Androgen receptor transactivates KSHV noncoding RNA PAN to promote lytic replication–mediated oncogenesis: A mechanism of sex disparity in KS

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Abstract

Kaposi’s sarcoma-associated herpesvirus (KSHV) preferentially infects and causes Kaposi’s sarcoma (KS) in male patients. However, the biological mechanisms are largely unknown. This study was novel in confirming the extensive nuclear distribution of the androgen receptor (AR) and its co-localization with viral oncoprotein of latency-associated nuclear antigen in KS lesions, indicating a transcription way of AR in KS pathogenesis. The endogenous AR was also remarkably higher in KSHV-positive B cells than in KSHV-negative cells and responded to the ligand treatment of 5α-dihydrotestosterone (DHT), the agonist of AR. Then, the anti-AR antibody-based chromatin immunoprecipitation (ChIP)-associated sequencing was used to identify the target viral genes of AR, revealing that the AR bound to multiple regions of lytic genes in the KSHV genome. The highest peak was enriched in the core promoter sequence of polyadenylate nuclear RNA (PAN), and the physical interaction was verified by ChIP–polymerase chain reaction (PCR) and the electrophoretic mobility shift assay (EMSA). Consistently, male steroid treatment significantly transactivated the promoter activity of PAN in luciferase reporter assay, consequently leading to extensive lytic gene expression and KSHV production as determined by real-time quantitative PCR, and the deletion of nuclear localization signals of AR resulted in the loss of nuclear transport and transcriptional activity in the presence of androgen and thus impaired the expression of PAN RNA. Oncogenically, this study identified that the AR was a functional prerequisite for cell invasion, especially under the context of KSHV reactivation, through hijacking the PAN as a critical effector. Taken together, a novel mechanism from male sex steroids to viral noncoding RNA was identified, which might provide a clue to understanding the male propensity in KS.
Although the incidence of Kaposi’s sarcoma (KS) is higher in men, little is known about the mechanisms by which male sex steroids contribute to this disparity. The present study confirmed the striking expression of the androgen receptor (AR) and its concordant nuclear distribution in KS tissues. High-throughput chromatin immunoprecipitation sequencing analysis showed that the AR had extensive binding sites in the KSHV genome, in which the highest enriched gene was PAN. PAN also exhibited the strongest upregulation of promoter activity and RNA transcription among various KSHV lytic genes after the male hormone treatment. Specifically, the effect was a result of the DNA-binding capability of AR to PAN promoter. Moreover, the AR induced dramatic cell invasion, especially under KSHV lytic replication, and the effect was greatly impaired by the inhibitory effect of siRNA on PAN RNA. This study provided a unique insight into the reason why KS occurred predominantly in men.

Introduction

Kaposi’s sarcoma (KS) occurs predominantly in men in the classic type or endemic population, and the male-to-female ratio ranges from 10:1 to 15:1 [1–3]. The male propensity is partly explained by sex-related differences in environmental influences and behaviors, for example, iron uptake and alcohol consumption, implying that host genetic factors are crucial determinants [4,5]. Researchers have made great efforts to elucidate the roles of host hormones in virus infection and/or pathogenesis, considering severe progression in male patients during the COVID-19 pandemic [6–8]. Regarding human oncoviruses, the hepatitis B virus (HBV) is currently the only virus that shows a positive regulatory circuit to male steroids [9–12]. The HBV X gene maintains the activation of AR, which in turn is associated with the enhancer I domain of HBV and promotes virus transcription and replication, thus increasing the risk of hepatocarcinogenesis [12]. The KS incidence and the infection by its causative agent, KS-associated herpesvirus (KSHV), also exhibited male propensity in a diverse cohort study. In Africa, the age-standardized incidence rate of KS was three times more in men than in women [13–15], and the KSHV virus load has been found to be higher in men [16,17]. However, the underlying mechanism needs further investigation.

It was once recognized that the latency in the biphasic life cycle of KSHV was majorly responsible for KS oncogenesis until highly active antiretroviral therapy implementation targeting KSHV lytic infection dramatically declined AIDS-KS incidence, indicating the pathological role of virus reactivation in KSHV oncogenesis [18–20]. In lytic-infected cells, the entire viral genomic transcription cascade occurred; 80% of this was referred to as a long noncoding transcript, polyadenylated nuclear RNA (PAN) [21]. As an emerging compartment to cellular noncoding RNA, the role of PAN RNA in KS pathogenesis and its crosstalk with cellular proteins remain obscure [22–25]. PAN RNA was demonstrated to be a structural scaffold for organizing demethylases UTX, JMJD3, or polycomb repressive complex 2 (PRC2) to KSHV lytic genes [23], while the related transcriptional events were unknown, especially from the host aspect.

To our knowledge, this was the first study to identify a striking expression of AR in the nuclei of KS spindle cells at a very high level, compared with the sporadic cytoplasm distribution from control tissues. As a notable steroid hormone receptor of the nuclear receptor family, the AR is classically thought to function as transcriptional regulators in the nucleus. Upon ligand binding, the intracellular AR translocates to the nucleus and then engages with specific androgen-responsive elements (AREs) to modulate gene transcription [26]. Therefore, the
restricted expression of AR in the nucleus of KS tissues indicates a possible transcription mechanism of AR for crosstalk with the oncogenic KSHV.

Using ChIP-seq analysis, extensive binding sites of AR to the KSHV genome of several lytic genes were observed, and PAN exhibited the highest enrichment. We further determined the direct interaction between AR and PAN promoter by electrophoretic mobility shift assay (EMSA) assay using the recombinant protein of AR mutant, whose DNA-binding domain was deleted. Functionally speaking, male steroid hormone treatment led to significantly increased KSHV lytic gene expression, especially PAN, and consequently the infectious progeny production, as determined by luciferase reporter assay and quantitative real-time polymerase chain reaction (qRT-PCR). Mechanically speaking, it was the ligand-activated nuclear transport of AR that determines the successful expression of downstream PAN RNA. Finally, the tumorigenic capability of AR overexpression was most significantly impaired by RNA interference of PAN upon KSHV lytic replication in the cell invasion assay. Taken together, the results of our study indicated that male hormones were functionally involved in the male propensity of KS by serving as a canonical transcription factor to viral noncoding RNA PAN.

