

The tissue-specific effects of different 17β -estradiol doses reveal the key sensitizing role of AF1 domain in ER α activity

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	Journal Pre-proof
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37	<u>HIGHLIGHTS</u>
38	• Some biological response to E2 displays biphasic dose relationship.
39	• ER α AF1 plays a key sensitizing role in ER α activity.
40	• Differences and even contradictory conclusions could be explain
41	differences of E2 doses.
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ABSTRACT

17β-Estradiol (E2) action can be mediated by the full-length estrogen receptor alpha (ERα66), 59 but also by the AF1 domain-deficient ER α (ER α 46) isoform, but their respective sensitivity 60 to E2 is essentially unknown. To this aim, we first performed a dose response study using 61 subcutaneous home-made pellets mimicking either metestrus, proestrus or a pharmacological 62 63 doses of E2, which resulted in plasma concentrations around 3, 30 and 600 pM, respectively. Analysis of the uterus, vagina and bone after chronic exposure to E2 demonstrated dose-64 dependent effects, with a maximal response reached at the proestrus- dose in wild type mice 65 expressing mainly $ER\alpha66$. In contrast, in transgenic mice harbouring only an $ER\alpha$ deleted in 66 AF1, these effects of E2 were either strongly shifted rightward (10 to 100-fold) and/or 67 attenuated, depending on the tissue studied. Finally, experiments in different cell lines 68 transfected with ER α 66 or 46 also delineated varying profiles of ER α AF1 sensitivity to E2. 69 Altogether, this work emphasizes the importance of dose in the tissue-specific actions of E2 70 and demonstrates the key sensitizing role of AF1 in ER α activity. 71

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- 73 Keywords: Estrogen receptor alpha (ERa), activation function (AF), dose response, tissue-
- 74 specific

1. Introduction

During her lifetime, a woman is exposed to varying levels of estrogens. At puberty, the onset of the menstrual cycle leads to a cyclical production of estrogens with large fluctuations in estrogen levels. Pregnancy is marked by an extensive increase in estrogens levels whereas menopause, the consequence of the cessation of estrogen production by ovaries in women, results in a sharp decrease in the main estrogen 17-β estradiol (E2) levels. These hormonal variations translate into changes not only in sex targets organs involved in sex secondary characters and reproduction, but also in many other non-reproductive tissues. In particular, estrogens play an important role in the growth and maturation of bone as well as in the regulation of bone turnover in adult. The importance of estrogen for skeletal health is well known and represents an important research area. Indeed, estrogen deficiency is a major risk for development of osteoporosis and leads to increased fracture risk, while estrogen treatment reduces this risk (Almeida *et al.*, 2017). Unfortunately, long-term estrogen therapy is also associated with serious side-effects, mainly increased risk of breast cancer and of deep venous thrombosis (Valera *et al.*, 2018).

Mouse models have already contributed to elucidation of the mechanisms of action of estrogens *in vivo* (Guillaume *et al.*, 2017; Hewitt and Korach, 2018; Rooney and van der Meulen, 2017). In general, mice are ovariectomized to eliminate the main source of endogenous estrogens, and to allow studying the effects of doses of exogenous E2 delivered by various devices, such as subcutaneous pellets. A large range of pellets that deliver estrogen from 0.01 mg/pellet (*i.e.*, 0.8 μg/kg/day) to 5 mg/pellet (*i.e.*, 2000 μg/kg/day) are available, and very different doses are used in the literature, often without justification of the dose employed. Furthermore, estrogens have frequently been reported to exert a hormetic or biphasic dose-response relationship, characterized by low-dose stimulation and high-dose inhibition (Strom *et al.*, 2011; Duarte-Guterman *et al.*, 2015). One example of such a

dichotomous effect was reported in ischemic stroke; whereas some studies have shown a neuroprotective effect of estrogen, others have reported neurotoxicity, the differences being attributed to the dose (Strom *et al.*, 2009).

The different responses to E2 are initiated by their binding to the estrogen receptors (ERs), ER α and ER β , which belong to the nuclear receptor superfamily and are structurally organized into 6 functional domains (A to F). The E domain supports hormone binding, which induces conformational changes that are required for ER transcriptional activity through the modulation of two activation functions (AFs), AF1 and AF2, located in the A/B and E domains, respectively. In addition to the full-length 66-kDa (ER α 66) isoform, a 46-kDa ER α -isoform (ER α 46), lacking the N-terminal portion (domains A/B), and thereby AF-1, has been reported to be expressed in various cell types such as human osteoblasts (Denger *et al.*, 2001), macrophages (Murphy *et al.*, 2009), and vascular endothelial cells (Murphy *et al.*, 2009), but also in cancer cells (Chantalat *et al.*, 2016). Mechanisms regulating both the expression of ER α 46 and its functions remain essentially unknown. ER α can be generated by either alternative splicing, proteolysis, or an alternative initiation of translation via an internal ribosome entry site (IRES) (Chantalat *et al.*, 2016).

ER-mediated transcriptional regulation involves either a direct interaction of ER with specific estrogen-responsive elements (EREs) located in enhancers or near the promoter region of target genes, or an indirect mechanism (tethered) via protein/protein interactions with other transcriptional factors (Hamilton *et al.*, 2017). ER α AF1 and ER α AF2 have been shown to play crucial roles in the transcriptional effects of ER α through the recruitment of coactivators. In cultured cells, the respective roles of ER α AF1 and ER α AF2 appear to depend on the state of cell differentiation (Merot *et al.*, 2004). Using mice selectively deficient in ER α AF1 or ER α AF2, we previously demonstrated that ER α AF2 is absolutely required for all nuclear effects of ER α in response to E2 whereas the role of ER α AF1 is highly tissue

126	dependent (Abot et al., 2013; Arnal et al., 2013; Billon-Gales et al., 2009; Borjesson et al.,
127	2011).
128	However, the potential role of the E2 dose on the different biological effects mediated by the

wild-type full length ERα-66 or the AF1/AB domain-deficient ERα has not been evaluated

nor taken into account in most in vivo studies.

In this work, we describe the effect of a 3-week exposure of ovariectomized mice to 3 doses of E2 mimicking metestrus (low dose), proestrus (medium dose) or a suprapharmacological dose on three major tissues: uterus, vagina and bone. This experimental paradigm allowed us to precisely compare tissue responses to these 3 doses. An additional very high dose of E2 was used in transgenic mice harbouring an ER α deleted for AF1 (ER α AF1 0) and in its littermate wild type mice.

