A Positive Feedback Signaling Loop between ATM and the Vitamin D Receptor Is Critical for Cancer Chemoprevention by Vitamin D

Huei-Ju Ting1,2, Sayeda Yasmin-Karim3, Shian-Jang Yan6, Jong-Wei Hsu2, Tzu-Hua Lin2, Weisi Zeng1, James Messing1, Tzong-Jeng Sheu4, Bo-Ying Bao5, Willis X. Li6, Edward Messing1,2, and Yi-Fen Lee1,2

Abstract

Both epidemiologic and laboratory studies have shown the chemopreventive effects of 1α,25-dihydroxyvitamin D3 (1,25-VD) in tumorigenesis. However, understanding of the molecular mechanism by which 1,25-VD prevents tumorigenesis remains incomplete. In this study, we used an established mouse model of chemical carcinogenesis to investigate how 1,25-VD prevents malignant transformation. In this model, 1,25-VD promoted expression of the DNA repair genes RAD50 and ATM, both of which are critical for mediating the signaling responses to DNA damage. Correspondingly, 1,25-VD protected cells from genotoxic stress and growth inhibition by promoting double-strand break DNA repair. Depletion of the vitamin D receptor (VDR) reduced these genoprotective effects and drove malignant transformation that could not be prevented by 1,25-VD, defining an essential role for VDR in mediating the anticancer effects of 1,25-VD. Notably, genotoxic stress activated ATM and VDR through phosphorylation of VDR. Mutations in VDR at putative ATM phosphorylation sites impaired the ability of ATM to enhance VDR transactivation activity, diminishing 1,25-VD-mediated induction of ATM and RAD50 expression. Together, our findings identify a novel vitamin D-mediated chemopreventive mechanism involving a positive feedback loop between the DNA repair proteins ATM and VDR. Cancer Res; 72(4); 1–11. ©2011 AACR.

Introduction

The chemopreventive role of vitamin D in numerous types of cancer, including colorectal, breast, and prostate cancer was first suggested by epidemiologic studies (1–3). Further studies showed that vitamin D deficiency is associated with risk of cancer development (4–6). Preclinical studies support the chemopreventive effect of vitamin D in carcinogen-induced animal tumor models (7–9). Moreover, vitamin D receptor (VDR)-deficient mice exhibit higher carcinogen-induced tumor incidence in numerous tissues (10). Therefore, vitamin D supplementation and the activation of the VDR signaling pathway protect organisms from malignant transformation.

Cells are constantly challenged by spontaneous errors as well as environmental insults that lead to DNA damage. Accumulated genomic mutations from improperly repaired DNA damages can lead to malignant transformation. Several studies indicate that vitamin D attenuates DNA damage levels. Vitamin D can reduce ultraviolet light irradiation–induced DNA photoproducts and chromosome aberrations in diethyl-nitrosamine-treated liver (11–13). This could result from decreasing sources of genotoxic stress, for example, the antioxidant effect of vitamin D protects cells against oxidative insults (13–16). Recently, accumulated evidence from gene profiling shows that vitamin D induces the expression of DNA repair genes (17, 18), suggesting that vitamin D could facilitate DNA repair pathways.

DNA double-strand breaks (DSB), mostly caused by exposure to reactive oxygen species (ROS), ionizing radiation (IR), or generated during replication of single-strand breaks, are susceptible to exonucleases that lead to loss of large genomic regions. Once DSBs occur, formation of the Mre11/Rad50/NBS complex recruits the DNA damage response (DDR) signaling kinase, ATM (ataxia telangiectasia mutated), to the DSB and then the H2A histone family member X (H2Ax) is phosphorylated by ATM. The formation of foci containing serine 139–phosphorylated H2Ax (γ-H2Ax) is required for retaining mediator proteins, TP53BP1, MDC1, BRCA1, and the Mre11/Rad50/NBS complex at the DSB. These mediator proteins facilitate assembly of the DNA repair machinery to conduct the repair of DSB (19). There are 2 DSB repair pathways, homologous recombination (HR) and non-homologous end joining (NHEJ). HR uses Holliday
junction formation to facilitate strand transfer exchange between sister chromatids and is therefore less error prone. NHEJ is an efficient but more error-prone repair pathway (20). Malfunctioned DDR signaling proteins and repair machineries can have catastrophic consequences that lead to premature aging and tumorigenesis (21, 22). Recent findings discovered that the DDR signaling cascades ATM/Chk2/p53 pathway is upregulated by oncogenic stress, and inhibition of ATM leads to large and invasive tumor development (23, 24). These studies conclude that the ATM signaling pathway is an anticancer barrier of early-stage tumorigenesis.

