Androgen Signaling in Esophageal Adenocarcinoma Cell Lines In Vitro

Helen M. Palethorpe¹ · Paul A. Drew¹,² · Eric Smith¹,³

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Abstract

Background We showed previously that nuclear localization of the androgen receptor (AR) and expression of the androgen-responsive gene FK506-binding protein 5 (FKBP5) in esophageal adenocarcinoma (EAC) tissues were associated with decreased patient survival, suggesting a role for androgens in this cancer.

Aim To investigate the effect of the AR ligand 5α-dihydrotestosterone (DHT) on AR-expressing EAC cell lines in vitro.

Methods and Results In tissue resection specimens from EAC patients, FKBP5 expression was positively associated with proliferation as measured by Ki-67 expression. We stably transduced AR into three AR-negative EAC cell lines, OE33, JH-EsoAd1, and OE19, to investigate androgen signaling in vitro. In the AR-expressing cell lines, 10 nM DHT, the concentration typically used to study AR signaling, induced changes in the expression of androgen-responsive genes and inhibited proliferation by inducing cell cycle arrest and senescence. At lower DHT concentrations near the half maximal inhibitory concentration (IC50), the AR-expressing cell lines proliferated and there were changes in the expression of androgen-responsive genes. In direct co-culture with cancer-associated fibroblast-like PShTert myofibroblasts, 10 nM DHT induced changes in the expression of androgen-responsive genes but did not inhibit proliferation.

Conclusions This is the first study to show that EAC cell lines respond to androgen in vitro. Proliferation together with the expression of androgen-responsive genes was dependent on the concentration of DHT, or the presence of a permissive microenvironment, consistent with observations in the tissues. These findings are consistent with a role for androgen signaling in EAC.

Keywords Esophageal adenocarcinoma · Androgen receptor · Fibroblast · FKBP5 · Dihydrotestosterone · Direct co-culture · In vitro

Abbreviations

ACTB Actin beta
CTV CellTrace violet
DHT 5α-Dihydrotestosterone
EAC Esophageal adenocarcinoma
E2F1 E2F transcription factor 1
FBXO32 F-box protein 32
FKBP5 FK506-binding protein 5
Given the association between AR localization, FKBP5 expression, and poor survival, we sought suitable cell lines to investigate the effect of androgen signaling on the behaviors of EAC cells. All available cell lines were AR negative, probably due to loss of AR expression during the establishment of cell lines from tissues [15]. We therefore stably transduced three EAC cell lines with AR. The aim of this study was to investigate factors that affect the growth and gene expression in AR-expressing EAC cell lines in response to androgen.

Methods

Immunohistochemistry for Ki-67 and FKBP5

Tissue microarrays (TMAs) composed of one or more representative cores of EAC were constructed as described previously [16]. Sequential sections consisting of cores from 74 cases were immunostained for FKBP5 [15] and Ki-67 [17]. Cores were scored as the percent of epithelial cells that expressed FKBP5 or Ki-67 as follows: 0, negative; 1, < 5% (rare); 2, < 25%; 3, > 25% < 75%; 4, > 75%. The median score for FKBP5 and Ki-67 (Ki-67 index) was determined for multiple cores from each case.

Cell Culture

The EAC cell lines OE33, JH-EsoAd1, and OE19 [18, 19] were obtained from the ATCC, Johns Hopkins University, and Sigma-Aldrich, respectively. They were stably transduced with AR and green fluorescent protein (GFP), or with GFP only [15]. At least six single-cell clones were established from each AR-transduced cell line, and the clone expressing the lowest amount of AR, as determined by western immunoblot, was used for all experiments unless otherwise stated. The cell lines expressing AR and GFP are referred to as OE33-AR, JH-AR, and OE19-AR, respectively. The EAC cell lines were maintained in androgen-depleted growth medium (stripped medium) consisting of phenol red-free RPMI-1640 containing l-glutamine (Life Technologies, Eugene, OR, USA), supplemented with 10% dextran charcoal stripped fetal bovine serum (Equitech-Bio, Inc., Kerrville, TX, USA), 200 U/mL penicillin, and 200 μg/mL streptomycin (Life Technologies). Stripped medium was used for experiments unless stated otherwise.

The PShTert myofibroblasts [20–22] were stably transduced with the SFG-RFP/Rluc construct to express red fluorescent protein (RFP) [23]. Neonatal foreskin fibroblasts (NFFs) and PShTert myofibroblasts were used between passages 10 and 20. Fibroblasts were maintained in DMEM containing l-glutamine (Life Technologies), supplemented with 10% fetal bovine serum (FBS; Sigma-Aldrich).

