Abnormal Mammary Gland Development and Growth Retardation in Female Mice and MCF7 Breast Cancer Cells Lacking Androgen Receptor

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Abstract
Phenotype analysis of female mice lacking androgen receptor (AR) deficient (AR−/−) indicates that the development of mammary glands is retarded with reduced ductal branching in the pre-pubertal stages, and fewer Cap cells in the terminal end buds, as well as decreased lobuloalveolar development in adult females, and fewer milk-producing alveoli in the lactating glands. The defective development of AR−/− mammary glands involves the defects of insulin-like growth factor I–insulin-like growth factor I receptor and mitogen-activated protein kinase (MAPK) signals as well as estrogen receptor (ER) activity. Similar growth retardation and defects in growth factor–mediated Ras/Raf/MAPK cascade and ER signaling are also found in AR−/− MCF7 breast cancer cells. The restoration assays show that AR NH2-terminal/DNA-binding domain, but not the ligand-binding domain, is essential for normal MAPK function in MCF7 cells, and an AR mutant (R608K), found in male breast cancer, is associated with the excessive activation of MAPK. Together, our data provide the first in vivo evidence showing that AR-mediated MAPK and ER activation may play important roles for mammary gland development and MCF7 breast cancer cell proliferation.

Key words: androgen receptor • knockout mice • mammary gland • breast cancer • MAPK

Introduction
The roles of androgen receptor (AR; reference 1), a transcripational factor that belongs to the nuclear receptor super-family (2), in female organs remains unclear. The lack of clarity could be due to the lack of in vivo models because male mice lacking functional AR are infertile; thus, they are unable to generate female AR knockout (AR−/−) offspring. Epidemiological analyses indicate some positive correlation between testosterone concentration and breast cancer incidence, although it is debatable that testosterone effects on breast cancer progression could also result from conversion to 17β-estradiol (E2) via aromatization in peripheral tissues (3, 4). However, other papers also suggested that androgens could negatively regulate the growth of mammary epithelial and breast cancer cells (5–7). AR is expressed in normal breast cells, and up to 85% of breast tumors are AR positive (8–10). Also, 25–82% of metastatic breast tumors, which are estrogen receptor (ER) and progesterone (P) receptor (PR) negative, still express a significant amount of AR.

Abbreviations used in this paper: AR, androgen receptor; BrdU, bromodeoxyuridine; CDS, charcoal dextran–stripped; DBD, DNA-binding domain; dprm, deletion of the proline-rich motif; E2, 17β-estradiol; Efp, estrogen-responsive finger protein; EGF, epidermal growth factor; ER, estrogen receptor; GFP, green fluorescent protein; HGF, hepatocyte growth factor; HI, heat-inactivated; HRG, heregulin; IGF-I, insulin-like growth factor I; LBD, ligand-binding domain; Luc, luciferase; MAPK, mitogen-activated protein kinase; np, natural promoter–driven; P, progesterone; PR, P receptor; siRNA, small interfering RNA; TBS, Tris-buffered saline; TEB, terminal end bud; UTR, untranslated region.
However, the mechanisms by which AR influences breast cancer progression and mammary gland development remain unclear. To answer these long-term puzzles and to dissect the molecular mechanism of AR in breast cells, we applied different strategies to knock out the AR gene in female mice and breast cancer MCF7 cells. Our results demonstrate that AR plays an important role in the mammary gland development and breast cancer cell growth.

Materials and Methods

**Generation of Female AR<sup>−/−</sup> Mice.** Construction of targeting vectors and generation of the chimera founder mice have been described previously (12). The strain of the mosaic founder was C57BL/6-129/SEVE. β-Actin is a housekeeping gene and is universally expressed in every tissue; therefore, the β-actin promoter-driven Cre (ACTB-Cre) will express and delete the floxed AR fragment in all the cells. The mating strategy is illustrated in brief in Fig. 1. The female AR<sup>−/−</sup> mice were genotyped by PCR, rather than by Southern blot analysis, as described in the figure legend and previously (12).

**Animal, Tissue Collection, and Real-time RT-PCR.** All animal experimentation was conducted in accordance with the standards of humane animal care. Tissues were fixed in buffered neutral formalin for 24 h. Fresh mammary gland tissues were frozen and stored in liquid nitrogen before RNA extraction. 3 µg of total RNA was reverse transcribed and subjected to real-time PCR using iCycle (Bio-Rad Laboratories), and the formulas used were described previously (13). Primer pair sequences were designed by Beacon Designer II software (Bio-Rad Laboratories). Primer pair sequences used for studying gene expression change are as follows: mouse PR, 5'-TGGCCTCAG-3' and 5'-GGTCTGCCTGTTCTGTCGTC-3'; and human Bcl-2, 5'-ACATCTGCCCTCTCTCGTG-3' and 5'-GGTGTGTTGTTGTTCTGTGTC-3'.

