent tamoxifen-resistant breast cancer cell lines.
have been reported (Xiong et al. 2016) and clinical trials are planned.

Secondly, studies using FDA approved low-dose estrogen (FDA approved!) and cocktails of precision survival inhibitors in LTED breast cancer can be confirmed to be effective in LTED MBC. The strategy would be in place for the evaluation of the best selective human estrogen receptor partial agonist (ShERPA) (Xiong et al. 2016) to then go into the adjuvant testing. Already liquid biopsies for breast cancer with low tumor burden are being advanced in clinical testing (Phallen et al. 2017). This technical advance will support adjuvant monitoring of apoptotic success.

Breast cancer treatment must evolve to improve women’s health. The view that ‘this is good enough’ must not take hold. A new SERM for adjuvant therapy must be the best to improve women’s health and the best to prevent breast cancer recurrences. Knowledge now exists for the eventual implementation of a new preemptive salvage therapy (Fig. 7) strategies based on the planned drug holidays in SOLE, and the use of a new SERM with a positive health pharmacology to replace AIs. Together, a new adjuvant SERM that destroys the ER with a SOLE adjuvant design of 3 month therapeutic windows, using precision medicines that kill micrometastatic breast cancer, would be an optimal strategy for breast cancer therapy to achieve. This would be cheap once patenting of precision medicines lapses, easy to administer orally and would address all current issues with current AI therapy. Nevertheless, it will be said that the plan could take decades so it cannot (should not) be attempted.

Forty years ago (Jordan 1978), there was no long-term adjuvant therapy, no understanding of dangerous side effects with tamoxifen, i.e. endometrial cancer, no AIs or SERMs for women’s health. The strategy of long-term adjuvant therapy in breast cancer was considered ‘mindless’ (Stoll 1991) at best and would not improve patient care, and dangerous at worst as it would encourage premature drug resistance. These medical opinions did not go unchallenged (Jordan 1991). Instead, the translational adjuvant strategies using antiestrogens conceived in the laboratory in the 1970/80s, gave patients fewer RR, unanticipated major decreases in mortality, fewer contralateral breast cancers, awareness of the link between tamoxifen and endometrial cancer, an understanding of the unique mechanisms of acquired resistance to antihormones that can be used to treat breast and prostate cancer and the new science of sex steroid-induced apoptosis. It is now prudent to plan for improvements in clinical care and build on past clinical advances.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of this review.

Funding
This work was supported by the National Institutes of Health NIH, MD Anderson’s Cancer Center support grant CA016672 and Susan G. Komen for the Cure Foundation under award number SAC100009, and Cancer Prevention Research Institute of Texas (CPRIT) for the STARS and STARS plus Awards. V C J thanks the benefactors of the Dallas/Ft. Worth Living Legend Chair of Cancer Research for their generous support. R C would like to thank the Romanian National Authority for Scientific Research and Innovation, Project no. 1.1.2/2017, CNCS – UEFISCDI, project number PN-II-RU-TE-2014-4-0422. Y H would like to thank the King Faisal Specialist Hospital & Research Centre (Gen. Org) (KFSh & RC-Jed) and the Royal Embassy of Saudi Arabia-Cultural Bureau in the USA for their financial support.

Acknowledgements
This article is submitted to celebrate Dr Jordan’s 70th birthday, the 40th anniversary of his proposal to use long-term adjuvant tamoxifen therapy clinically, and his discovery of the new group of medicines called SERMs, with tamoxifen as the pioneering SERM. His body of research work is recognized by the Endocrine Society with Dr Jordan’s selection as the 2018 Endocrine Society Laureate of the Gerald D Aubuch Award for Outstanding Translational Research.

References
Abderrahman B & Jordan VC 2016 Improving long-term adjuvant anti-oestrogenic therapy for breast cancer. Clinical Pharmacist. (https://doi.org/10.1211/CP.2016.20201203)
Abderrahman B & Jordan VC 2017 Rethinking extended adjuvant antiestrogen therapy to increase survivorship in breast cancer. JAMA Oncology [epub]. [https://doi.org/10.1001/jamaoncol.2017.351]
Ahmann FR, Citrin DL, deHaan HA, Guinan P, Jordan VC, Kreis W, Scott M & Trump DL 1987 Zoladex: a sustained-release, monthly luteinizing hormone-releasing hormone analogue for the treatment of advanced prostate cancer. Journal of Clinical Oncology 5 912–917. (https://doi.org/10.1200/JCO.1987.5.6.912)
Akakura K, Bruchovsky N, Goldenberg SL, Jennie PS, Buckley AR & Sullivan LD 1993 Effects of intermittent androgen suppression on androgen-dependent tumors. Apoptosis and serum prostate-specific antigen. Cancer 71 2782–2790. (https://doi.org/10.1002/1097-0142(19930501)71:9<2782::aid-CNCR2820719163;2.0.co;2-z)
Albertsen PC, Moore DF, Shih W, Lin Y, Li H & Lu-Yao GL 2011 Impact of comorbidity on survival among men with localized prostate cancer. Journal of Clinical Oncology 29 1335–1341. (https://doi.org/10.1200/JCO.2010.31.2330)
Alkner S, Bendahl PO, Ehinger A, Lovgren K, Ryden L & Ferno M 2016 Prior adjuvant tamoxifen treatment in breast cancer is linked to increased AIB1 and HER2 expression in metastrophic contralateral breast cancer. PLoS ONE 11 e0150977. (https://doi.org/10.1371/journal.pone.0150977)
Andersen RJ, Mawji NR, Wang J, Wang G, Haile S, Myung JK, Watt K, Tam T, Yang YC, Banuelos CA, et al. 2010 Regression of castrate-
recurrent prostate cancer by a small-molecule inhibitor of the amino-terminus domain of the androgen receptor. Cancer Cell 17 535–546. (https://doi.org/10.1016/j.ccr.2010.04.027)

Anderson KM & Liao S 1968 Selective retention of dihydrotestosterone by prostatic nuclei. Nature 219 277–279. (https://doi.org/10.1038/219277a0)

Anderson WE, Katki HA & Rosenberg PS 2011 Incidence of breast cancer in the United States: current and future trends. Journal of the National Cancer Institute 103 1397–1402. (https://doi.org/10.1093/jnci/djr257)

Anderson GL, Chlebowski RT, Aragaki AK, Kuller LH, Manson JE, Gass M, Bluhm E, Connelly S, Hubbell FA, Lane D, et al. 2012 Conjugated equine oestrogen and breast cancer incidence and mortality in postmenopausal women with hysterectomy: extended follow-up of the Women’s Health Initiative randomised placebo-controlled trial. Lancet Oncology 13 476–486. (https://doi.org/10.1016/S1470-2045(12)70075-X)

Andrieu G, Bostwick D, Brawley O, Gomella L, Marberger M, Tindall D, Breed S, Somerville M, Rittmaster R & Group RS 2004 Chemoprevention of prostate cancer in men at high risk: rationale and design of the dutasteride of prostate cancer events (REDUCE) trial. Journal of Urology 172 1314–1317. (https://doi.org/10.1097/01.ju.0000139320.78673.2a)

Ariazi EA, Cunliffe HE, Lewis-Wambi JS, Slifker MJ, Willis AL, Ramos P, Tapia C, Kim HR, Yerrum S, Sharma CGN, et al. 2011 Estrogen induces apoptosis in estrogen deprivation-resistant breast cancer through stress responses as identified by global gene expression across time. PNAS 108 18879–18886. (https://doi.org/10.1073/pnas.1115188108)

Arora VK, Schenkein E, Murari R, Subudhi SK, Wonvipat J, Balbas MD, Shah N, Cai L, Estathiou E, Logothetis C, et al. 2013 Glucocorticoid receptor confers resistance to antiandrogens by bypassing androgen receptor blockade. Cell 155 1309–1322. (https://doi.org/10.1016/j.cell.2013.11.012)

Attard G & Antonarakis ES 2016 Prostate cancer: AR aberrations and design of the reduction by dutasteride of prostate cancer events (REDUCE) trial. Journal of Urology 196 1516–1524. (https://doi.org/10.1097/JU.0000000000001910)

Attia DM & Ederveen AG 2012 Opposing roles of ERalpha and ERbeta in the genesis and progression of adenocarcinoma in the rat ventral prostate. Prostate 72 1013–1022. (https://doi.org/10.1002/pros.21507)

Barbieri R, Salvi S, Rubagotti A, Boccardo S, Spina B, Truini M, Attia DM & Ederveen AG 2012 Opposing roles of ERalpha and ERbeta in the genesis and progression of adenocarcinoma in the rat ventral prostate. Prostate 72 1013–1022. (https://doi.org/10.1002/pros.21507)

Beatson GT 1896 On the treatment of inoperable cases of carcinoma of the mamma: suggestions for a new method of treatment, with illustrative cases. Lancet 2 104–107. (https://doi.org/10.1016/0140-6736(05)72307-4)