In the current study, we show a cross-talk between DDR and vitamin D signaling in protecting DNA from genotoxic insults which is one mechanism mediating the chemopreventive effect of vitamin D against tumorigenesis.

Materials and Methods

Plasmids and reagents

NHEJ reporter, GFP-Pem1-Ad2, was a generous gift from Dr. Vera Gorbunova (University of Rochester, Rochester, NY). pDsRed-N1 was purchased from Clontech. Plasmids for GFP-based homologous recombination assay system, pDR-GFP and pCβAsce, were generous gifts from Dr. Maria Jasim (Memorial Sloan-Kettering Cancer Center, New York). The plasmids pGEX-KG-VDR-L, pGEX-KG-VDR-L1, and pGEX-KG-VDR-L2 were constructed by PCR amplifying VDR fragments with oligomers containing BamHI and XbaI sites, which were then inserted into pGEX-KG vectors (Promega). pcDNA3-flag-ATM and pcDNA-flag-ATMΔκd were generous gifts from Dr. Michael Kastan (St. Jude Children’s Research Hospital, Memphis, TN). pcDNA-flag-VDR was constructed by PCR amplifying VDR cDNA using oligomers containing BamHI and XbaI sites and then inserted into pcDNA-flag plasmids. pcDNA-flag-mutant VDRs were constructed by QuikChange Site-Directed Mutagenesis kit (Stratagene). Antibodies of embryonic inhibitor, KU55933, was purchased from Seltek. Mouse anti-human phospho-ATM (H-81), ATM, and (serine 1981) were purchased from Millipore; H2Ax was VDR (H-81), ATM, and

Xenograft mouse tumor model

The study was approved by the University of Rochester Committee on Animal Resources, and the mice were kept in a specific pathogen-free environment at the animal facility of the University of Rochester Medical Center. Young adult male athymic NCr-nu/nu mice (NCr-Frederick, Frederick, MD) at 8 to 10 weeks of age were subcutaneously injected with N-nitroso-N-methylurea (NMU)-transformed BPH-1 series cell lines into the dorsal lateral flank. Tumors were allowed to grow, measured weekly with calipers, and tumor volumes were calculated using the formula 0.532 × r1² × r2 (r1 < r2). Mice were weighed every week. Once they reached the endpoint (tumor size > 1 cm³), mice from all groups were euthanized by CO₂ and cervical dislocation.

Gene profiling

After NMU-induced transformation was completed, gene expression profiles were identified by the Oligo GEArray Human Cancer Microarray (SABioscience) according to manufacturer’s manual. Briefly, RNA harvested from cells was used as template for generating biotin-labeled probes. After hybridization of membranes containing cancer pathway-related genes with probes, membranes were washed and then developed for chemiluminescence imaging (VersaDoc, BioRad). After analysis using SABioscience online quantification software, genes with expression altered ≥1.5-fold were identified. By using GoMiner online software, altered genes between groups were functionally categorized and analyzed for statistical significance (P < 0.05).

GFP-based NHEJ and HR assay

These assays are described in the literature for NHEJ (25) and HR reporter assay (26). In brief, for NHEJ assay, BPH-1 cells were treated with EtOH or 1,25-VD for 24 hours and then transfected with GFP expression plasmids (HindIII digested; 0.5 μg/10⁵ cells) that express GFP only after repair of digested breaks and pDsRed-N1 (0.5 μg/10⁵ cells) for transfection control. Cells were harvested 2 days after transfection. The efficiency of NHEJ repair was analyzed using flow cytometry to quantify GFP-positive cells, which represent successful NHEJ DNA repair, and normalized by DsRed-positive cells for transfection efficiency. For HR assay, BPH-1 cells were treated as described above and transfected with plasmids for homologous recombination assay (pDR-GFP and pCβAsce, both at 0.5 μg/10⁵ cells). Cells transfected with pDR-GFP plasmid were selected by puromycin (2 μg/mL) for 2 days and then harvested at 6 days after transfection. Cells were subjected to analysis of GFP-positive cells under flow cytometry. Transfection was conducted by the Neon Transfection System (Invitrogen). Flow cytometry was carried out using FACSCanto-II (BD) and analyzed by Flowjo software.