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**Whole-mount Staining of Mammary Glands.** Whole mammary glands were spread on glass slides, fixed overnight in Carnoy's solution (Ethanol:CHCl<sub>3</sub>/acetic acid, 6:3:1), sequentially rehydrated by 100, 95, 70, and 50% ethanol and tap water, and stained in carmine red solution overnight until the whole gland became red. After staining, the tissue slides were dehydrated, cleared with xylene, and mounted.

**Immunohistochemistry and Bromodeoxyuridine (BrdU) Staining.** Mammary glands were fixed overnight in buffer neutral formalin (VWR Scientific Products) at room temperature. The tissues were dehydrated by passing through 70, 85, 95, and 100% ethanol, cleared in xylene and 1:1 xylene/paraffin for 45 min, and embedded in paraffin. Tissue sections were cut at a 5–7-µm thickness and mounted onto Probe-On Plus charged slides (Fisher Scientific).

For immunohistochemistry, sections were heated at 55°C for at least 2 h, deparaffinized in xylene, rehydrated, and washed in Tris-buffered saline (TBS), pH 8.0. For antigen retrieval, slides were microwaved in 0.01 M sodium citrate, pH 6.0, immersed with 1% hydrogen peroxide in methanol for 30 min, and blocked with 20% normal goat serum in TBS for 60 min. After washing with PBS, sections were incubated for 90 min in different antibodies diluted 1:200 to 1:500 in TBS containing 1% BSA, followed by goat anti-rabbit biotinylated secondary antibody diluted 1:300 in TBS containing 1% BSA. Sections were incubated with avidin-biotin–peroxidase complex solution for 30 min, followed by development with diaminobenzidine substrate kit (Vector Laboratories) for 5 min. Slides were counterstained with hematoxylin for 30 s, dehydrated, cleaned in xylene, and mounted. Primary antibody was replaced with normal rabbit IgG or 1% BSA in TBS for negative controls. Both of anti-phospho–mitogen-activated protein kinase (MAPK) and total MAPK antibodies were obtained from Cell Signaling Technology. BrdU labeling reagents and BrdU staining kit were purchased from Zymed Laboratories.

**Steroid Hormone RIA.** For characterization of hormonal profiles, sera were collected by the intracardiac method from AR<sup>−/−</sup> and AR<sup>+/+</sup> mice under ketamine and xylazine anesthesia (Sigma-Aldrich). Concentrations of E2 and P were determined using Coat-a-Count kits (Diagnostic Instruments).

**Construction of the AR Targeting Vector and Generation of AR<sup>−/−</sup> MCF7 Cells.** The targeting vector for generating AR<sup>−/−</sup> MCF7 cells was constructed by replacing the Smal–KpnI segments within the AR exon 1 with a promoterless neomycin cassette and inserting two flanking sequences, 5'- extending 1.1 kb into the human AR 5' untranslated region (UTR) and 3'- extending 6.2 kb into the AR intron 1, on a pGem-T easy vector (Promega). This promoterless neomycin cassette, containing a termination codon and a polyadenylation signal, was inserted in frame with AR, ATG start codon. The flanking homologous sequences were generated by PCR using the genomic DNA from human LN Cap cells as a template. For generation of AR<sup>−/−</sup> MCF7 cells, parental MCF7 cells were transfected with the AatII-linearized AR targeting vector using SuperFect (QIAGEN) and selected with 400 µg/ml neomycin. The genotypes of surviving clones were screened by Southern blot analyses. The homozygous clones (AR<sup>++/−</sup>) were picked up and subjected to the second gene targeting experiment using the same targeting vector. Clones were selected with a higher concentration of neomycin (1.25 mg/ml). The genotypes of surviving clones were screened again by Southern blot analyses.