Beaver JA & Park BH 2012 The BOLERO-2 trial: the addition of everolimus to exemestane in the treatment of postmenopausal hormone receptor-positive advanced breast cancer. Future Oncology 8 651–657. (https://doi.org/10.2217/fon.12.49)

Beck JT, Hortobagyi GN, Campone M, Lebrun F, Deulei I, Rugo HS, Pistilli B, Masuda N, Hart L, Melicher B, et al. 2014 Everolimus plus exemestane as first-line therapy in HR(+), HER2(−) advanced breast cancer in BOLERO-2. Breast Cancer Research and Treatment 143 459–467. (https://doi.org/10.1007/s10549-013-2814-5)

Bentrem DJ, Darde RS, Liu H, Macgregor-Schaffer J, Zapf JW & Jordan VC 2001 Molecular mechanism of action at estrogen receptor alpha of a newly clinically relevant antiestrogen (GW7604) related to tamoxifen. Endocrinology 142 838–846. (https://doi.org/10.1210/endo.142.2.7932)

Berstein LM, Wang JP, Zheng H, Yue W, Conaway M & Santen RJ 2004 Long-term exposure to tamoxifen induces hypersensitivity to estradiol. Clinical Cancer Research 10 1530–1534. (https://doi.org/10.1158/1078-0432.CCR-04-0333)

Bisson WH, Abagyan R & Cavasotto CN 2008 Molecular basis of agonicity and antagonicity in the androgen receptor studied by molecular dynamics simulations. Journal of Molecular Graphics and Modelling 27 452–458. (https://doi.org/10.1016/j.jmgm.2008.08.001)

Bohl CE, Gao W, Miller DD, Bell CE & Dalton JT 2005 Structural basis for antagonism and resistance of bicalutamide in prostate cancer. PNAS 102 6201–6206. (https://doi.org/10.1073/pnas.0500381102)

Bohl CE, Miller DD, Chen J, Bell CE & Dalton JT 2005b Structural basis for accommodation of nonsteroidal ligands in the androgen receptor. Journal of Biological Chemistry 280 37747–37754. (https://doi.org/10.1074/jbc.M507464200)

Bolla M, van Poppel H, Collette L, van Cangh P, Vekemans K, Da Pozzo L, de Reijke TM, Verhaegs M, Bossert JF, van Velthoven R, et al. 2005 Postoperative radiotherapy after radical prostatectomy for randomised controlled trial (EORTC trial 22911). Lancet 366 572–578. (https://doi.org/10.1016/S0140-6736(05)67101-2)

Bosland MC 2016 Is there a future for chemoprevention of prostate cancer? Cancer Prevention Research 9 642–647. (https://doi.org/10.1158/1940-6207.CAPR-160088)

Bostwick DG, Grignon DJ, Hammond ME, Amin MB, Cohen M, Crawford D, Gospodorowicz M, Kaplan RS, Miller DS, Montironi R, et al. 2000 Prognostic factors in prostate cancer. College of American Pathologists Consensus Statement 1999. Archives of Pathology and Laboratory Medicine 124 995–1000. (https://doi.org/10.1043/0033-9152(00)01249-5/PPHC=2.0.CO;2)

Boyd S 1990 On oophorectomy in cancer of the breast. British Medical Journal 1900 1161–1162

Breast International Group (BIG) I-98 Collaborative Group, Thrulmman B, Keshaviah A, Coates AS, Mouridsen H, Maurici L, Forbes JF, Paridaens R, Castiglione-Gertsch M, Gelber RD, et al. 2005 A comparison of letrozole and tamoxifen in postmenopausal women with early breast cancer. New England Journal of Medicine 353 2747–2757. (https://doi.org/10.1056/NEJMoa052558)

Brodie AM, Schwarzel WC, Shalik AA & Brodie HJ 1977 The effect of an amino-terminus domain of the androgen receptor.

Brodie AM, Schwarzel WC, Shaikh AA & Brodie HJ 1977 The effect of an amino-terminus domain of the androgen receptor. New England Journal of Medicine 366 520–529. (https://doi.org/10.1056/NEJMoa110653)

Brodie AM, Schwarzel WC, Shaikh AA & Brodie HJ 1977 The effect of an amino-terminus domain of the androgen receptor. New England Journal of Medicine 366 520–529. (https://doi.org/10.1056/NEJMoa110653)

Brodie AM, Schwarzel WC, Shaikh AA & Brodie HJ 1977 The effect of an amino-terminus domain of the androgen receptor. New England Journal of Medicine 366 520–529. (https://doi.org/10.1056/NEJMoa110653)
Brodie AM, Marsh D & Brodie HJ 1979 Aromatase inhibitors – IV. Regression of hormone-dependent, mammary tumors in the rat with 4-acetoxy-4-androstene-3,17-dione. Journal of Steroid Biochemistry 10 423–429. (https://doi.org/10.1016/0022-4731(79)90330-3)

Bruchovsky N & Wilson JD 1968 The intranuclear binding of testosterone and 5-alpha-androst-17-beta-ol-3-one by rat prostate. Journal of Biological Chemistry 243 5593–5596.

Bruchovsky N, Klotz LH, Sadar M, Crook JM, Hoffart D, Godwin L, Warkentin M, Gleave ME & Goldenberg SL 2000 Intermittent androgen suppression for prostate cancer: Canadian Prospective Trial and related observations. Molecular Urology 4 191–199.

Bubley GJ & Balk SP 2017 Association between androgen receptor splice variants and prostate cancer resistance to abiraterone and enzalutamide. Journal of Clinical Oncology 35 2103–2105. (https://doi.org/10.1200/JCO.2017.72.8808).

Buchanan G, Greenberg NM, Scher HI, Harris JM, Marshall VR & Tilley WD 2001 Colocalization of androgen receptor gene mutations in prostate cancer. Clinical Cancer Research 7 1273–1281.

Butun SE, Sharda G, Rink J, Sharma S & Simpson ER 1996 Distribution of aromatase P450 transcripts and adiopose fibroblasts in the human breast. Journal of Clinical Endocrinology and Metabolism 81 1273–1277. (https://doi.org/10.1210/jcem.81.8.8772611)

Busch S, Sims AH, Stal O, Ferno M & Landberg G 2015 Loss of TGFbeta receptor type 2 expression impairs estrogen response and confers tamoxifen resistance. Cancer Research 75 1457–1469. (https://doi.org/10.1158/0008-5472.CAN-14-1583)

Butt AJ, McNeill CM, Musgrove EA & Sutherland RL 2005 Downstream targets of growth factor and oestrogen signalling and endocrine resistance: the potential roles of c-Myc, cyclin D1 and cyclin E. Endocrine-Related Cancer 12 Supplement 1 547–559. (https://doi.org/10.1677/erc.1.00993)

Cai C, Chen S, Ng P, Bubley GJ, Nelson PS, Mostaghel EA, Marck B, Matsumoto AM, Simon NI, Wang H, et al. 2011 Intranuclear de novo steroid synthesis activates androgen receptor in castration-resistant prostate cancer and is upregulated by treatment with CYP17A1 inhibitors. Cancer Research 71 6503–6513. (https://doi.org/10.1158/0008-5472.CAN-11-0532)

Carey LA & Perou CM 2015 Palbociclib – taking breast-cancer cells out of gear. New England Journal of Medicine 373 273–274. (https://doi.org/10.1056/NEJMra1506800)

Catalano S, Giordano C, Panza S, Chemi F, Bonofoglio D, Lanzino M, Rizza P, Romiti F, Fuqua SA, Maggioni MP, et al. 2014 Tamoxifen through GPER regulates aromatase expression: a novel mechanism sustaining tamoxifen-resistant breast cancer cell growth. Breast Cancer Research and Treatment 146 273–285. (https://doi.org/10.1007/s10549-014-0107-4)

Chang KH, Encole CE & Sharifi N 2014 Androgen metabolism in prostate cancer: from molecular mechanisms to clinical consequences. British Journal of Cancer 111 1249–1254. (https://doi.org/10.1038/bjc.2014.268)

Chen T, Wang LH & Farrar WL 2000 Interleukin 6 activates androgen receptor-mediated gene expression through a signal transducer and activator of transcription 3-dependent pathway in LNCaP prostate cancer cells. Cancer Research 60 2123–2125.