Establishing siRNA-targeted VDR knockdown stable cell lines

Retrovirus-based plasmids (pSM2c) carrying siRNA-targeting VDR (siVDR) and control siRNA (SC) were purchased from Open Biosystem. Stable cell lines expressing these siRNAs
were generated according to the manufacturer’s manual (OligoEngine).

**Chromatin immunoprecipitation assay**

Chromatin immunoprecipitation assays were conducted according to a previous publication (27). DNA fragments containing VDREs in ATM and RAD50 genes were amplified by specific primer pairs (sequence available upon request).

**In vitro kinase assay**

ATM kinase assay was conducted according to a previous publication (28). Glutathione S—transferase (GST)-conjugated VDR fragment proteins were isolated according to the manufacturer’s manual (Promega).

**Immunoprecipitation and detection of phosphoserine**

BPH-1 cells on 10-cm dishes were treated with KU55933 for 2 hours and then exposed to 1 mmol/L H2O2 for 30 minutes and then changed to fresh normal medium. Proteins were harvested 3 hours after H2O2 challenge by radioimmunoprecipitation assay (RIPA) buffer containing protease inhibitor (cOmplete mini; Roche) and phosphatase inhibitor (1 mmol/L NaF and 1 mmol/L NaO,). Proteins were incubated with anti-phosphoserine and peroxidase-conjugated IgG fraction monoclonal mouse anti-rabbit IgG, light chain specific (Jackson ImmunoResearch Lab).

**Cell viability assay, quantitative PCR analysis, Western blotting assay, transient transfection and luciferase assays, and DNA pull-down assay**

Cell viability assay, quantitative PCR (Q-PCR) analysis, Western blotting assay, transient transfection and luciferase assays, and DNA pull-down assays were conducted according to previous publications (27, 29). Sequence of oligomers for amplifying actin, ATM, and RAD50 in Q-PCR assay are available upon request.

**Results**

**Vitamin D is chemopreventive in NMU transformation model**

To further investigate the chemopreventive effect of vitamin D supplementation, an *in vitro* model using NMU to transform prostate epithelial cells was applied (30). BPH-1 cells, a non-malignant human prostate epithelial cell line (31), were subjected to 3 repeated cycles of NMU treatment and then subcultured for another 6 passages to select clones with significant growth advantage (Fig. 1A).

In an anchorage-independent colony-forming assay, BPH-1 (NMU) cells formed many colonies indicating malignant transformation (Fig. 1B, middle). Importantly, 1,25-VD treatment can reduce NMU-induced BPH-1 malignant transformation with fewer colonies formed (Fig. 1B, bottom). The tumorigenicity was further confirmed in xenografted nude mice. The result showed that 60% of NMU-treated BPH-1 cells formed tumors (*n* = 10), but none of the dimethyl sulfoxide (DMSO)-treated BPH-1 cells (*n* = 16) or 1,25-VD–pretreated NMU-BPH-1 cells (*n* = 6) formed growing tumors (Fig. 1C). Tumor forming frequency between BPH-1 (NMU) and BPH-1 (VD + NMU) is 60% versus 0% (*P* = 0.034; Fisher exact test).

**DNA damage signaling genes and DNA repair are promoted by 1,25-VD**

To investigate the underlying molecular mechanism, we conducted gene profiling analysis using the human Cancer Pathway SuperArray (Supplementary Fig. S1A). We found that 63 genes were altered at a magnitude ≥1.5-fold by vitamin D in the NMU-treated groups, including 23 upregulated and 40 downregulated among a total of 440 genes. These genes were distributed in 4 statistically significant Gene Ontology (GO) categories (Supplementary Fig. S1B).