**Plasmids.** GAL4-Elk1, constitutively activated MEK1 (MEK-CA), Ras (Ras-CA), Raf (Raf-CA), dominant-negative Ras (Ras-DN) and Raf (Raf-DN), and MEK phosphatase CL-100 plasmids were provided by K.-L. Guan (University of Michigan Medical School, Ann Arbor, MI; reference 14). Constitutively activated Rac plasmid was a gift from J. Chernoff (Fox Chase Cancer Center, Philadelphia, PA; reference 15). The reporter genes (ARE)-4-luciferase (Luc), ERE-Luc, and pG5-Luc used for monitoring AR, ER, and GAL4-Elk transactivation, respectively, were described previously (16). FLAG-tagged AR deletion mutant expression plasmids were constructed by inserting PCR-generated AR cDNA fragments into pcDNA3 vector (Invitrogen) containing FLAG tag. The natural promoter–driven (np)-AR plasmid was constructed by inserting 3.6 kb hAR promoter, the entire hAR 5’-UTR, full-length AR cDNA, and 310-bp 3’-UTRs followed by 280-bp bovine GH poly(A) signals into pBlueScript sk− (vector) (Stratagene). The AR small interfering RNA (siRNA) expression vector that expresses an siRNA-targeting AR in mammalian cells was constructed by digesting and inserting a double-stranded polynucleotide 5’-GTCGGGCCCCT-
ATCC CA1GTTTGTGACAGA-GTTTGGGACTGGAAG

**Analysis of BrdU Staining Signal in Female Mice**

Observe the BrdU staining signal in both 4- and 6-wk-old female mice. Extension of mammary ducts, as compared with age-matched males, was observed. The sequence 5'-H11032

**Allele and Sequence**

With the sequence 5'-H11001

**Ductal Development of Mammary Gland**

In immature female mice, treatment with 0.1 nM E2 or 0.2% HI-FBS for treatment with 1 ml RPMI 1640 supplemented with 10% CDS-FBS for treatment with 0.4% low melting agarose (FMC Corp.) were layered on top of 1 ml of 0.8% agarose in 6-well culture plates. Cells were incubated 0.2% of heat-inactivated (HI)–FBS for treatment with 0.1 nM E2 or 0.2% HI-FBS for treatment with 100 ng/ml IGF-1 or 50 ng/ml epidermal growth factor (EGF). The cells were collected at indicated days for MTT assay according to the manufacturer’s instructions (Sigma-Aldrich).

**Soft-agar Colony Formation Assay**

2 × 10⁴ cells suspended in 0.4% low melting agarose (FMC Corp.) were layered on top of 1 ml of 0.8% agarose in 6-well culture plates. Cells were incubated with 1 ml RPMI 1640 supplemented with 10% CDS-FBS for treatment with 0.1 nM E2 or 0.2% HI-FBS for treatment with 100 ng/ml heregulin-α (HRG-α). After 4 wk of incubation, the colonies were visualized by incubating with 1 mg/ml INT (Sigma-Aldrich) for 24 h and counted with VersaDoc Imaging System (Bio-Rad Laboratories).

**Reporter Gene Assays**

Cells were plated in 96-well plates and plasmids at 0.5 μg per well were transfected into cells using SuperFect (QIAGEN). The medium was changed 2 h after transfection, and cells were cultured in media containing 10% CDS-FBS or 0.2% HI-FBS for 16 h, followed by treatment with 50 ng/ml EGF, 100 ng/ml IGF-1, 100 ng/ml HRG-α, 0.1 nM E2, or 1 nM DHT for another 16 h. 5 ng pRL-TK per well was used as internal control. Cells were harvested, and the Luc activity was analyzed using Dual-Luc Reporter Assay System (Promega).

**Western Blots**

Cells were lysed with radioimmunoprecipitation assay buffer containing 0.5% NP-40 and Western blotted with anti-AR (NH-27), anti-ER, antiactin, antimi–green fluorescent protein (GFP; Santa Cruz Biotechnology, Inc.), anti–total MAPK, and anti–phospho-MAPK (Cell Signaling Technology).

**Statistical Analysis**

All data were analyzed by one-way analysis of variance using minitab statistical software (State College). Mean separation was accomplished using Fisher’s pairwise comparison. Differences were considered significant at P < 0.05.