2. Material and Methods

2.1 Mice

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All procedures involving experimental animals were performed in accordance with the principles and guidelines established by the National Institute of Medical Research and were approved by the local Animal Care and Use Committee. The investigation conforms to the directive 2010/63/EU of the European parliament. Mice were housed in cages in groups of four to six and kept in a specific pathogen-free and temperature-controlled facility on a 12hour light to dark cycle. Each experimental group included at least six animals. Female C57BL/6J mice were purchased from Charles River Laboratories. ER\alpha AF-1⁰ mice were generated as previously described (Billon-Gales et al., 2009). Mice were anesthetized by intraperitoneal injection of ketamine (25 mg/kg) and xylazine (10 mg/kg), and ovariectomized or sham operated at 4 weeks of age, before puberty, to avoid any endogenous estrogen exposure. After 2 weeks of recovery, mice were implanted subcutaneously with pellets releasing either vehicle (cholesterol) or various amounts of E2 for 21 days. To deliver increasing doses of estradiol over prolonged periods of time, various amounts of E2 (E2758 Sigma-Aldrich) and cholesterol (C3045 Sigma-Aldrich) were thoroughly mixed in powder form and compacted to obtain pellets (20 mg total) that were then implanted subcutaneously into ovariectomized mice as previously described (Hsieh et al., 1998). Low (LD-E2), medium (MD-E2), high (HD-E2) and very high (VHD-E2) doses correspond respectively to weight/weight ratio of E2/cholesterol: 1/20000, 1/2500, 1/250 and 1/25.

2.2 Hormone assays

Plasma E2 levels were assayed by GC/MS, as described previously (Giton *et al.*, 2015) with minor modifications. Saline water / twice charcoal dextran stripped aged female rabbit serum (1/1, v/v) was used as the matrix for calibrators and quality control (QC) standards. Briefly, each sample (serums, calibration standards, quality controls, and blank matrix) was collected

in an 8 ml borosilicate tube. A spiking solution of deuterated steroid internal standard (IS) (50 163 ul containing 10 pg of E2-d4, except for blank matrix) (CDN Isotopes, Inc., Point-Claire, 164 Canada), and 3 ml of 1-chlorobutane were added to each sample. After fast centrifugation, the 165 upper organic phase was collected on conditioned Hypersep SI 500mg SPE minicolumn 166 (Thermo Scientific, Rockwood, USA). The column and adsorbed material were then washed 167 with ethyl acetate / hexane (6 ml: 1/9, v/v). The second fraction containing E2 was eluted 168 using ethyl acetate / hexane (4 ml; 1/1, v/v), then evaporated at 60°C to dryness. E2 was 169 170 derivatized with pentafluorobenzoyl chloride (PFBC) (103772-1G, Sigma-Aldrich, Steinheim, Germany). Final extracts were reconstituted in isooctane, then transferred into conical vials 171 for injection into the GC system (GC-2010 Plus, Shimadzu, Japan) using a 50% 172 phenylmethylpolysiloxane VF-17MS capillary column (20m x 0.15mm, internal diameter, 173 0.15µm film thickness) (Agilent Technologies, Les Ulis, France). A TQ8050 (Shimadzu, 174 175 Japan) triple quadrupole mass spectrometer equipped with a chemical ionization source and operating in Q3 single ion monitoring (SIM) mode was used for detection. The reagent gas for 176 the NCI detection was methane. The GC was performed in pulsed splitless mode with a 1 min 177 pulsed splitless-time. The oven temperature was initially 150°C for 0.50 min, further 178 increased to 305°C at 20°C/min and held at 305°C for 3.60 min, and then to 335°C at 179 30°C/min and held at 335°C for 1.7 min. The injection port and transfer line temperatures 180 were respectively 290 and 280°C. The flow-rate of helium (carrier gas) was maintained 181 constant at 0.96 ml/min. The mass spectrometer CI source temperature was 220°C. The 182 linearity of steroid measurement was confirmed by plotting the ratio of the steroid peak 183 response / internal standard (IS) peak response to the concentration of E2 for each calibration 184 standard. Accuracy, target ions, corresponding deuterated internal control, range of detection, 185 low limit of quantification (LLOQ), and intra & inter assay CVs of the quality control are 186 reported in Supplemental Table 1. 187

2.3 Determination of total plasma cholesterol

- Mice were anesthetized and blood samples were collected from the retro-orbital venous
- 190 plexus. Total plasma cholesterol was assayed using the CHOD-PAD kit (Horiba ABX,
- 191 Montpellier, France).

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2.4 Histological analysis

- 193 Paraffin-embedded transverse sections (4 µm) from formalin-fixed uterine or vaginal
- specimens were stained as previously described with anti–Ki67 (Abot et al., 2013) (RM-9106;
- 195 Thermo-scientific). Sections were examined after digitization using a NanoZoomer Digital
- Pathology[®]. To examine the proliferative effects of each treatment, the ratio of Ki67–positive
- 197 epithelial cell/total cell number from two microscopic fields of measurement at x20
- 198 magnification for each uterine and vaginal section was evaluated. For uterus, the luminal
- epithelial height (LEH) of endometrium is the mean of 10 measurements.

2.5 Peripheral quantitative computed tomography (pQCT):

- 201 Computerized tomography scans were performed using the pQCT XCT RESEARCH M
- 202 (version 4.5B, Norland Stratec, Germany) with a voxel size of 70 μm, as previously described
- 203 (Windahl et al., 1999). To measure trabecular volumetric bone marrow density (BMD), the
- scan was positioned in the distal metaphysis of the femur at a distance corresponding to 3.4%
- of the total femur length in the proximal direction from the distal growth plate. This area
- 206 contains trabecular and cortical bone and the trabecular region was defined as the inner 45%
- of the total cross-sectional area. The cortical bone parameters were analyzed in the mid-
- diaphyseal region of femur (Vidal et al., 2000).

2.6 Cell culture and transfection

- 210 HeLa, HepG2 and MDA-MB231 cell lines were maintained in DMEM (Invitrogen)
- supplemented with 10% fetal calf serum (FCS) (Biowest) and antibiotics (Invitrogen) at 37 °C in
- 212 5% CO₂. Transfections were carried out using jetPEI reagent according to manufacturer's

instructions (Polyplus transfection). One day before transfection, cells were plated in 24-well plates at 50% confluence. One hour prior to transfection, the medium was replaced with phenol red-free DMEM (Invitrogen) containing 2.5% charcoal-stripped FCS (Biowest). Transfection was carried out with 100 ng of the reporter gene C3-LUC, 100 ng of CMV-βGal internal control and 50 ng of expression vectors pCR-ERα or pCR-ERαAF-1⁰ (Merot *et al.*, 2004). Following an incubation overnight, cells were treated for 24 h with ethanol (vehicle control) or increasing concentration of E2. Cells were then harvested and luciferase and β-galactosidase assays were performed as previously described (Merot *et al.*, 2004).

2.7 Western Blot

One μg of pCR-ErαWT or pCR-ERαAF1⁰ plasmid were transcribed and translated using TNT[®] quick system Promega. Total proteins were separated on a 10% SDS/PAGE gel and transferred to a nitrocellulose membrane. Primary antibody against ERα was purchased from Santa Cruz Biotechnology (SP1 ABCAM, 1/500). Revelation was performed using an HRP-conjugated secondary antibody and visualized by enhanced chemoluminescence (ECL) detection according to the manufacturer's instructions (Amersham Biosciences/GE Healthcare), using ChemiDoc Imaging System (Bio-Rad).

2.8 Statistics

Statistics analysis were performed using GraphPad Prism software (version 6, GraphPad Software, Inc,). To test the different doses of E2, a 1-way ANOVA was performed, followed by a Fisher's LSD's multiple comparison post test. To test the respective roles of E2 treatment and genotype, a 2-way ANOVA was performed. When a significant interaction was observed between the 2 factors, the difference between genotype and treatment was evaluated by a Fisher's LSD's multiple comparison post test. *P*<0.05 was considered statistically significant.

3. Results

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(Table 1).

3.1 Dose-response profile to E2 is tissue specific in wild type mice.

We first studied in vivo the tissue-specific response to different doses of long-term E2 administration (3 weeks) on ovariectomized C57Bl/6 mice (Figure 1A). Plasma estradiol measurement allowed us to determine the E2/cholesterol ratio that mimicked either metestrus (low dose, LD-E2 ~1 pg/ml: 3pM) and proestrus (medium dose, MD-E2~10 pg/ml: 30pM) as previously reported (Nilsson et al., 2015), or a pharmacological dose (high dose HD-E2 ~200 pg/ml: 600pM) (Figure 1B). In mice on a chow diet, E2 treatment had no impact on body mass (Figure 1C) or plasma cholesterol content (Figure 1D), regardless of the dose administered. The pharmacological dose led to high level of E2 especially in view of the absence of SHBG in mice. Importantly, in mice receiving the high dose of E2, fraction of E2 was converted into estrone (E1) and became detectable in the plasma (Supplementary Table 2). By contrast, the level of estrone in mice supplemented with LD-E2 or MD-E2 is below the threshold of 10 pg/ml. Wet weight of the uterus represents the classic endpoint for measuring estrogenicity. As expected, ovariectomy led to uterine atrophy compared to intact mice, and all E2 doses increased uterine weight (Figure 2A). The maximal response in terms of weight gain was reached with the MD-E2, with an attenuated response observed using the HD-E2 (Figure 2A). We next evaluated the level of uterine epithelial proliferation using Ki67 staining. Epithelial proliferation in response to E2 was maximal at the medium dose, whereas the pharmacological dose of E2 led to a much lower epithelial proliferation; this represented a much greater decline than that reflected by uterine weight (Figures 2B and C). Accordingly, analysis of the height increase of uterine epithelial cells (LEH) as well as stromal proliferation and density confirmed that the medium dose is required to obtain strongest E2 response

Although the evaluation of morphological changes in the vagina is less used to assess estrogenic exposure in the mouse compared to measurements of uterine parameters, this tissue is another major target for estrogenic action. A maximal response of vagina in terms of weight gain was observed using the medium dose of E2, and it then plateaued with the medium or high dose of E2 (**Figure 2D**). By contrast, a low dose was sufficient to induce maximal epithelial proliferation (**Figures 2E and F**).

We next analyzed dose response to E2 treatment in bone, a major non-reproductive target of estrogens. As expected, ovariectomy significantly altered both cortical (**Figure 3A**) and trabecular (**Figure 3B**) bone compartments, but did not impact bone length (**Figure 3C**). Treatment with the lowest dose of E2 was sufficient to restore cortical thickness and cortical bone mineral content (**Figure 3A**) as well as trabecular bone mineral density (**Figure 3B**). As observed in reproductive tissues, the medium dose of E2 induced maximal tissue response thereafter reaching a plateau, with no additional elevation using the pharmacological dose.

Altogether, these results demonstrated that using pellets designed to release E2 levels similar to those encountered during metestrus (low dose) and proestrus (medium dose), the low dose of E2 stimulated 3 target tissues (*i.e.*, vagina, uterus and bone), whereas the medium dose was necessary to elicit a maximal effect.

3.2 Involvement of ERaAF1 action in mediating E2 response is dose dependent

Our results highlight that response to E2 is dose-dependent, with differences between tissues. In addition, we previously demonstrated that the role of ER α AF-1 is also tissue-specific, with a crucial role played in uterus and trabecular bone, but not in cortical bone (Abot *et al.*, 2013; Borjesson *et al.*, 2011). We first compared the dose response to E2 on an ERE-driven reporter gene (C3-LUC) in cells expressing either the wild type receptor (ER α 66) or the isoform having a deletion of AF1 (ER α AF1⁰ = ER α 46) (**Figure 4A**) that were introduced into three different cell lines by transient transfection (**Figures 4B-D**). Expression vectors encoding

wild type $ER\alpha$ and $ER\alpha AF1^0$ were expressed at similar level after transfection in these 287 different cell types (Supplementary Figure 1). In addition to the previously reported cell-288 specific differences in the transactivation efficiency between both receptors (Merot et al., 289 2004), we show here the need for higher doses of E2 for ERαAF-1⁰ than for ERαWT to 290 induce the activity of the reporter gene. The EC₅₀ was about 10 to 100 fold higher for ERαAF-291 1^0 than for ER α WT, depending of the cell-type (**Figures 4B-D**). In addition, even in higher 292 dose, the maximum response was also much lower using ERαAF-1⁰ (**Figure 4B, C**), except 293 for MDA-MB-231 cells (Figure 4D). 294 We then decided to systematically assess the role of ER α AF-1 in the estrogenic response. To 295 this end, we used ERαAF1⁰ mice expressing a 49-kDa truncated protein lacking the A domain 296 and motifs constituting ERa AF-1 in the B domain (Figure 5A) (Billon-Gales et al., 2009). 297 Since some previous studies have used high doses of estrogens in vivo, until 5 mg E2/ pellet 298 299 for 60 day release (Pedram et al., 2014), we have decided to add a very high dose (VHD-E2: ~2000pg/ml i.e 2048 pg/ml \pm 242) to evaluate the role of AF-1 function in vivo. We 300 previously reported that ERαAF1⁰ mice treated with pellets from Innovative Research (0.1 301 mg E2/pellet: 60 day release) presented a moderate uterine hypertrophy that was essentially 302 due to stromal edema, with minimal epithelial proliferation compared to WT mice (Abot et 303 al., 2013). Here, we confirm that no uterine hypertrophy was observed in ERαAF1⁰ mice in 304 response to LD and MD of E2 (Figure 5B). However, HD-E2 or VHD-E2 induced similar 305 increase of uterine weight in $ER\alpha AF1^0$ and $ER\alpha WT$ mice, demonstrating that the deletion of 306 the ERa AF1 function can be fully compensated by a 10 to 100-fold increase in the E2 dose 307 for some parameters such as uterine weigh (Figure 5B). For several other parameters, such as 308 endometrium epithelial proliferation (Figure 5C) and vagina weight gain (Figure 5D), E2 309 310 action was not only shifted rightward but the magnitude of the response was also decreased, remaining below the full effect observed in control mice, even at the VHD-E2 311

pharmacological dose. In addition, we demonstrate for the first time a role of ER α AF1 in vaginal epithelial proliferation, which as for uterine epithelial proliferation, is highly dependent of E2 dose (**Figure 5E**). Finally, we reconsidered the question of the role of ER\alpha F1 in bone, and, as previously described, we demonstrated that ER\alpha F1 is dispensable for cortical response to E2 regardless of the dose used, as revealed by the absence of interaction between genotype and E2 treatment (Figures 6A and B). It should be however noted a significant effect of genotype due to a smaller cortical thickness and a smaller cortical BMC in ERαAF1⁰ mice, independently on E2. By contrast, ERαAF1 is required for the increase of bone mineral density (BMD) in response to E2 in the trabecular compartment regardless of the dose (Figure 6B). We confirmed that E2 has no impact on bone length in both WT and ER\alphaAF10 mice but interestingly in ovariectomized mice, bone lengths from ERαAF1⁰ were smaller than in WT mice, as previously reported in older ERαAF1⁰ mice (Borjesson *et al.*, 2012) (**Figure 6C**). Altogether, these data definitely demonstrate that ERαAF1 is necessary for the full E2 response in uterus, vagina and the trabecular bone compartment. However, depending on the phenotypic feature, ERaAF1 deficiency can be partially or fully overcome by using high E2 levels.

4. Discussion

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To understand the physiology of estrogens, as well as to optimize the use of female sex hormones in medicine, it is crucial to precisely delineate the sensitivity of the tissues to estrogen levels. It is a common practice to ovariectomize mice to eliminate the main source of endogenous estrogens, and subsequently to administer E2 exogenously. Subcutaneous implantation of slow-release pellets is the most convenient and efficient way to deliver estrogen to mice; it avoids the stress associated with daily subcutaneous injections. In our study, the low dose of E2 (metestrus level) stimulated vagina, uterus and bone tissues in

ovariectomized mice, whereas the medium dose (proestrus level) was necessary to elicit a maximal effect. However, use of a supraphysiological dose (HD-E2) did not confer any further stimulation compared to the medium dose in these tissues. The HD of E2 even induced a lesser response than the MD of E2 in the uterus, but not in bone nor in the vagina. There are numerous examples of estrogen hormesis, *i.e.* lowering activity levels while increasing doses, concerning a wide variety of endpoints, including cerebral (Strom *et al.*, 2011; Strom *et al.*, 2009; Strom *et al.*, 2010), mammary gland differentiation (Vandenberg *et al.*, 2006) and insulin sensitivity (Gonzalez *et al.*, 2002). High concentrations of E2 can also lead to increased levels of metabolites, such as estrone (**Supplementary Table 2**), which can also influence the biological responses through either dependent or independent ER mechanisms (Thomas and Potter, 2013).

Remarkably, despite a significant effect on cortical thickness and BMC as well as on trabecular BMD, E2 even at a very high dose had no impact on bone length in our experimental conditions. This result differs from previous studies that reported a reduced bone length and growth plate width using high dose of E2 in older mice (Borjesson *et al.*, 2012; Borjesson *et al.*, 2010). These apparent discrepancies underline the importance of the age of ovariectomy, age of mouse studied, and timing of E2 supplementation.

E2 effects are mediated by hormone binding to ER α , ER β or GPR30 (Miller *et al.*, 2017). ER α also exists as two isoforms, designated ER α 66 and ER α 46, based on their respective molecular weight, and both isoforms were reported to regulate E2-mediated proliferation in the mouse uterus (Abot *et al.*, 2013). The expression of ER α 46 in the rat uterus was demonstrated even before the use of antibodies against ER α (Faye *et al.*, 1986). Importantly, we recently demonstrated that ER α 46 is frequently and sometimes abundantly expressed in ER α -positive breast tumors, but the mechanisms of its generation as well as pathophysiological significance of its expression are essentially unknown (Chantalat *et al.*,

2016). Furthermore, the specific signalling events associated with this AF1-deficient isoform as well as its interactions with the full length ERα66 remain unclear. Given the complexity of estrogen-mediated actions, which goes far beyond the single-receptor situation on which the linear dose-response model is based, it is not surprising that E2 can elicit hormetic responses. In particular, varied affinities for distinct receptors having different intrinsic activity levels would essentially lead to a "mixture" effect, where the summation of their individual dose-response curves would lead to composite behaviour that could be non-monotonic (Miller *et al.*, 2017). In particular, we previously proposed that ERα46 counteracts ERα66 on E2-induced cell proliferation (Abot *et al.*, 2013; Penot *et al.*, 2005).

The structural difference between ERα66 and ERα46, according to our current knowledge, resides mainly in the absence of AF1 in ER046 (Chantalat et al., 2016). Accordingly, both receptors exhibit different transactivation efficiencies in a cell-specific manner (Merot et al., 2004). It has been proposed that ERα46 could have a repressive role in AF1-permissive contexts through interference with ERα66 binding to DNA (Penot et al., 2005). The present study deepens the comparison between both ERα forms by revealing a major difference in their transcriptional sensitivity to the dose of E2. To reach the maximal level of ligand-dependent transcriptional activity, ER α 46 requires an E2 dose 10 to 100 times higher than does ER α 66. Several studies have shown that N-terminal truncated forms of ER α , including ER\alpha46, have an affinity for E2 which is similar to that found for the wild-type ERα66, indicating that the hormone binding domain is able to function independently of the N-terminal half of the ER (Kumar et al., 1986; Carlson et al., 1997; Lin et al., 2013). However, differences in ERa binding to DNA and in coactivator and/or corepressor recruitment could explain the distinctively lower sensitivity of ERα46 versus ERα66 to the dose of E2. We previously reported that although the qualitative overall recruitment of cofactors is quite similar between ERa46 and ERa66, there is a quantitative differential

modulation with some preference for cofactor binding to ER α 66 over ER α 46 (Chantalat *et al.*, 2016). ER α 46 is also able to recruit transcriptional repressors in a ligand-independent manner (Metivier *et al.*, 2004). In accordance with the specific binding of coregulators, we reported that some genes were specifically regulated by either the ER α 46 or ER α 66 isorforms, demonstrating that the absence of ER α AF1 affects specific transcriptional programs rather than leading to an overall decreased activity (Chantalat *et al.*, 2016).

Using mice harbouring an ER α deleted for AF1 (ER α AF1⁰), we previously demonstrated that ER α AF1 is necessary to induce uterine hypertrophy, suggesting that ER α 46 is not able to mediate E2 responses in uterus (Abot *et al.*, 2013). Here, we demonstrate that with a HD of E2, the increase in uterine weight is similar between ER α AF1⁰ and ER α WT, although an attenuated response is observed in terms of epithelial proliferation. By contrast, physiological LD or MD doses of E2 leading to levels encountered during the estrous cycle were unable to induce a uterine response in ER α AF1⁰ mice in contrast to ER α WT mice. These results demonstrate that the involvement of ER α AF1 in the E2 response is highly dependent on ligand dose, and that different signalling pathways regulate epithelial proliferation and uterine weight gain. Similarly, the response to E2 in vagina and trabecular bone are dependent of ER α AF-1, since this response is not only shifted rightward but also decreased in intensity, suggesting that these tissues can be used as an index of ER α AF-1-dependent estrogenicity.

There is also evidence that amino acid phosphorylation located in the ER α AF-1 domain, regulates numerous aspects of ER α function, including protein stability, hormone binding, receptor dimerization, DNA binding, protein-protein interactions, and transcription activity (Orti *et al.*, 1992; Lannigan, 2003; Weigel and Moore, 2007). In particular, Ser118 has been implicated in ER α mediated transcriptional regulation since a partial or complete

attenuation of E2-induced transcription has been observed in ERαS118A transfected cells (Ali *et al.*, 1993; Duplessis *et al.*, 2011). Appropriate transgenic mouse studies should be developed to determine the importance of ERα Ser118 phosphorylation *in vivo*.

5. Conclusion

We describe here the dose-effect of E2 on sexual targets and bone thanks to a simple, reliable and unexpansive slow release device. Our results demonstrate that the estrogenic effect on tissues is highly dependent on the dose used. Although the precise role of ER α AF1 both in physiology and in pathology remains to be clarified, the present work reveals that ER α AF1 provides an important layer of regulation through the sensitization of the response to estrogens. The role of ER α AF1 can be even more spectacular in the action of some selective ER modulators (SERMs) (Arao and Korach, 2019). For instance, tamoxifen selectively activates ER α AF1, and this activation is required for the beneficial effects of this SERM on arteries (Fontaine *et al.*, 2013) and metabolism (Guillaume *et al.*, 2017), while at the same time it prevents the recurrence of breast cancer. Whereas antagonist activity of tamoxifen is essentially through the ER α AF2 blockade, molecular mechanism of its partial agonist activity that involved ER α AF1 activation is not fully understood (Arao and Korach, 2019). Here, we report that importance of ER α AF1 domain in estrogenic response depend on both considered cell type/ tissues and ligand dose, two parameters which could be determinant in the agonist activity of E2 or SERM that control ER α AF-1-mediated physiological response.

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447	

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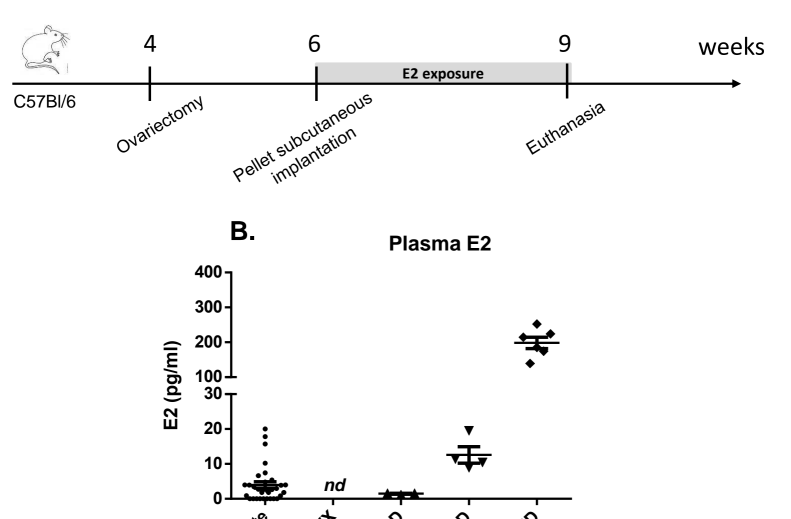
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588	FIGURE LEGEND
589	Figure 1: Plasma estradiol (E2) and cholesterol concentrations in mice implanted with
590	E2/cholesterol pellets.
591	Pellets releasing low dose (LD), medium dose (MD) or high dose (HD) of E2 were implanted
592	subcutaneously into ovariectomized mice for a period of 3 weeks (A). Plasma E2 (B), body
593	weight (C) and cholesterol content (D) of the mice were evaluated.
594	Figure 2: E2 dose response analysis of uterine and vaginal parameters
595	Pellets releasing low dose (LD), medium dose (MD) or high dose (HD) of E2 were implanted
596	subcutaneously into ovariectomized mice for a period of 3 weeks and uterus (A, B, C) and
597	vagina (D, E, F) were analyzed. Uterine (A) and vagina (D) weight. Percentage of uterine (B)
598	and vagina (E) Ki67-positive epithelial cells. Results are expressed as means \pm SEM. To test
599	effect of E2 exposure, one-way ANOVA and a Fisher's LSD's post test were performed (E2
600	exposure versus placebo *P < 0.05, ***P < 0.001, difference in E2 concentration ††† P<0.001
601	n=10/12 mice/group). Representative Ki-67 detection in transverse uterus (C) and vagina (F)
602	sections.
603	Figure 3: E2 dose response analysis of bone parameters
604	Pellets releasing low dose (LD), medium dose (MD) or high dose (HD) of E2 were implanted
605	subcutaneously into ovariectomized mice for a period of 3 weeks and femurs were analyzed.
606	Cortical thickness and cortical bone mineral content (BMC) (A) in the mid-diaphyseal part of
607	femur. Trabecular bone mineral density (BMD) in the distal metaphyseal part of femur (B)
608	and Bone length (C). Results are expressed as means \pm SEM. To test effect of E2 exposure,
609	one-way ANOVA and a Fisher's LSD's post test were performed (E2 exposure versus placebo
610	** $P < 0.01$, *** $P < 0.001$, difference in E2 concentration $^{\dagger\dagger\dagger}P < 0.001$, n=6/7 mice/group).
611	Figure 4: Dose response of E2 transcriptional activity in cultured cells expressing ERa66
612	or ERα46.