Two DSB repair genes, *ATM* and *RAD50*, were found to express higher in BPH-1 (VD + NMU) than in BPH-1 (NMU; Supplementary Fig. S1B and S1C). We suspected that 1,25-VD promotes or maintains DNA DSB damage repair.
capacity that is altered by carcinogen and other genotoxic challenges. We examined whether 1,25-VD treatment can also promote cell recovery from NMU-induced DSB by a γ-H2AX kinetic assay. Although NMU generates DNA alkylation that is repaired mostly by base excision repair, it is also known that DSB occurs upon alkylating reagents challenge (32) and that DSB repair protects cells from genotoxicity of alkylating agents (33). We found that NMU exposure increased the levels of γ-H2AX starting from 30 minutes, to a maximum at 2 hours in vehicle-treated cells, but the NMU-induced γ-H2AX induction was significantly reduced in the 1,25-VD–treated cells (Fig. 2A). Measuring γ-H2AX foci number found that 1,25-VD can reduce NMU-induced foci numbers at 2 hours after NMU exposure (Fig. 2B). This was not due to 1,25-VD altering cell-cycle distribution (Supplementary Fig. S2A). We also observed that the protective effect of 1,25-VD in H2O2 induced DSB at 6 hours (Supplementary Fig. S2B). These findings that γ-H2AX foci numbers and ATM activation (Supplementary Fig. S2C) by genotoxic insults were lower in 1,25-VD–treated groups suggest that 1,25-VD decreased DSB levels. RWPE-1 is another nonmalignant prostate epithelial cell line in which the DNA-protective effects of 1,25-VD were also observed (Supplementary Fig. S3). This could be a result of 1,25-VD reducing ROS generation as shown in our previous publication (29) and/or promoting DSB DNA repair when cells are faced with genotoxic insults.

To further understand whether 1,25-VD can promote DSB DNA repair, we examined the effects of 1,25-VD on 2 DSB repair pathways: NHEJ and HR repair. Treatment of BPH-1 cells with 1,25-VD significantly promoted the HR DNA repair capacity, as well as the NHEJ DNA repair capacity but to a lesser degree (Fig. 2C). Together, these results suggested that 1,25-VD protects cells from genotoxic insults through inducing DNA repair genes’ expression to promote the repair of DSB.

Figure 2. 1,25-VD protects cells from DSBs and promotes DSB DNA repair. A, 1,25-VD promotes cells’ recovery from NMU-induced DSB. BPH-1 cells were incubated with vehicle or 100 nmol/L 1,25-VD for 24 hours and then exposed to NMU (100 μg/mL) for 1 hour. DNA DSB marker γ-H2AX was detected at designated time points by Western blotting. After being normalized with H2AX, the relative intensity of γ-H2AX to 0 hours was calculated and plotted. **, P < 0.01 compared with EtOH-treated group at the same time point. B, BPH-1 cells were seeded at 10^5 cells per well on chamber slide and then treated with EtOH or 100 nmol/L 1,25-VD the next day. After 24 hours, cells were treated with or without H2O2 (0.1 mmol/L) for 30 minutes and then replenished with fresh medium containing EtOH or 1,25-VD. Cells were fixed at indicated time points and processed for staining with an antibody for γ-H2AX. Pictures were taken by confocal microscopy and representative pictures are shown (scale bar, 15 μm). γ-H2AX foci numbers were counted and then average foci numbers per cell ± SD were plotted. **, P < 0.01. C, 1,25-VD promotes NHEJ and HR repair. BPH-1 cells were treated with EtOH or 100 nmol/L 1,25-VD for 24 hours and then cotransfected with HindIII-digested NHEJ reporter and DsRed-expressing vector (NHEJ assay) or pD-βd-GFP and pCtδαSce plasmids (homologous recombination assay). Cells were continuously treated with EtOH or 1,25-VD until harvest. The percentage of GFP^+ and DsRed^+ cells were determined by fluorescence-activated cell-sorting (FACS) analysis. The relative repair efficiency was calculated by comparing with EtOH-treated group. *, P < 0.05; **, P < 0.01, compared with EtOH group (n = 3). DAPI, 4′,6-diamidino-2-phenylindole.
VDRs protect cells from genotoxicity and tumorigenesis