Materials and methods

Ethics statement

All experiments in the present study were conducted according to the principles mentioned in the Declaration of Helsinki. The use of the clinical sections of classic KS tissue specimens, normal skin, and non-KS angioma tissues was reviewed and approved by the Institutional Ethics Committee of the First Teaching Hospital of Xinjiang Medical University (Urumqi, Xinjiang, China; Study protocol # 20130216–53). Written informed consent was obtained from all participants, and all samples were anonymized. All tissue specimens had a pathological diagnosis except normal skin samples. All samples were collected from Xinjiang province, northwestern China.

Cell lines, antibodies, reagents, and plasmids

KS-derived endothelial SLK cells and 293T cells were maintained in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum (HyClone, Cytiva, USA). A derivative of SLK (termed iSLK) expressing a doxycycline-inducible replication and transcription activator (RTA) transgene was latently infected with a recombinant KSHV.219 virus, referred to as iSLK.219 cells, which constitutively expressed puromycin N-acetyl-transferase and green fluorescent protein (GFP); red fluorescent protein (RFP) was expressed during lytic replication [27]. The iSLK.219 cells were cultured as SLK cells but maintained using 1% penicillin–streptomycin, 1 μg/mL puromycin, 250 μg/mL G418, and 1 mg/mL hygromycin B. KSHV-negative B cells [BJAB (Epstein-Barr virus (EBV)-negative B lymphoma cell line) and DG75 (EBV-negative primary abdominal B lymphoma cell line)] and KSHV-positive primary effusion lymphoma cells [KSHV stably transfected BJAB (KSHV-BJAB) cells, BCBL1 (body-cavity-based lymphoma cell line)], BC3, and JSC1 (naturally EBV and KSHV co-infected) were generously provided by Dr. Erle S Robertson (University of Pennsylvania, USA) and maintained in Roswell Park Memorial Institute 1640 medium supplemented with 10% FBS (HyClone). Androgen-sensitive human prostate adenocarcinoma cells (LNCap) (TCHu173) and androgen-independent prostate cancer cells (PC3) (TCHu158) were purchased from cell bank/stem cell bank of Shanghai Institutes of Biological Sciences, Chinese Academy of Sciences (Shanghai, China). Charcoal-stripped fetal bovine serum (CD-FBS) (Sigma), from which endogenous hormones and growth factors were removed, was used for cell culture before DHT treatment.

The antibodies and reagents used were as follows: anti-AR antibody (Abcam, ab74272), anti-latency-associated nuclear antigen (LANA) monoclonal antibody (ABI, LN53), anti-
immunoglobulin G (IgG) antibody (Abcam, ab48386), anti-hemagglutinin (HA) antibody (Cell Signaling Technology, #3724), anti-Flag antibody (Cell Signaling Technology, #8146), and anti-glyceraldehyde-3-phosphate dehydrogenase (GAPDH) antibody (Cell Signaling Technology, #2118). The secondary antibodies (Thermo Fisher Scientific) included goat anti-rabbit antibodies conjugated with Alexa Fluor 488 [A-11094], 555 [A27017], and 680 [A27020], and goat anti-mouse antibodies conjugated with Alexa Fluor 488 [A-11001], 555 [A-21422], and 680 [A-28183]). DHT (Sigma–Aldrich, D-073), protease inhibitor cocktail set III (Millipore, 539134), phosphatase inhibitor cocktail (Santa Cruz Biotechnology, sc-45044), doxycycline hyclate (Sigma–Aldrich, D9891-25G-9), hygromycin (Sigma–Aldrich, V900372-1G), puromycin (Sigma–Aldrich, OG5541-5UG), and G418 disulfate salt (Sigma–Aldrich, A1720-5G) were used. Control siRNA (fluorescein isothiocyanate conjugate)-A (Santa Cruz Biotechnology, sc-36869), AR siRNA (Santa Cruz Biotechnology, sc-29204), Lipofectamine 3000 (Thermo Fisher Scientific, L3000001), 4',6-diamidino-2-phenylindole (DAPI) (Beyotime, c1002), anti-Flag M2 affinity gel (Sigma–Aldrich, A2220-5 mL), recombinant protein A/G agarose (Invitrogen, 5948-014/15920-010), glutathione sepharose 4B (GE Healthcare, 17-0756-01), SYBR Green Master Mix Kit (Toyobo, QPK-201), Corning Transwell polycarbonate membrane cell culture inserts (Corning, CLS3422-48EA), collagen type I cell ware coverslips (BD Biosciences, 354089), Dual-Luciferase Reporter Assay System (Promega, E1910), and Cell Counting Kit 8 (ab228554, Abcam) were also used in the study.

Plasmids: Flag-tagged full-length AR (1–919 aa, ref [M23263.1]) and its truncated constructs [△DNA-binding domain (DBD)+Hinge+ligand-binding domain (LBD), 1–537 aa; △NTD+DBD, 669–919 aa] were constructed by cloning the corresponding fragments into the pCDH-CMV-IRES-sf3-blast vectors (GE Healthcare). The glutathione S-transferase (GST)-fused truncated constructs of AR (NTD+DBD, 1–625 aa; NTD, 1–537 aa) were obtained by cloning the corresponding fragments into the pGEX-4T-1 vector. The reporter plasmids pGL3.0-RTAp, pGL3.0-LANAp, and pGL3.0-PANp were described previously [28,29]. A series of truncated reporter plasmids spanning ~2000 to +1 bp in the PAN promoter, each at ~200-bp interval and schematically shown in Fig 5A, were constructed by amplifying corresponding DNA fragments from the genomic DNA of induced iSLK.219 cells as the template and cloned into the pGL3.0 vector. All of the primers are listed in S1 Table.

**Immunohistochemical analysis**

Twelve KS tissues, one normal skin tissue, and six non-KS angioma tissues were collected. All the samples were formalin-fixed, paraffin-embedded, and immunostained with anti-AR or anti-LANA antibodies at a ratio of 1:200 as previously described [30]. The results were processed using Image-Pro plus 6.0 image analysis system (Media Cybernetics, MD, USA).