613	Schematic representation of the wild-type $ER\alpha 66$ ($ER\alpha AF1^{+/+}$) and $ER\alpha$ ($ER\alpha AF1^0$) isoforms
614	and expression of these isoforms obtained from in vitro translation pCR-ER α WT or pCR-
615	ERαAF1 ⁰ vectors performed with the TNT Coupled Reticulocyte Lysate System (A). HepG2
616	(B), HeLa (C) and MDA-MB231 (D) cells were transiently transfected with the C3-Luc
617	reporter constructs in the presence of pCR-ERαWT, pCR-ERαAF1 ⁰ , or empty pCR vectors.
618	Cells were treated with the indicated dose of E2 or vehicle (Ctrl) for 24 h. Normalized
619	luciferase activities were expressed as fold increase above values measured with empty pCR
620	and vehicle. Data correspond to the mean values \pm SEM of at least three separate transfection
621	experiments. Two-way ANOVA followed by Fisher's LSD's multiple comparison test were
622	$performed \; (ER\alpha AF1^{^{+/+}} \; \textit{versus} \; ER\alpha AF1^0 : \; *P < 0.05, **P < 0.01, ***P < 0.001).$
623	Figure 5: The role of ER α AF1 in uterine and vagina is dependent of E2 dose
623 624	Figure 5: The role of ER α AF1 in uterine and vagina is dependent of E2 dose Schematic representation of the wild-type ER α AF1 $^{+/+}$ and ER α AF1 0 mice constructs (A).
624	Schematic representation of the wild-type $ER\alpha AF1^{+/+}$ and $ER\alpha AF1^0$ mice constructs (A).
624 625	Schematic representation of the wild-type ERαAF1 ^{+/+} and ERαAF1 ⁰ mice constructs (A). Pellets releasing low dose (LD), medium dose (MD), high dose (HD) or very high dose
624 625 626	Schematic representation of the wild-type $ER\alpha AF1^{+/+}$ and $ER\alpha AF1^0$ mice constructs (A). Pellets releasing low dose (LD), medium dose (MD), high dose (HD) or very high dose (VHD) of E2 were implanted subcutaneously into $ER\alpha AF1^0$ ovariectomized mice during 3
624625626627	Schematic representation of the wild-type ERαAF1 ^{+/+} and ERαAF1 ⁰ mice constructs (A). Pellets releasing low dose (LD), medium dose (MD), high dose (HD) or very high dose (VHD) of E2 were implanted subcutaneously into ERαAF1 ⁰ ovariectomized mice during 3 weeks. Uterus (B , C) and vagina (D , E) were analyzed. Uterine (B) and vagina (D) weight.
624 625 626 627 628	Schematic representation of the wild-type ERαAF1 ^{+/+} and ERαAF1 ⁰ mice constructs (A). Pellets releasing low dose (LD), medium dose (MD), high dose (HD) or very high dose (VHD) of E2 were implanted subcutaneously into ERαAF1 ⁰ ovariectomized mice during 3 weeks. Uterus (B , C) and vagina (D , E) were analyzed. Uterine (B) and vagina (D) weight. Percentage of uterine (C) and vagina (E) Ki67-positive epithelial cells. Representative Ki-67
624625626627628629	Schematic representation of the wild-type $ER\alpha AF1^{+/+}$ and $ER\alpha AF1^0$ mice constructs (A). Pellets releasing low dose (LD), medium dose (MD), high dose (HD) or very high dose (VHD) of E2 were implanted subcutaneously into $ER\alpha AF1^0$ ovariectomized mice during 3 weeks. Uterus (B , C) and vagina (D , E) were analyzed. Uterine (B) and vagina (D) weight. Percentage of uterine (C) and vagina (E) Ki67-positive epithelial cells. Representative Ki-67 detection in transverse uterus sections. Results are expressed as means \pm SEM. Two-way

634	Figure 6: E2 dose response analysis of trabecular and cortical bone parameters in wild-
635	type $ER\alpha AF1^{+/+}$ and $ER\alpha AF1^0$ mice.
636	Pellets releasing low dose (LD), medium dose (MD), high dose (HD) or very high dose
637	(VHD) of E2 were implanted subcutaneously to ERαAF1 ⁰ ovariectomized mice for a period
638	of 3 weeks and femurs were analyzed. Cortical thickness and cortical bone mineral content
639	(BMC) in the mid-diaphyseal part of femur. 2-way ANOVA test revealed no significant
640	interaction, the overall effect of the treatment and genotype was analyzed (A). Trabecular
641	bone mineral density (BMD) in the distal metaphyseal part of femur (B) and Bone length (C).
642	Results are expressed as means \pm SEM. Since two-way ANOVA test revealed a significant
643	interaction between the 2 factors (E2 exposure and genotype), a Fisher's LSD's multiple
644	comparison test were performed (ER α AF1 $^{+/+}$ versus ER α AF1 0 : **P < 0.01, ***P < 0.001,
645	n=6/8 mice/group) (B , C).
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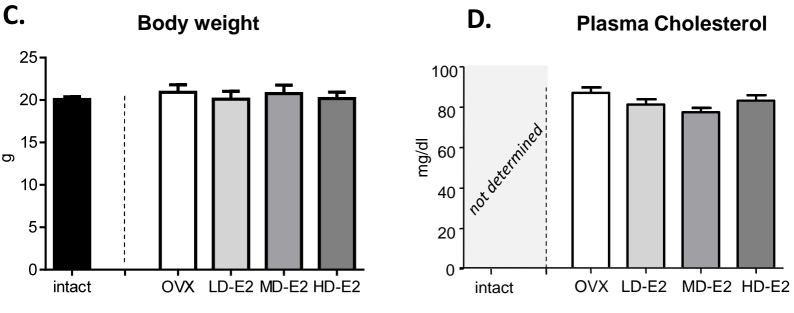


