THEMATIC REVIEW

90 YEARS OF PROGESTERONE

Progesterone and progesterone receptors in breast cancer: past, present, future

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Abstract

Progesterone and progesterone receptors (PR) have a storied albeit controversial history in breast cancers. As endocrine therapies for breast cancer progressed through the twentieth century from oophorectomy to antiestrogens, it was recognized in the 1970s that the presence of estrogen receptors (ER) alone could not efficiently predict treatment responses. PR, an estrogen regulated protein, became the first prognostic and predictive marker of response to endocrine therapies. It remains today as the gold standard for predicting the existence of functional, targetable ER in breast malignancies. PRs were subsequently identified as highly structured transcription factors that regulate diverse physiological processes in breast cancer cells. In the early 2000s, the somewhat surprising finding that prolonged use of synthetic progestin-containing menopausal hormone therapies was associated with increased breast cancer incidence raised new questions about the role of PR in ‘tumorigenesis’. Most recently, PR have been linked to expansion of cancer stem cells that are postulated to be the principal cells reactivated in occult or dormant disease. Other studies establish PR as dominant modulators of ER activity. Together, these findings mark PR as bona fide targets for progestin or antiprogestin therapies, yet their diverse actions have confounded that use. Here we summarize the early history of PR in breast cancer; debunk the theory that progesterone causes cancer; discuss recent discoveries that PR regulate cell heterogeneity; attempt to unify theories describing PR as either good or bad actors in tumors; and discuss emerging areas of research that may help explain this enigmatic hormone and receptor.

Key Words:
- progesterone
- progesterone receptor
- progestins
- cancer stem cells
- biomarker

Introduction

Progesterone is a small lipophilic hormone that plays a fundamental role in normal female biology and medicine. In premenopausal women, progesterone is primarily synthesized in a cyclical manner in the ovaries, with additional synthesis in peripheral tissues including adrenal glands, the nervous system and brain (Giatti et al. 2015, Africander & Storbeck 2018). At menopause, circulating progesterone levels decline sharply, but whether local tissue production continues is unknown. The breast is a major target of progesterone, where it regulates development of
the branched ductal epithelium and expansion of milk-secreting alveoli during lactation. Medically, bioidentical progesterone formulations or synthetic compounds termed progestins are taken by women of different ages for reasons that span birth control, to menopausal hormone therapies (MHT), to treatment of Alzheimer’s disease. However, exposure to exogenous progestins is associated with increased breast cancer incidence and/or disease progression. Nevertheless, progestins continue to be tested as treatments or preventatives for breast cancers. Some of the breast effects are clearly paradoxical and much remains to be learned about this hormone.

The major effector of progesterone and the target of progestins are the progesterone receptors (PR). PR are highly structured multi-domain proteins that upon ligand binding transmit their signals primarily through regulation of gene transcription. In humans, there are two main PR isoforms expressed from a single gene located at chromosome 11q22.1 – a 933 amino acid PR-B and a truncated 769 amino acid PR-A transcribed from an internal start-site (Kastner et al. 1990). PR share conserved functional domains with other members of the steroid/nuclear receptor family of transcription factors. These include an N-terminal domain that is highly modified post-translationally and contains transcriptional activation functions, a central DNA binding domain consisting of two cysteine-anchored zinc fingers, and a C-terminal ligand binding domain (Mangelsdorf et al. 1995, Takimoto et al. 2003). The PR gene is activated by estrogens, with both PR-A and PR-B expressed in approximately one-third of luminal epithelial cells of the normal breast, though there is some evidence for PR expression in basal epithelial cells as well (Hilton et al. 2012). The two PR isoforms are co-expressed in breast cancer cells, but often unequally, with a heightened PR-A:PR-B ratio correlating with poor prognosis (Graham et al. 1995, Hopp et al. 2004, Rojas et al. 2017). For the purposes of this review, we focus mainly on the collective activity of PR, recognizing that most PR+ breast cancers contain both PR-A and PR-B in varying ratios.

While early studies focused on PR structure and function, the last decade has seen analyses of PR-regulatory activities and biological end points. The demonstration that PR regulate breast cancer cell heterogeneity coincided with the reemergence of the cancer stem cell (CSC) theory that proposed a hierarchical mechanism by which rare pre-existing cancer cells avoid drug killing and perpetuate the re-population of bulk tumor cells (Reya et al. 2001). CSCs are currently recognized as a plastic state that can be achieved by genetic and/or phenotypic adaptation and can be influenced by the cellular microenvironment and environmental signals (Meacham & Morrison 2013). Multiple groups have described progestin and PR regulation of populations of cells with CSC properties (reviewed in Axlund & Sartorius 2012, Finlay-Schultz & Sartorius 2015, Simoes et al. 2015, Cenciari & Proietti 2019). The consequence of PR regulation of CSCs is unclear. It has been suggested that CSCs contribute to the long-term dormancy of estrogen receptor (ER)+ PR+ breast cancers; accelerate tumor progression upon development of endocrine therapy resistance; or conversely, that they impart cytostasis on estrogen-driven cells. Here we discuss the transition of PR as an important factor in breast cancer prognosis, to its role as a regulator of tumor cell plasticity and heterogeneity, and its future as a target of therapy.