Introduction

The incidence of esophageal adenocarcinoma (EAC) has increased rapidly over recent decades in Western countries [1–5]. It has a dismal prognosis, with around 65% of patients unsuitable for surgery at the time of diagnosis and an overall five-year survival rate of less than 15% [6, 7]. The major risk factors for EAC are gastro-esophageal reflux disease and obesity, leading to the only described precursor lesion for this cancer, Barrett’s esophagus. This cancer has one of the highest male-to-female ratios reported for cancers of non-reproductive organs, ranging from 7–10 to 1 [1, 2, 4, 6, 8–11], significantly higher than for the major risk factors. The gender difference appears to result from an approximate 20-year delay in onset in females of Barrett’s esophagus [12] and EAC [13]. These observations are consistent with a role for the sex steroid hormones in the biology of this cancer, with their concentrations differing between males and females, and changing over the lifespan.

The most important sex hormones in males are the androgens, and the most predominant androgen is testosterone. Testosterone passes through the cell membrane and into the cytoplasm where it, or its more physiologically effective metabolite 5α-dihydrotestosterone (DHT), binds to and activates the androgen receptor (AR). Activated AR translocates from the cytoplasm into the nucleus and binds to androgen response elements in the genome, influencing the transcription of androgen-responsive genes. The nature of the response can be modified by the relative abundance of multiple co-regulators (both co-activators and corepressors) [14].

We previously reported the immunostaining of EAC tissues for AR and the androgen-responsive gene FK506-binding protein 5 (FKBP5) [15]. We detected AR in the cancer cells of 75 of 77 cases, and in 70 it was nuclear. The expression of FKBP5 was observed in 64% of cases and only when the AR was nuclear. There was a significant association between nuclear AR and FKBP5 expression and decreased survival.

| Term       | Definition                                      |
|------------|------------------------------------------------|
| GFP        | Green fluorescent protein                      |
| HMOX1      | Heme oxygenase 1                               |
| IC50       | Half maximal inhibitory concentration          |
| NDRG1      | N-myc downstream regulated 1                   |
| NFF        | Neonatal foreskin fibroblast                   |
| qRT-PCR    | Quantitative reverse transcription polymerase chain reaction |
| RFP        | Red fluorescent protein                        |
| SA-β-gal   | Senescence-associated beta-galactosidase       |
| SD         | Standard deviation                             |
| TERT       | Telomerase reverse transcriptase                |
| TMA        | Tissue microarray                              |

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Aldrich, St Louis, MO, USA), 200 U/mL penicillin, and 200 μg/mL streptomycin (Life Technologies).

**Direct Co-culture**

The NFFs and PShTert myofibroblasts were cultured in stripped medium overnight, seeded at $4 \times 10^5$ cells per well into six-well plates (BD Biosciences, Franklin Lakes, NJ, USA), and then incubated for 48 h to form confluent monolayers. Next, OE33-ARs were seeded at $1 \times 10^5$ cells per well either in monoculture or in direct co-culture with the fibroblasts. The following day (day 0), and every 48 h thereafter, the medium was replaced with stripped medium supplemented with vehicle (0 nM DHT; 0.1% ethanol) or 10 nM DHT. Cells were harvested on day 6 of treatment, unless stated otherwise.

**Translocation of Androgen Receptor**

To establish direct co-cultures, fibroblasts were cultured in stripped medium overnight, then seeded at $8 \times 10^3$ fibroblasts per well in eight-well Lab-Tek Chamber Slides (Thermo Fisher Scientific, Rochester, NY, USA), and incubated for 48 h. Next, OE33-ARs ($2 \times 10^3$ cells per well) were added to the wells, followed by overnight incubation. The medium, supplemented with vehicle or 10 nM DHT (day 0), was replaced then and every 48 h for 6 days.

Following treatment, the cells were washed in Dulbecco’s phosphate-buffered saline (DPBS; Life Technologies), fixed in methanol on ice for 5 min, and air-dried. The cells were blocked with 10% normal goat serum (Dako, Glostrup, Denmark) in DPBS for 20 min and labeled with rabbit antihuman AR polyclonal IgG (clone N-20; 1 μg/mL in 1.5% goat serum; Santa Cruz Biotech Inc., Santa Cruz, CA, USA) for 1 h, followed by incubation with Alexa Fluor 568 goat anti-rabbit IgG (2 μg/mL in 1.5% goat serum; Molecular Probes by Life Technologies) for 45 min. Nuclei were stained with 1 μg/mL 4’,6-diamidino-2-phenylindole dihydrochloride (DAPI; Sigma-Aldrich) in DPBS for 15 min. Slides were mounted in fluorescent mounting medium (Dako) and stored at 4 °C in darkness. Images were captured using a Zeiss LSM 700 confocal microscope.