**Results**

**Generation and Phenotype of Female AR−/− Mice**

Using a Cre-lox conditional knockout strategy by mating the floxed AR male mice with AR+/− ACTB Cre+ females, we were able to generate female AR−/− mice, genotyped as AR−/− ACTB Cre+ (Fig. 1, A and B). Adult female mice with homologous deletion of AR appear healthy and develop normal external genitalia. Gross anatomical examination did not reveal obvious differences in the morphology of most organs between the AR+/+ and AR−/− litters. The body weights are similar between the AR+/+ and AR−/− mice, but the thymus of female AR−/− mice is

**Figure 1.** The generation and characterization of immature female AR−/− mice. (A) Gene targeting strategies. To generate female AR−/− mice, a Cre-lox strategy for conditional knockout was applied. The Cre-lox system uses the expression of P1 phage Cre recombinase to catalyze the excision of DNA located between flanking lox sites. (B) Breeding strategy of female AR−/− mice and genotyping of female AR−/− mice. Using the Cre-lox strategy, the targeted exon 2 of AR is not disrupted but floxed in the male mice. X and RI represent XbaI and EcoRI restriction enzyme sites, respectively. Thus, the AR functions normally in male mice, which can be bred with female AR+/− ACTB Cre+ mice and, thereby, generate homozygous female AR−/− mice. For examining the X chromosome with floxed AR, primers select and 2-3 are used. Select is located in the intron 1 with sequence 5′-GT-TGATACCTTAACCTCTGC-3′. This pair of primers will amplify a product with ∼510 bp for floxed AR and ∼460 bp for wt AR. For examining the AR knockout locus, primers select and 2-9 were used. 2-9 is located in intron 2 with the sequence 5′-CCTACATGACTGTGAGAGG-3′. The PCR product size from this pair of primers would be ∼270 bp for AR knockout allele and ∼600 bp for wt AR allele. The expression of Cre and internal control IL-2 was also confirmed by PCR genotyping. (C) The defects in the ductal development of mammary gland in immature female AR−/− mice. Whole breast mounts from 4-wk-old female AR−/− mice show lessened extension of mammary ducts, as compared with age-matched AR+/+ mice. (D) The decreases in the percentage of BrdU-positive staining (brown) are observed in both 4- and 6-wk-old AR−/− mice. (E) Statistic analyses of the distance of ductal extension in female AR−/− and AR+/+ mice (left). Statistic analyses of BrdU staining signal in female AR+/+ and AR−/− mice (right, n = 5 for each group). (F) The number of Cap cells (arrows) in TEB of AR−/− mice is less than that in the AR+/+ mice.
bigger than that of female AR<sup>+/+</sup> or AR<sup>+/−</sup> mice. Several estrogen target organs, including mammary gland, ovary, oviduct, and uterus, were collected from 4-, 6-, and 12-wk-old mice. The weights of these organs are 15–23% less in female AR<sup>−/−</sup> mice as compared with their age-matched littermates.

Defects in Mammary Gland Development in Prepubertal and Pubertal Stages in Female AR<sup>−/−</sup> Mice. First, we compared the morphology of mammary glands between immature (4- and 6-wk old) virgin AR<sup>+/+</sup> and AR<sup>−/−</sup> mice. By the fourth to sixth week, the ductal system has ~30–50% less extension in female AR<sup>−/−</sup> mice with reduced numbers and size of the terminal end buds (TEBs; Fig. 1, C and E). BrdU staining also revealed a 50% lower proliferation of AR<sup>−/−</sup> mammary glands, as compared with that of AR<sup>+/+</sup> mice (Fig. 1, D and E). The number of Cap cells (18), which are responsible for the ductal extension from TEBs, were also reduced in female AR<sup>−/−</sup> mice (Fig. 1 F). Together, our results indicated that the mammary gland development is retarded in prepubertal and pubertal stages in female AR<sup>−/−</sup> mice.

**Figure 2.** AR<sup>−/−</sup> mammary glands show the defects of the terminal branching and alveologenesis during maturity and pregnancy. Whole breast mounts from 8-, 16-, and 20-wk-old mature and 8-wk-old pregnant female mice were examined. (A–C) Less secondary and tertiary terminal branching in AR<sup>−/−</sup> mice, compared with AR<sup>+/+</sup> and AR<sup>+/−</sup> mice. (C) Early degeneration occurs in AR<sup>−/−</sup> mammary glands in 20-wk-old mice. (D) The decreased milk-producing lobuloalveolar development in the 8-wk-old pregnant AR<sup>−/−</sup> mice. (E) Using hematoxylin and eosin staining, the results indicate that the shrunken ductal space occurs in some AR<sup>−/−</sup> mammary glands in 16–20-wk-old mice (n = 4 for each group).