Past: PR as a biomarker and prognostic factor in breast cancer

Hormonal control of breast cancers was first demonstrated in the late nineteenth century when metastatic tumors of patients regressed following ovariectomy (Beaton 1896) (see Fig. 1 for a time-line of major discoveries relevant to progesterone and PR). By the early 1970s, it was known that approximately 30% of tumors were responsive to therapies involving either ablation of endocrine glands or addition of a variety of hormones or their inhibitors (Dao 1972, McGuire et al. 1974). Such tumors were classified generically as ‘hormone responsive’. The major ovarian hormones upon which studies focused were the estrogens, which accumulate in reproductive organs (Glascott & Hoekstra 1959, Jensen & Jacobson 1960). Experimental rat mammary tumors induced by the carcinogen DMBA were found to be estrogen target tissues (King et al. 1966). Development of the MCF-7 human breast cancer cell line at the Michigan Cancer Foundation and the demonstration of ER therein (Soule et al. 1973) laid the foundations for human ER research. Clinically, hormone dependent breast cancers were shown to accumulate more radioactive estrogens than autonomous ones, and this uptake was due to the presence of ER (McGuire 1973, McGuire et al. 1976). Elwood Jensen postulated that ER marked the hormone dependent tumor subset, and indeed, an international workshop convened in 1974 correlated the data from several trials in 380 patients, which showed that, regardless of treatment type, 55–60% of ER+ tumors regressed in response to endocrine therapies, while only 8% of ER− tumors did so (McGuire & Chamness 1973).

Progesterone is of course the other major ovarian hormone. Its importance in experimental mouse mammary
tumors was documented by the early studies of Huggins et al. (Huggins et al. 1962). However, to this day, the issue of whether progesterone is stimulatory or inhibitory in breast disease remains controversial with observations that vary depending on the models used, study of physiological vs pharmacologic doses, use of progesterone vs synthetic progestins, the presence or absence of estrogens or carcinogens, clinical data from MHT, and the like. We discuss this further subsequently. The first conclusive evidence that progesterone bound to PR used the estrogen-primed chick oviduct (O’Malley et al. 1970). In the early 1970s, B. O’Malley and his key collaborators, including M. Sherman, W. Schrader, D. Toft, T. Spelsberg and A. Means, showed in a series of elegant studies and multiple publications that the liganded receptors exist as dimers, compartment into both cytoplasm and nucleus, bind chromatin at specific sites, and regulate transcription (reviewed in Schrader & O’Malley 1978). Similar studies in mammals including human tissues proved to be difficult, however, due to progesterone’s relative low receptor binding affinity, rapid metabolism, and lack of specificity. This problem was solved by the synthesis and tritium labeling of the progestin R5020 at Roussel-Uclaf (Philibert & Raynaud 1974). [3H]R5020 in ligand binding assays (LBA) of human tumor biopsy extracts detected PR when radiolabeled progesterone failed to do so (Horwitz & McGuire 1975b). The availability of MCF-7 cells allowed for the first demonstration that ER and PR can coexist in one tumor, possibly in the same cell (Horwitz et al. 1975).

The clinical utility of these facts was readily apparent. As discussed previously, it was noted in the 1970s that, at best, 50–60% of ER+ tumors respond to endocrine therapies. Response failures were ascribed to flawed ER proteins or to errors in downstream ER signaling or transcription. It was learned in both the chick oviduct and the guinea pig uterus (Freifeld et al. 1974) that estrogen regulates a progesterone ‘receptor’. Horwitz et al. (1975a) reasoned that an ideal marker of hormone responsiveness in ER+ tumors would be a measurable product of estrogen action, and PR filled that need. We postulated that PR would be rare in tumors that lacked or failed ER signaling, but that PR positivity would mark an ER+ tumor capable of regulating at least one end product and would be hormone sensitive. Initial analysis of 50 tumor cytosols by LBA using [3H]R5020 found 0/14 (0%) to be ER−/PR+, but 20/36 (56%) to be ER+/PR+, a number close to the expected responders. We reported response to hormone therapies in nine patients. Objective remissions were restricted to tumors that were ER+PR+. Patients with ER+PR− or ER−PR+ tumors failed to respond. A larger set of 521 random tumors were 7% ER−PR−, 9% ER−PR+ and 74% ER+PR+, with a markedly higher likelihood of response associated with PR positivity (McGuire et al. 1977). Even before official publication of the PR paper (1975a), the White House learned of it in September 1974 when First Lady Betty Ford was diagnosed with breast cancer (Wu 2012, Sept 27). She underwent a radical mastectomy and her tumor was sent to us for analysis. We found it to be exceptionally PR rich (disclosed with Mrs Ford’s permission). In retrospect, she would have been an excellent candidate for minimal surgery and hormone therapies; the standard of care today. PR analysis was
quickly adopted. Since 1975, millions of patient tumor samples have been assessed for ER and PR, which has spared many women extensive mastectomies in favor of lumpectomies and hormone therapies. LBAs have been replaced by simple and reliable immunohistochemical assays, and the predictive value of well-validated PR assays in both the adjuvant setting and for advanced disease has been solidly documented (Osborne 1998, Hammond et al. 2010).