**Immunoblotting**

The cell lysates were prepared in radioimmunoprecipitation assay (RIPA) buffer (50mM Tris-HCl [pH 7.4], 150 mM NaCl, and 0.5% Triton X-100) containing protease and phosphatase inhibitors. Proteins were separated by sodium dodecyl sulfate–polyacrylamide gel electrophoresis and transferred to polyvinylidene difluoride membranes for immunoblotting with the indicated antibodies.

**Immunofluorescence assay**

The cells were fixed with 4% paraformaldehyde for 30 min at room temperature, permeabilized with 0.5% Triton X-100, blocked with 20% normal goat serum (Life Technologies), and reacted with indicated antibodies followed by fluorescent dye-conjugated secondary antibodies.
(1:1000). The dilution factor for individual primary antibodies was generally 1:200. The cover-slips were mounted with an anti-fade mounting medium (Beyotime) and photographed using a digital camera and software (Olympus FV-1200).

**Chromatin immunoprecipitation assay (ChIP)**

The ChIP protocol was adapted from the Rockland website with some modifications [31]. The cells were crosslinked in the medium with 1% formaldehyde for 15 min at room temperature and quenched with 0.125M glycine. After crosslinking, the cells were washed with phosphate-buffered saline twice and resuspended in 1 mL of buffer A (10mM Tris-HCl, pH 7.4, 10mM NaCl, 3mM MgCl₂, 0.2% Triton X-100, 1mM dithiothreitol, 0.5mM EDTA, and 0.2mM phenylmethylsulfonyl fluoride (PMSF) for 10 min at 4°C. The extracted nuclei were pelleted by centrifugation at 1300g for 5 min. The nuclei were lysed in sodium dodecyl sulfate (SDS) lysis buffer (50mM HEPES, 1mM EDTA, 1% SDS, and 1mM PMSF) for 10 min on ice. The lysates were subjected to sonication to obtain 200- to 500-bp fragments of DNA (Sonics; cycle, three pulses of 3 s; amplitude, 30%–35%) and then centrifuged at 12,000 rpm at 4°C for 10 min to obtain the supernatants. The samples were precleared with pretreated protein A or G beads (1 mg/mL bovine serum albumin, 1 mg/mL sperm DNA, 20% beads) for 2 h at 4°C. Then, 5% of the supernatants were kept as the input, and the remainder was divided into groups according to the experiment. The aliquots were incubated with pretreated protein A or G beads and 5 μg of anti-AR antibody or control rabbit immunoglobulin IgG antibody overnight at 4°C. After extensive washing with RIPA buffer, wash buffer (20mM Tris-HCl, pH 8.0, 1 mM EDTA, 250 mM LiCl, 0.5% NP-40, and 1 mM PMSF), and Tris-EDTA (TE) buffer (10 mM Tris-HCl, pH 8.0, 1 mM EDTA) (four times each), the beads were resuspended in TE buffer. The resuspended beads were subjected to RNase A and proteinase K digestion, and the crosslinking was reversed at 65°C for 8–10 h. DNA was recycled with a DNA purification kit (Qiagen, Shanghai, China). The primer sequences are summarized in S1 Table.

**ChIP-seq**

The procedures for ChIP for sequencing (ChIP-seq) were similar to those for ChIP described earlier, except for the pretreated beads to which sperm DNA was not added. More cells were needed for ChIP-seq, and 5 × 10⁷ iSLK.219 cells were harvested for the experiment. The lysates were sonicated to obtain DNA fragments of ~200 bp. The enriched DNA was subjected to library preparation for high-throughput sequencing. The ChIP-seq library was prepared according to Illumina instructions. The size selection range for the library was 100–400 bp.

**Bioinformatics analysis of ChIP-seq data**

The ChIP-seq data were aligned with the human genome (hg19) and the KSHV genome (HQ404500 plus 35 copies of the TR [U75699.1]) using Bowtie2 [32]; only one mismatch was allowed. The output files were subjected to peak calling with model-based analysis of ChIP-seq (MACS), as described previously [33,34]. The data aligned to the human genome were analyzed to obtain a model because the KSHV genome was too small to build a model for analysis. All the parameters were default, except that the parameter of the fold range was changed from 8 to 30. The default P value cutoff for peak detection was 10⁻⁷. However, the d values (distances between the modes of the Watson and Crick peaks) in the calculated models of AR were <60 bp; therefore, the analysis was done without the MACS model (-nomodel) following the protocol. The parameters were customized to analyze data aligned to the KSHV genome. Because of the smaller size of the KSHV genome, the IgG group was not used as a control for its strong bias; instead, the chromatin input was used as a control in each group. The parameters were
set as follows: —bw (bandwidth) 100, —nomodel (without the MACS model), —shiftsize (shift tags by an arbitrary number) 50, and —nolambda (use of a global lambda). The P value cutoffs at $10^{-3}$ and $10^{-5}$ were both analyzed. The results were visualized with the integrative genomics viewer (IGV) software [35]. For data validation, more reliable results were chosen ($P < 10^{-5}$).

**Dual-luciferase reporter assay**

All the procedures were performed as described in the manual of the luciferase assay system (Promega). The luciferase reporter plasmid was transfected into cells, with and without DHT treatment. Alternatively, the reporter plasmid was co-transfected individually with the plasmid expressing AR-pCDH-CMV-sf3-blast or ARΔNTD+DBD-pCDH-CMV-sf3-blast. The total amount of DNA was normalized with an empty vector in transfection. The cells were harvested 36 h after transfection for the luciferase assay. The fold change was calculated by normalizing the internal expression of Renilla luciferase and calibrated by control treatment.