**Cell Proliferation**

To measure cell proliferation, $1 \times 10^3$ cells per well were seeded in 96-well plates and cultured for 48 h. The cells were then treated with either vehicle or various concentrations of DHT for 6–12 days, depending on the cell line. The cells were next fixed in 10% neutral buffered formalin for 30 min, stained for 10 min with 1% crystal violet (Sigma-Aldrich) in 2% ethanol, washed eight times in distilled water, and then air-dried overnight. The crystal violet was eluted using 10% acetic acid and gentle rotation of the plates. The absorbance of the eluent was measured at 595 nM using a FLUOstar Optima microplate reader (BMG Labtech, Ortenberg, Germany). To determine whether growth inhibition with DHT was mediated via AR, OE33-ARs were seeded into six-well plates ($1 \times 10^5$ cells per well) and cultured for 48 h. The cells were treated for 6 days with 10 nM DHT and either vehicle (0.15% dimethyl sulfoxide; Sigma-Aldrich) or 15 μM enzalutamide (MedChem Express, Princeton, NJ, USA).

Cell division of OE33-AR was measured by dye dilution, using CellTrace Violet (CTV; Life Technologies). Cells were seeded in six-well plates ($3 \times 10^4$ cells per well) and incubated for 24 h. Wells for time 0 were then fixed in 4% paraformaldehyde. In the other wells, the medium was replaced daily, supplemented with either vehicle or DHT at around the half maximal inhibitory concentration (IC50; 0.06 and 0.1 nM) or 10 nM. Other wells were fixed with 4% paraformaldehyde 1, 3, or 5 days following. The amount of CTV in the cells was measured using a FACScanto II (BD Biosciences) with BD FACSDiva software. Cell doublets were excluded by doublet discrimination, based on nonlinearity of forward scatter and side scatter area versus height plots. Cells were gated based on GFP-positivity, and the median CTV intensity of this population determined using FlowJo software version 8.8.7 (Ashland, OR, USA).

**Quantitative Real-Time Reverse Transcription PCR (qRT-PCR)**

Total RNA was isolated from cells using TRIzol, and 1 μg was reverse-transcribed using the iScript cDNA Synthesis Kit (Bio-Rad Laboratories, Hercules, CA, USA) in a final reaction volume of 20 μL. Gene expression was determined by quantitative real-time reverse transcription polymerase chain reaction (qRT-PCR) using iQ SYBR Green Supermix (Bio-Rad Laboratories) in a final reaction volume of 10 μL, containing 0.1 μL of cDNA, and a final concentration of 0.2 μM of each forward and reverse primer (Supplementary Table S1). Reactions were performed using a CFX96 (Bio-Rad Laboratories) at 95 °C for 3 min, then 40 cycles of 95 °C for 15 s, 60 °C for 15 s, and 72 °C for 30 s, followed by a final extension of 72 °C for 1 min. Products were melted to confirm specificity. Normalized fold expression was calculated using actin beta (ACTB) as the reference gene.

**Cell Cycle Analysis**

OE33-ARs were seeded at $1 \times 10^5$ cells per well in stripped medium in six-well plates and incubated for 24 h.
The medium was then replaced daily, supplemented with either vehicle or 0.06, 0.1, or 10 nM DHT. Wells were harvested at 0, 24, 48, or 72 h. The cells were washed, resuspended in DPBS, and fixed with a final concentration of 70% ice-cold ethanol. The cells were pelleted, resuspended with 0.25% Triton X-100 (Sigma-Aldrich) in DPBS, and incubated for 2 h with 25 μg/mL propidium iodide (Sigma-Aldrich) and 40 μg/mL bovine pancreas ribonuclease A (Sigma-Aldrich) in DPBS. The DNA content of single cells was measured using a FACSCanto II. The percentages of cells in each cell cycle phase and in sub-G1 were calculated using BD FACSDiva software.