**Progestin drugs and breast ‘tumorigenesis’**

Because of its essential role in controlling human reproduction, bioidentical progesterone formulations and synthetic progestins that bind and activate PR have been developed for more than half a century. The first chemically synthesized progestin was norethindrone in the 1950s (Djerassi 1966), which became the first FDA-approved oral contraceptive in combination with estrogen in the US (1960) and Europe (1961). Today, progestrone or progestins are widely and safely used for contraception, treatment of infertility, endocrine disorders, and menopausal hormone therapy (MHT) with several on the WHO list of essential medicines (progesterone, medroxyprogesterone acetate (MPA), norethisterone). The benefits of progestins are, however, counterbalanced by their negative impact on rates of breast cancer diagnoses, particularly by users of MHT.

It was originally assumed that, for MHT, progestins would counteract any tumor-promoting effects of estrogens in the breast, akin to their protective effects in the uterus (Gambrell et al. 1983). This was despite studies in rodents which suggested otherwise (Huggins 1965, Lydon et al. 1999). For instance, mice given chronic long-term MPA develop mammary tumors with high frequency (Lanari et al. 2009). The protective hypothesis was debunked by two large studies in the US (Women’s Health Initiative) and UK (Million Women Study) that reported a significant increase in breast cancer incidence in women taking combined estrogen plus progestin compared to women taking estrogen alone (Chlebowskiet al. 2010, Beral et al. 2011). A 2019 meta-analysis by the Collaborative Group on Hormonal Factors in Breast Cancer confirmed the increased risk of breast cancer for MHT containing MPA, norethindrone acetate, or levonorgestrol, compared to never users or estrogen-only users (Collaborative Group on Hormonal Factors in Breast 2019). This was especially pronounced for long-term (>10 year) progestin users, who had twice the risk of developing breast cancer. Notably, this meta-analysis did not include bioidentical progesterone formulations, which had either no additional risk or even decreased breast cancer risk (discussed in Piette 2018). This has led to speculation that the androgenic and glucocorticoid activity of MPA and other progestins are responsible for the increased breast cancer risk (Carroll et al. 2017, Piette 2018). Other hypotheses argue that it is the expansion of occult malignant cells, pre-existent in the breasts of some women of menopausal age, that is stimulated by the progestins (Horwitz & Sartorius 2008). In this scenario, progestins promote occult disease or activate dormant tumor cells, while perhaps suppressing established disease. Furthermore, it is important to distinguish between progestins and natural progesterone. Currently these tend to be lumped together leading to the view that progesterone is ‘carcinogenic’ (i.e. cancer causer). It is our opinion that natural progesterone does not ‘cause’ breast cancer but can expand it (see subsequent section). Hence, despite widespread linkage between the terms ‘progestins’ and ‘carcinogenesis’, we suggest that care must be taken with these ideas, as with the term ‘bioidentical’, until solid data are available, in women, differentiating between the natural hormone and any biosynthetic ones.

**PR as therapeutic targets in advanced breast cancers**

In the century following the discovery that oophorectomy slowed progression of breast cancers (Beatson 1896, Boyd 1900), it became clear that estrogens are the main mitogens for about three quarters of tumors. Accordingly, most endocrine therapies in use today target the ER signaling axis in one manner or another. This has evolved from surgical and/or pharmacological blockade of ovarian estrogen production; to development of antiestrogens or ‘selective estrogen receptor modulators’ (SERMs) such as tamoxifen that bind to and alter the activity of ER; to newer agents such as fulvestrant that degrade or downregulate ER (SERDs) (Palmieri et al. 2014). Another approach is to inhibit tissue estrogen production in postmenopausal women by use of aromatase inhibitors (AI) such as anastrozole, exemestane or letrozole. Tamoxifen and other SERMs are now usually reserved for premenopausal women with ER+ disease, for cancer prevention in high risk women, and for patients intolerant of AI. Adjuvant endocrine therapies can be curative or provide long-term stabilization for many patients. Unfortunately, it is estimated that between 10% to more than 40% of women will experience a recurrence depending on initial disease stage and grade, and this risk persists for more than
20 years following successful initial treatment (Pan et al. 2017). Newer inhibitors that target CDK4/6 or mTOR used in combination with endocrine therapies modestly improve overall survival (Turner et al. 2018). Alternative endocrine therapies utilizing progestins, androgens, and glucocorticoids have been tested and used since the 1940s to supplement the estrogen inhibitors (Henderson & Canellos 1980).