**Protein purification and in vitro binding assay of the EMSA**

*Escherichia coli* strain BL21 (DE3) expressing GST or GST-fusion proteins was grown in Luria broth medium at 37˚C to the exponential phase and cultured for 12 h at 30˚C after induction with isopropyl-thiogalactopyranoside. The cells were harvested and resuspended in ice-cold lysis buffer (250mM NaCl, 2mM EDTA, 50mM Tris-HCl, 1% v/v Triton X-100, 10% v/v glycerol, with protease inhibitors, pH 8.0), followed by sonication lysis (Sonic; cycle, 3-s on, 3-s off pulses; amplitude, 30%). The cell lysates were centrifuged at 12,000 rpm to obtain the supernatant, which was combined with Sepharose 4B glutathione resin (GE Healthcare) for affinity purification following the manufacturer’s protocols. EMSA was conducted as described previously, with minor modifications [36]. Briefly, a 200-bp DNA sequence from 28,088 to 28,288 bp in the PAN promoter was prepared as the detective probe, because the sequence covered the summit at 28,188 bp located between the ChIP peak spanning from 27,580 to 29,193 bp, as identified in ChIP-seq and schematically shown in Fig 4B. A pair of fluorescein Alexa 700-conjugated primers was used to amplify the fragment from the genomic DNA template, which was extracted from the induced iSLK.219 cells. The sequences of the probes and primers are listed in S1 Table. The PCR product was purified by agarose gel extraction and quantified. AR–DNA complexes were prepared by adding the respective recombinant proteins or GST to a solution of diethyl pyrocarbonate-sterilized water containing 150mM NaCl, 50mM Tris-HCl (pH 8.0), 2mM MgCl$_2$, and 1mM dithiothreitol. Then, the DNA fragment was added, followed by incubation for 30 min at 25˚C. For each sample, free DNA and complexes were separated on a 6.5% native polyacrylamide gel run for 2 h at 120 V and 5 W at 4˚C and visualized using an Odyssey Fc Imaging System (LI-COR).

**Cell transfection**

iSLK.219 cells were seeded into 6- or 12-well plates and transfected at ~80% confluence with siRNA pools from Santa Cruz Biotechnology targeting the AR or the scramble control. siRNAs were transfected using Lipofectamine 3000 (Thermo Fisher Scientific) following the manufacturer’s protocols. The optimized concentration of siPAN for iSLK.219 was 200nM and for siAR was 3 or 6nM. The cells were cultured at 37˚C for 12 h, washed, and maintained for another 36 h. Recombinant expression plasmids were transfected into 293T cells using polyethyleneimine for 12 h, and the cells were continually cultured in a fresh medium for 36 h before collection.
Cell Counting Kit-8 assay

A CCK-8 assay for cell viability was conducted following the manufacturer’s protocols (Qia- gen, Shanghai, China). Briefly, 1000 cells were seeded per well in 96-well plates as indicated with the respective siRNA of AR, PAN, or siCtrl, or left untreated. After 48 h, 10 μL of Cell Counting Kit-8 (CCK-8) was added and mixed thoroughly. The cells were incubated for 2 h at 37˚C, and the absorbance was measured at 450 nm.

Quantification of KSHV DNA levels and RNA transcription in cells

KSHV DNA was extracted following the manufacturer’s protocol (Qiagen, Shanghai, China). A total of 200 ng of DNA from each sample was used in real-time DNA PCR using KSHV LANA gene-specific primers. LANA gene cloned into the pGL2.0 vector (Promega) was used as the external standard [28]. Known amounts of the LANA plasmid were used in the amplification reactions along with the test samples. The cycle threshold values were used to generate the standard curve and calculate the relative copy numbers of viral DNA in the samples. The KSHV DNA amount was normalized to the amount of purified cellular DNA as determined by the primers of GAPDH.

The cells were collected and lysed in TRIzol buffer (Life Technologies), and RNA was isolated following the manufacturer’s protocols. Reverse transcription was performed with a cDNA Reverse Transcription Kit (Toyobo). Real-time RT-PCR was performed with an SYBR green Master Mix kit (Toyobo). The relative mRNA levels were normalized to actin and calculated by the ΔΔCT method. The primer sequences are summarized in S1 Table.

Transwell invasion assay and cell counting

Transwell invasion assays were performed using Corning Transwell polycarbonate membrane cell culture inserts (8.0 μm). iSLK.219 cells were treated with siAR or scrambled siCtrl for 48 h, prior to doxycycline induction at 10μM for 24 h. Alternatively, the AR-pCDH-CMV-sf3-blast recombinant plasmid was transiently transfected into iSLK.219 cells for 24 h, followed by siPAN treatment for another 24 h and then subjected for doxycycline induction at the same concentration for 24 h. Typically, the cells were seeded in the upper chamber with a basic medium at 2 × 10^5 cells/well. The invasion was allowed to proceed for 24 h at 37˚C, and the cells were fixed with 70% methanol and stained with 0.5% crystal violet solution. The number of invaded cells was determined by counting stained cells from multiple randomly selected microscopic visual fields using ImageJ software (open source from National Institutes of Health, USA). Photographs were obtained, and independent experiments were performed in triplicate.

Statistical analysis

Experiments were carried out independently at least three times, and representative results were presented. Data were analyzed using the Student t test. A P value <0.05 was considered to be significant (two tailed). Error bars represented the standard error of the mean (SEM). The analysis was performed with GraphPad Prism software (GraphPad Software, Inc., CA, USA).

Results

AR was highly expressed in KS lesions and concentrated in the nuclei

Previous studies demonstrated that KS occurred disproportionately in men, suggesting a possible role of male hormones in KSHV infection and pathogenesis. Therefore, this study first explored the expression pattern of the AR in KS tissues. Twelve KS specimens and control
tissues, consisting of one normal skin sample and six non-KS angiomas, were collected from male patients. The AR expression was found in all specimens, contrary to a previous report showing the absence of sex-hormone receptors in KS [37]. Although the AR was expressed at low levels in the cytoplasm of the basal cells in control tissues (Fig 1B and 1C), it was extensively distributed in the cell nuclei in KS lesions; one tissue sample had both cytoplasmic and nuclear expression (Fig 1A). The AR was specifically distributed in the spindle areas of the KS tissues, which were characterized by LANA-positive staining (Fig 1A). Spindle cells were commonly recognized as KS tumor cells with an endothelial origin [38], and the results indicated a possible role of AR in KS pathogenesis.