Chen EJ, Sowalsky AG, Gao S, Cai C, Voznesensky O, Schafer R, Loda M, True LD, Ye H, Troncoso P, et al. 2015 Abiraterone treatment in castration-resistant prostate cancer selects for progesterone responsive mutant androgen receptors. Clinical Cancer Research 21 1273–1280. (https://doi.org/10.1158/1078-0432.CCR-14-1220)

Chodak G, Sharifi R, Kasimis B, Block NL, Macramalla E & Kennealey GT 1995 Single-agent therapy with bicalutamide: a comparison with medical or surgical castration in the treatment of advanced prostate carcinoma. Urology 46 849–855. (https://doi.org/10.1016/S0090-4295(99)00356-2)

Chu IM, Hengst L & Slingerland JM 2008 The Cdk inhibitor p27 in human cancer: prognostic potential and relevance to anticancer therapy. Nature Reviews Cancer 8 253–267. (https://doi.org/10.1038/nrc2347)

Chu VP, Konttinen JM, Hlippakia RA, Fukuchi J, Lin HP, Lin CY, Huo C & Su LC 2011a Androgens as therapy for androgen receptor-positive castration-resistant prostate cancer. Journal of Biomedical Science 18 53. (https://doi.org/10.1186/1423-0127-18-63)

Chu VP, Konttinen JM, Hlippakia RA, Fukuchi J, Lin HP, Lin CY, Huo C, Su LC & Liao S 2011b Androgen suppresses proliferation of castration-resistant LNCaP 104-R2 prostate cancer cells through androgen receptor, Skp2, and c-Myc. Cancer Science 102 2022–2028. (https://doi.org/10.1111/j.1349-7006.2011.02043.x)

Cocconi G 1994 First generation aromatase inhibitors – aminoglutethimide and testolactolone. Breast Cancer Research and Treatment 30 57–80. (https://doi.org/10.1007/BF00682741)

Colleoni M, Lao W, Karlsson P, Chirgwin JH, Aebi SP, Jerusalem GH, Neven P, Hitte E, Graas M-P, Simoncini E, et al. 2017 SOLE (Study of Letrozole Extension): a phase III randomized clinical trial of continuous vs intermittent letrozole in postmenopausal women who have received 4-6 years of adjuvant endocrine therapy for lymph node-positive, early breast cancer (BC). Journal of Clinical Oncology 35 503–503. (https://doi.org/10.1200/JCO.2017.35.6_suppl.503)

Coombes RC, Goss P, Dowsett M, Gazet JC & Brodie A 1984 4-Hydroxyandrostenedione in treatment of postmenopausal patients with advanced breast cancer. Lancet 2 1237–1239. (https://doi.org/10.1016/S0140-6736(84)92795-8)

Crawford ED, Eisenberger MA, Mcleod DG, Spaulding JT, Benson R, Dorr FA, Blumenstein BA, Davis MA & Goodman PJ 1989 A controlled trial of leuprolide with and without flutamide in prostate-carcinoma. New England Journal of Medicine 321 419–424. (https://doi.org/10.1056/NEJM198908173210702)

Culig Z, Hobisch A, Cronauer MV, Radmayr C, Trapani J, Hittmair A, Bartsch G & Klocker H 1994 Androgen receptor activation in prostatic tumor cell lines by insulin-like growth factor-I, keratinocyte growth factor, and epidermal growth factor. Cancer Research 54 5474–5478.

Culig Z, Bartsch G & Hobisch A 2002 Interleukin-6 regulates androgen receptor activity and prostate cancer cell growth. Molecular and Cellular Endocrinology 197 231–238. (https://doi.org/10.1016/S0303-7207(02)00266-0)

Cummins SR, Ensrud K, Delmas PD, LaCroix AZ, Reid DM, Goldstein S, Sirianun P, Lee A, Thompson J, et al. 2010 Lasoxifene in postmenopausal women with osteoporosis. New England Journal of Medicine 362 686–696. (https://doi.org/10.1056/NEJMoa0808692)

Cuzick J 2015 Statistical controversies in clinical research: long-term follow-up of clinical trials in cancer. Annals of Oncology 26 2363–2366. (https://doi.org/10.1093/annonc/mdv392)

Cuzick J, Forbes JF, Sestak I, Cawthorn S, Hamed H, Holli K, Howell A & International Breast Cancer Intervention Study II 2007 Long-term results of tamoxifen prophylaxis for breast cancer – 96-month follow-up of the randomized IBIS-I trial. Journal of the National Cancer Institute 99 272–282. (https://doi.org/10.1093/jnci/djk049)

Cuzick J, Sestak I, Baum M, Buzdar A, Howell A, Dowsett M, Forbes JF & Investigators AL 2010 Effect of anastrozole and tamoxifen as adjuvant treatment for early-stage breast cancer: 10-year analysis of the ATAC trial. Lancet Oncology 11 1135–1141. (https://doi.org/10.1016/S1470-2045(10)70257-6)

Cuzick J, Wickerham L & Powles T 2016 Differing perspectives on breast cancer chemoprevention. JAMA Oncology 2 276–277. (https://doi.org/10.1001/jamaoncol.2015.4406)

Davies C, Godwin J, Gray R, Clarke M, Darby S, McGale P, Wang YC, Peto R, Godwin J, Pan HC, et al. 2011 Relevance of breast cancer hormone receptors and other factors to the efficacy of adjuvant
tamoxifen: patient-level meta-analysis of randomised trials. *Lancet* **378** 771–784. (https://doi.org/10.1016/S0140-6736(11)60993-8)

Davies C, Pan HC, Godwin J, Gray R, Arriagada R, Raina V, Abraham M, Alencar VHM, Badran A, Bonfill X, et al. 2013 Long-term effects of continuing adjuvant tamoxifen to 10 years versus stopping at 5 years after diagnosis of oestrogen receptor-positive breast cancer: ATLAS, a randomised trial. *Lancet* **381** 805–816. (https://doi.org/10.1016/S0140-6736(13)61693-1)

De Bono JS, Oudard S, Ozuguroglu M, Hansen S, Machiels JP, Kokc i, Gravis G, Bodersg I, Mackenzie MJ, Shen L, et al. 2010 Prednisonse plus cabazitaxel or mitoxantrone for metastatic castration-resistant prostate cancer progressing after docetaxel treatment: a randomised open-label trial. *Lancet* **376** 1147–1154. (https://doi.org/10.1016/S0140-6736(10)61389-X)

De Bono JS, Logothetis CJ, Molina A, Fitzzi K, North S, Chu L, Chi KN, Jones RJ, Goodman OB Jr, Saad F, et al. 2011 Abiraterone and increased survival in metastatic prostate cancer. *New England Journal of Medicine* **364** 1995–2005. (https://doi.org/10.1056/NEJMo1a14618)

degraffenried LA, Friedrichs WE, Russell DH, Donzij ES, Middleton AK, Silva JM, Roth RA & Hidalgo M 2004 Inhibition of mTOR activity restores tamoxifen response in breast cancer cells with aberrant Akt Activity. *Clinical Cancer Research* **10** 8059–8067. (https://doi.org/10.1158/1078-0432.CCR-04-0035)

Dehm SM, Schmidt LJ, Heemers HV, Vessella RL & Tindall DJ 2008 Splicing of a novel androgen receptor exon generates a constitutively active androgen receptor that mediates prostate cancer therapy resistance. *Cancer Research* **68** 5469–5477. (https://doi.org/10.1158/0008-5472.CAN-08-0594)

Dosik M & Kaufman R 2004Raloxifene rebound regression. *Cancer Investigation* **22** 718–722. (https://doi.org/10.1081/CNV-200032978)

Dowsett M, Goss PE, Powles TJ, Hutchinson G, Brodie AM, Jefferoe SL & Coombs RC 1987 Use of the aromatase inhibitor 4-hydroxyandrostenedione in postmenopausal breast cancer: optimization of therapeutic dose and route. *Cancer Research* **47** 1957–1961.