The most common progestins used for breast cancer treatment are megestrol acetate (Megace) and MPA. Multiple clinical studies through the 1990s found that they were as effective as tamoxifen at improving progression-free survival of ER+ disease (reviewed in Carroll et al. 2017). Megace can be used at lower doses than MPA with equal or better efficacy and fewer toxicities (reviewed in Santen et al. 1990) and has therefore been the preferred progestin in use for late stage disease refractory to estrogen/ER targeted therapies. Although other targeted agents are now approved for treatment of endocrine refractory late stage breast cancer, progestins remain listed as options in ASCO guidelines (Rugo et al. 2016, Cardoso et al. 2018). Mifepristone (RU486), a compound that binds both PR and glucocorticoid receptors (GR) with both antiprogestin and antiglucocorticoid activity, has also been tested, but it achieved only partial remission in some patients and has unacceptable side effects (reviewed in Santen et al. 1990). Two ongoing window of opportunity trials in the UK and Australia are testing prometrium (micronized progesterone) in combination with either tamoxifen or letrozole in ER+PR+ disease (ISRCTN23662758; ACTRN12618000928213). The goal of these studies is to see if progesterone reduces proliferation below that obtained with ER targeted therapies alone. Onapristone (Apristor), a PR antagonist, is currently in Phase II trials in combination with fulvestrant for women who fail on CDK4/6 inhibitors (Context Therapeutics). Thus, whether targeting PR has efficacy in treating advanced breast cancer remains an open question, with ligands that both activate and deactivate the receptors in clinical use or trials.

Therapeutic targeting of PR in advanced breast cancers has both benefits and limitations. Formulations of natural progesterone, and to a lesser extent progestins, are generally well tolerated. Overall, PRs are expressed to varying degrees in ~75% of ER+ breast cancers at diagnosis (McGuire et al. 1986). However, up to 30% of advanced ER+ breast cancers have loss of heterozygosity at the PR locus, with many of these tumors losing PR expression (Tomlinson et al. 1996). ER+ tumors that have completely lost PR during tamoxifen therapy have worse prognoses (Cui et al. 2005). This suggests that, despite ER-positivity, PR loss is associated with development of resistance. However, threshold PR expression levels for efficient targeting are unknown. Tumors with low (1–10%) ER+ cells do not respond well to endocrine therapies (Yi et al. 2014); those that retain sufficient PR positivity (>10% PR+) may benefit from progestin or antiprogestin therapies. Some advanced breast cancers express high PR levels including those harboring mutations in the ER gene (ESR1), in which ER target genes such as PR are constitutively upregulated (Dustin et al. 2019). Whether PR could be co-targeted in these ER mutant cancers is unknown. There is also speculation that, if PR are retained, they can be an alternative driver in endocrine resistant tumors (Knutson & Lange 2014). Recent studies suggest that progestins could compromise immune surveillance by decreasing expression of interferon stimulated genes (Walter et al. 2017, Goodman et al. 2019). Thus, deciphering the context-dependent actions of PR in breast cancer is the next step in deciding whether to positively or negatively target the receptors.

**PR as drivers of normal breast differentiation and tumor-cell heterogeneity**

**Normal mouse mammary stem cells (MaSCs)**

Normal breast epithelium is maintained by a hierarchy of self-renewing MaSCs giving rise to progenitor cells that produce the bulk of terminally differentiated luminal and basal/myoepithelial cells (Villadsen et al. 2007, Fu et al. 2014). Progesterone is a key hormone controlling MaSC subpopulations. As this has been reviewed previously (Axlund & Sartorius 2012, Hilton et al. 2018), we highlight here the major findings. Two studies in mice showed that progesterone is necessary for regulating the number and function of MaSCs (Asselin-Labat et al. 2010, Joshi et al. 2010). Using flow cytometry with defined markers, Asselin-Labat et al. (2010) observed that, while ovariectomy did not reduce total MaSC numbers, it impaired their ability to repopulate a functional mammary gland upon transplantation. Restoration of fully functional MaSCs required supplementation with both estrogens and progesterone. Joshi et al. (2010) reported that murine MaSCs are located in a specialized niche in the basal epithelium. Their numbers are highest at diestrus when progesterone levels peak and higher in pregnant than in nulliparous animals. Estrogen appears to be necessary to induce PR, and progesterone then stimulates MaSC self-renewal. Since murine MaSCs are ER−PR−, progesterone upregulates them via paracrine factors such as Wnt4 and RANKL secreted from the
luminal PR+ cells, which then drive expansion of the basal MaSCs (Asselin-Labat et al. 2010, Joshi et al. 2010). RANK receptor inhibitors or targeted RANK antibodies impair MaSC function (Gonzalez-Suarez et al. 2010, Schramek et al. 2010). Overall, during the reproductive cycle of mice, progesterone plays a dynamic role in activating adult MaSCs within the mammary stem cell niche.

**Normal human MaSCs**

Hormonal regulation of epithelial cell hierarchy in the normal human breast has been studied in *ex vivo* organoid cultures from reduction mammoplasties. These form lobular structures that retain progenitor cells, plus committed luminal and myoepithelial cells (Graham et al. 2009b). In these models, progesterone increases proliferation indices, total cell number of organoids (Graham et al. 2009a), mammosphere formation, and aldehyde dehydrogenase (ALDH); all measures of stem/progenitor cell activity (Dontu et al. 2003, Ginestier et al. 2007). RANKL is an important PR target gene. In *ex vivo* organoids, well-defined lineage markers show that progesterone increases the levels of RANKL expression in PR+ luminal cells, and this is associated with an increase in ER−PR− progenitor cells (Tanos et al. 2013). RANKL itself is proliferative in the breast (Tanos et al. 2013); its role in regulating human MaSCs, tumorigenesis, and in tumor cell expansion is under extensive study.

