Evidence for an Epistatic Effect between TP53 R72P and MDM2 T309G SNPs in HIV Infection: A Cross-Sectional Study in Women from South Brazil

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Abstract

Objective: To investigate the associations of TP53 R72P and MDM2 T309G SNPs with HPV infection status, HPV oncogenic risk and HIV infection status.

Design: Cross-sectional study combining two groups (150 HIV-negative and 100 HIV-positive) of women.

Methods: Data was collected using a closed questionnaire. DNA was extracted from cervical samples. HPV infection status was determined by nested-PCR, and HPV oncogenic risk group by Sanger sequencing. Both SNPs were genotyped by PCR-RFLP. Crude and adjusted associations involving each exposure (R72P and T309G SNPs, as well as 13 models of epistasis) and each outcome (HPV status, HPV oncogenic risk group and HIV infection) were assessed using logistic regression.

Results: R72P SNP was protectively associated with HPV status (overdominant model), as well as T309G SNP with HPV oncogenic risk (strongest in the overdominant model). No epistatic model was associated with HPV status, but a dominant (R72P over T309G) protective epistatic effect was observed for HPV oncogenic risk. HIV status was strongly associated (risk factor) with different epistatic models, especially in models based on a visual inspection of the results. Moreover, HIV status was evidenced to be an effect mediator of the associations involving HPV oncogenic risk.

Conclusions: We found evidence for a role of R72P and T309G SNPs in HPV status and HPV oncogenic risk (respectively), and strong associations were found for an epistatic effect in HIV status. Prospective studies in larger samples are warranted to validate our findings, which point to a novel role of these SNPs in HIV infection.

Introduction

Infection susceptibility variability among individuals is observable in many infectious diseases. Understanding such variation and identifying its causal factors may have important implications for clinical practice and population health. Although intra and inter-population variation of viral infection susceptibility is attributable to several factors, host genetics likely plays an important role. Regarding HIV infection, a common example of genetic resistance is the CCR5 32-bp deletion. This allele, when transcribed and translated, results in a non-functional receptor, thus providing resistance to HIV infection [1]. Importantly, these findings are currently being incorporated into therapeutic approaches [2,3], thus evidencing that understanding the genetic basis of viral infection susceptibility has practical implications for human health. In this regard, several genome-wide association studies (GWAS) have been conducted to investigate the roles of host genetics in HIV load and/or disease progression [4–14], and, more recently, some GWAS focused on genetic factors associated with HIV acquisition in different populations [15–20]. Such attempts indicate the importance given to host genetics regarding HIV pathogenesis.

Considering that HIV infection is considered a major risk factor for HPV infection, the identification of host genetic factors involved with HIV infection may also have implications for HPV-related outcomes. Given the well-established roles of HPV in cancer, the majority of the studies involving HPV and host
genetics are related with cancer development and/or progression associated with infection by oncogenic HPV strains. In this context, p53 pathway genes have been extensively studied given that oncogenic HPV E6 orchestrates, in association with E6AP, p53 degradation by the ubiquitin-proteasome system[21]. Theoncogenic effects of HPV have also been investigated epide

icantly (in different populations) using genetic variants in p53 pathway genes, including the TP53 R72P SNP (rs1042522) [22–24], which was evidenced to interfere with p53 apopotic and transcriptional functions [25–27]. More recently, the MDM2 T309G SNP (rs2279744) [28–30], which influences p53 activity by affecting MDM2 tran

scription, have also been investigated in similar contexts [31,32].

In a recent review, different biological mechanisms by which interfering with p53 pathway may impact viral infection (or virus persistence after exposure to it) were proposed [33]. In addition, there are substantial amounts of in vitro evidence supporting a functional relationship between p53 and different HIV proteins (with evidence for implications for p53-mediated apoptosis), as well as some evidence regarding MDM2. These functional relationships with the p53 pathway have been evidenced for gp120 [34,35], Rev, Tat [36] and Vpu [37] proteins. Considering the roles of R72P and T309G SNPs in p53 apoptosis and regulation (respectively) and the evidence for a role of p53 pathway in HPV and HIV biology, we aimed to investigate the association of these SNPs with HPV infection status, HIV oncogenic risk and HIV infection status.

Materials and Methods

Study Design and Participants

We performed a cross-sectional study, selecting two independent groups of women based on HIV status (determined by medical diagnosis). From May 2010 to May 2011, 250 (150 HIV-negative and 100 HIV-positive) women seeking gynecologic care at the gynecological ambulatory clinic of Faculty of Medicine of Federal University of Pelotas (South Brazil) that fulfilled eligibility criteria (not pregnant, sexually active, and not menstruating) and agreed to participate were sequentially included in the study. Data were collected using an adapted version of a closed questionnaire [38], which was applied by a trained female interviewer. Routine gynecological exams (cervicitis indicators, visual inspection with acetic acid and Lugol’s iodine), were performed and included in the questionnaire, as well as the patient’s recorded information (last Pap test result).

Ethics Statement

The study was approved by the Ethics Committee of the Faculty of Medicine of Federal University of Pelotas (June 2009). Written informed consent was obtained from all participants. All procedures were performed in accordance with the Helsinki Declaration guidelines.

DNA Collection

Cervical samples were collected with a cytobrush and placed into 1.5 ml microtubes containing 300 ml of Cell Lysis Solution (Puregene™ DNA Extraction Kit, Gentra Systems Minneapolis, MN). The material was enzymatically digested using 1.5 μl of proteinase K (10 mg/ml, New England Biolabs, MA) and incubated overnight at room temperature. DNA was extracted according to manufacturer’s specifications.