To explore whether VDR mediates the protective role of 1,25-VD against genotoxicity, we established siRNA VDR knocked down (BPH-1siVDR) versus scramble control (BPH-1SC) BPH-1 cell lines. VDR knockdown efficiency was 70% and VDR transactivity measured by the induction of CYP24, ATM, and RAD50 expression was reduced (Fig. 3A). The protective effect of 1,25-VD against H2O2 and IR-induced cell death was lost in BPH-1siVDR cells compared with BPH-1SC cells (Fig. 3B; Supplementary Fig. S4A). Moreover, the effect of 1,25-VD facilitating DS recovery could only be seen in BPH-1SC but not in BPH-1siVDR cells (Fig. 3C). All these data support role of VDR in protecting cells from genotoxicity.

Next, we examined the role of VDR in the chemopreventive effect of 1,25-VD. Knockdown of VDR sufficiently induced colony formation, representing tumorigenicity, even without NMU treatment (Fig. 3D, DMSO-treated BPH-1siVDR vs. BPH-1SC). This is observed from 2 independent stable clones and is therefore unlikely to result from insertion of transgenes into critical genome sites. This result strongly supports the tumor-suppressive role of VDR. However, 1,25-VD treatment can still reduce NMU-induced tumorigenicity in BPH-1siVDR cells. (The representative photos of colonies are shown in Supplementary Fig. S4B.)

These results suggest that 1,25-VD has VDR-independent chemopreventive effects or can still function through the residual VDR signal amplified by ATM-VDR signaling loop during carcinogen challenges. To clarify whether VDR is essential in mediating the chemopreventative effect of 1,25-VD, we obtained VDR-null MEFs from VDR knockout mice (VDRko) to examine whether 1,25-VD can still suppress the tumorigenesis process in the absence of VDR. At the eleventh passage, VDRko MEFs spontaneously formed colonies in soft agar assays whereas VDR heterozygous deletion (VDRhet) MEFs formed very few colonies. Treatment of 1,25-VD can still reduce NMU-induced tumorigenicity in BPH-1siVDR cells.

To test whether VDR is a downstream target of DDR phosphorylation sites in GST-VDR-L and GST-VDR-L1 fragment to generate GST-dmVDR-L and GST-dmVDR-L1. The result showed that the phosphorylation was reduced but not completely abolished in these mutated fragments. VDR transactivation activity was enhanced by ATM but not by ATM dead mutant (ATMkd; Fig. 4C). The single serine-mutated VDRs (VDRs208g and VDR s222a) and double serine mutant VDR (dmVDR) can respond to 1,25-VD–induced transactivity (Fig. 4C). However, only the dmVDR completely lost response to ATM (Fig. 4C). These results indicate that ATM enhances VDR activity through phosphorylation of 2 amino acids, ser208 and ser222. Similarly, overexpression of ATM promotes endogenous VDR transactivity in BPH-1 cells (Fig. 4D).

We next found that ATM-specific inhibitor, KU55933, inhibited H2O2-enhanced VDR phosphorylation and transactivity (Fig. 4E and F). Therefore, DNA damage induced phosphorylation and transactivation of VDR through ATM. We further compared the DNA-binding ability of VDR and dmVDR. The result showed that H2O2 and 1,25-VD-induced DNA binding ability of wild-type (wt) VDR. On the other hand, dmVDR bound VDRE but did not respond to 1,25-VD and H2O2 stimulation as strongly as wtVDR (Fig. 4G). Taken together, these data suggest that DNA damage signals enhance VDR activity through phosphorylation at s208 and s222 by ATM.

ATM phosphorylates and activates VDR transactivity

ATM kinase is a key molecule that senses DNA damage and activates the DDR signaling cascade. H2O2 exposure quickly activates ATM within 30 minutes at concentrations starting at 0.1 mmol/L (Supplementary Fig. S5A). Interestingly, we observed that the 1,25-VD–induced VDR transactivity was further enhanced by H2O2 challenge (Supplementary Fig. S5B and S5C).

To test whether VDR is a downstream target of DDR signaling kinases upon induction of genotoxic stress stimuli, we screened for the phosphorylation motif on VDR using the web tool Scansite (http://scansite.mit.edu/). Three putative ATM target sites were identified (Fig. 4A).