Dowsett M, Cuzick J, Ingle J, Coates A, Forbes J, Bliss J, Buyse M, Baum M, Buzdar A, Colleoni M, et al. 2010 Meta-analysis of breast cancer outcomes in adjuvant trials of aromatase inhibitors versus tamoxifen. *Journal of Clinical Oncology* **28** 509–518. (https://doi.org/10.1200/JCO.2009.23.1274)

Duck CB, Jones A, Bohl CE, Dalton JT & Miller DD 2011 Unexpected binding orientation of bulky-B ring anti-androgens and implications for future drug targets. *Journal of Medicinal Chemistry* **54** 3973–3976. (https://doi.org/10.1021/jm200097)

Early Breast Cancer Trialists’ Collaborative Group, Dowsett M, Forbes JF, Bradley R, Ingle J, Alhara T, Bliss J, Boccardo F, Coates A, Coombs RC, et al. 2015 Aromatase inhibitors versus tamoxifen in early breast cancer: patient-level meta-analysis of the randomised trials. *Lancet* **386** 1341–1352. (https://doi.org/10.1016/S0140-6736(15)60174-1)

EBCTCG 1998 Tamoxifen for early breast cancer: an overview of the randomised trials. Early Breast Cancer Trialists’ Collaborative Group. *Lancet* **351** 1451–1467

Edlind MP & Hsieh AC 2014 PI3K-AKT-mTOR signaling in prostate cancer progression and androgen deprivation therapy resistance. *Asian Journal of Andrology* **16** 378–386. (https://doi.org/10.4103/1008-682X.122876)

Ellis MJ, Gao F, Dehdashti F, Jeffe DB, Marcom PK, Carey LA, Dickler MN, Silverman P, Fleming GF, Kommedaddy A, et al. 2009 Lower-dose vs high-dose oral estradiol therapy of hormone receptor-positive, aromatase inhibitor-resistant advanced breast cancer: a phase 2 randomized study. *JAMA* **302** 774–780. (https://doi.org/10.1001/jama.2009.1204)

Fan P, McDaniel RE, Kim HR, Clagett D, Haddad B & Jordan VC 2012 Modulating therapeutic effects of the c-Src inhibitor via oestrogen receptor and human epidermal growth factor receptor 2 in breast cancer cell lines. *European Journal of Cancer* **48** 3488–3498. (https://doi.org/10.1016/j.ejca.2012.04.020)

Fan P, Agboke FA, Cunliffe HE, Ramos P & Jordan VC 2014a A molecular model for the mechanism of acquired tamoxifen resistance in breast cancer. *European Journal of Cancer* **50** 2866–2876. (https://doi.org/10.1016/j.ejca.2014.08.011)

Fan P, Agboke FA, McDaniel RE, Sweeney EE, Zou X, Creswell K & Jordan VC 2014b Inhibition of c-Src blocks oestrogen-induced apoptosis and restores oestrogen-stimulated growth in long-term oestrogen-deprived breast cancer cells. *European Journal of Cancer* **50** 457–468. (https://doi.org/10.1016/j.ejca.2013.10.001)

Fan P, Cunliffe HE, Griffith OL, Agboke FA, Ramos P, Gray JW & Jordan VC 2014c Identification of gene regulation patterns underlying both oestrogen- and tamoxifen-stimulated cell growth through global gene expression profiling in breast cancer cells. *Molecular Cancer Research* **13** 1367–1376. (https://doi.org/10.1158/1541-7786.MCR-14-0494)

Fanning SW, Mayne CG, Dharmarajan V, Carlson KE, Martin TA, Novick SJ, Toy W, Green B, Panchamukhi S, Katzenellenbogen BS, et al. 2016 Estrogen receptor alpha somatic mutations Y537S and D538G confer breast cancer endocrine resistance by stabilizing the activating function-2 binding conformation. *Elife* **5** e12792. (https://doi.org/10.7554/eLife.12792)

Fasolo A & Sessa C 2008 mTOR inhibitors in the treatment of cancer. *Expert Opinion on Investigational Drugs* **17** 1717–1734. (https://doi.org/10.1517/13543784.17.11.1717)

FDA 2011 FDA drug safety communication: S-alpha reduce inhibitors (S-ARIs) may increase the risk of a more serious form of prostate cancer. Rockville, MD, USA: FDA (available at: https://www.fda.gov/Drugs/DrugSafety/ucm258314.htm)

Feldman BJ & Feldman D 2001 The development of androgen-independent prostate cancer. *Nature Reviews Cancer* **1** 34–45. (https://doi.org/10.1038/35094009)

Finn RS, Crown JP, Lang I, Boer K, Bondareno IM, Kulyk SO, Ettl J, Patel R, Pinter T, Schmidt M, et al. 2015 The cyclin-dependent kinase 4/6 inhibitor palbociclib in combination with letrozole versus letrozole alone as first-line treatment of oestrogen receptor-positive, HER2-negative, advanced breast cancer (PALOMA-1/TRIO-18): a randomised phase 2 study. *Lancet Oncology* **16** 25–35. (https://doi.org/10.1016/S1470-2045(14)71159-3)

Fisher B, Costantino JP, Wickerham DL, Cecchini RS, Cronin WM, Robidoux A, Bevers TB, Kavanah MT, Atkins JN, Margolese RG, & et al. 2005 Tamoxifen for the prevention of breast cancer: current status of the National Surgical Adjuvant Breast and Bowel Project P-1 study. *Journal of the National Cancer Institute* **97** 1652–1662. (https://doi.org/10.1093/jnci/dji372)

Flesner N, Keane TE, Lawton CA, Mulders PE, Payne H, Taneja SS & Morris T 2008 Adjuvant androgen deprivation therapy augments cure and long-term cancer control in men with poor prognosis, nonmetastatic prostate cancer. *Prostate Cancer and Prostatic Diseases* **11** 46–52. (https://doi.org/10.1038/sj.pcan.4500982)

Fradeit A, Bouchet M, Delliaux C, Gervais M, Kan C, Benelotto C, Fantano P, Vargas G, Bouazza L, Crosst M, et al. 2016 Estrogen related receptor alpha in castration-resistant prostate cancer cells promotes tumor progression in bone. *Oncotarget* **7** 77071–77086.
Fujimoto N, Mizokami A, Harada S & Matsumoto T 2001 Different expression of androgen receptor coactivators in human prostate. *Urology* **58** 289–294. (https://doi.org/10.1001/0000-9495(2001)01117-7)

Furr BJ & Jordan VC 1984 The pharmacology and clinical uses of tamoxifen. *Pharmacology and Therapeutics* **25** 127–205. (https://doi.org/10.1016/0163-7258(84)90043-3)

Goss PE, Ingle JN, Martino S, Robert NJ, Muss HB, Piccart MJ, Castiglione M, Tu D, Shepherd LE, Pritchard KI, et al. 2003 A randomized trial of letrozole in postmenopausal women after five years of tamoxifen therapy for early-stage breast cancer. *New England Journal of Medicine* **349** 1793–1802. (https://doi.org/10.1056/NEJMoa032312)

Gottardis MM, Wagner RJ, Borden EC & Jordan VC 1987 Development of advanced postmenopausal breast cancer with an aromatase inhibitor, 4-hydroxysterondienone: phase II report. *Cancer Research* **46** 4823–4826.

Gottardis MM, Wagner RJ, Borden EC & Jordan VC 1989 Development of advanced postmenopausal breast cancer with an aromatase inhibitor, 4-hydroxysterondienone: phase II report. *Cancer Research* **49** 4090–4093.

Gottardis MM, Wagner RJ, Borden EC & Jordan VC 1989b Differential ability of antiestrogens to stimulate breast cancer cell (MCF-7) growth in vivo and in vitro. *Cancer Research* **49** 4765–4769.

Gray RG, Rea D, Handley K, Bowden SJ, Perry P, Earl HM, Poole CJ, Bates T, Chettyawardana S, Dewar JA, et al. 2013 aTTom: long-term effects of continuing adjuvant tamoxifen to 10 years versus stopping at 5 years in 6,953 women with early breast cancer. *Journal of Clinical Oncology* **31** (15 Supplement) abstract 5. (https://doi.org/10.1200/jco.2013.31.15_suppl.5)

Gregory CW, He B, Johnson RT, Ford OH, Mohler JL, French FS & Wilson EM 2001b Androgen receptor-mediated prostate cancer recurrence after androgen deprivation therapy. *Cancer Research* **61** 4315–4319.

Gregory CW, Johnson RT Jr, Mohler JL, French FS & Wilson EM 2001a A mechanism for androgen receptor-mediated prostate cancer recurrence after androgen deprivation therapy. *Journal of Clinical Oncology* **19** 3396–3403. (https://doi.org/10.1200/JCO.2002.10.057)

Higgins C & Hodges CV 1914 Studies on prostatic cancer - I The effect of castration, of estrogen and of androgen injection on serum phosphatases in metastatic carcinoma of the prostate. *Cancer Research* **1** 293–297.