Figure 1

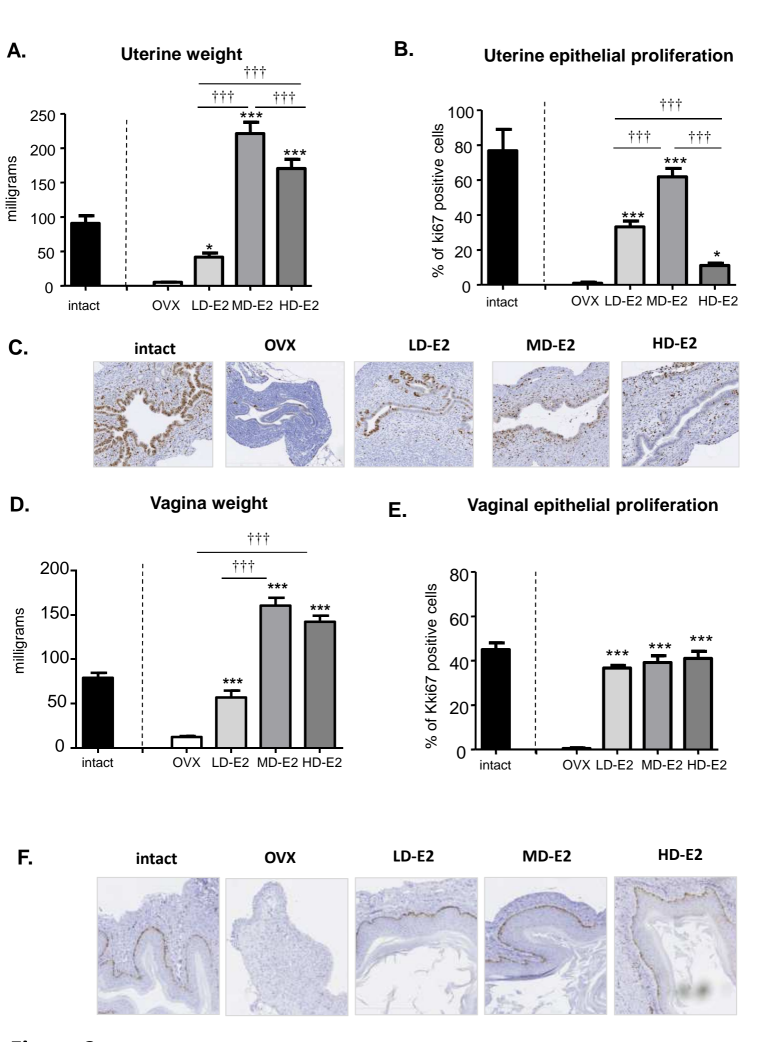
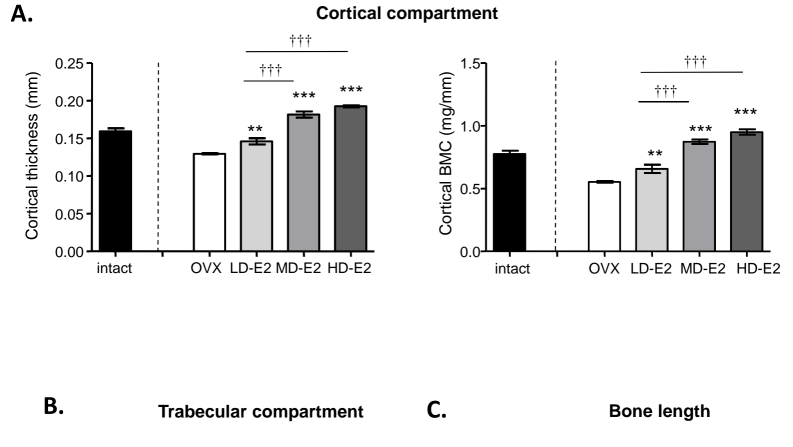


Figure 2



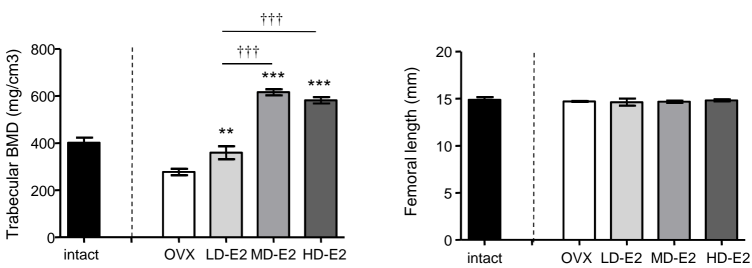


Figure 3

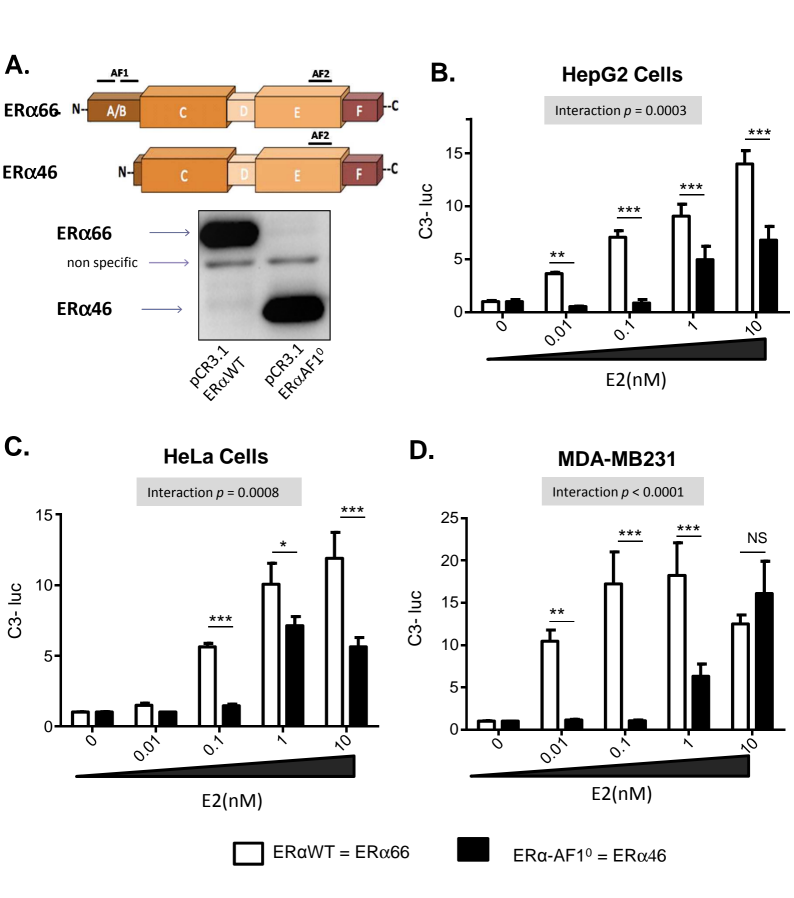
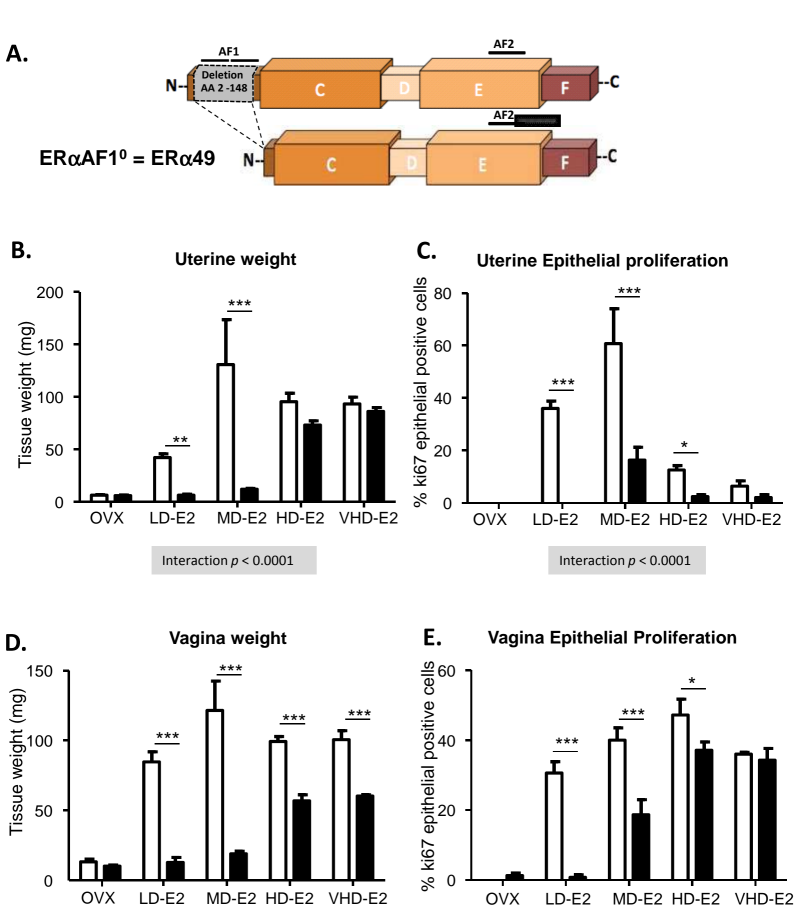