HPV Detection and Genotyping

HPV detection was performed using nested-PCR in two rounds: amplification of a 450 bp fragment using the MY09/11 primer pair [39] and amplification of a 140 bp fragment using the GP5/6 primer pair [40]. MY09/11 and GP5/6 PCRs (final reaction volume of 25 μl) were performed as follows: initial denaturation for 9 min at 95°C; 40 cycles of denaturation (for 1 min at 95°C and for 30 s at 94°C, respectively), primer annealing (for 1 min at 55°C and 30 s at 45°C, respectively), and extension (for 1 min and for 30 s, respectively, at 72°C); and final extension for 5 min at 72°C [41,42]. PCR amplificates were visualized on 2.0% agarose gels stained with GelRed™ (Biotium Inc., CA). HPV-positive amplicates (from the second nested-PCR round) were purified using Gel Band purification Kit (GE Healthcare, USA) according to manufacturer’s instructions. To determine HPV oncogenic risk group (i.e., HPV genotype), Sanger sequencing was performed in a MegaBACE 1000 DNA sequencer (GE Healthcare, USA) using Dynamic ET-terminator technology. Chromatograms were assembled and analyzed using the ContigExpress module of the Vector NTI 10.0 suite (Invitrogen, USA). The assembled sequences were submitted to BLAST alignment (www.ncbi.nlm.gov/BLAST) against sequences available in GenBank.

Both SNPs were genotyped by PCR-RFLP using GoTaq qPCR Master Mix (Promega, USA) [in 12 μl reactions] with primers, restriction enzymes and PCR conditions described previously [43–45]. Briefly, for the R72P SNP, the 199 bp amplicate was cleaved using BspUI (New England Biolabs, MA) and loaded on 2.5% agarose gel stained with GelRed™ (Biotium Inc., CA). Genotyping was performed as follows: one fragment of 199 bp corresponds to R72P genotype; three fragments of 199 bp, 113 bp and 86 bp correspond to R72P genotype; and two fragments of 113 bp and 86 bp correspond to R72R genotype. For the T309G SNP, the 157 bp amplicate was cleaved by MspI (New England Biolabs, MA) and loaded on 2.5% agarose gel stained with GelRed™. Genotyping was performed as follows: one fragment of 157 bp corresponds to T309T genotype; three fragments of 157 bp, 109 bp and 48 bp correspond to T309G genotype; and two fragments of 109 bp and 48 bp correspond to G309G genotype.

Statistical Analyses

Analyses were performed in R (version 3.0.1, http://www.r-project.org/). Descriptive analyses were stratified according to HIV infection status for both SNPs [including assessing Hardy-Weinberg equilibrium (HWE) by Fisher’s exact test using the “genetics” R package: http://cran.r-project.org/web/packages/genetics/], status of HPV infection (positive or negative), HPV oncogenic risk (high or low) and potential confounding variables (i.e., skin color, achieved schooling in years, family income in minimum salaries and age). Skin color was a categorical variable defined by interviewer’s observation. Achieved schooling in years was categorized in illiterate, 1–4, 5–8, 9–11 and 11 or more according to the formal educational system that the recruited women frequented [composed of: 8 years or Primary Education, which changes substantially after the first 4 years; Secondary Education, composed of 3 years (in a total of 11 years); and Higher Education]. Age was categorized in groups of (approximately) 5 years (adjusting the age limits of some groups to avoid categories with very few individuals) for a more detailed comparison between HIV groups. Crude comparisons between HIV strata were performed by either chi-squared or Fisher’s exact test (using the “gmodels” R package: http://cran.r-project.org/web/packages/gmodels/index.html).

Importantly, any confounding effect of age, education and family income (after adjusting for skin color) on the association
between a genetic factor and a given trait is expected to occur by chance (since an individual's genotype is not influenced by environmental factors), thus imposing a difficulty to establish an adequate conceptual framework to select confounding variables. Therefore, we used a statistical-oriented approach, although skin color was invariable adjusted for. The reason for doing so is the well-known possibility of confounding in SNP-outcome analyses due to population stratification. Since both socioeconomic status (in several populations including Brazil) and genotypic frequencies of the vast majority of SNPs vary substantially according to skin color, the last can create spurious associations between genetic factors and outcomes influenced by socioeconomic factors. This is relevant for this manuscript since the last are known to have profound implications for sexually transmitted diseases.

The confounding variable selection was performed by stepwise backwards selection (critical P = 0.2 according to likelihood-ratio chi-squared test using the “car” R package: http://cran.r-project.org/web/packages/car/index.html). Since skin color was invariable adjusted for due to conceptual considerations, it was not subjected to removal. This process was performed having, as the independent variable, each outcome of the study (i.e., HPV status, HPV oncogenic risk and HIV status) and each main exposure (R72P and T309G genotypes, as well as the genotypes combined). These analyses were performed by logistic and multinomial logistic regression (using the “nnet” R package: http://cran.r-project.org/web/packages/nnet/index.html), respectively. The variables that remained in the final models of the outcome and the genetic exposure of interest were considered confounding factors for the associations involving such exposure-outcome pair.

Associations involving R72P and T309G SNPs and study outcomes were assessed in crude and adjusted logistic regression [estimating odds ratio (OR)] models. Five genetic models (obtained using the “SNPassoc” R package: http://cran.r-project.org/web/packages/SNPassoc/index.html) were tested: codominant or genotypic (i.e., each genotype is coded as a distinct category), additive (i.e., the SNPs are coded numerically according to the number of variant alleles), overdominant (i.e., homozygous genotypes = 0 and heterozygous genotype = 1), dominant (i.e., homozygous wild = 0; heterozygous and homozygous variant genotype = 1) and recessive (i.e., homozygous wild and heterozygous genotypes = 0 and homozygous variant genotype = 1), with the category corresponding to (or containing) the wild-type