In vitro kinase assays revealed that ATM phosphorylated the VDR-L and VDR-L1 fragments both containing s208 and s222 but not the VDR-L2 (Fig. 4B, bottom). In the sample containing ATM and VDR-L1, 3 phosphorylated bands were found including ATM, VDR-L1 (†), and an unknown band (#) that might be degraded VDR. ATM autophosphorylation and BRCA1 phosphorylation bands were observed as positive control.

We further mutated the putative phosphorylation sites in GST-VDR-L and GST-VDR-L1 fragment to generate GST-dmVDR-L and GST-dmVDR-L1. The result showed that the phosphorylation was reduced but not completely abolished in these mutated fragments. VDR transactivation activity was enhanced by ATM but not by ATM dead mutant (ATMkd; Fig. 4C). The single serine-mutated VDRs (VDRs208g and VDR s222a) and double serine mutant VDR (dmVDR) can respond to 1,25-VD–induced transactivity (Fig. 4C). However, only the dmVDR completely lost response to ATM (Fig. 4C). These results indicate that ATM enhances VDR activity through phosphorylation of 2 amino acids, ser208 and ser222. Similarly, overexpression of ATM promotes endogenous VDR transactivity in BPH-1 cells (Fig. 4D).

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ATM-modified VDR is required for VDR cellular protective effects against H2O2 challenge

On the basis of our results showing that 1,25-VD treatment can protect cells against DNA damage insult (Fig. 2) and VDR is a downstream target of the ATM cascade (Fig. 4), we then hypothesized that DNA-protective effect of 1,25-VD-VDR is modulated by ATM, such that phosphorylation of VDR by ATM is critical for this protective effect.

To test this hypothesis, we compared cell survival upon H2O2 challenge in cells overexpressing wtVDR and dmVDR. Because the transactivity of dmVDR cannot be stimulated by ATM (Fig. 4C), dmVDR serves as a tool to delineate the cellular function of ATM-phosphorylated VDR. BPH-1 cells were transiently over-expressed with wtVDR or dmVDR, treated with 1,25-VD for 24 hours, and then exposed to H2O2 to determine cell survival. The results showed that cell survival was increased in BPH-1 cells overexpressing wtVDR and treatment with 1,25-VD promoted cell survival.

In contrast, cells overexpressing dmVDR had a similar basal survival rate as compared with vector control cells (EHO group) but the protective effect of 1,25-VD was lost in these dmVDR cells (Fig. 5A). The overexpression level of wtVDR and dmVDR mRNA and proteins were confirmed to be equivalent (Fig. 5B). In addition, we examined the expression of RAD50 induced by 1,25-VD in cells expressing various types of VDR. The expression of ATM and RAD50 was induced by 1,25-VD in BPH-1 (vector) cells and further induced in BPH-1 (VDR) cells which were overexpressed with wtVDR. Interestingly, the
Figure 3. The protective effect of 1,25-VD against genotoxic challenges in VDR-depleted cells. A, 1,25-VD induced gene expression in VDR-depleted cells. BPH-1SC and BPH-1siVDR cells were treated with EtOH or 100 nmol/L 1,25-VD for 24 hours and then harvested for RNA extraction. The expression of indicated genes was measured by Q-PCR. The relative expression compared with BPH-1SC treated with EtOH was calculated and mean ± SD plotted. B, the protective effect of 1,25-VD against genotoxicity in VDR-depleted cells. BPH-1SC and BPH-1siVDR cells were treated with 100 nmol/L 1,25-VD for 24 hours and then exposed to different doses of \( \text{H}_2\text{O}_2 \). After 6 days, MTT assays were conducted. Percentages of surviving cells in each treatment group compared with no \( \text{H}_2\text{O}_2 \) exposure were calculated (*, \( P < 0.05 \) compared with EtOH with same dosage of \( \text{H}_2\text{O}_2 \)-treated group; \( \text{n} = 3 \)). C, the effects of 1,25-VD on genotoxic agent–induced DSBs. BPH-1SC or BPH-1siVDR cells were incubated with EtOH or 100 nmol/L 1,25-VD for 24 hours and then exposed to \( \text{H}_2\text{O}_2 \) (0.1 mmol/L) for 30 minutes. At designated time points, proteins were harvested for detecting \( \gamma\)-H2AX level. The relative intensity of \( \gamma\)-H2AX (normalized by actin) compared with 0 hours was calculated and plotted (*, \( P < 0.05 \) compared with EtOH-treated groups at the same time points). D, the chemopreventive effect of 1,25-VD in VDR-depleted cells. NMU-induced malignant transformation was conducted in BPH-1SC and BPH-1siVDR cells as in Fig. 1A. Soft agar colony formation assay was conducted for examining tumorigenicity. Colony numbers were counted and averaged from 9 different fields of each well under microscope at 50× magnification. Average colony numbers from 2 independent experiments of 2 independent clones of BPH-1SC and BPH-1siVDR stable cell lines were calculated and mean ± SD plotted. E, MEF cells from VDRhet and VDRko mice were analyzed for their tumorigenicity after EtOH or 100 nmol/L 1,25-VD treatment for 24 hours three times. Colony numbers were counted and accumulated from 10 different fields of each well under microscope. Average colony numbers from 3 independent counting were calculated and mean ± SD plotted.
induction of ATM and RAD50 by 1,25-VD is abolished in the BPH-1 (dmVDR) cells (Fig. 5C).