Ignatov A, Ignatov T, Roessner A, Costa SD & Kalinski T 2010 Role of GPR30 in the mechanisms of tamoxifen resistance in breast cancer MCF-7 cells. *Breast Cancer Research and Treatment* **123** 87–96. (https://doi.org/10.1007/s10549-009-0624-6)

Ingle JN, Ahmann BL, Green SJ, Edmonson JH, Bresalier RS, Kools IK, Nichols WC, Creagan ET, Hahn RG, Rubin J, et al. 1981 Randomized
clinical trial of diethylstilbestrol versus tamoxifen in postmenopausal women with advanced breast cancer. New England Journal of Medicine 304 16–21. (https://doi.org/10.1056/NEJM198101010340104)
Jarrard DF, Kinoshita H, Shi Y, Sandefur C, Hoff D, Meinsen LE, Chang C, Herman JG, Issacs WB & Nasif N 1998 Methylation of the androgen receptor promoter CpG island is associated with loss of androgen receptor expression in prostate cancer cells. Cancer Research 58 5310–5314.
Jensen EV & Jacobson HI 1962 Basic guides to the mechanism of estrogen action. Recent Progress in Hormone Research 18 387–414.
Jeselsohn R, Yelensky R, Buchwalter G, Frampton G, Mercier-Bernstam F, Gonzalez-Angulo AM, Ferrer-Lozano J, Perez-Fidalgo JA, Cristofanilli M, Gomez H, et al. 2014 Emergence of constitutively active estrogen receptor-alpha mutations in pretreated advanced estrogen receptor-positive breast cancer. Clinical Cancer Research 20 1757–1767. (https://doi.org/10.1158/1078-0432.CCR-13-2332)
Jiang SY, Wolf DM, Yingling JM, Chang C & Jordan VC 1992 An estrogen-receptor positive Mcf-7 clone that is resistant to antiestrogens and estradiol. Molecular and Cellular Endocrinology 90 77–86. (https://doi.org/10.1016/0303-7207(92)90104-E)
Joly-Pharboiz MO, Ruffion A, Roch A, Michel-Calemard L, Andre J, Chantepie J, Nicolas B & Panaye G 2000 Inhibition of growth and induction of apoptosis by androgens of a variant of LNCaP cell line. Journal of Steroid Biochemistry and Molecular Biology 73 237–249. (https://doi.org/10.1016/S0960-0760(00)00076-5)
Jordan VC 1976 Effect of tamoxifen (ICI 46,474) on initiation and growth of DMBA-induced rat mammary carcinomas. European Journal of Cancer 12 419–424. (https://doi.org/10.1016/0014-2295(76)90030-X)
Jordan VC 1978 DMBA-induced rat mammary carcinoma system for the evaluation of tamoxifen treatment as a potential adjuvant therapy. Reviews on Endocrine-Related Cancer (October Supplement) 49–55.
Jordan VC 1984 Biochemical pharmacology of antiestrogen action. Pharmacological Reviews 36 245–276.
Jordan VC 1991 Prolonged adjuvant tamoxifen: a beginning not the end. Annals of Oncology 2 481–484. (https://doi.org/10.1093/oxfordjournals.annonc.a005799)
Jordan VC 2003 Tamoxifen: a most unlikely pioneering medicine. Nature Reviews Drug Discovery 2 205–213. (https://doi.org/10.1038/nrd1031)
Jordan VC 2006 Tamoxifen (ICI46,474) as a targeted therapy to treat and prevent breast cancer. British Journal of Pharmacology 147 (Supplement 1) S269–S276. (https://doi.org/10.1038/sj.bjp.0706399)
Jordan VC 2008 The 38th David A Karnofsky lecture: the paradoxical the tamoxifen tale. European Journal of Cancer 44 239–251. (https://doi.org/10.1016/j.ejca.2008.09.056)
Jordan VC, Dix CJ & Allen KE 1979 The effectiveness of long-term tamoxifen treatment in a laboratory model for adjuvant hormone therapy of breast cancer. In Adjutant Therapy of Cancer II, pp 19–26. Eds SE Salmon & SE Jones. New York, NY, USA: Grune and Strattin.
Jordan VC, Naylor KE, Dix CJ & Prestwich G 1980 Anti-oestrogen action in experimental breast cancer. Recent Results in Cancer Research 71 30–44.
Jordan VC, Fritz NE, Langan-Fahey S, Thompson M & Tormey DC 1991 Alteration of estrogen endocri parameters in premenopausal women with breast cancer during long-term adjuvant therapy with tamoxifen as the single agent. Journal of the National Cancer Institute 83 1488–1491. (https://doi.org/10.1093/jnci/83.20.1488)
Jordan VC, Curpan R & Maximov PY 2015 Estrogen receptor mutations found in breast cancer metastases integrated with the molecular pharmacology of selective ER modulators. Journal of the National Cancer Institute 107 djv075. (https://doi.org/10.1093/jnci/djv075)
Jordan VC, Fan P, Abderrahman B, Maximov PY, Hawsawi YM, Bhattacharya P & Pokharel N 2016 Sex steroid induced apoptosis as a rational strategy to treat anti-hormone resistant breast and prostate cancer. Discovery Medicine 21 411–427.
Kaplan CP, Haas JS, Perez-Stable EJ, Des Jarlais DC & Gregorich SE 2005 Factors affecting breast cancer risk reduction practices among California physicians. Preventive Medicine 41 7–15. (https://doi.org/10.1016/j.ypmed.2004.09.041)
Kattan MW, Wheeler TM & Scardino PT 1999 Postoperative nomogram for disease recurrence after radical prostatectomy for prostate cancer. Journal of Clinical Oncology 17 1499–1507. (https://doi.org/10.1200/JCO.1999.17.5.1499)
Katzenellenbogen BS, Kendra KL, Norman MJ & Berthois Y 1987 Proliferation, hormonal responsiveness, and estrogen receptor content of MCF-7 human breast cancer cells grown in the short-term and long-term absence of estrogens. Cancer Research 47 4355–4360.
Kawata H, Ishikura N, Watanabe M, Nishimoto A, Tsunenari T & Aoki Y 2011 Administration of tamoxifen and prevention of breast cancer. Cancer Research 71 1580–1586.
Kawara H, Ishikura N, Watanabe M, Nishimoto A, Tsunenari T & Aoki Y 2011 Administration of tamoxifen and prevention of breast cancer. Cancer Research 71 1580–1586.
Kokontis J, Takakura K, Hay N & Liao S 1994 Increased androgen receptor expression and androgen hypersensitivity. Prostate 20 745–754. (https://doi.org/10.1002/pros.21107)
Kennedy BJ 1965 Hormone therapy for advanced breast cancer. Cancer 18 1551–1557. (https://doi.org/10.1002/1097-0142(196512)18:12<1551::AID-CNCR2820181206>3.0.CO;2-1)
Kirk M, Hirsh C & Crawford ED 2017 Characterising the castration-resistant prostate cancer population: a systematic review. International Journal of Clinical Practice 71 1180–1192. (https://doi.org/10.1111/ijc.12414.2011.02799.x)
Klotz L & Higano CS 2016 Intermittent androgen deprivation therapy-an important treatment option for prostate cancer. JAMA Oncology 2 1531–1532. (https://doi.org/10.1001/jamaoncol.2016.3138)
Kokontis J, Takakura K, Hay N & Liao S 1994 Increased androgen receptor expression and altered c-myc expression in prostate cancer cells after long-term androgen deprivation. Cancer Research 54 1566–1573.
Kote-Jaral Z, Leongamornlert D, Saunders E, Tymrakiewicz M, Castro E, Mahmud N, Guy M, Edwards S, O’Brien L, Sawyer E, et al. 2011 BRC2A is a moderate penetrance gene contributing to young-onset prostate cancer: implications for genetic testing in prostate cancer patients. British Journal of Cancer 105 1230–1234. (https://doi.org/10.1038/bjc.2011.383)

Labrie F, Dupont A, Giguere M, Borsary J-P, Belanger A, Lacourciere Y, Emond J & Monfette G 1986 Advantages of the combination therapy in previously untreated and treated patients with advanced prostate cancer. Journal of Steroid Biochemistry 25 877–883. (https://doi.org/10.1016/0022-4731(86)90319-5)

Legare S & Basik M 2016 Minireview: the link between ERAlpha corepressors and histone deacetylates in tamoxifen resistance in breast cancer. Molecular Endocrinology 30 965–976. (https://doi.org/10.1210/me.2016-1072)

Lemmo W 2016 Anti-estrogen withdrawal effect with raloxifene? A case report. Integrative Cancer Therapies 15 245–249. (https://doi.org/10.1177/1534735416658954)

Lemmon MR, Mahmud N, Tymrakiewicz M, Saunders E, Dadaev T, Castro E, Goh C, Govindasami K, Guy M, O’Brien L, et al. 2012 Germline BRC1A mutations increase prostate cancer risk. British Journal of Cancer 106 1697–1701. (https://doi.org/10.1038/bjc.2012.146)

Lerner LJ & Jordan VC 1990 Development of antiestrogens and their use in breast cancer: eighth Cain memorial award lecture. Cancer Research 50 4177–4189.