Single cell RNA sequencing of normal human breast epithelial cells detects ER and PR transcripts in two luminal epithelial cell types termed ‘secretory’ and ‘hormone responsive’ (Nguyen et al. 2018). Other studies of cells isolated by flow cytometry detect PR transcripts in the basal epithelial cell fraction and in occasional PR+ myoepithelial cells (SMA+p63+) (Hilton et al. 2012, 2018), suggesting that progesterone may also have direct autocrine effects on a subset of PR+ myoepithelial cells (Hilton et al. 2018). Given these heterogeneities, a definitive picture of progesterone’s role in generating the physiological and complex cellular architecture of the normal human breast requires further study.

**Luminal breast CSCs in cell lines.**

The term ‘CSC’ has undergone evolving definitions. It was originally thought that tumors, from their origin, contain small populations of cells that share properties of normal stem cells including quiescence, self-renewal capacity, and the ability to generate differentiated progeny (Reya et al. 2001). Studies in multiple cancer types redefined CSCs as a functional state, that can exist *de novo* and/or be acquired through environmental cues or therapeutic pressure as tumors evolve (Meacham & Morrison 2013). Generically, for breast disease, CSCs were initially defined (Al-Haaj et al. 2003) as having the surface marker signature CD44+CD24−/lowEpCAM+, with increased ALDH activity (Ginestier et al. 2007). It is unclear whether these factors define the luminal ER+PR+ CSCs subtype. It has been postulated that phenotypic adaptation through transcriptional and epigenetic modulation may be the dominant force dictating ER+ tumor heterogeneity and evolution (Patten et al. 2018) and that, as in the normal tissues, hormones are key factors controlling CSC number and function in luminal disease. Liganded PR, especially PR-A (Truong et al. 2019), increase populations of cells with CSC properties and tumoursphere formation capacity (reviewed in Axlund & Sartorius 2012, Finlay-Schultz & Sartorius 2015, Simoes et al. 2015, Cenciarini & Proietti 2019) (Cittelly et al. 2013, Hilton et al. 2014, Finlay-Schultz et al. 2015, Knutson et al. 2017, Truong et al. 2019). Additionally, several genes implicated in CSC number and function are targets of liganded PR, including the pluripotent transcription factor KLF4 (Cittelly et al. 2013), the co-activator FOXO1 (Truong et al. 2019), the transcriptional repressor BCL6 (Sato et al. 2014), and micro-RNAs (miR)-29 and miR141 (Cittelly et al. 2013, Finlay-Schultz et al. 2015). Thus, progesterone and progestins increase populations of self-renewing CSCs in all models tested and the signaling intermediates are being defined since they could serve as therapeutic targets.

**CK5+ luminal CSCs**

Molecular profiling of breast cancers led to their classification into five major subtypes (Perou et al. 2000), including Luminal A (ER and PR rich), Luminal B (trending to lower ER, low or no PR, high proliferation rate), HER2+ER−, normal-like, and ER−PR− basal-like. This has been updated to include additional TNBC subtypes including claudin-low (mesenchymal-like) with the note that basal-like breast cancers can span into HER2+ and ER+ subtypes (Prat & Perou 2011). Interestingly, luminal tumors have relatively low or no CD44+ and ALDH+ CSCs compared to TNBC, leading to speculation that luminal tumors have alternate CSCs. Analyses of luminal breast cancer cell xenografts treated with progestins show increased transcript levels of basal cytokeratins (CK) (CK5, CK6, and CK17) and decreased levels of luminal CKs (CK8, CK18, and CK19) (Sartorius et al. 2005). CK5 is especially
interesting since it is a marker of luminal progenitor cells (Lim et al. 2009) and a well-established indicator of poor prognosis (Malzahn et al. 1998, van de Rijn et al. 2002, Cheang et al. 2008). Of note, progestin-induced breast cancer cells lose ER and PR while gaining CK5. Such cellular heterogeneity is not surprising since, clinically, one-third to one-half of all ER+ breast cancers contain ER−PR−CK5+ cell subpopulations (Horwitz et al. 2008, Haughian et al. 2012, Joensuu et al. 2013). Compared to CK5− cells, CK5+ cells possess all the hallmarks of CSCs including tumor initiation capacity, quiescence, and resistance to chemo- and endocrine therapies (Horwitz et al. 2008, Kabos et al. 2011, Axlund et al. 2013, Sato et al. 2014, Goodman et al. 2016). Further, as CK5+ cells lose hormone responsiveness, they gain expression of mesenchymal transcription factors such as Slug and Twist and increase Wnt and Notch signaling (Haughian et al. 2012). Progestins target the CK5 gene directly by rapidly recruiting PR to the proximal CK5 promoter at two progesterone response elements (Fettig et al. 2017).