**Reduced Ductal Morphogenesis in the Mammary Glands of the Mature AR<sup>−/−</sup> Mice.** At maturity (8-, 16-, and 20-wk), we could see AR<sup>−/−</sup> mammary glands were filled with large bloated ducts terminating with bloated ends. It is also obvious that AR<sup>−/−</sup> mammary glands have fewer secondary and tertiary ductal branches as compared with age-matched AR<sup>+/+</sup> and AR<sup>+/−</sup> mice (Fig. 2, A–C). During the pregnancy stage, the retarded ductal branches in AR<sup>−/−</sup> mice are partially restored, yet compared with AR<sup>+/+</sup> mice, the AR<sup>−/−</sup> mice mammary glands still have less milk-producing alveoli (Fig. 2 D). In agreement with these findings, we also observed shrunken ductal spaces in some AR<sup>−/−</sup> mice mammary glands in 16-wk-old or older mice (Fig. 2 E), and reluctant nursing behavior in some AR<sup>−/−</sup> mothers. The decreased milk-producing alveoli and shrunken ductal spaces may result in the lessened capacity for AR<sup>−/−</sup> mice to feed their offspring. In 20-wk-old mice, we found that mammary glands in some AR<sup>−/−</sup> mice underwent degeneration earlier than those of the AR<sup>+/+</sup> mice (Fig. 2 C). Overall, Fig. 2 demonstrates that the lack of AR in female mice may retard the mammary gland development and affect the capacity of female AR<sup>−/−</sup> mice to feed their offspring.

Defects of MAPK Activity and Insulin-like Growth Factor I (IGF-I)–IGF-IR Pathway in AR<sup>−/−</sup> Mammary Glands. Early works indicated that E2/ER, P/PR, and paracrine growth factors/MAPK signals may contribute to the growth and development of mammary glands (19–21). First, we examined the MAPK activity in 4-wk-old mice. The immunohistochemical staining data show that the overall phospho–MAPK expression is weaker in the mammary cells from AR<sup>−/−</sup> mice, as compared with AR<sup>+/+</sup> mice (Fig. 3 A), although we could observe some ductal mammary cells with intensive phospho–MAPK staining in AR<sup>−/−</sup> mammary glands. The total MAPK protein expression is similar between AR<sup>−/−</sup> and AR<sup>+/−</sup> mice, as shown in Fig. 3 B. We examined the upstream regulators of MAPK signals, IGF-I–IGF-IRs. We found that IGF-IR, but not IGF-I, mRNA expression is reduced by 46% in immature AR<sup>−/−</sup> mammary glands (Fig. 3 C), suggesting that the IGF-I–IGF-IR→MAPK signaling pathway may be defective in female AR<sup>−/−</sup> mice. We investigated the expression of the downstream target, cyclin D1, in AR<sup>−/−</sup> and AR<sup>+/+</sup> mice using real-time quantitative PCR. The cyclin D1 mRNA expression was significantly reduced in female AR<sup>−/−</sup> mice (Fig. 3 C). Together, these data showing the reduction of IGF-IR, cyclin D1, and MAPK activity may suggest that the defects in the AR→IGF-I–IGF-IR→MAPK→cyclin D1 signaling pathway might result in the retarded mammary gland development in female AR<sup>−/−</sup> mice. This is in agreement with early papers showing that IGF-I–IGF-IR is an important paracrine growth factor for mammary gland development (22, 23), and cyclin D1 is a downstream mediator of growth factor–induced mammary gland proliferation (24).

Before systematic hormone function that occurs after puberty, growth factors, such as IGF-I, are the major con-
Reduced ER Activity in AR−/− Mammary Glands. Early analyses indicated that MAPK could also influence ER function (25), and that cyclin D1 could also be a downstream target gene for ER (26). Defects in MAPK and cyclin D1 may suggest that ER signals could also be impaired in the female AR−/− mouse. Therefore, we ovariectomized 4-wk-old mice, treated them with E2 for 2 d, and harvested the mammary glands for the comparison of the ER activity by examining ER target gene expression between AR−/− and AR+/+ mice. We found that E2-induced Efp (27) and HGF (28) were down-regulated in female AR−/− mice as compared with AR+/+ mice (Fig. 3 D). Early works also indicated that both Efp and HGF were important factors for breast cell growth (20, 29). Interestingly, we found that PR expression in mammary glands was similar between AR−/− and AR+/+ (Fig. 3 D). This finding is consistent with a previous paper showing that PR expression is E2/ER-independent in 5-wk-old or younger mice (30). Nevertheless, we found that the serum levels of PR’s ligand, P, was reduced in 12- to 16-wk-old adult female AR−/− mice (Fig. 3 E), which may result in the reduction of P/PR activity in mature mice. As the P/PR signal pathway plays important roles for the tertiary ductal branching and alveolar development (19), the lower P/PR activity in AR−/− mice may contribute to the retarded branching and lobuloalveolar formation in the development of mature stage mammary glands.