**Confocal Microscopy**

To assess morphology, OE33-AR (1.72 × 10^3), JH-AR (5.73 × 10^3), or OE19-AR (5.73 × 10^3) cells were seeded into 96-well μ-plates (Ibidi, Martinsried, Germany) in stripped medium supplemented with either vehicle or 10 nM DHT replaced daily for 3 days. For direct co-cultures, NFFs were labeled using the CellTrace Violet (CTV) Cell Proliferation Kit according to the manufacturer’s protocol (Life Technologies). Fibroblasts were seeded at 1.14 × 10^5 per well, and OE33-ARs were seeded overnight at 1.43 × 10^5 cells per well, in either monoculture or overlying fibroblasts followed by treatment with vehicle or 10 nM DHT for 6 days, with medium replaced every 48 h. Images were captured using a Zeiss LSM 700 confocal microscope with Zen2012 SP1 (black edition) software version 8.1.

**Senescence-Associated Beta-Galactosidase Assay**

Cells were seeded in 24-well plates at 2 × 10^3 cells per well for OE33 and OE33-AR and 3 × 10^3 cells per well for JH-AR and OE19-AR, followed by 48-h incubation. The culture medium was then replaced daily with fresh medium supplemented with vehicle or DHT (IC50s and 10 nM). On days 0, 1, 3, and 5 for OE33 and OE33-AR and 0, 4, 6, and 8 for JH-AR and OE19-AR, wells were stained with the Senescence Cells Histochemical Staining Kit (Sigma-Aldrich). The percentage of senescence-associated beta-galactosidase (SA-β-gal) positive cells was calculated from a count of 200 cells.

**Statistics**

The statistical software used was Prism 6.0d for Macintosh (GraphPad Software, San Diego, CA, USA). Proliferation dose–response curves were fitted, and the IC50 determined by nonlinear regression analysis. Data are the mean ± SD of a single experiment reproduced in triplicate with p values determined by parametric unpaired Student’s t test assuming equal standard deviations (SD), unless stated otherwise. Differences were considered significant when the two-tailed p value was ≤ 0.05.

**Results**

**EAC Tissues with a High Percentage of FKBP5-Positive Cells Had a High Proliferation Index**

Previously, we reported that the expression of FKBP5, a surrogate marker for androgen signaling, was associated with reduced survival in EAC [15]. To determine the relationship between AR signaling and tumor growth in vivo, we measured the percentage of FKBP5 positive cells and the Ki-67 proliferation index in immunostained EAC resection specimens (Supplementary Fig. S1). There was a positive correlation between the FKBP5 expression and the proliferation index (nonparametric Spearman correlation \( r = 0.37, 95\% \) confidence interval 0.1533–0.5628; \( p = 0.0010 \)), with the proliferation index significantly higher in those tissues with higher FKBP5 expression (Fig. 1; \( p = 0.0002 \)).

**DHT Inhibited Proliferation of AR-Expressing EAC Cell Lines In Vitro**

Expression of AR protein was confirmed in the three EAC cell lines stably transduced with AR, OE33-AR, JH-AR, and OE19-AR, by western immunoblot (Supplementary Fig. S2a) and immunocytochemistry (Supplementary Fig. S2b). In the absence of DHT, AR immunoreactivity

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**Fig. 1** Ki-67 proliferation indices in EAC resection tissues with low (≤ 2) or high (> 2) FKBP5 expression. Tissue microarrays were immunostained and scored as the percentage of epithelial cells that expressed FKBP5 or Ki-67 as follows: 0, negative; 1, < 5% (rare); 2, < 25%; 3, > 25% < 75%; 4, > 75%. Data are the median score of cores from each case. p value by Mann–Whitney U test
was seen by confocal microscopy to be moderate in the cytoplasm and mild to moderate in the nucleus. Exposure to 10 nM DHT induced complete nuclear localization of the AR, confirming that the transduced AR was functionally responsive to androgen.

In the AR-negative cell lines, exposure to DHT at concentrations up to 100 nM did not significantly alter proliferation (data not shown). The dose–response curves for each of the AR-expressing lines, given a single dose of DHT at the start of culture, are shown in Fig. 2a. The concentration of DHT used in most reported studies of AR signaling in vitro is 10 nM, which completely inhibited proliferation of OE33-AR and JH-AR, and almost completely of OE19-AR. The IC50s were 0.09, 0.26, and 1.3 nM for OE33-AR, JH-AR, and OE19-AR, respectively. To determine whether the differences in the DHT dose–response curves between the cell lines were due to the amount of AR expressed, we compared, within each of the transduced lines, clones with the highest and lowest expression of AR and found no significant differences in the DHT dose–response curves (data not shown). We also observed the same or similar dose–response for the uncloned OE33-AR. The addition of the AR antagonist enzalutamide (15 μM) completely blocked the growth inhibition of the AR-expressing cells induced by 10 nM DHT, confirming that the anti-proliferative effect was mediated by the AR (p < 0.0001; Fig. 2b).