We further verified the enrichment of AR at the protein level by involving various KSHV-positive primary effusion lymphoma cells of BCBL1 and BC3 cells and the negative B cells of DG75 and BJAB. The results showed that the AR was considerably expressed in KSHV-infected cells, when compared with the undetectable level in DG75 and BJAB (Fig 1D). Since BJAB was not an ideal control for BCBL1 and BC3 cells, we further compared the expression...
of AR in BJAB and KSHV stably transfected BJAB (KSHV-BJAB) cells. We found that the AR was expressed in KSHV-BJAB cells, but in much lower amounts than in BCBL1 cells (Fig 1E). Regarding the ligand effect, androgen treatment strongly increased the expression of AR in a dose-dependent manner in KSHV-BJAB and BCBL1 cells, compared with MeOH solvent treatment (Fig 1E). The effect was more significant in BCBL1 cells and was validated by approximately two times transcriptional upregulation of prostate-specific antigen (PSA), the classical target gene of AR. PSA expression in androgen-sensitive prostate cancer cells of LNCaP was 5.2 times higher than that in solvent control, indicating the potency of DHT treatment (Fig 1F). The specificity of the anti-AR antibody and DHT efficacy were validated by the abundance of AR in the 110-kDa full-length isoform. A dramatic increase was observed upon DHT treatment in LNCaP cells, but not in nonsensitive cells of PC3 (S1A Fig). In addition, an AR transcript was detected in B cells and control cells of LNCaP and PC3. We surprisingly found that the AR could not be detected at the mRNA level in B cells and PC3 cells, while it could only be successfully amplified from LNCaP cells; the Ct value of the AR and reference gene of actin was 24.6 and 19.6, respectively, while the values of AR in other cells were more than 31 (S1B Fig). The results were in high concordance with the data set in which the AR transcript was hardly detected in versatile types of B cells, such as HDLM-2, Daudi, RPMI-8226, U-226-70, U-698, and Kapas-707 cell lines in the Human Protein Atlas (https://www.proteinatlas.org/). Taken together, the nuclear localization of AR in clinical samples and the ligand-dependent transactivity of AR in KSHV-positive BCBL1 cells indicated a mode of genomic regulation of AR to the KSHV genome by acting as a canonical transcription factor.

**High throughput of ChIP-sequencing revealed that the AR bound to KSHV genome**

All the genome-wide binding sites of AR were mapped using ChIP-seq to determine whether the AR bound to KSHV genome and the interacting domains. iSLK.219 cells were pretreated with 10μM DHT or its solvent control MeOH for 48 h, without doxycycline induction. The cell lysates were subjected to immunoprecipitation with the anti-AR antibody or the control anti-IgG antibody. The ChIP-enriched DNA with the anti-AR antibody and the chromatin input in the iSLK.219 cells treated with DHT or MeOH solvent were analyzed by deep sequencing. The sequence reads for each sample were mapped to the KSHV genome (HQ404500) and human genome (hg19) using Bowtie 2, and the aligned files were subjected to peak calling using MACS. The peak models built using MACS based on human genome alignment indicated that both DNA input and α-AR-immunoprecipitated (IP) samples had broad peaks. The peaks of the four samples ($P < 10^{-5}$) on the whole KSHV genome are shown in Fig 2A, and the information of AR to the human genome is shown in S1 Data. Compared with the ChIP peaks induced by solvent control treatment, the most significant peaks upon DHT ligand treatment were concentrated on the known promoter regions of K1, PAN/K7, ORF69, and ORF75, which represented the AR-binding sites, accompanied by smaller peaks scattering throughout the genome (Fig 2A). The highest enrichment signals recruited by the AR upon ligand treatment were localized in the promoter regions of the PAN gene (Fig 2B). The annotated information on the AR peaks ($P < 10^{-3}$) is shown in Fig 2C. The highest peak in ChIP-seq ranging from 27,580 to 29,193 bp covered the promoters of both K7 gene and PAN. The detailed schematic illustration about the overlapping region between the PAN promoter and the coding sequence of the K7 gene is shown in Fig 2D. Thus, which is the key viral gene between PAN and K7 responsive to male sex steroid treatment needs complete differentiation. Taken together, these results suggested a possible physical interaction between AR and KSHV genome in the herpesvirus field.
Association of AR with the promoter of PAN and, additionally, that of RTA

Five pairs of primers at the individual promoter were designed to determine the binding so as to verify further the specific binding sites of AR on the aforementioned lytic genes (Fig 3A and 3B). Similarly, iSLK.219 cells were treated with DHT and its solvent, without the induction of
doxycycline, and subjected to ChIP detection using an anti-IgG or anti-AR antibody. The AR showed significant binding to the PAN and ORF75 promoters (Fig 3A). The binding of AR in the presence or absence of DHT was determined to demonstrate the role of DHT treatment in AR recruitment, that is, whether the recruitment of AR to the viral promoter regions was dependent on the activation of the whole signaling pathway. It showed that the enrichment of AR in the PAN promoter region was detected at the highest level, compared with no effect on the K7 promoter (Fig 3B). The immunoprecipitated DNA in the ChIP assays targeting different sites on the PAN promoter was subjected to qPCR analysis and subsequent DNA gel confirmation (Fig 3C).

The viral genes were all lytically expressed, and K1, PAN, and ORF69 showed a replication and transcription activator (RTA)-responsive element (RRE) or the lytic origin of replication (ori-Lyt) domain in their promoters, which were essential for RTA binding and ori-Lyt-dependent DNA replication [22,39,40]; these might suggest a possible role of RTA in AR-mediated
KSHV lytic gene expression. As RTA is the only viral lytic protein sufficient to disrupt latency and promote complete lytic cascade [28], the possible regulatory effect of AR on KSHV RTA was additionally analyzed. Although it was not shown in peaks by ligand treatment in ChIP-seq, the direct recruitment of AR to the RTA promoter was actually detected only in the context of ectopic AR (Fig 3D). The immunoprecipitation products for the various binding sites on the RTA promoter were comprehensively analyzed (Fig 3D). iSLK.219 cells were left untreated or transduced with the lentivirus of AR, without doxycycline, and then subjected to immunoprecipitation with the anti-AR antibody. Collectively, these results indicated that the AR might physically bind with KSHV lytic genes, especially with PAN.