| Variable                   | Categories | HIV status          | P-value |
|----------------------------|------------|---------------------|---------|
|                            |            | Negative (n = 250)  | Positive (n = 100) |
| HPV status                 | Positive   | 176 (70.4%)         | 32 (32.0%)     | <0.001* |
|                            | Negative   | 74 (29.6%)          | 68 (68.0%)     |         |
| HPV oncogenic risk         | Low-risk   | 7 (10.1%)           | 38 (66.7%)     | <0.001* |
|                            | High-risk  | 62 (89.9%)          | 19 (33.3%)     |         |
| Age groups                 | 18–24      | 43 (17.2%)          | 3 (3.0%)       | <0.001* |
|                            | 25–30      | 55 (22.0%)          | 10 (10.0%)     |         |
|                            | 31–35      | 45 (18.0%)          | 17 (17.0%)     |         |
|                            | 36–39      | 50 (20.0%)          | 17 (17.0%)     |         |
|                            | 40–45      | 57 (22.8%)          | 53 (53.0%)     |         |
| Skin color                 | White      | 164 (65.6%)         | 51 (51.0%)     | 0.005*  |
|                            | Black      | 31 (12.4%)          | 26 (26.0%)     |         |
|                            | Brown      | 55 (22.0%)          | 23 (23.0%)     |         |
| Family income              | ≤1: 50     | 50 (26.6%)          | 27 (30.7%)     | 0.389*  |
|                            | >1, ≤2     | 80 (42.6%)          | 33 (37.5%)     |         |
|                            | >2, ≤3     | 46 (24.5%)          | 18 (20.5%)     |         |
|                            | ≥4         | 12 (6.4%)           | 10 (11.4%)     |         |
| Achieved schooling         | Illiterate | 4 (1.6%)            | 0 (0.0%)       | <0.001* |
|                            | 1–4        | 32 (13.0%)          | 22 (23.2%)     |         |
|                            | 5–8        | 93 (37.8%)          | 43 (45.3%)     |         |
|                            | 9–10       | 36 (14.6%)          | 0 (0.0%)       |         |
|                            | ≥11        | 81 (32.9%)          | 30 (31.6%)     |         |
| R72P                       | R72R       | 27 (10.8%)          | 18 (18.0%)     | 0.200*  |
|                            | R72P       | 118 (47.2%)         | 43 (43.0%)     |         |
|                            | P72P       | 105 (42.0%)         | 39 (39.0%)     |         |
| T309G                      | T309T      | 35 (14.0%)          | 16 (16.0%)     | 0.543*  |
|                            | T309G      | 101 (40.4%)         | 45 (45.0%)     |         |
|                            | G309G      | 114 (45.6%)         | 39 (39.0%)     |         |

*Fisher’s exact test.
†Chi-square test.
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The homozygous genotype as the reference group (as described elsewhere [46]).

For the analyses of epistasis, a total of 13 epistatic models were tested using crude and adjusted logistic regression models. Eleven of them were described elsewhere [46]. Briefly, the following epistatic models were tested: dominant epistasis [both R72P over T309G (reference: R72R T309T; category 1: R72R G309_; category 2: P72P _309) and vice-versa (reference: R72R T309T; category 1: R72R G309G; category 2: _72_ G309_); recessive epistasis [both R72P over T309G (reference: R72_ T309_; category 1: R72_ G309G; category 2: P72P _309) and vice-versa (reference: R72_ T309_; category 1: R72R T309_; category 2: _72_ G309G); dominant and recessive epistasis [both R72P over T309G (reference: remaining genotypic combinations; category 1: P72_ G309G) and vice-versa (reference: remaining genotypic combinations; category 1: P72P G309_); double dominant

**Table 2.** Crude and adjusted associations [showing OR (95% CI) and p-values] between R72P and T309G SNPs and HPV status.

| Genetic model | R72P Crude | R72P Adjusted | T309G Crude | T309G Adjusted |
|---------------|------------|---------------|------------|---------------|
| Codominant    | P = 0.071  | P = 0.065     | P = 0.786  | P = 0.859     |
| A/A           | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| A/a           | 0.63 (0.40–1.00) | 0.62 (0.39–0.99) | 1.18 (0.74–1.87) | 1.14 (0.71–1.84) |
| a/a           | 1.16 (0.59–2.28) | 1.14 (0.57–2.28) | 1.12 (0.58–2.12) | 1.08 (0.54–2.11) |
| Overdominant  | P = 0.024  | P = 0.021     | P = 0.542  | P = 0.612     |
| A/A-a/a       | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| A/a           | 0.61 (0.39–0.94) | 0.60 (0.39–0.93) | 1.14 (0.74–1.76) | 1.12 (0.72–1.74) |
| Additive      | P = 0.649  | P = 0.602     | P = 0.602  | P = 0.707     |
| N° of “a” alleles | 0.93 (0.68–1.27) | 0.92 (0.66–1.27) | 1.08 (0.80–1.47) | 1.06 (0.77–1.46) |
| Dominant      | P = 0.146  | P = 0.125     | P = 0.500  | P = 0.601     |
| A/A           | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| A/a-a/a       | 0.73 (0.47–1.12) | 0.71 (0.46–1.10) | 1.16 (0.75–1.79) | 1.13 (0.72–1.77) |
| Recessive     | P = 0.227  | P = 0.229     | P = 0.924  | P = 0.991     |
| A/A-A/a       | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| a/a           | 1.47 (0.78–2.77) | 1.48 (0.78–2.82) | 1.03 (0.56–1.87) | 1.00 (0.53–1.87) |

**Table 3.** Crude and adjusted associations [showing OR (95% CI) and p-values] between R72P and T309G SNPs and HPV oncogenic risk.