For subsequent experiments, we used two concentrations of DHT near the IC50 (0.06 and 0.1 nM for OE33-AR, 0.25 and 0.5 nM for JH-AR, and 0.5 and 1.0 nM for OE19-AR), as well as 10 nM. We next examined the possibility that proliferation may differ with daily

Fig. 2  Effect of DHT on the proliferation of AR-expressing EAC cell lines.  a  Dose–response curves for the proliferation of AR-expressing EAC cells grown for 6–12 days with vehicle or tenfold serial dilutions of DHT. Proliferation was measured by crystal violet assay. Data are the mean ± SD of six replicates, and the corresponding nonlinear regression curve, from a representative experiment for each cell line.  b  The effect of 15 μM enzalutamide on the proliferation of OE33-AR treated with 10 nM DHT

Fig. 3  Effect of single compared to daily doses of DHT on OE33-AR growth and FKBP5 expression.  a  Dose–response curves for the proliferation of OE33-AR grown for 5 days with vehicle or twofold serial dilutions of DHT given as a single dose or replaced daily. Data are the mean ± SD of six replicates, and the corresponding nonlinear regression curve, from a representative experiment.  b  Normalized fold FKBP5 expression after 3 days culture with a single dose or daily replacement of vehicle or DHT. Data are the mean ± SD of triplicate reactions for three biological replicates
replenishment of DHT compared to a single treatment at the start of the culture. The results in Fig. 3a show the DHT dose–response curves for OE33-AR. These were similar whether the DHT was given as a single treatment or daily in fresh medium, with IC50s of 0.08 and 0.06 nM, respectively.

DHT Induced Androgen-Responsive Gene Expression in AR-Expressing Cell Lines

In preliminary studies, we found that cell lines cultured with a single dose of 10 nM DHT expressed high levels of the androgen-responsive gene FKBP5, even though proliferation was inhibited. At lower concentrations of DHT, around the IC50, there was partial inhibition of growth but no, or very low, FKBP5 expression. This appeared to conflict with our immunostains of resection tissues where we frequently measured FKBP5 and Ki-67 expression together.

We therefore compared the effect of a single dose to daily replenishment on the induction of the androgen-responsive gene FKBP5 in OE33-AR (Fig. 3b). A single dose induced an increase in FKBP5 expression compared to vehicle of 1.2-fold for 0.06 nM DHT ($p = 0.002$), 2.2-fold for 0.1 nM ($p = 0.0004$), and 22-fold for 10 nM ($p = 0.0005$) after 3 days of culture. In contrast, when the DHT was replenished daily, the increase in FKBP5 expression compared to vehicle was twofold for 0.06 nM ($p = 0.03$), fourfold for 0.1 nM ($p < 0.0001$), and 16-fold for 10 nM ($p = 0.0002$). The increases in FKBP5 expression for daily compared to single dosing were significantly greater for 0.06 and 0.1 nM DHT ($p = 0.02$ and $p < 0.0001$, respectively), but not for 10 nM ($p = 0.639$). Daily dosing, which would be expected to more closely mimic in vivo conditions, with concentrations of DHT around the IC50, permitted growth and induced significant expression of FKBP5.

Next, we measured the expression of known androgen-responsive genes in each of the three AR-expressing cell lines at the different concentrations of DHT. We found that the pattern of response was similar between the cell lines, although there were differences in the fold increases (Supplementary Fig. S3).

DHT Inhibited Cell Division and Induced Cell Cycle Arrest and Cell Senescence in AR-Expressing Cells

To understand better the inhibition of growth, we analyzed the effect of DHT on cell division, cell cycle arrest, and cell senescence. Cell division in OE33-AR, as measured by the intracellular dilution of CTV, was inhibited after 3 and 5 days of culture (Fig. 4a). The median CTV content at day 0 was 25,724 fluorescence units (FU). After 5 days of treatment with vehicle, it was reduced to 535 FU. In contrast, after 5 days of treatment with 0.06, 0.1, and 10 nM DHT the median CTV content was 621 FU, 930 FU, and 5760 FU, respectively, indicating that there was less cell division as the concentration of DHT increased.