Lewis JS, Mekee K, Osipo C, Ross EA, Kidawi N, Li T, Bell E, Chandel NS & Jordan VC 2005a Intrinsic mechanism of estradiol-induced apoptosis in breast cancer cells resistant to estrogen deprivation. Journal of the National Cancer Institute 97 1746–1759. (https://doi.org/10.1093/jnci/dji400)

Lewis JS, Osipo C, Mekee K & Jordan VC 2005b Estragon-induced apoptosis in a breast cancer model resistant to long-term estrogen withdrawal. Journal of Steroid Biochemistry and Molecular Biology 94 131–141. (https://doi.org/10.1016/j.jsbmb.2004.12.032)

Li J, Yen C, Liaw D, Podosyanika K, Bose S, Wang SL, Puc J, Miliaresis C, Rodgers L, McCombie R, et al. 1997 PTEN, a putative protein tyrosine phosphatase gene mutated in human brain, breast, and prostate cancer. Science 275 1943–1947. (https://doi.org/10.1126/science.275.5308.1943)

Lin HK, Yeh S, Kang HY & Chang C 2001 Akt suppresses androgen-receptor signaling in prostate cancer cells. PNAS 98 7200–7205. (https://doi.org/10.1073/pnas.121172982)

Lipton A & Santen RJ 1974 Proceedings: medical adrenalectomy using cortisol, fludrocortisone and Dexamethasone. JAMA 219 792–804. (https://doi.org/10.1001/jama.1974.033300300272x)

Locke JA, Guns ES, Lubik AA, Adomat HH, Hendy SC, Wood CA, Ettinger SL, Gleave ME & Nelson CC 2008 Androgen levels increase by intratumoral de novo steroidogenesis during progression of castration-resistant prostate cancer. Cancer Research 68 6407–6415. (https://doi.org/10.1158/0008-5472.CAN-07-5997)

Lu-Yao GL, Albertsen PC, Moore DF, Shih W, Lin Y, DiPaola RS, Lipton A & Santen RJ 1974 Proceedings: medical adrenalectomy using cortisol, fludrocortisone and Dexamethasone. JAMA 219 792–804. (https://doi.org/10.1001/jama.1974.033300300272x)

Luo Y, Albersen PC, Moore DF, Shih W, Lin J, DiPaola RS, Barry MJ, Zietman A, O’Leary M, Walker-Corkery E, Barry MJ, Zietman A, O’Leary M, Walker-Corkery E, et al. 2009 Outcomes of localized prostate cancer following conservative management. JAMA 302 1202–1209. (https://doi.org/10.1001/jama.2009.1348)

Mainwaring WI 1969 A soluble androgen receptor in the cytoplasm of rat prostate. Journal of Endocrinology 45 531–541. (https://doi.org/10.1677/joe.0.0450531)

Marcelli M, Ittmann M, Mariani S, Sutherland R, Nigam R, Murthy L, Zhao Y, DiConcini D, Puxedu E, Esen A, et al. 2000 Androgen receptor mutations in prostate cancer. Cancer Research 60 944–949.

Marques RB, Dits NF, Erkens-Schulze S, van Weerden WM & Jenster G 2010 Bypass mechanisms of the androgen receptor pathway in therapy-resistant prostate cancer cell models. PLoS ONE 5 e13500. (https://doi.org/10.1371/journal.pone.0013500)

Masamura S, Santner SJ, Hcjtian DF & Santen RJ 1995 Estrogen deprivation causes estradiol hypersensitivity in human breast cancer cells. Journal of Clinical Endocrinology and Metabolism 80 2918–2925. (https://doi.org/10.1210/jcem.80.8.7559875)

Mathew P 2008 Prolonged control of progressive castration-resistant metastatic prostate cancer with testosterone replacement therapy: the case for a prospective trial. Annals of Oncology 19 395–396. (https://doi.org/10.1093/annonc/mdm568)

Maximov PY, Lee TM & Jordan VC 2013 The discovery and development of selective estrogen receptor modulators (SERMs) for clinical practice. Current Clinical Pharmacology 8 135–155. (https://doi.org/10.2174/1574884711308020006)

McGuire WL, Carbone PP, Vollmer EP, United States National Cancer Institute & Breast Cancer Treatment Committee 1975 Estron Receptors in Human Breast Cancer. New York: Raven Press.

Melisko ME, Goldberg ME, Hwang J, De Luca A, Fang S, Esserman LJ, Chien AJ, Park JW & Rugo HS 2017 Vaginal testosterone cream vs estradiol vaginal ring for vaginal dryness or decreased libido in women receiving aromatase inhibitors for early-stage breast cancer: a randomized clinical trial. JAMA Oncology 3 313–319. (https://doi.org/10.1001/jamaoncol.2016.3904)

Miller TW 2013 Endocrine resistance: what do we know? American Society of Clinical Oncology Educational Book. (https://doi.org/10.1200/EdBook_AM.2013.33.e37)

Mishra S, Tai Q, Gu X, Schmitz J, Pouillard A, Fajardo R, Mahalingham D, Chen Z, Zhu X & Sun LZ 2015 Estron and estrogen receptor alpha promotes malignancy and osteoblastic tumorigenesis in prostate cancer. Oncotarget 6 44388–44402. (https://doi.org/10.18632/oncotarget.6317)

Mo Z, Liu M, Yang F, Luo H, Li Z, Tu G & Yang G 2013 GPR30 as an initiator of tamoxifen resistance in hormone-dependent breast cancer. Breast Cancer Research 15 R114. (https://doi.org/10.1186/bcr3581)

Mulholland DJ, Tran LM, Li Y, Cai H, Morein A, Wang S, Plaisier S, Giarraway IP, Huang J, Graeber TG, et al. 2011 Cell autonomous role of PTEN in regulating castration-resistant prostate cancer growth. Cancer Cell 19 792–804. (https://doi.org/10.1016/j.ccr.2011.05.006)

Nettles KW, Lazzaro OP, Fernandez L, Chesa N & Diaz-Chico BN 2002 Transition to androgen-independence in prostate cancer. Journal of Steroid Biochemistry and Molecular Biology 81 191–201. (https://doi.org/10.1016/S0960-0760(02)00064-X)

Neri R, Florence K, Kozioł P & Van Cleave S 1972 A biological profile of a nonsteroidal antiandrogen, SCH 13521 (4-nitro-3 trifluoromethoxybenzyl)boronanilide). Endocrinology 91 427–437. (https://doi.org/10.1210/endo-91-2-427)

Neffels KW, Bunring JB, Gil G, Nowak J, Sharma SK, Ahm B, Kulp K, Hochreg RB, Zhou H, Katzenellenbogen JA, et al. 2008 NKappaB selectivity of estrogen receptor ligands revealed by comparative crystallographic analyses. Nature Chemical Biology 4 241–247. (https://doi.org/10.1038/nchembio.76)

Ni L, Yang CS, Gioeli D, Frierson H, Toft DO & Paschal BM 2010 FKBP51 promotes assembly of the Hsp90 chaperone complex and regulates androgen receptor signaling in prostate cancer cells. Molecular and Cellular Biology 30 1243–1253. (https://doi.org/10.1128/MCB.018632-16)

Nishiyama T, Hashimoto Y & Takahashi K 2004 The influence of androgen deprivation therapy on dihydrotestosterone levels in the prostatic tissue of patients with prostate cancer. Clinical Cancer Research 10 7121–7126. (https://doi.org/10.1158/1078-0432.CCR-04-0913)
Steroid-induced apoptosis in postmenopausal women with metastatic breast cancer. An updated analysis. Breast Cancer Research and Treatment 54 117–122. (https://doi.org/10.1023/A:1006185805079)

Penson DE, Armstrong AJ, Concepcion R, Agarwal N, Obsson C, Karsh L, Dunschee C, Wang E, Wu K, Krivoshik A, et al. 2016 Enzalutamide versus bicalutamide in castration-resistant prostate cancer: the STRIVE trial. Journal of Clinical Oncology 34 2098–2106. (https://doi.org/10.1200/JCO.2015.64.9285)

Perez-Tenorio G, Berglund T, Eguerra Merca A, Nordenskjold B, Rutqvist LE, Skoog I & Stal O 2006 Cytoplasmic p21WAF1/CIP1 correlates with Akt activation and poor response to tamoxifen in breast cancer. International Journal of Oncology 28 1031–1042. (https://doi.org/10.3892/ijo.28.5.1031)

Pether M, Goldenberg SL, Bhagrath K & Gleave M 2003 Intermittent androgen suppression in prostate cancer: an update of the Vancouver experience. Canadian Journal of Urology 10 1809–1814.

Phallen J, Sausen M, Adleff V, Leal A, Hruban C, White J, Anagnostou V, Fiksel J, Cristiano S, Papp E, et al. 2017 Direct detection of early-stage cancers using circulating tumor DNA. Science Translational Medicine 9.

Pink JJ, Jiang SY, Fritsch M & Jordan VC 1995 An estrogen-independent MCF-7 breast cancer cell line which contains a novel 80-kilodalton estrogen receptor-related protein. Cancer Research 55 2583–2590.