| Genetic model | R72P Crude | R72P Adjusted | T309G Crude | T309G Adjusted |
|---------------|------------|---------------|------------|---------------|
| Codominant    | P = 0.680  | P = 0.553     | P = 0.024  | P = 0.019     |
| A/A           | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| A/a           | 1.43 (0.65–3.21) | 1.54 (0.69–3.55) | 0.35 (0.15–0.78) | 0.32 (0.13–0.74) |
| a/a           | 1.19 (0.40–3.85) | 1.48 (0.46–5.22) | 0.91 (0.28–3.27) | 0.81 (0.24–2.99) |
| Overdominant  | P = 0.410  | P = 0.381     | P = 0.006  | P = 0.005     |
| A/A-a/a       | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| A/a           | 1.37 (0.65–2.99) | 1.41 (0.66–3.09) | 0.36 (0.17–0.75) | 0.34 (0.16–0.73) |
| a/a           | 0.545 (0.30–0.95) | 0.350 (0.18–0.63) | 0.321 (0.16–0.67) | 0.244 (0.12–0.47) |
| Additive      | P = 0.011  | P = 0.033     | P = 0.021  | P = 0.021     |
| N° of “a” alleles | 1.17 (0.70–2.01) | 1.30 (0.75–2.32) | 0.77 (0.45–1.30) | 0.73 (0.42–1.24) |
| Dominant      | P = 0.410  | P = 0.277     | P = 0.033  | P = 0.021     |
| A/A           | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| A/a-a/a       | 1.36 (0.65–2.84) | 1.53 (0.71–3.33) | 0.43 (0.19–0.94) | 0.40 (0.17–0.87) |
| a/a           | 0.969 (0.60–1.56) | 0.761 (0.47–1.24) | 0.344 (0.20–0.60) | 0.394 (0.23–0.60) |
| Recessive     | P = 0.964  | P = 0.761     | P = 0.344  | P = 0.394     |
| A/A-A/a       | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| a/a           | 1.02 (0.36–3.16) | 1.19 (0.40–3.90) | 1.67 (0.59–5.49) | 1.60 (0.56–5.29) |

“*A*” and “a” correspond to wild-type (i.e., either R72 or T309) and variant alleles (i.e., either P72 or G309), respectively. doi:10.1371/journal.pone.0089489.t002
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Table 4. Crude and adjusted associations [showing OR (95% CI) and p-values] between R72P and T309G SNPs and HIV status.

| Genetic model | R72P | T309G |
|---------------|------|-------|
|               | Crude | Adjusted | Crude | Adjusted |
| **Codominant** |       |         |       |         |
| A/A           | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| A/a           | 0.98 (0.59–1.63) | 0.90 (0.54–1.52) | 1.30 (0.79–2.17) | 1.60 (0.91–2.84) |
| a/a           | 1.79 (0.88–3.61) | 1.45 (0.69–2.99) | 1.34 (0.66–2.65) | 1.85 (0.83–4.07) |
| **Overdominant** |       |         |       |         |
| A/A-a/a       | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| A/a           | 0.84 (0.53–1.34) | 0.82 (0.51–1.31) | 1.21 (0.75–1.93) | 1.35 (0.81–2.25) |
| **Additive**  |       |         |       |         |
| P = 0.205     | P = 0.517 | P = 0.305 | P = 0.073 |
| N$^*$ of “a” alleles | 1.25 (0.89–1.75) | 1.12 (0.79–1.60) | 1.19 (0.86–1.64) | 1.41 (0.97–2.05) |
| **Dominant**  |       |         |       |         |
| P = 0.606     | P = 0.984 | P = 0.259 | P = 0.064 |
| A/A           | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| A/a-a/a       | 1.13 (0.71–1.83) | 1.00 (0.62–1.64) | 1.31 (0.82–2.11) | 1.65 (0.97–2.86) |
| **Recessive** |       |         |       |         |
| P = 0.077     | P = 0.216 | P = 0.634 | P = 0.347 |
| A/A-A/a       | 1 (Reference) | 1 (Reference) | 1 (Reference) | 1 (Reference) |
| a/a           | 1.81 (0.94–3.45) | 1.53 (0.77–2.97) | 1.17 (0.60–2.19) | 1.42 (0.68–2.87) |

$^*$“A” and “a” correspond to wild-type (i.e., either R72 or T309) and variant alleles (i.e., either P72 or G309), respectively.

(a) For R72P SNP, only skin color was considered a covariate for all outcomes. For T309G SNP, in addition to skin color, age was also considered a covariate for HPV status and HIV status (tables S1 and S2). Crude and adjusted analyses for the associations of R72P and T309G SNPs with the outcomes are shown in tables 2–4. For HPV status, no significant associations with T309G SNP were observed (P>0.500). Regarding R72P SNP, a marginal association was observedfor the codominant model [crude (genotypes R72P and T309G); adjusted: OR (95% CI), 0.63 (0.40–1.00) and 1.16 (0.59–2.28); P = 0.071; adjusted: OR (95% CI), 0.62 (0.39–0.99) and 1.14 (0.57–2.28); P = 0.065] and there

Results

Sample Description

The characteristics of the sample (stratified according to HIV infection status) are shown in table 1. There were statistically significant differences between HIV-positive and HIV-negative regarding the two HPV-related outcomes, age, schooling (P<0.001) and skin color (P = 0.005). Family income did not significantly differ between HIV strata (P = 0.389). In this initial analysis, no significant differences were observed between HIV strata regarding the genotypic frequencies of R72P (P = 0.200) and T309G (P = 0.543) SNPs. Moreover, there was no evidence for departures from HWE regarding R72P and T309G SNPs in the total sample (P>0.999 and P = 0.102, respectively) and within skin color strata (white: P = 0.642 and P = 0.474; black: P>0.999 and P>0.999; brown: P = 0.797 and P = 0.331).

The observed differences are in the expected direction regarding HPV infection prevalence (70.4% in HIV-positive and 32.0% in HIV-negative women), age (prevalence of age groups: 18–24, 17.2% and 3.0%; 25–30, 22.0% and 10.0%; 31–35, 18.0% and 17.0%; 36–39, 20.0% and 17.0%; and 40–45, 22.8% and 53% in HIV-positive and HIV-negative groups, respectively), skin color (prevalence of skin color groups: white, 65.6% and 51.0%; black, 12.4% and 26.0%; and brown, 22.0% and 23.0% in HIV-positive and HIV-negative women, respectively) and achieved schooling in years (prevalence of schooling categories: illiterate, 1.6% and 0.0%; 1–4, 13.0% and 23.2%; 5–8, 37.8% and 45.3%; 9–10, 14.6% and 0.0%; and 11 or more, 32.9% and 31.6%, in HIV-positive and HIV-negative groups, respectively). Although the genotypic frequencies of none of the SNPs significantly differed between HIV strata, the heterogeneity of these groups was considered relevant for the association analyses.