Next, we measured the cell cycle phase distribution by flow cytometry following 3 days of culture. The results are shown in Fig. 4b. There were no significant changes in the proportion of cells in each phase of the cell cycle following treatment with 0.06 nM DHT. Compared to vehicle, there was a 19% (p = 0.006) and 52% increase (p = 0.0004) in the proportion of cells in the G0/G1 phase following treatment with 0.1 and 10 nM DHT, respectively, a 34% (p = 0.0004) and 47% (p = 0.003) decrease in cells in the G2/M phase, and no difference and a decrease of 51% (p = 0.009) in the S phase. There was no sub-G1 population with any of the concentrations, suggesting that DHT did not induce cell death. The expression of E2F1, a transcriptional activator necessary for progression through the G1/S transition, was significantly inhibited with 0.1 nM DHT, which was least pronounced in OE33-AR and least in OE19-AR (Fig. 4c).

Using time-lapse confocal microscopy, we observed that each of the three AR-expressing cell lines underwent extensive morphological changes over 3 days in monolayer culture with 10 nM DHT (Fig. 4d). The cells became dis cohesive, enlarged, and flattened, with the appearance of many large cytoplasmic holes. There was no microscopic evidence of extensive cell death. Because these changes were suggestive of senescence, we stained cultures for the senescence marker, senescence-associated beta-galactosidase (SA-β-gal). There was a concentration and time-dependent increase in the percentage of SA-β-gal stained cells over the duration of culture in each of the three cell lines, which was most pronounced in OE33-AR and least in OE19-AR (Fig. 4e). Together these results demonstrated that DHT inhibited the proliferation of AR-expressing EAC cell lines in vitro by inducing growth arrest and senescence in a dose-dependent manner.
The Effect of Direct Co-culture with Fibroblasts

We next explored the possibility that fibroblasts, the major cell population within the tumor microenvironment, which are known to influence the response of tumor cells to drugs [24], might modify the response of AR-expressing EAC cells to androgens. Normal neonatal foreskin fibroblasts (NFFs) and PShTert myofibroblasts representative of cancer-associated fibroblasts were used.

The morphology of OE33-AR in monoculture and direct co-culture with NFFs, vehicle, or 10 nM DHT is shown in Fig. 5a. With vehicle, there was no apparent microscopic difference between OE33-AR grown in monoculture or in direct co-culture with NFFs. They formed numerous clusters of cells with distinct cell borders, polygonal shape, and well-defined nuclei. With 10 nM DHT, growth of OE33-AR was inhibited in both the monoculture and the NFF direct co-culture. The OE33-AR were enlarged and contained numerous refractile, round bodies devoid of obvious structural content under phase microscopy and lacking GFP under fluorescence microscopy. This effect of DHT on OE33-AR growth in monoculture and direct co-culture with NFFs was also reflected in the cell counts shown in Fig. 5b.

In relation to the co-culture of OE33-AR with cancer-associated fibroblast-like PShTert myofibroblasts, there were no apparent microscopic differences between the monoculture and the direct co-culture with the myofibroblasts, either with vehicle or with 10 nM DHT (Fig. 5a), and no differences between the counts of OE33-AR grown with vehicle compared to 10 nM DHT in the direct co-culture (Fig. 5b). Similar effects of the myofibroblasts on morphology (Supplementary Fig. S4) and cell counts (Supplementary Fig. S5) in the presence of 10 nM DHT were observed for the other two AR-expressing EAC cell lines, JH-AR and OEt9-AR. Increasing the DHT concentration to 100 or 1000 nM resulted in complete inhibition of OE33-AR proliferation in direct co-culture with the myofibroblasts ($p < 0.0001$) (Supplementary Fig. S6). The addition of enzalutamide to cultures blocked the DHT mediated inhibition of OE33-AR growth in monoculture ($p = 0.0001$), but did not significantly alter the outcome of direct co-culture with the myofibroblasts ($p = 0.544$) (Fig. 5c).
Translocation of AR in Direct Co-cultures

We investigated whether nuclear translocation of AR was altered in OE33-AR in direct co-culture. The results in Fig. 6 show that the OE33-AR treated with vehicle, in either monoculture or direct co-culture, had mild AR immunoreactivity in the cytoplasm and weak immunoreactivity in the nucleus, consistent with a lack of AR activation. With 10 nM DHT, there was complete translocation of AR to the nucleus and no AR in the cytoplasm of OE33-AR in monoculture or direct co-culture with NFFs. In contrast, there was a DHT dose-dependent distribution of AR in OE33-AR directly co-cultured with the myofibroblasts. With 10 nM DHT, there was moderate immunoreactivity in the nucleus, and mild to moderate immunoreactivity in the cytoplasm. With 100 nM DHT, there was complete nuclear translocation in about 50% of OE33-AR, with mild cytoplasmic and moderate nuclear immunoreactivity in remaining cells, and with 1000 nM DHT, there was complete nuclear translocation in all cells. The finding of both nuclear and cytoplasmic AR in OE33-AR directly co-cultured with myofibroblasts was similar to our findings in EAC resection samples where AR was expressed in both the nucleus and cytoplasm in the majority of tissues.