The AR−/− MCF7 Cells Exhibit Severe Defects in Growth and Colony Formation. To further dissect the mechanisms of AR roles in breast tissue at molecular and cellular levels, we applied the homologous recombination by using a targeting vector carrying a promoterless neomycin cassette to generate AR-deficient (AR−/−) MCF7 cells (Fig. 4 A). Two AR−/− MCF7 clones have been successfully obtained, and the targeted loci were confirmed by Southern blot analysis (Fig. 4 B). In these two homologous clones, the expression and the ligand-activated transcriptional activity of AR were indeed abrogated (Fig. 4 C and D). We found that AR−/− MCF7 cells exhibit a severe impairment in proliferation when cultured in media containing normal, steroid-deprived or 10−10 M E2-treated serum (Fig. 4 E). The soft-agar colony formation assay also showed that the
with np-AR. Full length of the 110-kD AR was detected by anti-AR antibody (NH27) using Western blotting assays. Cells were cotransfected with pEGFP-C1 vector for normalization of transfection efficiency. GFP expression was detected with anti-GFP antibody. (H) The mRNA expression of Ki67 and c-myc, but not Bcl-2, was reduced in AR+/− MCF7 cells transfected with AR siRNA using electroporation, compared with the cells transfected with vector alone. Electroporation was performed using 0.4-cm cuvettes and Gene Pulser II set at 280 V and 950 μF.

The Growth Factor–mediated Proliferation and MAPK Activation Is Impaired in AR+/− MCF7. Next, we examined whether the loss of AR impairs the growth factor–mediated proliferation and MAPK activation in AR+/− MCF7 cells. Treatment of AR+/+ MCF7 cells with IGF-I, EGF, or HRG-α could stimulate cell proliferation and activate GAL4-Elk1, a direct target of MAPK, in a low serum-containing medium (Fig. 5, A and B). In contrast, the growth factor–stimulated cell proliferation and MAPK activation were largely impaired in the AR+/− MCF7 cells (Fig. 5, A and B). Using another strategy by transection of AR siRNA into AR+/+ MCF7 cells, we found that suppression of AR expression could also diminish IGF-I/EGF/HRG-α–induced MAPK activation (Fig. 5 B). Moreover, the reduced transcriptional activity of GAL4-Elk1 in AR+/− MCF7 cells could be rescued by transfection of np-AR (Fig. 5 C). Interestingly, adding EGF with np-AR could synergistically enhance the transactivation of GAL4-Elk1 in AR+/− MCF7 cells compared with the cells treated with EGF alone (Fig. 5 C), suggesting a significant involvement of AR in the growth factor signaling pathway. Furthermore, the AR-activated GAL4-Elk1 activity could be diminished by MAPK phosphatase-1 (CL-100) or a specific inhibitor U0126, as well as Ras-DN or Raf-DN (Fig. 5 D; reference 14). Also, the reduction of MAPK activation by AR siRNA could be recovered by MEK-CA, Ras-CA, or Raf-CA, but not by Rac-CA (15) or PI3K (Fig. 5 E, p110 catalytic subunit). These results indicate that AR is an important upstream regulator of the Ras/Raf/MAPK cascade.

The Transcriptional Activity of ER Is Defective in AR+/− MCF7 Cells. Using an ERE-Luc reporter, we further compared the ER activity in AR+/+ and AR+/− MCF7 cells. The transcriptional activities of ER were reduced by 58.8, 53.8, and 55.0% in AR+/− MCF7 cells in the presence of E2 at 10−12, 10−10, and 10−8 M, respectively (Fig. 5 F). The reduced transcriptional activity of ER in AR+/− MCF7 cells could be restored by transfection of np-AR.
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These results are consistent with the data in Fig. 3 C showing that the ER target gene expression is reduced in AR-/−/− MCF7 cells.

The NH2 Terminus/DNA-binding Domain (DBD) of AR Is Required for MAPK Activation.