To further validate that ATM-phosphorylated VDR is critical for 1,25-VD protective effect against genotoxicity, the different VDRs (wt vs. dmVDR) were reexpressed in BPH-1siVDR cells (Fig. 5D). The result showed that 1,25-VD has no protective effect on BPH-1siVDR cells (Fig. 5D, left), and wtVDR reexpression rescued the protective effect of 1,25-VD (middle) but not dmVDR (right). In summary, we conclude that activation of VDR by ATM phosphorylation is essential for the DNA-protective effect of 1,25-VD and this 1,25-VD/VDR protective effect provides one functional mechanism mediating antitumorigenic effect of vitamin D.
Discussion

Here, we investigate the mechanisms underlying the chemopreventive effect of vitamin D in tumorigenesis. These results show that 1,25-VD treatment can protect cells from carcinogen-induced genotoxic stress via VDR-mediated transcriptional upregulation of DNA repair genes, ATM and RAD50, and thereby facilitate DSB repair. Reciprocally, DDR signaling kinase ATM phosphorylates and enhances the transactivity of VDR. Figure 6 illustrates that activation of the ATM-VDR–positive signaling loop upon carcinogen/oncogenic stress contributes to the protective effect of 1,25-VD.
1,25-VD against genomic insult–induced cell death and malignant transformation.

Currently, the effect of vitamin D supplementation in human cancer prevention is still under evaluation. One major concern for using vitamin D supplements in tumor prevention is the hypercalcemia side effect due to high dosage vitamin D uptake. In this study, we found that a high concentration of 1,25-VD (100 nmol/L) is required to prevent BPH-1 from malignant transformation. It is estimated that more than 4,000 IU/kg vitamin D supplement is required to reach such dosage. However, the current physiologic recommended upper limit dosage of vitamin D supplementation is 2,000 IU. One possibility for this high dosage requirement in BPH-1 is that it is immortalized by SV40 T antigens which disrupts vitamin D signaling pathways by squelching p53 and/or downregulating VDR expression (34, 35). We predict that the physiologic concentration of 1,25-VD will be sufficient to activate the antitumorigenic effect of VDR in normal cells that contain functional p53 and Rb. Therefore, the optimal dosage of vitamin D supplement required for preventing cancer needs to be further studied.

Phosphorylation of ATM putative target sites, s208 and s222, in VDR can enhance VDR transactivity and that mutation of both serines in VDR (dmVDR) impairs ATM-VDR signaling loop. ATM enhances VDR transactivity by increasing the DNA-binding ability of VDR as evidenced by VDRE-binding studies showing that genotoxic stress induces VDR-VDR complexes of wtVDR but not of dmVDR (Fig. 4G; Supplementary Fig. S5C). As stated in the literature, casein kinase II can also phosphorylate VDR s208 (36). Phosphorylation of VDR s208 does not affect ligand binding, DNA binding, or RXR heterodimerization of VDR but does promote the recruitment of DRIP205 as noted by the enhancement of VDR transactivity (36, 37). Therefore, we expect that DRIP205 recruitment is another mechanism by which ATM enhances VDR transactivity.