Associations of R72P and T309G SNPs with HPV Outcomes and HIV Status

For R72P SNP, only skin color was considered a covariate for all outcomes. For T309G SNP, in addition to skin color, age was also considered a covariate for HPV status and HIV status (tables S1 and S2). Crude and adjusted analyses for the associations of R72P and T309G SNPs with the outcomes are shown in tables 2–4. For HPV status, no significant associations with T309G SNP were observed (P>0.500). Regarding R72P SNP, a marginal association was observed for the codominant model [crude (genotypes R72P and T309G); adjusted: OR (95% CI), 0.63 (0.40–1.00) and 1.16 (0.59–2.28); P = 0.071; adjusted: OR (95% CI), 0.62 (0.39–0.99) and 1.14 (0.57–2.28); P = 0.065] and there

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was a significant association for the overdominant model [crude: OR (95% CI), 0.61 (0.39–0.94); P = 0.024; adjusted: OR (95% CI), 0.60 (0.39–0.93); P = 0.021]. Indeed, the OR patterns observed for the codominant model indicate a protective overdominant effect.

For HPV oncogenic risk, the results were somehow opposite. No significant associations were observed for R72P SNP (P > 0.277). Regarding T309G SNP, significant associations were observed for codominant [crude (genotypes T309G and G309G, respectively): OR (95% CI), 1.03 (0.53–1.96) and 0.96 (0.49–1.88); P = 0.024; adjusted: OR (95% CI), 0.96 (0.49–1.88) and 0.81 (0.44–2.48); P = 0.019], overdominant [crude: OR (95% CI), 0.36 (0.17–0.75); P = 0.006; adjusted: OR (95% CI), 0.34 (0.16–0.73); P = 0.005] and dominant [crude: OR (95% CI), 0.43 (0.19–0.94); P = 0.033; adjusted: OR (95% CI), 0.40 (0.17–0.87); P = 0.021] models. Again, the codominant model already indicated a protective overdominant effect.

In contrast to HPV outcomes, only marginal associations were observed for HIV status. There were neither significant nor marginal associations for R72P SNP (P > 0.205). Regarding T309G SNP, marginal associations were observed for additive [crude: OR (95% CI), 1.19 (0.86–1.64); P = 0.305; adjusted: OR (95% CI), 1.41 (0.97–2.05); P = 0.073] and dominant [crude: OR (95% CI), 1.31 (0.82–2.11); P = 0.259; adjusted: OR (95% CI), 1.65 (0.97–2.86); P = 0.064] models. Although marginal, these associations lead to the speculation that HIV status could be an (at least partial) effect mediator of the associations between T309G SNP and HPV oncogenic risk. In this regard, the analysis of association between this SNP and HPV oncogenic risk were repeated with the inclusion of HIV status as a covariate (table S3). By doing so, only the overdominant model remained, although attenuated, statistically significant [OR (95% CI), 0.38 (0.14–0.96); P = 0.040]. The same was performed for the analysis of association between R72P SNP and HPV status, but no substantial differences were observed [overdominant model: OR (95% CI), 0.60 (0.38–0.96); P = 0.032], further evidencing an effect mediation role in HIV status in the associations between T309G SNP and HPV oncogenic risk.

### Table 5. Associations between 11 epistatic models and HPV status.

| Epistatic model | Genotypic combination | Crude OR (95% CI) | P-value | Adjusted OR (95% CI) | P-value |
|----------------|-----------------------|------------------|---------|---------------------|---------|
| Dominant (1.1) | R72R T309T            | 1 (Reference)    | 0.192   | 1 (Reference)       | 0.129   |
|                | R72R G309_            | 1.45 (0.74–2.87) | 1.44    | 0.72–2.88           | 0.019   |
|                | _72_ _309_            | 0.91 (0.50–1.67) | 0.83    | 0.45–1.55           | 0.006   |
|                | _72_ G309_            | 1.11 (0.61–2.03) | 1.04    | 0.57–1.93           | 0.021   |
| Recessive (2.1)| R72_ T309_            | 1 (Reference)    | 0.479   | 1 (Reference)       | 0.378   |
|                | R72_ G309G            | 1.03 (0.53–1.96) | 0.96    | 0.49–1.88           | 0.019   |
|                | _72_ G309G            | 1.48 (0.78–2.80) | 1.58    | 0.82–3.07           | 0.044   |
| Recessive (2.2)| R72_ T309_            | 1 (Reference)    | 0.533   | 1 (Reference)       | 0.494   |
|                | R72R T309_            | 1.47 (0.74–2.90) | 1.53    | 0.76–3.07           | 0.066   |
|                | _72_ G309G            | 1.08 (0.58–1.99) | 1.05    | 0.55–1.97           | 0.021   |
| Dominant and recessive (3.1) | P72_ G309G | 0.75 (0.33–1.64) | 0.75    | 0.32–1.66           | 0.044   |
| Dominant and recessive (3.2) | P72G309G | 0.97 (0.41–2.21) | 1.06    | 0.44–2.48           | 0.096   |
| Double dominant (no effect accumulation) (4) | Other | 1 (Reference) | 0.482 | 1 (Reference) | 0.475 |
| Double recessive (no effect accumulation) (5) | Other | 1.24 (0.76–2.01) | 1.24    | 0.75–2.03           | 0.433   |
| Double dominant (with effect accumulation) (6) | Other | 1.15 (0.63–2.12) | 1.11    | 0.61–2.08           | 0.021   |
|                | P72_ G309_            | 0.89 (0.47–1.72) | 0.80    | 0.41–1.57           | 0.062   |
| Double recessive (with effect accumulation) (7) | Other | 1.22 (0.74–2.00) | 1.20    | 0.72–2.00           | 0.021   |
|                | P72P G309G            | 1.55 (0.82–2.51) | 1.92    | 0.34–11.01          | 0.089   |
| Quantitative (8) | N’ of P72 and G309 alleles | 1.01 (0.81–1.26) | 0.950 | 0.78–1.24 | 0.893 |

*The epistatic models were numbered as described previously [46]. The "_" indicates that the effect is irrespective of the allele. E.g., R72P G309_ represents the genotypic combinations R72P T309G - R72P G309G.

Other: 1. _72_ T309_ - R72R _309_. 2. R72_ _309_ - _72_ T309T. 3. _72_ G309_ - P72_ _309_. 4. P72_ _309_ - _72_ G309_ . 5. P72P _309_ - _72_ G309G. 6. R72R G309_ - P72_ T309T. 7. P72P T309_ - R72_ G309G.

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Epistatic Models Were Associated with HPV Oncogenic Risk and HIV Status

In addition to skin color, age was considered a covariate for HPV status and HIV status (table S4). Crude and adjusted analyses for the associations of epistatic models and study outcomes are shown in tables 5–7. No associations were observed for HPV status (P = 0.192 and P = 0.129, respectively). Regarding HPV oncogenic risk, a significant association was observed for dominant epistasis with R72P SNP overcoming the effects of T309G SNP [model 1.1 (genotypic combinations R72P G309_ and R72_ G309G, respectively); crude: OR (95% CI), 0.24 (0.06–0.77) and 0.49 (0.13–1.53); P = 0.038; adjusted: OR (95% CI), 0.22 (0.06–0.73) and 0.54 (0.14–1.72); P = 0.060]. In addition, marginal associations were observed for dominant epistasis with T309G SNP overcoming the effects of R72P SNP [model 1.2 (genotypic combinations P72 T309T and _72_ G309G, respectively); crude: OR (95% CI), 0.59 (0.14–2.22) and 0.31 (0.08–0.94); P = 0.077; adjusted: OR (95% CI), 0.67 (0.15–2.63) and 0.31 (0.08–0.94); P = 0.060], dominant and recessive epistasis with R72P having a dominant effect given that G309G occurs [model 3.1; crude: OR (95% CI), 4.82 (0.84–90.98) and 1.11 (0.39–3.46); P = 0.082; adjusted: OR (95% CI), 4.75 (0.83–89.77) and 1.25 (0.42–4.11); P = 0.076]

Table 6. Associations between 11 epistatic models and HPV oncogenic risk.

| Epistatic Model* | Genotypic Combination | Crude OR (95% CI) | P-value | Adjusted OR (95% CI) | P-value |
|------------------|-----------------------|-------------------|---------|---------------------|---------|
| Dominant (1.1)   | R72R T309T            | 1 (Reference)     | 0.038   | 1 (Reference)       | 0.024   |
|                  | R72P G309_            | 0.24 (0.06–0.77)  | 0.22    | (0.06–0.73)         |         |
|                  | P72 _309_             | 0.49 (0.13–1.53)  | 0.54    | (0.14–1.72)         |         |
| Dominant (1.2)   | R72R T309T            | 1 (Reference)     | 0.077   | 1 (Reference)       | 0.060   |
|                  | P72 T309T             | 0.59 (0.14–2.22)  | 0.67    | (0.15–2.63)         |         |
|                  | _72_ G309G            | 0.31 (0.08–0.94)  | 0.31    | (0.08–0.94)         |         |
| Recessive (2.1)  | R72_ T309_            | 1 (Reference)     | 0.608   | 1 (Reference)       | 0.649   |
|                  | R72_ G309G            | 1.81 (0.58–6.87)  | 1.70    | (0.54–6.50)         |         |
|                  | P72 _309_             | 1.11 (0.39–3.46)  | 1.25    | (0.42–4.11)         |         |
| Recessive (2.2)  | R72_ T309_            | 1 (Reference)     | 0.633   | 1 (Reference)       | 0.656   |
|                  | R72R T309G            | 1.09 (0.35–3.78)  | 1.24    | (0.38–4.53)         |         |
|                  | _72_ G309G            | 1.69 (0.59–5.59)  | 1.63    | (0.57–5.43)         |         |
| Dominant and Other* | Other               | 1 (Reference)     | 0.082   | 1 (Reference)       | 0.086   |
| recessive (3.1)  | P72 G309G             | 4.82 (0.84–90.98) | 4.75    | (0.83–89.77)        |         |
| Dominant and Other* | Other               | 1 (Reference)     | 0.876   | 1 (Reference)       | 0.768   |
| recessive (3.2)  | P72PG309G             | 1.12 (0.28–5.52)  | 1.24    | (0.30–6.27)         |         |
| Double dominant (no effect accumulation) (4) | R72R T309T | 1 (Reference)     | 0.069   | 1 (Reference)       | 0.074   |
| Double recessive (no effect accumulation) (5) | Other | 0.37 (0.10–1.08)  | 0.37    | (0.10–1.10)         |         |
| Double dominant (with effect accumulation) (6) | Other | 1.39 (0.60–3.37)  | 1.45    | (0.62–3.54)         |         |
| Double recessive (with effect accumulation) (7) | Other | 0.34 (0.09–0.92)  | 0.34    | (0.09–0.94)         |         |
|          | P72 G309G             | 0.43 (0.11–1.47)  | 0.44    | (0.11–1.52)         |         |
| Double dominant (with effect accumulation) (8) | Other | 1.41 (0.59–3.55)  | 1.45    | (0.61–3.70)         |         |
|          | P72PG309G             | 1.21 (0.11–26.53) | 1.36    | (0.12–30.58)        |         |
| Quantitative (8) | N° of P72 and G309 alleles | 0.95 (0.65–1.39) | 0.781   | 0.96 (0.66–1.42)    | 0.854   |

*The epistatic models were numbered as described previously [46].

*Other: 3.1: _72_ T309_ - R72R T309G; 3.2: R72_ _309_ - _72_ T309T; 4: P72 _309_ - _72_ G309G; 5: P72P _309_ - _72_ G309G; 6: R72R G309_ - P72 _309_ T7: P72P T309_ - _72_ G309G.

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Effects of R72P SNP [model 1.2 (genotypic combinations P72 T309T and _72_ G309_, respectively); crude: OR (95% CI), 3.03 (1.33–7.59) and 2.80 (1.32–6.71); P = 0.015; adjusted: OR (95% CI), 2.57 (1.06–6.78) and 3.09 (1.38–7.72); P = 0.020] and double dominant epistasis with cumulative effect [model 7 (genotypic combinations P72P T309_ - R72_ G309G and P72P G309G, respectively); crude: OR (95% CI), 3.26 (1.52–7.81) and 2.31 (1.02–5.78); P = 0.007; adjusted: OR (95% CI), 3.29 (1.47–8.20) and 2.32 (0.97–6.04); P = 0.011]. Marginal associations were observed for double recessive epistasis without cumulative effects [model 5; crude: OR (95% CI), 1.56 (0.93–2.59); P = 0.093; adjusted: OR (95% CI), 1.72 (0.98–3.02); P = 0.060] and quantitative [model 8; crude: OR (95% CI), 1.23 (0.96–1.56); P = 0.098; adjusted: OR (95% CI), 1.28 (0.98–1.67); P = 0.074] models. Since statistically significant associations of different epistatic models with HPV oncogenic risk and HIV status were observed, the adjusted analyses for the first were repeated, including HIV status as a covariate (table S5). The significant associations were not maintained (P=0.142), thus suggesting a mediation effect of HIV status.

Since there were significant associations of different epistatic models with HIV status, a visual inspection of OR (with 95% CI) resulting from the adjusted analyses between genotypic combinations of the two SNPs and HIV was performed to identify patterns and, possibly, elaborate further epistatic models. Larger (and similar to one another) OR values were observed for the genotypic combinations R72R G309G and P72PT309T than for the rest (figure 1). This pattern indicates that a different epistatic effect may be underlying the observed associations. Therefore, an additional model was elaborated to reflect the following epistatic relationship: a double dominant effect that both affects disease risk and blocks a “hidden” additional recessive effect, which is manifested in the presence of a homozygous-wild homozygous-variant genotypic combination (which could be called double dominant epistasis with blocked recessive effects). Such model was tested in categorical (assuming distinct effects for the following genotypic combinations: R72R T309T (reference); R72P _309_ - R72R T309_, R72_ T309_ - R72R _309_, R72R _309_ - R72_ G309_.

| Epistatic model | Genotypic combination | Crude OR (95% CI) | P-value | Adjusted OR (95% CI) | P-value |
|----------------|-----------------------|-------------------|---------|----------------------|---------|
| Dominant (1.1) | R72R T309T            | 1 (Reference)     | 0.009   | 1 (Reference)        | 0.004   |
|                | R72P G309_            | 3.52 (1.54–8.88)  |         | 4.34 (1.79–11.57)   |         |
|                | P72_ _309_            | 2.63 (1.23–6.29)  |         | 2.45 (1.09–6.10)    |         |
| Dominant (1.2) | R72R T309T            | 1 (Reference)     | 0.015   | 1 (Reference)        | 0.020   |
|                | P72_ T309T            | 3.03 (1.33–7.59)  |         | 2.57 (1.06–6.78)    |         |
|                | _72_ G309_            | 2.80 (1.32–6.71)  |         | 3.09 (1.38–7.72)    |         |
| Recessive (2.1) | R72_ T309_            | 1 (Reference)     | 0.166   | 1 (Reference)        | 0.139   |
|                | R72_ _309_            | 1.28 (0.62–2.50)  |         | 1.46 (0.67–3.10)    |         |
|                | P72P _309_            | 1.88 (0.96–3.61)  |         | 2.01 (0.95–4.20)    |         |
| Recessive (2.2) | R72R T309_            | 1.96 (0.97–3.92)  |         | 1.97 (0.90–4.25)    |         |
|                | _72_ G309G            | 1.29 (0.66–2.45)  |         | 1.53 (0.73–3.14)    |         |
| Dominant and recessive (3.1) | Other | 1 (Reference) | 0.140 | 1 (Reference) | 0.420 |
| Dominant and recessive (3.2) | Other | 1 (Reference) | 0.948 | 1 (Reference) | 0.903 |
| Double dominant (no effect accumulation) (4) | Other | 2.88 (1.38–6.78) | 0.201 | 1 (Reference) | 0.060 |
| Double recessive (no effect accumulation) (5) | Other | 1.56 (0.93–2.59) | 1.72 (0.98–3.02) | 0.011 |
| Double dominant (with effect accumulation) (6) | Other | 3.26 (1.52–7.81) | 3.29 (1.47–8.20) | 0.011 |
|                | P72_ G309_            | 2.31 (1.02–5.78)  |         | 2.32 (0.97–6.04)    |         |
| Double recessive (with effect accumulation) (7) | Other | 1.57 (0.92–2.64) | 1.69 (0.95–3.00) | 0.011 |
|                | P72P G309G            | 1.41 (0.19–7.40)  |         | 2.29 (0.25–15.36)   |         |
| Quantitative (8) | N° of P72 and G309 alleles | 2.13 (0.96–1.56) | 0.998 | 1.28 (0.98–1.67) | 0.074 |

The epistatic models were numbered as described previously [46]. The underscore indicates that the effect is irrespective of the allele. E.g., R72P G309_ represents the genotypic combinations R72P T309G - R72P G309G. Other: 3.1: _72_ T309_ - R72R _309_. 3.2: _72_ T309_ - _72_ G309_. 4: P72_ _309_ - _72_ G309_. 5: P72P _309_ - _72_ G309G. 6: R72R G309_ - P72_ T309T. 7: P72P T309_ - R72_ G309G.

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T309G and P72P G309G; and R72R G309G and P72P T309T] and in numeric (i.e., linear tendency) forms, hereafter referred to as models 9.1 and 9.2, respectively. Even stronger associations than the previous ones were observed for models 9.1 [genotypic combinations R72P _309G_ - _72_ T309G - P72P G309G and R72R G309G - P72P T309T, respectively; crude: OR (95% CI), 2.43 (1.15-5.77) and 6.87 (2.72-18.91); P<0.001; adjusted: OR (95% CI), 2.47 (1.12-6.06) and 7.08 (2.59-21.01); P = 0.001] and 9.2 (crude: OR (95% CI), 2.66 (1.67-4.35); P<0.001; adjusted: OR (95% CI), 2.68 (1.62-4.60); P<0.001).

**Discussion**

We investigated, for the first time in the epidemiological setting, the evidence linking p53 and MDM2 with HIV, as well as potential implications of SNPs in the p53 pathway for HPV-related outcomes other than cancer development and/or progression. Analyses of epistatic models provided evidence for a novel mechanism linking the p53 pathway with HIV infection status. Interestingly, we observed large effect sizes for the epistatic models (achieving notably low P-values in a relatively small sample). This contrasts with a recent GWAS involving 6300 cases and 7200 controls that tested approximately 8 million common variants, which suggested that host genetic influences on HIV acquisition are either rare or have very small effects (not detectable given the study power) [20]. In our study, all genotypic combinations of the epistatic model 9.1 had a relatively high prevalence [R72R T309T: 50 (20.0%); R72R G309G - P72P T309T: 20 (8.0%) and the remaining genotypic combinations: 180 (72.0%) in HIV-positive and R72R T309T: 8 (8.0%); R72R G309G - P72P T309T: 22 (22.0%) and the remaining genotypic combinations: 70 (70.0%) in HIV-negative women] and large effect sizes (OR, others: 2.47; R72R G309G - P72P T309T: 7.08). Indeed, when comparing our findings with the cited GWAS studies on HIV acquisition [15,20], our effect sizes are larger than all of the OR of statistically-significant SNPs reported in these studies, with one exception (OR, 9.51) that was observed in an exploratory phase of the analyses but was markedly decreased in the expanded and external validation analyses (2.67 and 1.69, respectively) [18]. This observation reinforces the importance of analyzing epistasis even in genome-wide scale studies [47,48].

An important consideration is that we found relatively weak associations of epistatic models with HPV-related outcomes (especially for HPV status), which would not be expected given their strong associations with HIV status. While very strong associations were found between some epistatic models and HIV status, there were no associations of models 9.1 and 9.2 with HPV status and HPV oncogenic risk (data not shown). It is conceptually difficult to concede a factor causally involved in HIV status but not with HPV status when this is not adjusted for in the analysis (unless HIV status does not have a causal effect on HPV status). However, this issue does not invalidate our findings, since they are unlikely to be caused by confounding (discussed below) or chance alone (the P-values for models 9.1 and 9.2 were very low).

An important limitation of our study was the sample size (which increase was not possible due to practical/logistic reasons). Power analyses showed that, as expected, there was generally low power (power <0.80) when HPV oncogenic risk was the dependent...
variable (power <0.80) for single-SNP and epistasis analyses (due to reduced sample size) alike. For the former (table S6), associations involving OR<2 and performed under the recessive model (of which the exposure prevalence is smaller than of other genetic effects) were also generally underpowered. For the latter (table S7), power was also reduced for associations involving OR<3 and for several models when HIV status was the dependent variable. These results, in addition to the partial redundancy of some genetic/epistatic models, are reflected in the several marginal associations observed. Although these observations illustrate that our analyses were underpowered under some circumstances, they support the causality of our associations, since the effect sizes were large enough to achieve statistical significance even in a small sample. This is particularly illustrated in the epistatic model 9.1: while power analyses indicate low power when HIV status was the dependent variable, very strong associations were observed due to the large OR values.

Another important consideration is that the study design was not optimal for causal inference. This is particularly relevant considering the hypothesis that the SNPs (in combination) might influence virus establishment/persistence after exposure to it, which would be a rather dynamic mechanism. However, it is well-known that associations involving germ-line genetic markers as independent variables are not subjected to reverse causation and are generally robust against confounding, as reviewed in the context of Mendelian randomization [49]. To further reduce the possibility of residual confounding caused by population stratification, the analyses were adjusted for skin color and additional covariates. Nonetheless, the optimal design would be a prospective study to validate our findings and understand their underlying mechanisms. Besides, the results for models 9.1 and 9.2 were very similar, thus requiring additional studies for validating not only the genotypic combinations R72P T309G - R72P G309G, but also the actual model epistatic model. Another issue is the true relationship among genetic markers and the study outcomes: it cannot be determined from our study whether the SNPs are truly causal or are in linkage disequilibrium with the causal variants. Nevertheless, the value of our findings regarding genetic predisposition to HIV persistence after viral exposure is independent of this issue. In summary, our results provide evidence for a role of the p53 pathway – involving R72P and T309G SNPs – in HIV status and HPV oncogenic risk, and strong associations were found for an epistatic effect on HIV status. Our results require validation in prospective cohort studies aiming at either elucidating HIV pathogenesis or developing new therapeutic/preventive strategies.

Supporting Information

Table S1 Likelihood-ratio chi-squared tests P-values of the selection of confounders based on association with each outcome.

Table S2 Likelihood-ratio chi-squared tests P-values of the selection of confounders based on association with each SNP.

Table S3 Adjusted associations [showing OR (95% CI) and P-values], including HIV status as a covariate, between R72P SNP and HPV status and between T309G SNP and HPV oncogenic risk.

Table S4 Likelihood-ratio chi-squared tests P-values of the selection of confounders based on association with combined genotypes.

Table S5 Adjusted associations (including HIV status as a covariate) between 11 epistatic models and HPV oncogenic risk.

Table S6 Statistical power of the single-SNP analyses for different OR values.

Table S7 Statistical power of the epistasis analyses for different OR values.

Methods S1 Power Analysis.

Author Contributions

Conceived and designed the experiments: LGE TC FKS MFS. Performed the experiments: LGE EMN FMR. Analyzed the data: FPH. Contributed reagents/materials/analysis tools: TC FKS MFS. Wrote the paper: FPH.

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