DHT Induced Expression of Androgen-Responsive Genes in Co-cultures

We next measured the effect of 10 nM DHT on the transcript levels of the androgen-responsive genes FKBP5, HMOX1 (heme oxygenase 1) and NDRG1 (N-myc downstream regulated 1) in OE33-AR in monoculture or direct co-culture. The OE33-ARs (GFP-positive) were sorted from NFFs (GFP-negative) or myofibroblasts (RFP-positive) in the direct co-cultures. The results in Fig. 7a show that the change in expression induced by DHT, induction, or repression was similar in the monoculture and co-cultures.

![Fig. 6](image-url) Nuclear translocation of AR induced by DHT in OE33-AR in direct co-culture with fibroblasts. Fluorescent micrographs of OE33-AR grown for 6 days in monoculture or direct co-culture with PShTerts with vehicle or 10, 100, or 1000 nM DHT, or in direct co-culture with NFFs with vehicle or 10 nM DHT. Cells were labeled with rabbit antihuman AR polyclonal IgG (clone N-20; 1 μg/mL in 1.5% goat serum) followed by secondary antibody Alexa Fluor 568 goat anti-rabbit IgG (2 μg/mL in 1.5% goat serum) and DAPI (1 μg/mL). Merged channel images were captured using a Zeiss confocal LSM 700 microscope. Scale bar is 75 μM.
although the extent of the change varied. This suggests that, although DHT did not inhibit OE33-AR growth in direct co-culture with myofibroblasts, there was still a change in the expression of androgen-responsive genes.

This response of OE33-AR to DHT in direct co-culture with myofibroblasts was also confirmed in two other AR-expressing EAC cell lines, JH-AR and OE19-AR (Supplementary Fig. S7). Co-cultured cells remained unsorted since the expression of androgen-responsive genes in myofibroblasts did not increase with DHT (FKBP5, \( p = 0.437 \); HMOX1, \( p = 0.244 \); NDRG1, \( p = 0.339 \)) (Supplementary Fig. S8). Unsorted OE33-AR co-cultures were used for comparison. FKBP5 was upregulated with 10 nM DHT in both monoculture and direct co-culture with myofibroblasts for all three AR-expressing EAC cell lines. HMOX1 and NDRG1 had limited androgen responsiveness for JH-AR, as shown previously in Supplementary Fig. S3. OE19-AR showed upregulation of HMOX1 and downregulation of NDRG1 with DHT in both monoculture and direct co-culture with myofibroblasts. While the use of unsorted cells may have reduced the significance of these changes, the results suggest that, for all three AR-expressing EAC cell lines, the myofibroblast blocked the growth inhibition but not the gene expression induced by DHT in monoculture.

To confirm that changes in gene expression were mediated through the AR signaling pathway, the anti-androgen enzalutamide was used. The results in Fig. 7b show the effect of enzalutamide on FKBP5 expression in OE33-AR directly co-cultured with myofibroblasts with 10 nM DHT. Again, we measured expression in unsorted cells because FKBP5 expressed by myofibroblasts did not increase with DHT. Enzalutamide reduced the expression both in monoculture (tenfold; \( p = 0.0002 \)) and in direct co-culture with the myofibroblasts (17-fold; \( p = 0.0003 \)), indicating that FKBP5 upregulation was mediated through AR. This confirms that, in co-culture conditions where DHT did not inhibit proliferation, DHT was still functional in regulating the expression of androgen-responsive genes through the AR signaling pathway.