Powles TJ, Ashley S, Tidy A, Smith IE & Dowsett M 2007 Twenty-year follow-up of the Royal Marsden randomized, double-blinded tamoxifen breast cancer prevention trial. Journal of the National Cancer Institute 99 283–290. (https://doi.org/10.1093/jnci/djk050)

Prat A & Perou CM 2011 Decomposing the molecular portraits of breast cancer. Molecular Oncology 5 23–23. (https://doi.org/10.1016/j.molonc.2010.11.003)

Pritchard KI, Chia SK, Simmons C, McLeod D, Paterson A, Provencher L & Rayson D 2017 Enhancing endocrine therapy combination strategies for the treatment of postmenopausal HR+/HER2- advanced breast cancer. Oncologist 22 12–24. (https://doi.org/10.1634/theoncologist.2016-0185)

Rathkopf DE, Morris MJ, Danilla DC, Slovin SE, Steinbrecher JE, Arauz G, Curley T, Rix PJ, Maneval EC, Chen I, et al. 2012 A phase I study of the androgen signaling inhibitor ARN-509 in patients with metastatic castration-resistant prostate cancer (mCRPC). Journal of Clinical Oncology 30 (15 Supplement) 4548–4548. (https://doi.org/10.1200/jco.2012.30.15_suppl.4548)

Rathkopf DE, Morris MJ, Fox JJ, Danilla DC, Slovin SE, Hager JH, Rix PJ, Chow Maneval EC, Chen I, Gonen M, et al. 2013 Phase I study of ARN-509, a novel antiandrogen, in the treatment of castration-resistant prostate cancer. Journal of Clinical Oncology 31 3525–3530. (https://doi.org/10.1200/JCO.2013.50.1684)

Rau KM, Kang HY, Cha TL, Miller SA & Hung MC 2005 The mechanisms and management of hormone-therapy resistance in breast and prostate cancers. Endocrine-Related Cancer 12 511–532. (https://doi.org/10.1677/erc.1.01026)

Ravdin PM, Fritz NE, Tormey DC & Jordan VC 1988 Endocrine status of premenopausal node-positive breast cancer patients following adjuvant chemotherapy and long-term tamoxifen. Cancer Research 48 1026–1029.

Raynaud JP, Bonne C, Bouton MM, Lagace I & Labrie F 1979 Action of a non-steroid anti-androgen, RU 23908, in peripheral and central tissues. Journal of Steroid Biochemistry 11 93–99. (https://doi.org/10.1016/0022-4731(79)90281-4)

Reeder-Hayes K & Muss HB 2017 Vaginal estrogens and aromatase inhibitors: how safe is safe enough? JAMA Oncology 3 305–306. (https://doi.org/10.1001/jamaoncol.2016.3934)

Riggins RB, Bouton AH, Liu MC & Clarke R 2005 Antiestrogens, aromatase inhibitors, and apoptosis in breast cancer. Vitamins and Hormones 71 201–237.
Endocrine-Related Cancer

Steroid-induced apoptosis in endocrine cancers

25:2

R111

Risbridger GP, Davis ID, Birdsell SN & Tilley WD 2010 Breast and prostate cancer: more similar than different. *Nature Reviews Cancer* **10** 205–212. (https://doi.org/10.1038/nrc2795)

Robertson JFR, Bondarenko IM, Trishkina E, Dvorkin M, Panasci L, Manikhas A, Shparyk Y, Cardona-Huerta S, Cheung KL, Philco-Salas MJ, et al. 2016 Fulvestrant 500 mg versus anastrozole 1 mg for hormone-receptor-positive advanced breast cancer (FALCON): an international, randomised, double-blind, phase 3 trial. *Lancet* **389** 2997–3005. (https://doi.org/10.1016/S0140-6736(16)32389-3)

Robinson DR, Wu YM, Vats P, Su E, Lonigro RJ, Cao X, Kalyana-Sundaram S, Wang R, Ying N, Hodges L, et al. 2013 Activating ERα11 mutations in hormone-resistant metastatic breast cancer. *Nature Genetics* **45** 1446–1451. ([https://doi.org/10.1038/ng.2823](https://doi.org/10.1038/ng.2823))

Robinson D, Van Allen EM, Wu YM, Schultz N, Lonigro RJ, Mosquera JM, Montgomery B, Taplin ME, Pritchard CC, Attard G, et al. 2015 Integrative clinical genomics of advanced prostate cancer. *Cell* **161** 1215–1226. ([https://doi.org/10.1016/j.cell.2015.05.001](https://doi.org/10.1016/j.cell.2015.05.001))

Ropke A, Erbersdobler A, Hammerer P, Palisaar J, John K, Stumm M, et al. 2008-2016) Molecular endocrine-mechanisms underlying hormone-dependent development of human breast cancer: a systematic review and meta-analysis. *Sterility* **22** 703–721. (https://doi.org/10.1016/S0013-0282(16)38580-6)

Schally AV, Comaru-Schally AM, Plonowski A, Nagy A, Halmos G & Rekasi Z 2000 Peptide analogs in the therapy of prostate cancer. *Prostate* **45** 158–166. (https://doi.org/10.1002/1097-0045(20001001)45:2<158::AID-PROS10>3.0.CO;2-K)

Scher HI, Beer TM, Higano CS, Andar A, Taplin ME, Efstathiou E, Rathkopf D, Shelley J, Yu EY, Alumkal J, et al. 2010 Antitumour activity of MDV3100 in castration-resistant prostate cancer: a phase 1-2 study. *Lancet* **375** 1437–1446. (https://doi.org/10.1016/S0140-6736(10)60729-2)

Schiff R, Massarweh SA, Shou J, Bharwani L, Molisin SK & Osborne CK 2004 Cross-talk between estrogen receptor and growth factor pathways as a molecular target for overcoming endocrine resistance. *Clinical Cancer Research* **10** 3315–3366. (https://doi.org/10.1158/1078-0432.CCR-031212)

Schrey MP & Patel KV 1995 Prostaglandin E2 production and metabolism in human breast cancer cells and breast fibroblasts. *Regulation by inflammatory mediators. British Journal of Cancer* **72** 1412–1419. (https://doi.org/10.1038/bjc.1995.523)

Schweizer MT, Antonarakis ES, Wang H, Ajiboye AS, Spitz A, Cao H, Luo J, Haffner MC, Yegnasubramanian S, Carducci MA, et al. 2013 Effect of bipolar androgen therapy for asymptomatic men with castration-resistant prostate cancer: results from a pilot clinical study. *Science Translational Medicine* **7** 269ra2. (https://doi.org/10.1126/scitranslmed.3010563)

Seidenfeld J, Samson DJ, Hasselblad V, Aronson N, Albertsen PC, Bennett CL & Witt TJ 2000 Single-therapy androgen suppression in men with advanced prostate cancer: a systematic review and meta-analysis. *Annals of Internal Medicine* **132** 566–577. (https://doi.org/10.17326/0003-4819-132-7.20000440-00009)

Shiplewy WU, Seiferheld W, Lukka HR, Major PP, Heny NM, Grignon DJ, Sartor O, Patel MP, Bahary JP, Zietman AL, et al. 2017 Radiation with or without antiandrogen therapy in recurrent prostate cancer. *New England Journal of Medicine* **376** 417–428. (https://doi.org/10.1056/NEJMoa1607529)

Shou J, Massarweh S, Osborne CK, Wakeling AE, Ali S, Weiss H & Schiff R 2004 Mechanisms of tamoxifen resistance: increased estrogen receptor-HER2/neu cross-talk in ER/HER2-positive breast cancer. *Journal of the National Cancer Institute* **96** 926–935. (https://doi.org/10.1093/jnci/djh166)

Siegel RL, Miller KD & Jemal A 2015 Cancer statistics, 2015. *CA: A Cancer Journal for Clinicians* **65** 5–29. (https://doi.org/10.3322/caac.21254)

Sledge GW, Mamounas EP, Hortobagyi GN, Burstein HJ, Goodwin PJ & Wolff AC 2014 Past, present, and future challenges in breast cancer treatment. *Journal of Clinical Oncology* **32** 1979–1986. (https://doi.org/10.1200/JCO.2014.55.4139)

Smith DC, Smith MR, Sweeney C, Elfyk AA, Logothetis C, Corn PG, Vogelzang NJ, Small EJ, Harzstark AL, Gordon MS, et al. 2013 Cabozantinib in patients with advanced prostate cancer: results of a phase II randomized discontinuation trial. *Journal of Clinical Oncology* **31** 412–419. (https://doi.org/10.1200/JCO.2012.45.0494)

Smith SG, Side L, Meisel SF, Horne R, Cuzick J & Wardle J 2016 Participant-reported symptoms and their effect on long-term adherence in the international breast cancer intervention study I (IBIS I). *Journal of Clinical Oncology* **34** 3873–3879. (https://doi.org/10.1200/JCO.2016.64.5341)

Smith SG, Sestak I, Howell A, Forbes J & Cuzick J 2017 Participant-reported symptoms and their effect on long-term adherence in the international breast cancer intervention study I (IBIS I). *Journal of Clinical Oncology* **35** 3873–3879. (https://doi.org/10.1200/JCO.2016.64.5341)
Steroid-induced apoptosis in endocrine cancers

Clinical Oncology 10 2666–2673. (https://doi.org/10.1200/JCO.2016.71.7439)

Song RXD, Mor G, Naftolin F, McPherson RA, Song J, Zhang ZG, Yue W, Wang JP & Santen RJ 2001 Effect of long-term estrogen deprivation on apoptotic responses of breast cancer cells to 17 beta-estradiol. Journal of the National Cancer Institute 93 1714–1723. (https://doi.org/10.1093/jnci/93.22.1714)

Span PN, Tjan-Heijnen VC, Manders P, Beex LV & Sweep CG 2003 Cylcwn-E is a strong predictor of endocrine therapy failure in human breast cancer. Oncogene 22 4898–4904. (https://doi.org/10.1038/sj.onc.1206818)

Stoll BA 1991 Overprolonged adjuvant tamoxifen therapy in breast cancer. Advances in Cancer Research 56 249–258. (https://doi.org/10.1016/S0065-3744(08)60452-7)

Thorpe JF, Jain S, Marczylo TH, Gescher AJ, Steward WP & Mellon JK 2005 Structural basis for an unexpected mode of SERM-mTOR antagonism. Molecular and Cellular Endocrinology 237 159–167. (https://doi.org/10.1016/j.mce.2005.04.014)

Tolis G, Ackman D, Stellos A, Mehta A, Labrie F, Fazeika AT, Comaruschally AM & Schally AV 1982 Tumor-Growth Inhibition in Patients with Prostatic-Carcinoma Treated with Luteinizing-Hormone-Releasing Hormone Agonists. Prostate 7 1696–1697. (https://doi.org/10.1002/pro.2120070311)

Tomiguchi M, Yamamoto Y, Yamamoto-Ikuzuki M, Goto-Yamaguchi L, Fujiki Y, Fujisawa S, Sueta A, Hayashi M, Takeshita T, Inao T, et al. 2016 Fibroblast growth factor receptor-1 protein expression is associated with prognosis in estrogen receptor-positive/human epidermal growth factor receptor-2-negative primary breast cancer. Cancer Science 107 491–498. (https://doi.org/10.1111/cas.12897)

Toy W, Shen Y, Won H, Green B, Sakr RA, Will M, Li Z, Gala K, Fanning S, King TA, et al. 2013 ESRI1 ligand-binding domain mutations in hormone-resistant breast cancer. Nature Genetics 45 1439–1444. (https://doi.org/10.1038/ng.2822)

Turner NC, Ro J, Andre F, Loi S, Verma S, Iwata H, Harbeck N, Loibl S, Huang Bartlett C, Zhang K, et al. 2015 Palbociclib in hormone-resistant advanced breast cancer. New England Journal of Medicine 373 209–219. (https://doi.org/10.1056/NEJMoai1505270)

Umemita Y, Hiyakka RA, Kokonis JM & Liao S 1996 Human prostate tumor growth in athymic mice: inhibition by androgens and stimulation by finasteride. Prostate 31 11802–11807. (https://doi.org/10.1002/pro.21231-4)

van de Velde CJ, Rea D, Seynaeve C, Putter H, Hasenburg A, Vancranel JM, Paardebens R, Markopoulos C, Heuzey Y, Hille ET, et al. 2013 Adjuvant tamoxifen and exemestane in early breast cancer (TEAM): a randomised phase 3 trial. Lancet 377 321–331. (https://doi.org/10.1016/S0140-6736(13)62112-4)

Vicier C, Dieci MV, Arnedos M, Delaloge S, Viens P & Andre F 2014 Clinical development of mTOR inhibitors in breast cancer. Breast Cancer Research 16 203. (https://doi.org/10.1186/bcr3618)

Waltering KK, Urbanucci A & Visakorpi T 2012 Androgen receptor (AR) aberrations in castration-resistant prostate cancer. Molecular and Cellular Endocrinology 360 38–43. (https://doi.org/10.1016/j.mce.2011.12.019)

Wang ZY & Yin L 2015 Estrogen receptor alpha-36 (ER-alpha36): a new player in human breast cancer. Molecular and Cellular Endocrinology 107 193–206. (https://doi.org/10.1016/j.mce.2015.04.017)

Watson PA, Chen YE, Balbas MD, Wongvipat J, Scooi ND, Viale A, Kim KS & Sawyer LS 2010 Constitutively active androgen receptor splice variants expressed in castration-resistant prostate cancer require full-length androgen receptor. Prostate 107 16759–16765. (https://doi.org/10.1016/j.pms.2014.02.002)

Welshons WV & Jordan VC 2015 Adaptation of estrogen-dependent MCF-7 cells to low estrogen (phenol red-free) culture. European Journal of Cancer and Clinical Oncology 23 1935–1939. (https://doi.org/10.1074/jbc.M111.265124)

Wilsson TM, Henke BR, Moonen THM, Charlison PS, Batchelor KW, Lubahn DB, Moore LB, Oliver BR, Sauls HR, Trantafiliou JA, et al. 1994 3-[1,2-Diphenylbut-1-enyl]phenyl]acrylic acid: a non-steroidal estrogen with functional selectivity for bone over uterus in rats. Journal of Medicinal Chemistry 37 1500–1522. (https://doi.org/10.1021/jm010037a002)

Wolf DM & Jordan VC 1993 A laboratory model to explain the survival advantage observed in patients taking adjuvant tamoxifen therapy. Recent Results in Cancer Research 127 23–33.

Wu YL, Yang X, Ren Z, McDonnell DP, Norris JD, Wilsson TM & Greene GL 2005 Structural basis for an unexpected mode of SERM-mediated ER antagonism. Molecular Cell 18 413–424. (https://doi.org/10.1016/j.molcel.2008.12.004)

Xiong R, Patel HK, Gutgesell LM, Zhao J, Delgado-Rivera L, Pham TND, Zhao H, Carlson K, Martin T, Katzenellenbogen JA, et al. 2016 Selective Human Estrogen Receptor Partial Agonists (ShERPAs) for tamoxifen-resistant breast cancer. Journal of Medicinal Chemistry 59 219–237. (https://doi.org/10.1021/acs.jmedchem.5b01276)

Yang X, Guo Z, Sun F, Li W, Alfano A, Shimelis H, Chen M, Brodie AM, Chen H, Xiao Z, et al. 2011 Novel membrane-associated androgen receptor splice variant potentiates proliferative and survival responses in prostate cancer cells. Journal of Biological Chemistry 286 36152–36160. (https://doi.org/10.1074/jbc.M111.265124)

Yao K, Lee ES, Bentrem DJ, England G, Schafer JJ, O’Regan RM & Jordan VC 2000 Antitumor action of physiological estradiol on tamoxifen-stimulated breast tumors grown in athymic mice. Clinical Cancer Research 6 2078–2086.

Yeh S, Miyamoto H, Shima H & Chang C 1998 From estrogen to androgen receptor: a new pathway for sex hormones in prostate. Prostate 95 5527–5532. (https://doi.org/10.1002/pro.21231-4)

Yu K, Toral-Barza L, Discafani C, Zhang WG, Skotnicki J, Frost P & Gibbons JJ 2001 mTOR, a novel target in breast cancer: the effect of CCI-779, an mTOR inhibitor, in preclinical models of breast cancer. Endocrine-Related Cancer 8 249–258. (https://doi.org/10.1677/erc.0.0080249)
Steroid-induced apoptosis in endocrine cancers

Yue W, Wang JP, Conaway MR, Li Y & Santen RJ 2003 Adaptive hypersensitivity following long-term estrogen deprivation: involvement of multiple signal pathways. *Journal of Steroid Biochemistry and Molecular Biology* **86** 265–274. (https://doi.org/10.1016/S0960-0760(03)00366-2)

Zhang L, Altuwaijri S, Deng F, Chen L, Lai P, Bhanot UK, Korets R, Wenske S, Lilja HG, Chang C, et al. 2009 NF-kappaB regulates androgen receptor expression and prostate cancer growth. *American Journal of Pathology* **175** 489–499. (https://doi.org/10.2353/ajpath.2009.080727)

Zhao XY, Malloy PJ, Krishnan AV, Swami S, Navone NM, Peehl DM & Feldman D 2000 Glucocorticoids can promote androgen-independent growth of prostate cancer cells through a mutated androgen receptor. *Nature Medicine* **6** 703–706. (https://doi.org/10.1038/76287)

Zhou Y, Yau C, Gray JW, Chew K, Dairkee SH, Moore DH, Eppenberger U, Eppenberger-Castori S & Benz CC. 2007 Enhanced NF kappa B and AP-1 transcriptional activity associated with antiestrogen resistant breast cancer. *BMC Cancer* **7** 59. (https://doi.org/10.1186/1471-2407-7-59)

Received in final form 17 November 2017
Accepted 21 November 2017
Accepted preprint published online 21 November 2017

This work is licensed under a Creative Commons Attribution 4.0 International License.