To further determine which functional domain of AR is required to restore the normal MAPK activity in AR-/−/− MCF7 cells, various AR fragments were reintroduced into AR-/−/− MCF7 cells (Fig. 6 A). We found that the NH2 terminus together with DBD (but not ligand-binding domain [LBD]), LBD with deletion of helix 12 domain (LBD-dH12), DBD alone, or NH2 terminus alone can restore the MAPK activation (Fig. 6 A).

Next, because an AR mutant (AR-R608K) has been suggested to be associated with male breast cancer (31), we were interested to know its effect on the MAPK activation. We found that in AR-/−/− MCF7 cells, AR-R608K had a higher induction fold on MAPK activation than AR-FL (Fig. 6 B), suggesting that the contribution of AR-R608K to breast cancer incidence may involve the excessive activation of MAPK. Next, the requirement of AR NH2 terminus/DBD, but not LBD, to restore the MAPK activation was further confirmed by using a double mutation AR (AR-R614H-dprm) with a deletion of the proline-rich motif (dprm) at the NH2-terminal region (32) and a point mutation on the second zinc-finger motif (R614H) at the DBD (33). Although AR with a single mutation, AR-R614H or AR-dprm, still partially retains the ability to activate MAPK, the double mutation AR-R614H-dprm almost loses the whole capacity to restore the MAPK activity, even though these AR mutants contain intact LBDs (Fig. 6 B).

Together, these restoration assays indicate that AR, through its NH2 terminus/DBD domains, plays an important role for the MAPK activation in MCF7 cells.

Discussion

Using three different approaches (Cre-lox conditional KO, siRNA, and homologous recombination) to abrogate the AR function in female mice and MCF7 breast cancer cells, we demonstrate that AR may go through interruption of MAPK activity and ER signaling to exert its important roles for the normal mouse mammary gland development and MCF7 breast cancer cell growth. These results are in agreement with early papers showing that both MAPK and ER are essential factors for mammary gland development and breast cancer growth. For example, female ER-/−/− mice exhibit undeveloped mammary glands similar to those of newborn mice, indicating the essential roles of ER in the ductal growth (21), and the loss of ER can significantly delay the onset of tumor induction in MMTV-
that loss of AR could disrupt the IGF-I–, EGF–, and H-Ras–induced mammary tumor growth (38). AR could contribute to the retardation of mammary gland development because AR abrogation may also occur in males, but not in females. Similar phenomena were observed in male mice lacking functional ER (36, 37). The MAPK inhibitor PD908059 could inhibit both the mammary gland alveolar morphogenesis (20) and Her2/Neu–, H-Ras–, or C-myc–initiated mammary tumor growth (38).

To dissect how AR influences MAPK activity, we found that loss of AR could disrupt the IGF-I–, EGF–, and HRG–α–induced MAPK activity (Fig. 4 B) and reduced IGF-IR expression in AR−/− mice (Fig. 3 C). Bonnette et al. (23) found that mice with defective IGF-IR have less branching and decreased cellular proliferation of TEBs in developing mammary glands, and these defects could be partly restored during pregnancy. Similar phenomena also occurred in AR−/− mice (Figs. 1 and 2), suggesting that the suppression of IGF-IR by the AR abrogation may contribute to the retarded mammary gland development in AR−/− mice. Consistent with data (Fig. 4 E and Fig. 5 B) showing that the loss of AR interrupts HRG–α–induced anchorage-independent cell growth and MAPK activation, Watson et al. (39) found that in transgenic rats with overexpression of HER2–Neu in the mammary gland, normal males, but not castrated males, developed mammary tumors. These results may suggest the potential cross-talk between androgen–AR and HER2–Neu signaling pathways in mammary tumor progression.

Further mechanism dissection studies indicate that AR–induced MAPK activation is via the Ras/Raf/MAPK cascade (Fig. 5 D), although the detailed mechanisms of how AR influences Ras-Raf activity remain unclear. Migliaccio et al. (32) reported that AR could activate MAPK via the interaction between the proline-rich motif of AR and the SH3 domain of c-Src. They demonstrated that this interaction occurred quickly and could be initiated in a short time by androgen or estrogen treatment. In contrast, Fig. 6 A demonstrates that AR NH2 terminus/DBD, without the LBD, or LBD–dH12 alone, could induce MAPK activity, suggesting that AR, but not androgen, is the major factor to activate MAPK activity. This raises an interesting question: what is the role of AR in the breast cancer development? Early analyses of androgen–AR roles in breast cancer mainly focused on the effect of androgen or estrogen treatment on the breast cancer showing that androgen is likely that AR signals may use multiple pathways, including the classic androgen/AR→AR target genes of genomic actions as well as AR→AR interaction proteins of nongenomic action to exert its roles in the breast cancer progression. This is in agreement with early papers showing that ER could also cross-talk to MAPK in breast cancer cells (25, 41). In addition to estrogens, ER could be activated via phosphorylation at Ser118 by MAPK to induce its target gene expression (25). In return, ER could also induce the Ras/Raf/MAPK cascade via nongenomic action (32). Therefore, our results showing that AR could influence both MAPK and ER signals suggest that the reduction of ER activity could be due to the reduced MAPK activity, and the reduced MAPK activity may be due to the reduced ER activity in female mice and MCF7 cells lacking AR.

Neu (Neu/ER−/−) or Wnt-1 (Wnt-1/ER−/−) transgenic mice lacking functional ER (36, 37). The MAPK inhibitor PD098059 could inhibit both the mammary gland alveolar morphogenesis (20) and Her2/Neu–, H-Ras–, or C-myc–initiated mammary tumor growth (38). To dissect how AR influences MAPK activity, we found that loss of AR could disrupt the IGF-I–, EGF–, and HRG–α–induced MAPK activity (Fig. 4 B) and reduced IGF-IR expression in AR−/− mice (Fig. 3 C). Bonnette et al. (23) found that mice with defective IGF-IR have less branching and decreased cellular proliferation of TEBs in developing mammary glands, and these defects could be partly restored during pregnancy. Similar phenomena also occurred in AR−/− mice (Figs. 1 and 2), suggesting that the suppression of IGF-IR by the AR abrogation may contribute to the retarded mammary gland development in AR−/− mice. Consistent with data (Fig. 4 E and Fig. 5 B) showing that the loss of AR interrupts HRG–α–induced anchorage-independent cell growth and MAPK activation, Watson et al. (39) found that in transgenic rats with overexpression of HER2–Neu in the mammary gland, normal males, but not castrated males, developed mammary tumors. These results may suggest the potential cross-talk between androgen–AR and HER2–Neu signaling pathways in mammary tumor progression.

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In conclusion, this paper provides the first in vivo evidence showing that AR may go through growth factors, MAPK, and ER/PR signals (summary in Fig. 6 C) to control the normal breast development, and modulate the breast cancer cell proliferation, especially in the conditions of absence of or reduced E2 (Fig. 4 E). Whether this would imply there is a positive correlation between AR expression and breast cancer incidence/progression in women with lower circulating E2, such as with the postmenopausal stages, remains an interesting question for future study. Epidemiological papers support this correlation suggesting that AR expression is more significantly associated with breast cancer in postmenopausal women than premenopausal women (9, 13, 42), and up to 50% of the AR-positive breast cancers are ER- and/or PR-negative (13, 43). Finally, as AR NH2 terminus/DBD, but not LBD, may play AR expression is more significant with breast cancer cell proliferation, especially in the conditions stages, remains an interesting question for future study. Epidemiological papers support this correlation suggesting that AR expression is more significantly associated with breast cancer in postmenopausal women than premenopausal women (9, 13, 42), and up to 50% of the AR-positive breast cancers are ER- and/or PR-negative (13, 43). Finally, as AR NH2 terminus/DBD, but not LBD, may play a role in the normal breast development, and modulate the breast cancer cell proliferation, especially in the conditions of absence of or reduced E2 (Fig. 4 E). Whether this would imply there is a positive correlation between AR expression and breast cancer incidence/progression in women with lower circulating E2, such as with the postmenopausal stages, remains an interesting question for future study. Epidemiological papers support this correlation suggesting that AR expression is more significantly associated with breast cancer in postmenopausal women than premenopausal women (9, 13, 42), and up to 50% of the AR-positive breast cancers are ER- and/or PR-negative (13, 43). Finally, as AR NH2 terminus/DBD, but not LBD, may play essential roles to modulate the growth factor signaling pathways in MCF7 breast cancer cells, targeting the function of AR NH2 terminus/DBD may represent a new potential therapeutic approach to the battle against breast cancer.

We thank J. Xu, H. Land, P. Keng, D.A. Pearce, P.A. di Sant’Agnese, K.-L. Guan, and J. Chernoff for helpful discussions and their reagents. We also thank K. Wolf for proofreading.

This work was supported by National Institutes of Health grants DK60905, Army DAMD17-02-1-0557, NYSDOH-C017947, and their reagents. We also thank K. Wolf for proofreading.

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