Recent evidence suggests that CK5 contributes functionally to the luminal CSC phenotype. Studies using knockdown (shRNA) or knockout (CRISPR-Cas9) of the CK5 found impaired progestin-induced tumorsphere formation (Fettig et al. 2017, McGinn et al. 2020). Chronic hormone-withdrawn luminal cells constitutively upregulate CK5 (Haughian et al. 2012). Such cells were used to identify non-cytoplasmic interactors of CK5 that influence cell phenotype (McGinn et al. 2020). One is β-catenin, a downstream effector of the Wnt signaling factor, an essential component of adherens junctions and an important regulator of normal and malignant stem cells (Reya & Clevers 2005). CK5+ luminal cancer cells, whether progestin-induced or constitutive, have reduced β-catenin and E-cadherin at the cell membrane; factors that influence cell polarity, migration, and invasion in association with poor prognosis. This agrees with the fact that, in breast cancers, invasive leader cells express basal CKs including CK5 (Cheung et al. 2013). CK5 is infrequently mutated in breast cancer (www.cbioportal.org) and unlikely to be a primary driver; however, cells that express CK5 are implicated as the origin of some basal-like breast cancers (Lim et al. 2009), and therefore, functional properties of CK5 could impact tumorigenesis. This work suggests that progestins transition a minor subpopulation of malignant cells from a proliferative but non-aggressive hormone sensitive ER+PR+CK5− state to a more invasive hormone resistant ER−PR−CK5+ state (see Fig. 2). If so, antiprogestins could serve as potent preventatives in moderate to high-risk premenopausal

Figure 2
Hypothesis of progesterone and progestin effects in early and late luminal breast cancer. For simplicity, progesterone and progestins are collectively termed ‘P’, although individual ligands may impart different magnitudes of effects. Left: In localized, micro ER+PR+ disease such as DCIS, P upregulate ER−PR− luminal CSCs (green), potentially by signaling from normal ER+PR+ (blue) or malignant ER+PR+ (red) via paracrine factors. Center: In detectable invasive carcinomas, P upregulate of CSCs promotes tumor invasiveness over proliferation, while estrogens (E) promote proliferation. Right: In established local or metastatic disease, endocrine therapies target estrogen (E)-driven growth. Addition of P suppress E-driven growth but upregulate CSCs. Such tumors may be indolent or enter long-term dormancy but remain capable of reactivation by P or other signals. Created, in part, using Biorender.com.
women; a theory currently under clinical investigation using ulipristal acetate in the UK (BC-APPS1).

**Progesterone, PR, tumor dormancy and metastasis**

In the absence of a chemical carcinogen it is difficult if not impossible to ‘cause’ breast cancer with progesterone. Yet we and others remain puzzled by the clear evidence that women prescribed MHT have a heightened incidence of breast cancer if a progestin is added to the estrogen. What is the progestin doing if not ‘causing’ disease? In the seminal studies of Huggins (Huggins et al. 1962, Huggins 1965) and others (Sivaraman et al. 2001) on effects of progesterone or pregnancy on tumor ‘induction’, animals were always pre-treated with a carcinogen, followed weeks later by endogenous or exogenous estrogen plus progesterone. Tumor development was recorded by palpation. Such studies found that high progestational states decreased the time to tumor formation or increased tumor number. Notably, when progesterone was included as a control without the carcinogen, no tumors developed (Jabara 1967). In hindsight, these studies document the failure of progesterone alone to ‘induce’ palpable disease and were technically incapable of detecting carcinogen-induced micro-disease prior to hormone addition. Instead, we suggest that they point to progesterone as a promoter of pre-established disease caused by the carcinogen and postulate that pre-existing occult, possibly dormant, micro-disease explains the promoting effects of progestins in women prescribed MHT. Thus, progesterone and progestins may differentially impact early and late breast cancer (Fig. 2).

We developed models to test these theories using multicolor fluorescence to track malignant cells in mice (Ogba et al. 2014). We showed that for weeks after hormone dependent ER+PR+CK5– luminal breast cancer cells are injected into ovariectomized mice there is no evidence of disease. However, some apparently disease-free mice harbor a reservoir of luminal CK8/18+ micro-metastases in lymph nodes and other sites that have low or no ER, PR or CK5. If mice are re-exposed to physiological hormones, fulminant disease emerges rapidly. And if the regimen includes progesterone, this is accompanied by extensive upregulation of CK5+ CSC-like cells. Clearly, in these models, hormones trigger expansion of microscopic occult cells into overt disease. Given this, it is not unreasonable to propose that some women of menopausal age unknowingly harbor pre-existent minimal disease, which expands upon hormone supplementation leading to a diagnosis of breast cancer. This scenario also argues strongly against prescribing MHT to breast cancer survivors. It is of considerable interest that it is in such early, clinically hidden cancers, in which progesterone increases CSCs, it also triggers cell migration leading to widespread metastatic dissemination in mouse models of early HER2-driven breast cancer (Hosseini et al. 2016).

**PR modulation of ER action**

The prevalence of ER+PR+ breast cancers increases with age; they constitute >80% of diagnoses in postmenopausal women. In this environment, locally produced estrogens are the major mitogens and the role of endogenous and exogenous progestins remains unclear (Giulianelli et al. 2013, Diep et al. 2015, Carroll et al. 2017). In general, there are two types of studies that either test progesterone and progestins in the absence of estrogens or assess their impact on estrogen mediated growth. Studies in breast cancer cells initially recognized that treatment with the progestin R5020 alone decreases 2D cell growth (Horwitz & Freidenberg 1985). This was later found to be due to inhibition of the cell cycle after a transient round of proliferation (Musgrove et al. 1991, Groshong et al. 1997). Other studies found that progestins increase growth of BT-474 breast cancer xenografts (Liang et al. 2010). Regarding PR isoforms, cells expressing PR-B increase proliferation in response to progestin, and this is attenuated by PR-A expression (Tung et al. 1993). Following short-term treatment of ex vivo breast tumor explants, progestins have variable effects on proliferation (Knutson et al. 2017), confirming that the impact of progestins on breast cancer growth is both tumor and context dependent.

In breast cancer, PR are co-expressed with ER and their activities are intertwined. This was first recognized using simple promoter-reporter constructs, where PR were found to repress ER transcriptional activity (Wen et al. 1994). Other studies found that exogenously expressed PR abolish estrogen-induced growth in MCF-7 cells (Zheng et al. 2005). Various immunoprecipitation assays have demonstrated direct interactions between ER and PR (Ballare et al. 2003, Giulianelli et al. 2012, Daniel et al. 2015, Mohammed et al. 2015), finding the two receptors in multi-protein complexes and/or co-localized in proximity on chromatin. Two more recent studies in breast cancer cell lines discovered that short-term (3–24 h) treatment with progesterone or a progestin reprograms ER DNA binding (Mohammed et al. 2015, Singhal et al. 2016).
Mohammed et al. (2015) found that PR are present at ER target genes in the absence of ligand and act as cofactors. However, in the presence of progesterone, PR redirect ER to a cistrome with decreased occupancy of oncogenic genes coinciding with reduced cell or tumor growth. Singhal et al. (2016) noted that, while there was some coordinate activity of the agonist R5020 and estrogen at target genes, the overall phenotypic effect was a progestin-induced reduction in estrogen-driven cell and tumor growth. Notably, they also found that the selective PR antagonist CDB4124 (telapristone acetate) suppressed estrogen-induced tumor growth. That is, either agonist- or antagonist-ligated PR were additive with tamoxifen. This could partially explain why both PR agonists and antagonists have seen some success in clinical trials. Overall, most studies find that liganded, activated PR attenuate ER activity and have a net repressive effect on estrogen mediated growth. Taken together, the data confirm the long-held notion that progesterone and progestins are schizophrenic with regard to breast disease: progestins are harmful in occult, early, low density lesions where they upregulate stemness and promote cell migration, proliferation and metastases. On the other hand, progesterone and progestins are beneficial in high density palpable tumors where they suppress these activities by directly suppressing ER proliferative effects, at the expense, however, of expanding the CSC reservoir (see Fig. 2).

New ER+PR+ human tumor models

Basic and preclinical studies on PR in breast cancer have relied heavily on several human breast cancer cell lines that were developed in the 1970s including MCF-7, T47D, ZR75, and BT-474 (Brooks et al. 1973, Engel et al. 1978, Lasfargues et al. 1978, Keydar et al. 1979). Among these, T47D cells have traditionally been the go-to cell line for PR research, due primarily to their unique constitutively high PR expression levels independent of exogenous estradiol supplementation (Horwitz et al. 1982), allowing analysis of the autonomous actions of progestins. While numerous other cell-line collections have been subsequently developed, the majority lack ER and/or PR at the protein level (Neve et al. 2006). A great deal of information regarding PR structure and function has been dissected using cell line models. They have also been extremely useful for initial studies of solid tumors since, with estrogen supplementation, ER+PR+ cell lines grow readily as xenografts in immune compromised mice (Clarke 1996). There are also numerous transgenic murine mammary tumor models. However, these are rarely ER+ (with a few models expressing ER/PR only transiently early in tumor development) and are not estrogen dependent (Pfefferle et al. 2013). With improved immune compromised mouse strains such as NOD-scid IL2Rgnull (NSG), a series of patient-derived xenografts (PDX) have been developed by direct transplantation of tumor fragments into mouse mammary glands. More than 500 such PDX are available worldwide (Dobrolecki et al. 2016), but only one-third are ER+ due to intrinsic limitations in generating them. Furthermore, in many ER+ PDX, PR are either not expressed or only expressed in a small cell fraction. Our group has generated several ER+PR+ PDX that are well-suited for study of ER and PR action (Kabos et al. 2012). Direct grafting of cancer cells into milk ducts is an improved method for growing PDX that maintain ER and PR positivity without the requirement for exogenous estrogen supplementation (Sflomos et al. 2016). Breast cancer specimens can also be adapted to patient-derived organoids (PDO) (Sachs et al. 2018) or patient-derived explants, in which tumor tissue is partitioned into a sponge culture and cultured with hormones for up to 72 h (Centenera et al. 2018). Each of these models has limitations including the fact that they are difficult to manipulate genetically. To get around this, we recently generated three ER+ PDX-derived breast cancer cell lines, two of which express PR, which are amenable to manipulations such as viral transduction (Alzubi et al. 2019). Collectively, the expanding repertoire of human breast cancer models is allowing the study of PRs in heterogeneous tumors and will broaden our understanding of their actions.

PR regulation of RNA polymerase III

We have used our ER+PR+ PDX tumor models to define how progestins influence tumor behavior and PR action in heterogeneous disease and find that chronic treatment with estrogen plus either progesterone or MPA slows estrogen-driven tumor growth (Finlay-Schultz et al. 2017), supporting earlier cell line data and reflecting clinical data. Using gene expression analyses, we noted that progesterone or MPA reverse expression of approximately half of estrogen-regulated genes by decreasing upregulated ones and increasing downregulated ones. In contrast, few transcripts were regulated by estrogen and progesterone/MPA in the same direction. We assessed ER chromatin occupancy by ChIP-seq and found that progesterone caused a ~20% loss or gain of estrogen-induced ER binding sites. Thus, akin to data in cell lines, in PDXs, progestins
impact ER gene regulation and attenuate estrogen driven growth. However, the ER chromatin shift is not as striking as in cell lines. This may be due to longer in vivo treatment times, a more complex hormone environment, and greater cellular heterogeneity.

Somewhat surprisingly, by assessing chromatin occupancy, we found that PR were localized at approximately 50% of tRNA genes, unlike ER which bind there only sparsely. We confirmed that progesterin co-treatment decreased unprocessed pre-tRNAs and their corresponding mature tRNAs. Furthermore, rapid IP-mass spectrometry of endogenous protein analysis displayed PR in association with RNA polymerase III (Pol III); the polymerase that transcribes all tRNA genes. Pol III transcription is a key target of the dynamic balance between cell growth and quiescence. Its direct or indirect regulation is necessary for breast cancer targeted therapies such as mTOR inhibitors that push cells into cytostasis. Interestingly, direct repression of Pol III is a survival mechanism for cells undergoing stress, inducing a quiescent state like one that maintains the longevity of normal tissue stem cells. Pol III can even increase organismal lifespan (Moir & Willis 2013, Filer et al. 2017). tRNAs are the most abundant product of Pol III and changes in their levels and type regulates cell phenotype through reduced and/or selective mRNA translation (Frenkel-Morgenstern et al. 2012). Thus, PR regulation of Pol III could simultaneously contribute to both growth inhibition and expansion of CSCs. Furthermore, this mechanism could impact ER+ tumors independent of direct effects on ER transcription.

**Future: where do progesterone and PR go from here?**

It is indisputable that, in luminal breast cancers, estrogens acting through functional ER are the major mitogenic drivers, explaining the success of drugs that target ER at several signaling stages. Therefore, it has long been clinically important to assess for ‘functional ER’, and PR expression, an end point of functional ER signaling, continues to serve that purpose. The issue before us is not just whether PR proteins, but whether functioning progesterone- or progestin-liganded PR proteins, play a role in breast cancers. There is no question that progesterone is a key hormone required for maturation of the normal breast, where it drives paracrine signaling from PR+ cells to PR− MaSCs, which then generate the organ’s complex cellular architecture and function. In short, in the normal breast, progesterone promotes differentiation at the expense of growth. Is it possible that in breast malignancies progesterone’s functions are similar? That progesterone targets stem cells and promotes cellular heterogeneity in cancers like it does in the normal breast? We address questions raised by this hypothesis subsequently.

First, we need to debunk the notion that progesterone ‘causes’ breast cancers. There is considerable experimental and clinical evidence that, alone and at physiological levels, progesterone is incapable of causing breast cancers so that its reputation as a ‘tumorigenic’ or ‘carcinogenic’ hormone is undeserved. It would be useful to have definitive proof of this once and for all and to eliminate use of these terms in reference to progesterone and the breast.

Further, we review data leading us to postulate that progesterone behaves in breast disease like it does in the normal gland – it targets stem cells. But in the case of breast cancers, these are ‘cancer’ stem cells located in pre-existing disease previously induced by other mechanisms, a carcinogen perhaps. Such a role for progesterone in transformed cells is of course highly consequential, and it is our contention that following-up on the myriad possible pathways is where the future lies. From these ideas flow the likelihood that progesterone plays a role in activating dormant disease; in generating tumor-cell heterogeneity; in enhancing aggressiveness of one or more tumor-cell subpopulations; and in promoting tumor-cell dissemination and metastasis. Extensive research is required in human models and the clinic with regard to progesterone’s role in these areas: (1.) Stem cells: What are the molecular markers of luminal CSCs (and are there several such cell types); how do they arise; how are they transformed, induced or activated; what is the role of progesterone therein; how are CSCs metabolically fueled and maintained; how do they receive and send signals; can CSCs or their signals be pharmacologically or immunologically suppressed; what is their role in resistance to hormone therapies? (2.) Aggressiveness: How and what heterogenous cell subpopulation(s) are induced or activated by progesterone; is this at an early or late stage of disease; are there leading-edge cell markers; how are cells targeted to one or more distant organs; can they be pharmacologically or immunologically suppressed at early and/or late stages? (3.) Minimal vs established disease: Can minimal disease be diagnosed; are CSCs identifiable in minimal disease; is tumor dormancy real and can it be sustained; what signals activation of dormant tumor-cells; is there a role for progesterone or antiprogestins therein; is progesterone a growth suppressor or a mitogen in
established cancers; does this differ across disease stages; are progesterone and synthetic progesterins similar or not; do synthetic progestins ‘cause’ cancer; if so, how? As new, human breast cancer models are developed and tested and existing or unimagined new technologies are applied, these questions can be answered. We believe that we are entering an important and exciting period in research on progesterone and PR in breast cancers and their future as targets of therapy.

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

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