AR bound to the PAN promoter through direct interaction
The AR complexed with the PAN promoter sequence in ChIP-qPCR detection, but whether it occurred through direct association needed further investigation. All nuclear receptors have the same basic structure, comprising a NH2-terminal transactivation domain, a DBD, a hinge domain, and a carboxyl-terminal LBD [41]. Two truncated constructs of AR were synthesized, consisting of AR-NTD+DBD (amino acids 1–625) and AR-NTD (amino acids 1–537), the latter of which lacking the DBD domain, as schematically shown in Fig 4A. Two GST-fused recombinant proteins of AR were induced from Escherichia coli and purified. Then, the PAN promoter probe–based in vitro binding assay of EMSA was performed. The summit information in ChIP peaks indicated that the highest enrichment of AR binding to the PAN promoter occurred at the summit of 28,188 bp, schematically shown in Fig 4B. Therefore, the covering 200-bp sequence from 28,088 to 28,288 bp was amplified, labeled using fluorescein, and then used as a DNA probe (Fig 4B). The PCR template was extracted from the genomic DNA of doxycycline-induced iSLK.219 cells using fluorescent Alexa-700-labeled primers at each 5’-terminus (Fig 4C). After incubation, the complexes between the GST and GST-fused protein of AR were assayed using EMSA. Fig 4D showed that both AR recombinant proteins led to a strong complex formation, and the bands that represented the complexes were significantly shifted in lanes 9 and 10. Expectedly, AR-binding shifts were observed and augmented by incubation with increasing amounts of 320 pmol of recombinant AR proteins, but not at the lower level of 40 or 160 pmol (Fig 4D). It was notable that the AR-NTD, which lacked the DNA-binding domain, also showed an impaired but still high capability of binding to the probe and formed the shift, compared with that of AR-NTD+DBD (Fig 4D). This might indicate that more extending sequences other than the traditional DBD domain might be responsible for the effect. Collectively, these results proved that, in addition to the RNA transcript of PAN bound with several epigenetic modifiers, the AR could be another subset of host factors that physically interacted with its DNA sequence.

AR transactivated the PAN promoter as a canonical transcription factor
The study explored whether AR binding to PAN necessarily led to the promoter activity of KSHV lytic genes using a dual-luciferase reporter assay. A schematic illustration of reporter plasmids of pPAN and a series of truncation plasmids of PAN promoter from pP1 to pP10, which extended to position ~2000 bp from the transcriptional start site, at continued intervals of ~200 bp is shown in Fig 5A. The major host factors to the promoter and RNA transcript of PAN were highlighted (Fig 5A). The study confirmed that DHT treatment significantly induced the promoter activity of PAN and RTA, which was 14.8 and 5.2 times higher than that of pGL3.0 vector control, respectively, but had no effect on the LANA promoter (Fig 5B). 293T cells were pretreated with DHT or its solvent control for 24 h, followed by transfection with the promoter plasmid for another 36 h. Strikingly, the ectopic expression of the full-length AR
transactivated the PAN at a very high level in a dose-dependent manner; the auto-activation of RTA to its own promoter was used as the positive control (Fig 5C). 293T cells were transfected with reporter plasmids of RTA, PAN, LANA promoter, or pGL3.0 vector or in combination with the expression plasmid of AR in a dose-dependent manner. A mutant of AR NTD+DBD plasmid was constructed from the wild type, from which the transactivation domains of N-terminus and DNA-binding domain were deleted, to illustrate whether the effects were dependent on the DBD domain. Correspondingly, a dramatic decline in the promoter activity of PAN and RTA was observed compared with that of wild-type AR, indicating that the AR activated PAN and RTA dependent on DNA-binding capability (Fig 5D). To fine map the interaction between AR with PAN promoter, the ectopic AR-induced promoter activity was

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Fig 4. AR interacted with the PAN promoter majorly through its DNA-binding domain. (A) Illustration of AR functional structure and truncated constructs used in this study. (B) Illustration of fluorescence-labeled DNA probes used in the EMSA assay, spanning from 28,088 to 28,288 bp of PAN promoter. The fragment covered the summit at 28,188 bp in the ChiP peak on the PAN promoter as identified from deep sequencing. (C) Alexa Fluor 700 dye was used to label the 5’ terminus of the primers. Two fusion proteins of AR both formed complexes with the PAN promoter, while the mutant lacked the DBD domain. Equal amounts of 320 pmol of DNA probe were used in each group. It was incubated with increasing amounts of GST-fused AR truncations or GST alone. The two GST-fused AR truncations comprised an N-terminus with a DNA-binding domain of AR (GST-NTD+DBD) and a shorter version of GST-NTD, which lacked the DBD domain. GST protein alone was used as a negative control. The lane with only DNA probe was used to indicate the size of free probes. Representative images are shown. Each reaction was repeated in, at least, triplicate.

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comprehensively compared between the 10 truncation plasmids showed in Fig 5A, using a pPAN plasmid as positive control. The results showed that only the promoter activity of pP8 reporter plasmid, which covered the sequence from 28,220 to 28,419 bp, was significantly increased by AR overexpression, compared with that of pCDH-sf3 (Fig 5E). The results were verified by the consistent upregulation of the pPAN plasmid, which covered precisely the whole insertion of pP8 (Fig 5A and 5E). More interestingly, the N-terminus of pP8 insertion was also overlapped by the summit sequence from 28,088 to 28,288 bp, which was identified in the ChIP peak on the PAN promoter (Fig 4B). Therefore, these results indicated that a minus AR-binding region in the PAN promoter was the 69-bp sequence from 28,220 to 28,288 bp. The luciferase activity was normalized by TK expression in all groups, and the fold change was determined by comparing AR-induced promoter activity with that of the vector.

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together, the results suggested that male sex steroid treatment transactivated the promoters of KSHV lytic genes, especially PAN and RTA, in a canonical transcription factor way.

**AR promoted lytic gene expression and KSHV production**

We further investigated the functional consequence of the promoter transactivation of PAN by male sex steroids. Upon AR overexpression, a large number of RFP-positive cells, representing increasing KSHV reactivation, were observed, especially after doxycycline treatment for 72 h (Fig 6A). The results were validated by quantitative analysis of the dramatically increased expression of PAN, RTA, and ORF57 and DNA copies of KSHV using RT-qPCR (Fig 6B and 6C). iSLK.219 cells were transduced with AR-expressing lentivirus or virus control for 24 h and then exposed to doxycycline for another 72 h before harvest. Surprisingly, spontaneous KSHV reactivation occurred with a very high dose of AR-expressing lentivirus, even without
the induction of doxycycline (Fig 6D). Ten times the amount of AR-expressing lentivirus (multiplicity of infection = 10) was used to infect iSLK.219 cells for 96 h without doxycycline. The effect was confirmed by >200 times higher expression of PAN RNA in RT-qPCR analysis, compared with the little effect on RTA and LANA expression (Fig 6E). Moreover, RNA interference of AR in iSLK.219 cells was used, and the knockdown efficiency of siAR was validated by decreasing the expression of endogenous AR compared with scrambled siControl treatment (Fig 6F). The cell viability was measured by a CCK-8 assay 48 h after siRNA transfection (S2 Fig). The transfection of AR siRNA attenuated the RFP signals and mRNA expression of PAN and RTA in a dose-dependent manner (Fig 6G and 6H). Collectively, the results indicated that the male hormone axis might facilitate the RNA transcription of multiple KSHV lytic genes, especially PAN and RTA, which were two critical effectors for the entire viral genomic transcription cascade.

**Nuclear import of AR was a requirement for transactivation to PAN in a ligand-dependent manner**

The AR has been found to contribute to lytic gene expression, especially to PAN. However, whether the effect is obtained by ligand-mediated nuclear localization of AR remains unknown. Mounting evidences indicate that AR is localized to the cytoplasm in the absence of androgen and translocates into the nuclei to activate gene expression in the presence of the ligand [26,41]. During the process, two nuclear localization signals (NLSs) of AR have been identified as responsible for translocation from the cytoplasm into the nucleus in the presence of the ligand, with the constitutively activated one located in the DBD and the hinge region at amino acids 617–633 of AR [42], and the other in the LBD [43]. The specific residues and functions of the latter NLS are unclear [43]. The effect of the ligand on PAN RNA expression was thus explored. DHT treatment led to the strongest upregulation of PAN RNA expression in a dose-dependent manner upon doxycycline induction (Fig 7A). iSLK.219 cells were treated with DHT or the solvent prior to doxycycline induction for 48 h. Whether the results were acquired by the androgen-mediated nuclear localization of AR through NLSs needed full illustration. A deletion mutant was constructed from which the DBD, hinge region, and LBD where NLSs were located were removed, as schematically shown in Fig 7B. Fig 7C (left panel) clearly demonstrates that the transfection of the mutant plasmid of ARΔDBD+Hinge+LBD-pCDH-sf3 led to the AR expression stringently outside of the nucleus compared with the disperse distribution of the full-length wild-type AR throughout the transfected cells; the specificity of AR expression was validated by the negative signal from the vector group detected using the anti-Flag antibody. Compared with the strong nuclear translocation of full-length AR upon androgen treatment without doxycycline, ARΔDBD+Hinge+LBD-pCDH-sf3 remained perinuclear in the cytoplasm, thus confirming the loss of function in the ligand-induced nuclear import of the mutant. The nuclear distribution in some cells in the mutant group indicated that other NLS existed besides the known two (Fig 7C, left panel). Regarding the effect of KSHV reactivation between the wild-type AR and the mutant, the ARΔDBD+Hinge+LBD-pCDH-sf3 mutant was confirmed to have a compromised capability to induce lytic gene expression represented by a significant less amount of RFP+ signals compared with that in wild-type AR, with or without DHT treatment [Fig 7C, left panel, consistent with the expression of PAN RNA in the same treatment detected by RT-qPCR in Fig 7D (left panel)]. The effects were considerably recapitulated using inducible doxycycline in the right panel of Fig 7C and 7D. It showed that the significant upregulation of RFP-positive signals by wild-type AR transfection was dramatically diminished in the group of NLSs-deleted mutant, especially under ligand treatment (Fig 7C, right panel). The results were verified by a nearly 54%
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(A) mRNA relative expression (LSK-219 cells).

(B) Schematic diagram showing the domain structure of the viral noncoding RNA PAN.

(C) Immunofluorescence images showing the effects of doxycycline on the expression of AR, GFP, and α-Flag(AR) in different cells.

(D) Bar graph showing the relative expression of PAN under different conditions.
decrease in PAN RNA transcript in the mutant, compared with that in the wild-type one (Fig 7D, right panel). Intriguingly, it seemed that doxycycline treatment led to the feedback regulation of the expression of AR and its cellular distribution (Fig 7C, right panel), which will be very interesting to explore in the future. Taken together, the results indicated that the ligand-activated nuclear transport of AR was a prerequisite for the successful transactivation to PAN.

AR-mediated PAN contributed to cell invasion

The present study attempted to understand whether AR-induced PAN was biologically correlated with KS pathogenesis. Surprisingly, siRNA-induced inhibition of AR remarkably reduced the number of invaded cells, especially on doxycycline induction, while the effect in the absence of induction was relatively minor (Fig 8A). The staining intensity representing the cell numbers in the aforementioned groups was quantitatively analyzed by involving randomly collected three to five fields from individual groups using ImageJ software (Fig 8B). iSLK.219 cells were transfected with siAR or the control prior to doxycycline treatment. Next, the study investigated whether the regulation of PAN RNA by the AR was a prerequisite for AR-mediated cell invasion. The results showed that AR overexpression elicited an increased cell invasion dependent on the use of doxycycline (Fig 8C). However, the effect was inhibited significantly by additional involvement of siPAN, especially under doxycycline induction (Fig 8C). The images in Fig 8C were quantitatively analyzed and addressed in Fig 8D. The decreasing number of cells in siPAN groups with or without doxycycline induction was 30% versus 8.7% (Fig 8D). The knockdown efficiency of RNA interference to PAN was measured by RT-qPCR, and it showed that approximately 40% of PAN transcript was reduced compared with that in siControl, upon induction (Fig 8E). iSLK.219 cells were transduced with siPAN or the control siRNA. Then, the cells were treated with either doxycycline or dimethyl sulfoxide (DMSO) for 24 h. Taken together, the results demonstrated that PAN RNA was necessary for AR-mediated cell invasion, especially in the context of active KSHV lytic replication.

