CCR5 promoter activity correlates with HIV disease progression by regulating CCR5 cell surface expression and CD4 T cell apoptosis

Anjali Joshi1, Erin B. Punke1, Melina Sedano1, Bethany Beauchamp2, Rima Patel2, Cassady Hossenlopp2, Ogechika K. Alozie3, Jayanta Gupta4,5, Deabrata Mukherjee3 & Himanshu Garg1

CCR5 is the major co-receptor for HIV and polymorphisms in the CCR5 gene as well as promoter region that alter cell surface expression have been associated with disease progression. We determined the relationship between CCR5 promoter polymorphisms and CD4 decline and other immunopathological features like immune activation and CD4+ T cell apoptosis in HIV patients. CCR5 promoter haplotype HHC was significantly associated with higher CD4 counts in patients. The relative promoter activity (RPA) of each haplotype was determined in vitro and combined promoter activity based on both alleles (CRPA) was assigned to each patients. Interestingly, CCR5 CRPA correlated inversely with CD4 counts and CD4:CD8 ratio specifically in viremic patients. In normal individuals, the CRPA correlated with the number of CCR5+CD4+ T cells in the peripheral blood suggesting an effect on CCR5 expression. In a subset of high viremic patients harboring R5 tropic HIV, there was a strong correlation between CCR5 CRPA and both CD4 counts and CD4 T cell apoptosis. Our study demonstrates that, CCR5 promoter polymorphisms correlate with CD4 T cell loss possibly by regulating CD4 T cell apoptosis in HIV patients. