Title: Activating ESRI mutations differentially impact the efficacy of ER antagonists

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Abstract

Recent studies have identified somatic \textit{ESR1} mutations in patients with metastatic breast cancer (MBC) and found some of them to promote estrogen-independent activation of the receptor. The degree to which all recurrent mutants can drive estrogen-independent activities and reduced sensitivity to ER antagonists like fulvestrant is not established. In this report, we characterize the spectrum of \textit{ESR1} mutations from over 900 patients. \textit{ESR1} mutations were detected in 10\%, with D538G being the most frequent (36\%), followed by Y537S (14\%). Several novel, activating mutations were also detected (e.g. L469V, V422del, Y537D). While many mutations lead to constitutive activity and reduced sensitivity to ER antagonists, only select mutants such as Y537S caused a magnitude of change associated with fulvestrant resistance \textit{in vivo}. Correspondingly, tumors driven by Y537S, but not D5358G, E380Q or S463P were less effectively inhibited by fulvestrant than more potent and bioavailable antagonists including AZD9496. These data point to a need for antagonists with optimal pharmacokinetic properties to realize clinical efficacy against certain \textit{ESR1} mutants.

Statement of Significance

A diversity of activating \textit{ESR1} mutations exist, only some of which confer resistance to existing ER antagonists that might be overcome by next generation inhibitors such as AZD9496.
Introduction

Therapeutic targeting of estrogen synthesis is a mainstay of therapy for the over 70% of breast cancers that feature estrogen receptor expression. These treatments markedly reduce the risk of recurrence from early stage disease and improve outcomes in those with advanced disease (1, 2).

Despite this efficacy, a significant subset of estrogen receptor positive (ER+) breast tumors ultimately develops resistance to anti-estrogen therapy. Recent work has identified a set of recurrent mutations in the estrogen receptor, ESR1, from patients with hormone-refractory metastatic breast cancer (3-7). These analyses from small populations of patients described mutations within the ligand-binding domain (LBD) of the receptor, with the majority of the mutations at residues Y537 and D538. It was not clear whether a greater repertoire of somatic LBD mutants might be identified by sequencing a larger population of metastatic breast cancers. Recurrent mutations at Y537 and D538 were characterized using biochemical, cellular, and structural analyses and were shown to promote (1) an apo-receptor conformation similar that to that of estradiol-bound receptor, (2) constitutive coactivator binding and transcriptional activity in the absence of estrogen, and (3) hormone-independent proliferation when expressed in hormone-dependent cells (3, 8, 9). Also evident from these studies was the potential for ER antagonists to potently inhibit mutant receptor activities. However, whether existing antagonists had sufficient potency and adequate pharmacokinetic properties to inhibit mutant receptors in vivo and thereby overcome hormone resistant phenotypes has not been clear.

In this study, we analyzed ESR1 DNA sequences from a large series of metastatic breast tumors and found several novel LBD mutations that constitutively activate the receptor
and promote breast cancer phenotypes. We further investigated the ability of ER antagonists to potently inhibit mutant receptor activities. We observed differential sensitivity of the LBD mutants to selective estrogen receptor degraders (SERDs). Among the mutants Y537S was the most constitutively active and required the highest drug concentrations to fully inhibit the receptor. This specific mutant proved to be less effectively antagonized in vivo by fulvestrant, a drug with suboptimal pharmacokinetic properties compared to a more potent and orally bioavailable SERD, AZD9496. Collectively, these data suggest that activating ESR1 LBD mutations differentially impact the efficacy of ER antagonists.
Results

Novel ESR1 LBD mutations in hormone-resistant breast cancer patients

With an expansion of our efforts to analyze mutations present in metastatic breast cancer using next generation sequencing (National Clinical Trials Registry #00897702), we now have a more comprehensive portrait of the diversity and frequency of ESR1 mutations in metastatic breast cancer (MBC) (Fig 1A). In this series, over 929 cases of breast cancer (including ER+, HER2+ and ER- tumors) were analyzed with 95 patients having somatic mutations in ESR1 (Table 1). Somatic mutations were found in the LBD in all but 1 case. Clinically, 85 out of 95 patients with ESR1 mutations had ER+/HER2- metastatic breast cancer, while 10 of them were ER+/HER2+. In terms of treatment in the metastatic setting, 67.4% of the ESR1 mutant patients had prior exposure to an aromatase inhibitor (AI), while only 18.8% of the WT patients had an AI as a treatment for metastatic disease (Supplementary Table 1). Among the metastatic sites with ESR1 mutations detected, liver and bone were the two most frequent while none were detected in brain metastasis biopsies. The most frequent mutations in this series were D538G (n=34), Y537S (n=13), E380Q (n=20), Y537C (n=6), Y537N (n=5), and L536H (n=4). A number of other mutations were also observed at low frequency (n≤2), most of which have not previously been described (Supplementary Table 2). Although these individual mutations are not common, in aggregate they represent 20% of the cases of LBD mutations in ESR1.

The presence of a somatic mutation in the LBD of ESR1 does not necessarily imply constitutive activity and thus we sought to characterize the activity of a set of these infrequent mutants and compare them to the activities of the better-described variants such as Y537S. We generated hemagglutinin (HA)-tagged ESR1 constructs with
mutations introduced by site-directed mutagenesis and determined their activity through luciferase assays performed in breast cancer cell lines MCF7 (ER+) and SKBr3 (ER-). First, to determine if the mutants had the ability to drive estrogen independent transcription in the cellular context in which they are typically found (ER+ breast cancer), we examined the ability of transient expression of mutant ER to promote transcription from an ERE-luciferase construct in MCF7 cells. We found that most of the mutant ERs had elevated ERE-luciferase activity in the absence of estradiol (E2) compared to wild type (WT) ER, although none of them displayed activity higher than Y537S (Fig 1B). There were two notable exceptions, however, in which the S432L and V534E mutants showed no increase in activity in the absence of E2, despite being expressed at relatively similar levels as that of WT (Supplementary Fig 1A). In addition to the augmentation of E2-independent activity, we examined whether any of the mutants had altered response to E2. ER with mutations in amino acid 537 showed little further induction with addition of estradiol, while all the other mutants, including S432L and V534E, showed induction with the addition of E2, as was the case for WT. Because the effect of E2 addition for these mutants could be ascribed to the presence of the endogenous WT ER, we also examined the effect of mutant ER expression in ER- SKBr3 cells (Supplementary Fig 1B). With the exception of Y537S, which is already maximally activated, the other ER mutants could be further activated by addition of E2 in SKBr3 cells. This includes the S432L and V534E mutants, demonstrating that while these mutations are not constitutively active, they are also not inactivating alterations. The E2-independent activities of the ER mutants were not restricted to the luciferase reporter, as demonstrated by the induction of endogenous ER dependent transcripts such as PGR and GREB1 (Fig
1C and Supplementary Fig 1C). As observed with the luciferase assays, most of the mutants could induce transcripts to levels above that from WT in the absence of E2. Once again, the S432L and V534E mutants showed little evidence of constitutive activity in the absence of E2 while the 536 and 537 mutants were the most active. In addition to regulating ER target genes directly through ER binding sites in the genome, ER modulates gene expression indirectly via interaction with other transcription factors, such as the activator protein-1 (AP-1). We thus examined the effect of several ER mutants on the induction of AP-1 target genes, namely CCND1 and MYC and found no significant changes in the levels of these two gene transcripts in comparing WT and mutant ERs (data not shown).

We looked for further evidence of biologic activation of these somatic LBD mutants beyond assays for transcriptional activation. The phosphorylation of Serine 118 and Serine 167 of ER has been correlated with receptor activation in the absence of ligand (10-12). Consistent with the transcriptional assays, all mutants except S432L and V534E showed increased steady state ER S118 and S167 phosphorylation compared to WT in the absence of E2 when expressed in MCF7 cells (Fig. 1D). The addition of E2 suppressed apparent differences between phosphorylation of WT and most of the mutants (Fig. 1D). There were slight differences in the mutants’ expression levels (HA and ER blots) despite our efforts to achieve equal expression. This may be partly due to differential effects of some mutants on receptor stability, which we have demonstrated previously with Y537S and D538G (3). Together, these data demonstrate the constitutive activity of most of the infrequent ESR1 mutations, and also reveal that at least two mutants (V534E and S432L) show no evidence of constitutive activation.
Given their constitutive activities, we hypothesized that these infrequent ER mutants might promote estrogen-independent tumor growth. We therefore generated doxycycline inducible cell lines that express four representative ER mutants (E380Q, S463P, L536R and Y537S) in the estrogen dependent MCF7 model and examined their proliferation in estrogen-deprived medium. The concentrations of doxycycline used to induce comparable levels of ER expression were determined by immunoblot analyses and quantitative PCR (Supplementary Fig 2A-B). We found that this set of mutants behaved similar to Y537S in their ability to promote E2 independent growth (Fig 1E). However, there were some differences observed in the degree of growth stimulation such as the E380Q mutant promoting growth to a lesser degree than the other mutants. Overall, the data point to several infrequent LBD mutants driving higher estrogen-independent receptor activity and thereby potentially promoting resistance to estrogen deprivation therapy.

The data together point to significant differences in the level of estrogen-independent activity of recurrent mutants with the well characterized mutants in the loop between helix 11-12 (Y537S, D538G) being more potent inducers than some of the less frequent mutants (e.g. S463P or E380Q). One hallmark of both ligand-bound ER as well as constitutively active mutants such as Y537S is their ability to induce a conformation that binds coactivator proteins. To characterize whether these less frequent mutants could induce this “agonist conformation” in vitro without estradiol, we performed a time-resolved Förster Resonance Energy Transfer (tr-FRET) assay with SRC3 nuclear receptor domain (NRD) titrated to a fixed amount of mutant ER-LBD (Y537S, D538G, S463P and E380Q) (13, 14). Steroid receptor coactivator 3 (SRC3) is a well-established ER co-
activator that is highly expressed in breast cancer cells and has a high affinity for ligand bound ER (15). As shown in Figure 1F, SRC3 NRD bound to unliganded Y537S with the highest affinity (18.2 nM). The D538G mutant was also able to bind SRC3 in the absence of E2 (137 nM) albeit somewhat less avidly than Y537S. By contrast, no binding was detected for S436P and E380Q LBD whatsoever in the absence of E2. When the ER mutants are pre-saturated with E2, all of them bound to SRC3 NRD with varying affinities. S463P and E380Q showed similar affinities (EC$_{50}$ = 3.9 nM and 3.4 nM) while Y537S and D538G had higher affinities (EC$_{50}$ = 0.3 nM and 1.1 nM). The results of a trypsin digestion FRET assay further demonstrated the differences between the conformations of the mutants. This assay tests accessibility of the region of surrounding K529 to trypsin access and cleavage. All mutants tested provide substantial to major conformational stabilization of this region, but there were significant differences between mutants (Supplementary Table 3). In the absence of ligand (Apo), the t½ of WT-ER is the shortest at 3 minutes, implying that this region of the WT-ER is very accessible to trypsin attack. In contrast, Y537S and S463P demonstrate the greatest stabilization (in the range of 15-25 fold, followed by D538G, E380Q and L536R with less than 5-fold stabilization. In the presence of ligand, marked stabilization is provided by the same two mutations (Y537S and S463P), as well as by D538G compared to WT. Although the degree of stabilization provided by the mutations does not fully correlate with their degree of constitutive activity, it provides evidence that the conformations of the mutants, at least in the region of helix 11, are quite different. Altogether, the data are consistent with mutants inducing constitutive activity through distinctive mechanisms and with Y537S appearing to be the most active.
Efficacy of SERDs against ER mutants

Previous work has suggested that ER antagonists are able to inhibit ER mutants but may be less potent than they are against WT ER (3). Given the differential activities of the LBD mutants, we asked whether different LBD mutants had comparable activity against the pure ER antagonist, fulvestrant (ICI). We examined the effect of selected ER mutants upon the dose dependent inhibition of ER driven transcription and proliferation by fulvestrant in MCF7 cells. Fulvestrant was able to inhibit the activity of all of the mutants; however, it appeared that the Y537S mutant required significantly higher levels of drug (70-fold higher IC$_{99}$ for Y537S compared to WT) to fully inhibit their activities (Fig 2A). In contrast, E380Q and S463P showed only slight resistance to fulvestrant with 2-fold higher IC$_{99}$ as compared to WT (Fig 2B). In terms of cell proliferation, Y537S required the highest concentration of fulvestrant (IC$_{50}$ = 55-fold higher than WT) for complete growth inhibition while several other mutants had modest effects (Y537C/N/D, D538G, S463P) (Fig 2C-D and Supplementary Figure 3A-D). This result is fairly concordant with the data from Figure 2A-B. One exception is that S463P mutant appeared to have a greater effect on fulvestrant sensitivity in the proliferation assay than the transcriptional assay. This is consistent with a greater impact of this mutant in promoting estrogen-independent proliferation than transcription (Fig 1E and 1C). The differential effects of fulvestrant upon ER driven transcription from different mutants were also evident by assessing GREB1 and PGR transcript levels by quantitative PCR (Supplementary Fig 3E-F). Taken together, the data indicate that certain ESR1 mutants can alter fulvestrant sensitivity in vitro.
To test if the partial resistance conferred by ER mutants was a class effect for SERDs, we compared the effects of three other SERDs, namely AZD9496 (16), RU58668 and GDC-0810, on the WT and mutant cell lines. While all three SERDs were capable of inhibiting cell proliferation by all mutants, significantly higher concentrations were consistently required for Y537S (Supplementary Fig 4A-F). These findings corroborate previously published reports where the Y537S mutation led to reduced rates of ligand association and greater resistance to disruption of ligand binding by urea compared to WT (17).

Because SERD potency was tested using a model involving transient expression of the mutants, we sought to confirm the findings regarding Y537S in a system in which the mutation was expressed from its endogenous genomic locus. We therefore generated an *ESR1* knock-in of Y537S into the MCF7 cell line, using CRISPR/Cas9-mediated genome editing. The insertion of Y537S into *ESR1* gene locus was confirmed by ddPCR analysis of the parental cells, pooled cells (cells before single cell sorting) and Y537S CRISPR knock-in cells (clone B3). These data demonstrated that the knock-in cell line is homozygous for the Y537S mutation, having an insignificant number of WT positive droplets detected (Supplementary Fig 5). Cells expressing the Y537S allele also demonstrated enhanced expression of progesterone receptor (PR) protein and proliferation in hormone depleted media compared to parental cells (Fig 2E-F). The potency of fulvestrant and AZD9496 in Y537S expressing cells was examined by assaying cell proliferation over a range of drug doses. As shown in Figure 2G, the growth of MCF7 WT cells was inhibited by fulvestrant and AZD9496 at an IC$_{50}$ concentration of 0.4 nM, whereas in the Y537S expressing cells, fulvestrant and AZD9496 inhibited growth with an IC$_{50}$ of 25 nM (Fig 2H). These reduced sensitivities in the CRISPR
knock-in Y537S cell line are consistent with the results seen with the doxycycline induced overexpression mutant cell lines.

Superior anti-tumor activity on ER mutant expressing xenografts achieved with AZD9496

The poor pharmacokinetic properties of fulvestrant are well documented both in mouse and humans (3, 18). Given the comparable efficacy of AZD9496 and fulvestrant in vitro and the high serum levels of AZD9496 achievable through oral administration, we sought to compare these drugs in xenograft models of ER mutant disease. We examined xenografts from the MCF7 cells engineered to express WT or mutant ER under a doxycycline inducible promoter (3). We have previously demonstrated that the growth of these tumors is entirely dependent on ER signaling and that complete ablation of ER is able to fully block tumor growth. Maximally effective doses of fulvestrant (200mg/kg) twice weekly via subcutaneous injection or AZD9496 (15mg/kg orally once daily) were administered to mice with tumors expressing WT, E380Q, S463P, Y537S, or D538G ER. Fulvestrant fully inhibited the growth of the WT, E380Q and S463P expressing tumors while nearly completely inhibiting the growth of D538G tumors (Fig 3A-E). The Y537S expressing tumors, however, continued to grow in the presence of fulvestrant, albeit more slowly than untreated controls. Whereas fulvestrant was only partially effective at suppressing the Y537S tumors, AZD9496 was able to completely inhibit their growth. The relative resistance of the Y537S tumors to fulvestrant was evident when we compared the sizes of the tumors at the end of the study by plotting them as scatter plots, which showed that the Y537S tumors were significantly larger than WT or D538G.
tumors in the fulvestrant arms (ICI) but not in the AZD9496 arms (Fig 3F-G). By contrast, tumors expressing the alternate Y537 mutations, Y537C/N as well as V422del, responded similarly to WT upon inhibition by either fulvestrant or AZD9496 (Supplementary Fig 6A-D). Similar observations were made when we tested the effect of another bioavailable SERD, GDC-0810, on the growth of ER mutant tumors, suggesting that the improved pharmacokinetic properties of the antagonists aid in efficient targeting of the mutant ER (Supplementary Fig 7A-C).

As further support of the potential for ER antagonists to target most ER mutant tumors, we examined the effect of AZD9496 and fulvestrant on the growth of a patient-derived xenograft (PDX), CTC-174 (19), that expresses a D538G ESR1 mutation (Supplementary Fig 8A-B). Both AZD9496 and fulvestrant were able to slow tumor growth in this model with a slightly enhanced efficacy seen for AZD9496 in this context similar to what was observed for the MCF7-D538G model (Fig 3H and 3C). It should be noted that unlike the MCF7 model, growth of this particular xenograft has never been shown to be fully dependent on ER signaling alone and this may be partially impacted by the PIK3CA N345K (activating mutation) and two ARID1A truncation mutations (E1776* and S705fs) (inactivating mutations) identified in sequencing of this model. Consistent with these results, assessment of ER-driven transcripts in the AZD9496 and vehicle-treated tumors shows marked reduction of ER activity from AZD9496 administration suggesting it was efficacious in blocking the ER-driven component of this disease (Fig 3I). Overall, the data from the two ER mutant models are consistent with the ability of AZD9496 to inhibit mutant ER driven tumor growth across a broader range of mutants than fulvestrant in vivo. In support of this, we examined the drug levels of AZD9496 and fulvestrant in
serum using pharmacokinetic modeling. We found that the average area under the curve (AUC) and Cmax for fulvestrant (AUC = 5.29 μM·h, Cmax = 0.65 μM) were markedly lower than those obtained for AZD9496 (AUC = 218 μM·h, Cmax = 59 μM)). This difference was observed despite dosing fulvestrant at levels approximately 5-fold higher than are administered to patients. Taken together, the data are consistent with fulvestrant and AZD9496 both inhibiting mutant ER activities in vitro but AZD9496 having superior pharmacokinetic properties enabling in vivo efficacy across a broader range of ER mutant models including those driven by Y537S.

Discussion

The two major strategies for therapeutic targeting of hormonal signaling in breast cancer are estrogen deprivation and direct antagonism of the estrogen receptor. Through large clinical investigations, estrogen deprivation has become the clinical standard for both the adjuvant and metastatic setting. Recently however, resistance to estrogen deprivation therapy has been characterized as frequently involving activating mutations in the estrogen receptor (20). Biochemical and structural studies of the most common mutations have shown them to promote an activated conformation in the absence of ligand. This conformation remains permissive for ligand binding, thus leading to the hypothesis that direct antagonism may be a rational strategy for these mutant receptors. However, the ability of different ER antagonists to effectively block all of the different ER mutants that occur in the clinic has not been established. In this study, we characterize the diversity of activating ESR1 mutations that are observed in the clinic and determine conditions needed for ER antagonists to be effective against the mutations.
Initial reports on the existence of somatic *ESR1* mutations in ER+ metastatic breast cancer came from somewhat smaller populations of tumors in the range of 11-76 tumors (3-6). Recurrent alterations were the focal point of the *in vitro* characterizations and it was demonstrated that mutations D538G, Y537S/N/C, and L536R all promoted estrogen-independent activation of the receptor (3-6). In addition, several of these reports assessed the ability of ER antagonists such as 4-hydroxytamoxifen and fulvestrant to block these mutant receptors (3-6). From these data, it appeared that the mutant receptors might be inhibited, albeit at higher drug concentrations (3, 4, 6, 21). Whether all *ESR1* mutants followed the pattern of these mutations in amino acids 536-538 was unknown. In the current study, we report the identification of *ESR1* LBD mutations from a large cohort of patients with metastatic breast cancer. The majority of mutations are in amino acids 538, 537, 380, and 536. However, we detected a number of low frequency novel mutations, which collectively comprised 20% of the mutations detected within this series. An analysis of the different mutations revealed a range of activities. Across a variety of assays surveying *in vitro* conformation, phosphorylation, transcriptional activity, and estrogen-independent proliferation, mutation in Y537S appeared to have the greatest effect. Mutation in neighboring residues 536 and 538 also led to high activity; however, in both cases these were often at or below the level observed with estradiol stimulation. It is also notable that mutation at the same 537 site to C, D and N caused receptor activation, but to a lesser degree than did the S mutant. Hence, the level of ER activation depends both on the site of mutation and the nature of the mutant residue. Beyond these mutations, all of the mutants outside this region show only modest activity in the absence of estradiol. In addition, two somatic mutations (S432L and V534E) showed no
activation in the absence of estrogen and so their role in promoting resistance to aromatase inhibition is not supported by these data. The basis for these differences in basal activation levels are likely to lie in the conformational changes that these mutations induce. Structures of D538G and Y537S ligand binding domains have now been reported and show similarities in the mechanisms whereby they induce ligand independent activation (22). However, even these two structures highlight important differences in hydrogen bonding and side chain packing that may well translate into the apparent differences in coactivator binding affinity and basal transcriptional activity. Alterations such as E380Q or S463P appear likely to promote hormone independence through still other mechanisms as evidenced by the lack of coactivator binding they induce in vitro in the absence of estradiol. These findings support ongoing efforts to characterize structures of all of the different recurrent ER mutants as they likely reveal the different constraints that prevent unliganded ER from activating transcription.

From a translational perspective, the varying activities observed from different ESR1 mutations raise several issues. First, it is not yet clear whether different mutations are more or less able to promote resistance to aromatase inhibitors. The data revealing the presence of ESR1 mutations in hormone-independent tumors and their association with poor outcomes has largely been obtained from small populations that are underpowered to look for differences in outcome due to different mutations. One recent report suggested that patients with D538G and Y537S mutant tumors may have slight differences in survival outcomes, but this again involved numbers too small to be conclusive on this point (20). Our data raise the possibility that some mutations may indeed be more effective in promoting resistance than others. Perhaps of even greater significance,
estrogen receptor antagonists might have differential efficacies as a function of mutation
type and activity, a possibility that we have investigated in our models.

Estrogen receptor antagonists such as fulvestrant appear broadly effective against ER
mutants in vitro, but important differences emerge when comparing the potencies against
individual mutants. Whereas several mutations had a modest effect on fulvestrant
efficacy, Y537S led to major changes in the concentration required to fully inhibit ER
activities. What accounts for the specific differences in drug potency is not yet clear but it
is notable that the Y537S mutant is the most activated in the absence of ligand and also
shows reduced ligand association rates and binding affinities in vitro. Perhaps this points
to a particularly active state of the receptor that might be targeted by a unique
pharmacologic strategy. However, despite this reduced affinity, antagonists such as
fulvestrant can ultimately inhibit ER mutants including Y537S in vitro, albeit at higher
doses. To address whether these differences in potency might have clinical implications
we examined the in vivo effects of these drugs in xenograft models.

Using these models, we found that fulvestrant was capable of fully inhibiting wild type,
E380Q, and S463P ER driven breast cancers. However, Y537S mutants were not fully
inhibited by fulvestrant despite dosing to higher levels than are typically achieved in the
clinic. The tumor model driven by D538G was nearly completely inhibited by fulvestrant
and so appeared to behave more like the E380Q and S463P mutants in vivo. The oral
SERDs, AZD9496 and GDC-0810, were able to completely block growth of WT and all
mutant ER-driven models including those driven by Y537S. These findings are consistent
with clinical observations that the major limitation of fulvestrant is its poor
pharmacokinetic properties. While fulvestrant is highly potent in vitro, receptor
occupancy in vivo is incomplete at the steady state serum concentrations reached with current dosing (18). However, higher peak and steady state levels of AZD9496 and GDC-0810 are achievable despite administering significantly less drug to mice. As several more bioavailable SERDs such as AZD9496 and GDC-0810 are now in early clinical trial testing, the value of higher drug levels can be formally evaluated. Our data indicate the need to include assessment of specific mutations in this evaluation because mutations in amino acids 537 and perhaps also 536 and 538, will likely necessitate higher drug levels to achieve complete inhibition.

Finally, it is notable that the patient-derived xenograft model we analyzed only demonstrated partial tumor growth inhibition with AZD9496. While the data for ESR1 mutations are consistent with these alterations being common and reducing tumor dependence on estrogen, they do not imply that all mutant tumors are exclusively dependent on ER signaling for their growth. Tumor genotyping of ESR1 mutant breast cancers also reveals recurrent alterations in the PI3K/AKT pathway, cyclin D1, and FGF receptors, among others (3). These alterations likely reduce tumor dependence on ER signaling. Such tumors are appropriate candidates for combinations of antiestrogens with inhibitors of PI3K, AKT, CDK4/6, FGFRs that are all in Phase 2/3 testing. Our data suggest that the specific hormonal drug used in such a combination is likely to matter significantly, a point further emphasized by our recent observation that inhibition of growth signals such as PI3K/AKT lead to further activation of and restored dependence upon ER signaling (23). Taken together, our data and the emerging literature suggest that more potent and bioavailable compounds to block ER signaling may play a key role in the management of ER+ metastatic breast cancer.
Methods

Reagents
17β-estradiol (E2) and RU-58668 were from Sigma-Aldrich (St Louis, MO) and Tocris Bioscience (R&D Systems, MN, USA) respectively. All the drugs were dissolved in DMSO. AZD9496 and Fulvestrant were kindly provided by AstraZeneca UK Limited (Cheshire, UK). GDC-0810 was a kind gift from Seragon Pharmaceuticals Inc (San Diego, CA). Hemagglutinin (HA)-tag (Cat. No: 2367/ Clone: 6E2), Progesterone receptor A/B (PR) (Cat. No: 8757S/ Clone: D8Q2J) and anti-β-Actin (Cat. No: 4970S/ Clone: 13E5) antibodies were purchased from Cell Signaling Technology.

Cell Culture
All cell lines were maintained at 37°C and 5% CO2 in humidified atmosphere. SKBr3 cell line was a kind gift from Dr Neal Rosen while MCF7 Tet-On® was obtained from Clontech in February 2013 and MCF7 was from DSMZ in 2012. SKBr3 cells were grown in DMEM/F12 supplemented with 10% FBS, 100 ug/ml penicillin, 100 mg/ml streptomycin, 4mM glutamine while MCF7 Tet-On® were grown in DMEM/F12 supplemented with 10% Tet System Approved FBS (Clontech), 100 ug/ml penicillin, 100 mg/ml streptomycin, 4mM glutamine and 100ug/ml of G418. MCF7 cells were cultured in phenol red-free RPMI-1640 media (Sigma, R7509) supplemented with 10% heat-inactivated fetal calf serum (FCS) or 5% charcoal/dextran-treated serum (CSS) to deplete hormones, 2mmol/L L-glutamine (Corning). All cell lines were tested negative for mycoplasma and authenticated by short-tandem repeat (STR) analysis in 2013.
Generation of Y537S CRISPR knock-in cell lines

MCF7 cells were transfected with a sgRNA CAS9 vector and a donor cassette as a non-digested plasmid at a 2:1 ratio using Fugene (Promega). The gRNA sequence used was ctccagcagcaggtcataga. The donor cassette contained a 800bp and 1kb homology regions for incorporation of the Y537S mutation via homologous directed repair (HDR). Between the homology regions, a Neomycin resistance gene was encoded, which was under the PKG promotor and used for selection of HDR events 48 hours post transfection. After two weeks of selection, single cell clones were generated and screened with ddPCR for evidence of Y537S mutation. The digital droplet PCR was performed (24) using ddPCR primers (cgggttgctctaagtagt and aatgcgatgaagtagagccc) and specific probes (BHQ_cc{C}ctc{tAt}gacc{t}g HEX and BHQ_CC{A}CTC{TCT}GAC{C}TG_FAM). The location of the insertion was confirmed using junction PCR with the following primer pairs,

1 (TTAGATCATGCTGTAGGCCCTG) + 2 (CTGGAACCCATGACC GGAAAG),
3 (GCAGATCCAGGGGGCATTTA) + 4 (GATGTGGAATGTGTGCGAGC),
2 (CTGGAACCCATGACC GGAAAG) 5 (GGATCAATTCTCTAGAGCTCGC).

Tide analysis (25) was used to confirm the frame shift mutation of the 2\textsuperscript{nd} ESR1 allele. Targeted locus amplification (TLA) sequencing (26) confirmed the genotype of the 3 ESR1 alleles (A2942C (Y537S) knock-in, inactivating single base insertion knock-out and inactivating 48bp deletion).
Patient-derived xenografts (PDX)

CTC-714 PDX model was derived from patient circulating tumor cells (CTC) cultures, consented according to the Human Biological Samples Policy and purchased from Conversant Biologics. The CTCs were obtained from a 63-year-old patient with stage IV ER+ breast cancer after 42 days of fulvestrant therapy and 26 days of eribulin therapy.

To generate the PDX, EpCAM+CD44+ cells were suspended in phosphate buffered saline (PBS) mixed with high concentration matrigel (BD Biosciences) at 10 mg/ml and ~ 650 cells were injected into the third mammary fat pad of a NOD/SCID (Cg-Prkdcscid Il2rgtm1Wjl/SzJ) mouse. In tumour transplantation study, 2 × 2 mm pieces of tumour tissue from CTC-derived tumour xenografts were implanted in the mammary fat pad of Beige Nude XID mice. All procedures were performed in accordance with US federal, state and Institutional guidelines in an AAALAC-accredited facility. Tumour growth was calculated weekly by bilateral caliper measurement (length x width) and mice randomised into vehicle or treatment groups with approximate mean start size of 0.2 to 0.4 cm³ for efficacy studies or 0.5 to 0.8 cm³ for PD studies. Mice were dosed once daily by oral gavage or subcutaneous (s.c.) injection for fulvestrant at the times and doses indicated for the duration of the treatment period. Tumour growth inhibition from start of treatment was assessed by comparison of the mean change in tumour volume for the control and treated groups.

Animal studies

6-8 week old nu/nu athymic BALB/c female mice were obtained from Harlan Laboratories, Inc. and maintained in pressurized ventilated caging. All studies were
performed in compliance with institutional guidelines under an IACUC approved protocol (MSKCC#12-10-016). MCF7 inducible HA-ER xenograft tumors were established in nude mice by subcutaneously implanting 0.18 mg sustained release 17β-estradiol pellets with a 10g trocar into one flank followed by injecting $1 \times 10^7$ cells suspended 1:1 (volume) with reconstituted basement membrane (Matrigel, Collaborative Research) on the opposite side 3 days afterwards. When the tumors reached a size of ~200 mm$^3$, the mice bearing tumors from each cell line were randomized into 3 treatment groups, fed with water containing 0.2-0.5 mg/ml of doxycycline (0.2 mg/ml for WT, 1.0 mg/ml for E380Q, 0.1 mg/ml for S463P and 0.5 mg/ml for vector control, Y537S and D538G respectively) and 0.1% sucrose for induction of ER expression, 24 hours before being treated with vehicle, 200mg/kg of Fulvestrant subcutaneously twice a week or 15mg/kg of AZD9496 via gavage once daily. Tumor dimensions were measured with vernier calipers and tumor volumes calculated ($\pi/6 \times $ larger diameter $\times$ (smaller diameter)$^2$). In this study, there was no blinding of the investigator as randomization of animals was done. Based upon our previous work measuring the variability in size and growth of MCF7 xenografts, we estimated 10 mice/group would allow us to detect tumor size differences of >200mm$^3$.

**Sequencing of tumor biopsies in the MSK-IMPACT series**

**Study population:** All the patients were enrolled in an institution-wide IRB approved umbrella protocol allowing us to perform genomic testing on their tumors. Informed consents were obtained from all participating patients. This study was conducted in accordance with the Declaration of Helsinki. Between 4/2014 and 6/2015, 929 patients
with confirmed metastatic breast carcinoma (631 with ER+/HER2- disease) underwent MSK-IMPACT testing (27). Detailed clinical information including treatment exposures, and subsequent clinical outcomes were collected from all patients.

**DNA extraction:** 15-20 unstained 10μm-thick FFPE sections were obtained and micro-dissected to ensure >85% tumor content. DNA was extracted using the QIAamp DNA Micro Kit (Qiagen) and standard protocols. Mononuclear cells from peripheral blood were used to extract patient-matched normal DNA.

**Sequencing:** Deep sequencing of targeted genes were performed utilizing the MSK-IMPACT assay (27). Briefly, MSK-IMPACT is a targeted sequencing assay that involves hybridization of barcoded libraries to custom oligonucleotides (Nimblegen SeqCap) designed to capture all protein-coding exons and select introns of 410 commonly implicated oncogenes, tumor suppressor genes, and members of pathways deemed actionable by targeted therapies. The captured pool was sequenced on an Illumina HiSeq 2500 as 2x100bp paired-end reads, resulting in approximately 500-1000 fold coverage per tumor.

Note: Please refer to the Supplementary Methods for the rest of the methods mentioned in the article.
**Author Contributions**


**Acknowledgements**

The authors thank Kinisha Gala, Allison Smith and Neal Rosen for their insightful comments on the manuscript and Marcello Maresca for the design and generation of the reagents used for the production of the Y537S ESR1 MCF7 cell line. SC and GG are supported by a Department of Defense Breast Cancer Research Project Grant (W81XWH-14-1-0359). SC is also supported by funds from Susan G. Komen and the Breast Cancer Research Foundation. The Marie-Josée and Henry R. Kravis Center for Molecular Oncology as well as NCI Cancer Center Support Grant (CCSG, P30 CA08748) supported genomic sequencing. WT was supported by a MSKCC Translational Research Oncology Training Fellowship made possible by the generous contribution of First Eagle Investment Management.
References


PubMed PMID: 25129690.

Table 1. Clinical characteristics of the tumor biopsies sequenced in the study

<table>
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<tr>
<th>Characteristics</th>
<th>Total N = 929</th>
<th>ESR1 Wild Type N = 834</th>
<th>ESR1 Mutated N = 95</th>
<th>P value</th>
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<tbody>
<tr>
<td><strong>Receptor Subtype</strong></td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>HR+/HER2-</td>
<td>139</td>
<td>546 (86.5%)</td>
<td>85 (13.5%)</td>
<td></td>
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<tr>
<td>HR+/HER2+</td>
<td>126</td>
<td>116 (92.1%)</td>
<td>10 (7.9%)</td>
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</tr>
<tr>
<td>HR-/HER2+</td>
<td>50</td>
<td>50 (100%)</td>
<td>0 (0%)</td>
<td></td>
</tr>
<tr>
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<td>122</td>
<td>122 (100%)</td>
<td>0 (0%)</td>
<td>&lt;0.0001</td>
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<td><strong>Histology</strong></td>
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<tr>
<td>Invasive Ductal</td>
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<td>626 (89.7%)</td>
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<tr>
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<tr>
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<tr>
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<tr>
<td>Liver</td>
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<td>126 (78.8%)</td>
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<tr>
<td>Lymph Node</td>
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<tr>
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<tr>
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<tr>
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<td>0 (0%)</td>
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<tr>
<td>Soft Tissue</td>
<td>25</td>
<td>22 (88%)</td>
<td>3 (12%)</td>
<td></td>
</tr>
<tr>
<td>Ovary</td>
<td>22</td>
<td>21 (95.5%)</td>
<td>1 (4.5%)</td>
<td></td>
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<tr>
<td>Skin</td>
<td>24</td>
<td>21 (87.5%)</td>
<td>3 (12.5%)</td>
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<tr>
<td>Pleura</td>
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<td>20 (87%)</td>
<td>3 (13%)</td>
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<tr>
<td>Other</td>
<td>51</td>
<td>38 (74.5%)</td>
<td>13 (25.5%)</td>
<td>0.0017</td>
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Figure 1. Newly detected \textit{ESR1} mutations exhibit a range of estrogen-independent activities.

(A) Diagram of \textit{ESR1} Ligand Binding Domain with somatic mutations identified from 929 breast tumors analyzed. Height of the circles correlates to the number of cases with that specific mutation. The color codes of the circles are as follow: green for missense mutations, red for truncating mutations (Nonsense, Nonstop, Frameshift deletion, Frameshift insertion, Splice site) and black for in frame mutations. (B) Activation of ER reporter gene. ER+ MCF7 cells were transfected with empty vector, HA-ER\textalpha wild type (WT) or indicated \textit{ESR1} mutation, ERE-luciferase and Renilla luciferase reporter constructs in hormone-depleted medium with 10 nM of E2 added for 24 hours where indicated. Firefly luciferase activity shows increased activity in absence of E2 or presence of E2 for certain mutations. Graphs were plotted with the mean ± SD of three biological replicates. *, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$; ****, $P < 0.0001$. (C) Activation of ER target genes. MCF7 cells were transfected with empty vector, HA-ER\textalpha WT or mutant in hormone-depleted medium and harvested 48 hours post-transfection for qRT-PCR analysis. Bars represent mean ± SD of three technical replicates normalized to actin (ACTB) expression. *, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$; ****, $P < 0.0001$. (D) Activation of ER phosphorylation in MCF7 cells. Expression level of the mutant HA-tagged ERs and their relative phosphorylation status at Serine118 and Serine 167, treated with or without 10 nM E2 for 24 hours by immunoblot analysis with specific antibodies as indicated. (E) Activation of hormone independent cell proliferation. Doxycycline inducible ER mutant receptors (E380Q, S463P, L536R and Y537S) expressing MCF7 cells were seeded in 96-well plates in hormone-depleted medium with or without the
addition of doxycycline and proliferation was assayed using resazurin reagent. Data show sufficiency of these 4 mutants to promote cell growth in the absence of estradiol. Each point in the graph represented mean ± SD of 6 technical replicates. (F) Binding of the SRC3 NRD to Y537S, D538G, E380Q or S463P ERα LBD in the absence or presence of E2. SRC3 was titrated into a fixed amount of ER-LBD-biotin and time-resolved Förster resonance energy transfer (tr-FRET) indicated that only Y537S and D538G were able to recruit SRC3 in the absence of E2 but not E380Q and S463P. LBD, ligand-binding domain.

Figure 2. Efficacy of SERDs against ER mutants.

Inhibition of WT and mutant driven ERE-luciferase (A-B) and proliferation (C-D) by fulvestrant (ICI). Doxycycline-induced WT and mutant ER expressing MCF7 cells were treated with various doses of each antagonist in regular medium, demonstrating that the more active mutants required higher level of antagonists for complete inhibition. Graphs were plotted with the mean ± SD of 2 technical or 6 biological replicates respectively. (Notes: SD values for Figure 3A-B are too low for error bars to be visible in the graphs)

(E) Detection of PR levels in MCF7 Y537S CRISPR knock-in cell by Western blot showed elevated PR levels in the Y537S knock-in cells in comparison to the parental line, indicating the expression of Y537S ER mutant receptors. (F) Proliferation assays of parental and Y537S CRISPR knock-in cell lines performed in hormone-depleted or regular media whereby Y537S knock-in cells showed estrogen-independent cell proliferation. (G-H) Proliferation assays of parental MCF7 and MCF7 Y537S CRISPR knock-in cells treated with various doses of fulvestrant, AZD9496 or 4OHT,
demonstrated significantly higher doses are required to cause growth inhibition of Y537S expressing cells by ER antagonists. Cell confluency was measured using the IncuCyte Zoom standard software over a few days. Graphs were plotted with the mean ± SD of three biological replicates.

Figure 3. AZD9496 demonstrates superior anti-tumor effects on ER mutant expressing xenografts.

Mice bearing MCF7 inducible HA-ER WT (A), Y537S (B), D538G (C), E380Q (D) or S463P (E) tumors were randomly assigned to treatment groups of either 15 mg/kg of AZD9496, daily orally or 200 mg/kg of Fulvestrant twice weekly, s.c. Tumors treated with AZD9496 showed greater growth inhibition as compared to those treated with Fulvestrant. The result was presented as average tumor volume measured for each group ± SD (n = 10 mice/group). *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001. (F-G) Scatter plots of the volumes of tumors expressing WT, Y537S or D538G treated with either fulvestrant (ICI) or AZD9496 taken at the end of the xenograft studies shown in A-C. T-tests comparing the volumes of mutant tumors to that of the WT indicated significant resistance of Y537S mutant tumors to fulvestrant treatment. *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001. (H) Patient-derived xenograft (PDX), CTC-714, with D538G mutation, showed greater growth arrest with 5mg/kg of AZD9496. The result was presented as average tumor volume measured for each group ± SEM. (I) Quantitative PCR detection of various ER target genes of AZD9496 treated CTC-714, the D538G PDX model, showed significant reduction in the transcript levels of GREB1, STC2, TFF1 and PGR when compared to the vehicle, indicating inhibition of ER.
signaling pathway by AZD9496. Graphs were plotted with the mean ± SD of three technical replicates.
Figure 1

A. Diagram of ESR1 mutations and domain locations.

B. ERE-Luciferase assay results for different mutations.

C. PGR expression levels normalized to ACTB.

D. Western blot analysis of ER and pER (S118) with vehicle and 10nM E2 treatment.

E. Fluorescence intensity over time for MCF7 Tet-ON HA-ER stables.

F. Normalized fluorescence intensity as a function of Log of SRC3 concentration for various mutations and treatments.
Figure 2
Figure 3
Activating ESR1 Mutations Differentially Impact the Efficacy of ER Antagonists

Weiyi Toy, Hazel Weir, Pedram Razavi, et al.

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