**Discussion**

We have previously reported that nuclear localization of AR and/or expression of the androgen-responsive gene FKBP5 was associated with decreased survival in EAC, suggesting a role for androgens in this cancer [15]. Here we have extended that study to show that EAC resection tissues with a higher percentage of FKBP5 positive cells also had a higher proliferation index, showing a positive relationship between androgen signaling and cancer cell growth. We have investigated the effects of androgen on...
the behavior of AR-expressing EAC cell lines in vitro. The three EAC cell lines that we stably transduced with full-length human AR cDNA were responsive to androgen, as shown by DHT induced nuclear localization of the receptor, and dose-dependent changes in cell proliferation, morphology, and gene expression. The commonly used concentration of DHT, 10 nM, markedly altered the expression of androgen-responsive genes, but completely inhibited cell proliferation by inducing cell cycle arrest and senescence, which appeared to be inconsistent with our findings in patient tissues of increased proliferation and upregulation of androgen-responsive gene expression. We then found that lower concentrations of DHT around the IC50 allowed proliferation and induced significant, although smaller, changes in the expression of androgen-responsive genes. We also found that in direct co-culture with an immortalized myofibroblast cell line, PShTert, the growth inhibitory effect of 10 nM of DHT was largely nullified, but the effect on expression of androgen-responsive genes was unaltered.

Studies of the effect of androgens on cell proliferation have generated conflicting results. Inhibition of proliferation has been reported often, in normal and cancer cell lines from a range of tissues, either naturally expressing or transduced with AR [25–40]. No change or an increase in cell proliferation has also been reported [34, 40–43]. Why androgens in some cells increase and in others decrease proliferation is unclear. Many of the reported studies only used a single concentration of DHT, most commonly 10 nM. When dose–response studies have been reported, the IC50 for DHT inhibition of proliferation has been of the same order as we measured [29, 30, 33, 40]. Our finding that growth inhibition was associated with cell cycle arrest and the induction of senescence is also consistent with other reports [30, 44–46].

This is the first comprehensive study of the expression of androgen-responsive genes across a range of DHT concentrations, in parallel with measurements of proliferation, in AR-expressing cells that are growth inhibited by DHT. We showed a DHT dose-dependent alteration of the expression of these genes. This could be measured from DHT concentrations around the IC50, significantly lower than the 10 nM most commonly used for in vitro studies [27, 47, 48]. Most reported studies use a single dose of DHT given at the start of the culture period. Single doses around the IC50 resulted in small, but significant, changes in the expression of the androgen-responsive genes. Daily replenishment of the DHT increased the magnitude of the gene expression response without significantly altering proliferation. Daily replenishment, compared to a single dose, would be expected to more closely mimic the delivery of hormone in vivo. Our observations that DHT concentrations around the IC50 were sufficient to increase FKBP5 expression and allow proliferation were consistent with the association between FKBP5 and Ki-67 expression we measured in patient EAC resection specimens, which suggest our in vitro findings are clinically relevant.

The tumor microenvironment is an important determinant of the response of cancer cells to molecules such as hormones and drugs [24, 49]. We therefore examined the effect of androgens on our AR-expressing EAC cell lines in direct co-culture with fibroblasts, the primary cellular component of the microenvironment. The PShTert myofibroblast line, an immortalized activated fibroblast line which has properties typical of cancer-associated fibroblasts [20–22, 50–53], allowed the growth of AR-expressing EAC cells at concentrations of DHT that inhibited the growth of cells in monoculture, without affecting the gene expression changes induced by DHT in monoculture. These are the first reported studies of EAC cell lines in co-culture with fibroblasts or myofibroblasts. There are a number of studies of the effect on prostate cancer cell lines of co-culture with PShTert myofibroblasts, but none have investigated how fibroblasts affect the inhibition of cancer cell proliferation by androgen [20–22]. In our study, the myofibroblasts do not appear to be depleting androgen from the culture medium since there are still changes in the expression of androgen-responsive genes that can be blocked by the anti-androgen enzalutamide. This suggests that the microenvironment has the potential to modify the response of AR-expressing EAC cells to androgen in vivo. The mechanisms for this are unknown, warranting further investigation that may lead to new targets for manipulating the progression of EAC and possibly other androgen-responsive cancers.

This is the first study to show a positive association between androgen signaling and cancer cell proliferation in EAC, and that AR-expressing EAC cell lines respond to androgens in vitro. Proliferation of AR-expressing EAC cells in monoculture was inhibited by higher concentrations of DHT. Our in vitro findings suggest that, in cancer tissues in vivo, AR-expressing EAC cells at lower concentrations of DHT, or in the presence of activated fibroblasts in the microenvironment, would proliferate and DHT would alter the expression of androgen-responsive genes. Our findings are consistent with a role for androgen signaling in EAC.

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Author’s contribution HMP performed experiments. All authors contributed to the conception and design of the experiments, data analysis, and the writing of the manuscript.

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