Figure 4



Interaction p = 0.0002

ERα-AF1⁰

Figure 5

Interaction p < 0.0001

 $ER\alpha$ -AF1+/+

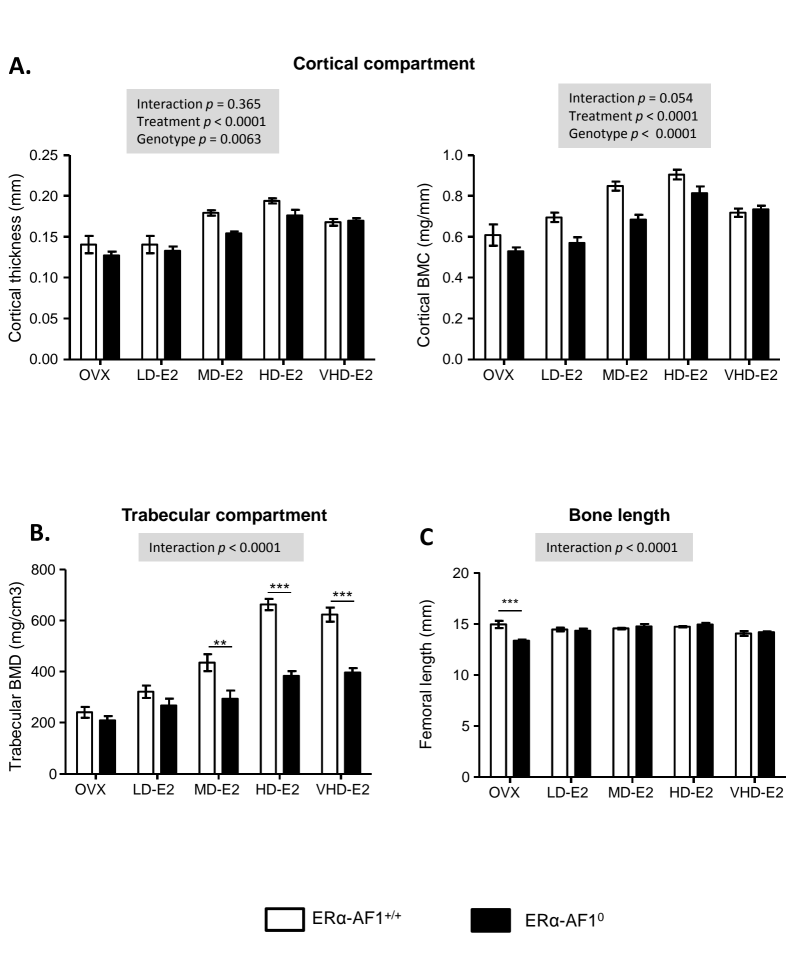


Figure 6

	OVX	LD-E2	MD-E2	HD-E2
Uterine LEH (µm)	12.1 +/- 0.6	20.2 +/- 1.2 **	37.6 +/- 1.9 ***	40.5 +/- 2.1 ***
Uterine Stromal Density	257 +/- 1.14	116.7 +/- 44.6 ***	87 +/- 35.5 ***	72.7 +/- 34.5 ***
Uterine Stromal proliferation (%Ki67positive cells)	0.3 +/- 0.5	8 +/- 5.6	42.1 +/- 11 ***	21.6 +/- 8 ***

Results are expressed as means \pm SEM. To test the respective roles of each treatment, one-way ANOVA and a Fisher's LSD's *post test* were performed (treatment versus placebo **, P < 0.01, ***, P < 0.001)

Table 1: Analysis of uterine parameters in C57BI/6 mice

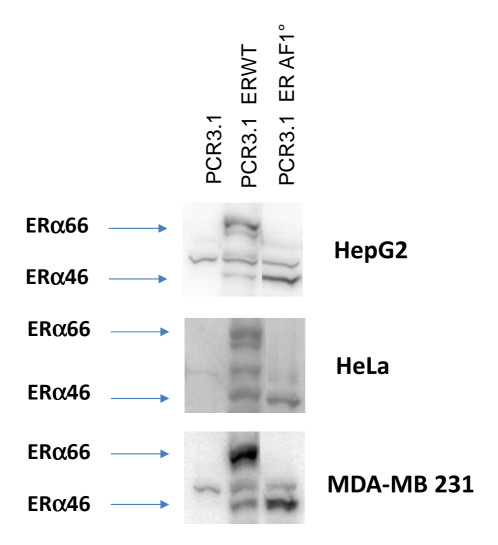
Accucary (%)	Analyte	Target ion analyte /IS (amu)	Range (pg/ml) R ²	Mean (pg/ml) 17 runs Intra- & Inter assay CVs (%)			
				LLOQ	Low QC	Middle QC	High QC
94-107	E2	660/664	0.2- 56.0 0.9994	0.22 17.3 -19.6	2.87 2.9-4.2	6.07 2.6-3.7	12.88 2.3-3.33

LLOQ: low limit of quantification; QC: quality control

	OVX	LD-E2	MD-E2	HD-E2
Estrone (E1)	< 10pg/ml	< 10pg/ml	< 10pg/ml	22,5 +/- 5,4

Results are expressed as means ± SEM.

Supplementary Table 2: GC/MS plasma Estrone (E1) dosage



Supplementary Figure 1: Western blot analysis was performed on an equal amount of cellular extracts prepared from cells transfected with empty vectors (lane 1) expression vectors encoding wild type (lane 2), or $ER\alpha AF1^0$ mutant (lane 3).

HIGHLIGHTS

- Some biological response to E2 displays biphasic dose relationship.
- ER α AF1 plays a key sensitizing role in ER α activity.
- Differences and even contradictory conclusions could be explained by differences of F2 doses