Discussion

As the most common disease in patients with AIDS, KS is found in approximately 50% of tumor cases in the population [44]. KS occurs predominantly in men, consistent with the male-associated seroprevalence of KSHV, at least in Africa [13–17]. A previous study demonstrated that the membrane-localized AR organized a series of signaling events to activate the virus receptor of ephrin type-A receptor 2 (EphA2) during KSHV early infection [45]. It
explained that KSHV seroprevalence and virus DNA copies from male patients were higher than those in female patients. However, the underlying mechanism for the male propensity in KS oncogenesis remains elusive. This study was novel in demonstrating that the AR acted as a transcription factor to exert the genomic regulation of KSHV noncoding RNA PAN, thereby

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facilitating lytic replication–mediated oncogenesis, which implied a novel mechanism for sex disparity in KS. The study highlighted that the AR orchestrated the androgen-regulated transcriptome to virus-originated noncoding RNA. It demonstrated that the AR might also be exploited by KSHV in virus-induced malignancy, as in the case of HBV [10–12]. It indicated the striking expression of AR in KS spindle cells and the potency of androgen treatment in KSHV-positive B cells. As spindle cells are KS tumor cells of endothelial origin characterized by nuclear staining for the KSHV oncoprotein LANA [38], these data might indicate a novel role of male hormones in KS malignancy.

Similar to other steroid hormone receptors, the AR is located in the cytoplasm in the absence of androgens, forming a complex with chaperones and co-chaperones such as heat shock proteins and immunophilins. Upon androgen binding, the AR undergoes conformational changes and translocates to the nucleus, where it binds to androgen response elements as a homodimer and then activates the expression of AR target genes [26,41]. This mode of action is the classical nuclear receptor signaling pathway, also known as the genomic action, had being reported to noncoding RNA of cellular HOTAIR by the estrogen receptor [46]. In the present study, the PAN RNA was confirmed as the first microbial noncoding RNA that responded to male sex steroid regulation. Different from the previous findings of chromatin modifiers of histone H3K27 demethylases UTX, JMJD3, and lysine methyltransferase MLL2 binding to PAN RNA [23], we identified a unique way of AR directly interacting with the promoter sequence of PAN, thus highlighting a novel function of AR in viral infectious disease. This study showed that the AR was a new transcriptional regulator for PAN RNA transcript. Interestingly, the effect was acquired even without targeting the classical motif of androgen-responsive element in PAN, which might extend the current knowledge of the regulatory events to KSHV PAN. Similar to how estrogen receptor functions on cellular noncoding RNA, ER binds strongly to a genomic region of about 14.5 kb upstream to the transcription start site of the HOTAIR gene [46]. We demonstrated that a minus AR-binding site from 28,220 to 28,288 bp in the PAN promoter was indispensable for AR transactivation. Combined with the finding that the nuclear translocation of AR in the ligand-dependent fashion was critical for increased PAN RNA, the results indicated a canonical way of AR to transcriptionally regulate the viral noncoding gene.

In common with other herpesviruses, the γ-herpesviruses remain latent in their natural host, and only a fraction of the viral genes are expressed [5,15,20,40]. A subset of these latent viruses can reactivate and enter the lytic cycle [47]. Although viral genes transcribed during latency are generally considered essential for the transformation process and the development of viral-induced neoplasia, it is now clear that a subset of lytic genes can also contribute to the development of viral-induced cancers through a paracrine mechanism [20,47,48]. The role of lytic replication in KS pathogenesis is supported by the fact that interrupting lytic replication with immune reconstitution or using anti-lytic herpes antivirals, such as gancyclovir, can also inhibit or prevent KS development [19,20]. The PAN, K1, and ORF69, as identified using ChIP-seq analysis, were found to have RTA-responsive elements (RRE) for RTA binding on their promoters [21,22,39,40]. As RTA is the major viral protein governing the transition of the KSHV genome from the latent to the lytic phase [28,47], this study further verified the physical interaction of AR with RTA promoter by ChIP-PCR detection but only under the condition of AR overexpression. Accordingly, ectopic AR dramatically increased the promoter activity and mRNA expression of RTA. In addition, we surprisingly found that another known early-lytic gene of ORF57 also was transactivated by AR overexpression. Besides, a two-repeat classical ARE motif of “AGAACA” [49] was identified in the ORF57 promoter, which was AGAA\^A\^AGA\^C. Therefore, the significance of male sex steroids in KSHV lytic replication was emphasized.
Recent studies have found that AR-targeted gene type II transmembrane serine protease (TMPRSS 2) is responsible for the viral spike (S) protein priming and the subsequent entry via binding of its S protein to angiotensin-converting enzyme 2 (ACE 2) receptor of coronavirus infectivity [8,50,51]. Hence, male hormones are relevant to the male predominance in certain infectious diseases through versatile mechanisms [50–54]. In research on the herpesvirus, however, the understanding of the reasons for this male predominance is limited. This study demonstrated that the AR bound and transactivated viral noncoding RNA PAN, thus suggesting a novel mechanism for sex disparity in KS. Considering that viral noncoding RNA is a common feature of herpesvirus family, but a neglected field of research in biology, the findings of the present study have scientific relevance for other types of viral diseases.

Supporting information

S1 Fig. (A) Specificity of the anti-AR antibody and DHT efficacy were validated by the abundance of AR at 110-kDa full-length isoform, which dramatically increased upon DHT treatment in LNCaP cells, but not in nonsensitive cells of PC3. (B) AR transcript could not be detected at the mRNA level in the B cells and PC3 cells, while it could only be successfully amplified from LNCaP cells; the Ct values of AR and reference gene of actin were 24.6 and 19.6, respectively, which those in other cells were all more than 31.

(TIF)

S2 Fig. RNA interference of AR and PAN did not affect the cell viability of cells. A total of 1000 iSLK.219 cells were seeded per well in 96-well plates and transfection with 6nM siAR, 200nM siPAN, or siCtrl, or left untreated for 48 h.

(TIF)

S1 Table. Primers and probes.

(DOCX)

S1 Data. Identified AR peaks in human genome (hg19).

(XLS)

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