The positive regulatory loop constituted by ATM-VDR led us to predict that 1,25-VD treatment will act synergistically with DDR signaling in suppressing tumorigenesis. Our study showed that once the ATM-VDR loop is broken by either siVDR or dmVDR, the 1,25-VD regulation of ATM and RAD50 expression and protective effect of 1,25-VD against genotoxic challenges are lost (Figs. 3 and 5). These results support a critical role of the intact ATM-VDR loop in mediating DNA-protective effect of vitamin D. Interestingly, the chemopreventive effect of 1,25-VD is not abolished by siVDR (Fig. 3). This suggests the DNA-protective effect of 1,25-VD might not be the only chemopreventive mechanism of 1,25-VD. Other cancer pathways identified in array can also contribute to the antitumorigenic effect of 1,25-VD (Supplementary Fig. S1B). Remaining ATM-VDR signaling is possibly sufficient in mediating 1,25-VD regulation of other cancer pathways. Indeed, complete depletion of VDR by using VDRko MEF abolishes the tumor-suppressive effect of 1,25-VD. This supports the essential role of ATM-VDR loop in mediating the antitumorigenic effect of vitamin D. In addition, we challenged MEFs with NMU (3 cycles and one passage) but failed to induce more colonies (data not shown). More passages might be required to enrich the malignant population to examine the chemopreventive effect of vitamin D during carcinogen-induced malignant transformation. Meanwhile, we also immortalized MEFs by SV40LT. Interestingly, these immortalized MEFs from VDRko and VDRhet form colonies spontaneously. Once MEFs are being transformed by SV40LT, vitamin D is no longer able to reduce their colony-forming ability (data not shown). Therefore, cells should be treated with vitamin D prior malignant transformation to show its chemopreventive activity. On the other hand, we expect that if ATM-phosphorylated VDR is constitutively expressed, the chemopreventive effect of 1,25-VD could be amplified. Exploiting the ATM-VDR signaling loop by combination of 1,25-VD and ATM activators is therefore potentially a novel strategy to prevent cancer while mitigating the hypercalcemia side effects of vitamin D.

Overall, our efforts in mechanistic studies support the role of vitamin D in guarding genomic integrity through regulation of genes involved in anti-oxidation (29) and DNA repair. Other potential pathways including inducing Ras signaling and suppressing antiapoptotic genes demand future study. Another gene profiling study of 1,25-VD–treated RWPE-1 cells identifies the WNT, Notch, NF-kB, insulin—like growth factor 1, and inflammation signaling, those are also potential chemopreventive mechanisms of 1,25-VD worth pursuing (38). In patients with cancer, the DNA-protective effect of vitamin D can compromise the efficacy of therapies targeting DNA, such as radiotherapy and chemotherapy. Therefore, the status of vitamin D

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**Figure 6.** Schematic diagram of DNA protective and antitumorigenic mechanism of ATM-VDR signaling axis. Carcinogen and other oncogenic stress can stimulate ATM triggering DDR signaling cascade to activate p53 and other factors in forming anticancer barrier in early-stage cancer initiation. Activated ATM can also phosphorylate and enhance VDR transactivity in regulating expression of ATM and RAD50, hence promoting DSB repair guarding genome integrity. Other anticancer pathways, such as senescence and apoptosis, are potentially regulated by 1,25-VD-VDR signaling.
signaling, including the availability of ligands and receptor, could serve as a prognostic biomarker for predicting the response of patients to radiotherapy and chemotherapy. On the other hand, vitamin D supplementation might be one supportive treatment for protection against DNA damage caused by radiation exposure in healthy individuals. Finally, a novel vitamin D–based chemopreventive strategy could be developed on the basis of the ATM-VDR–positive signaling loop.

References


A Positive Feedback Signaling Loop between ATM and the Vitamin D Receptor Is Critical for Cancer Chemoprevention by Vitamin D

Huei-Ju Ting, Sayeda Yasmin-Karim, Shian-Jang Yan, et al.

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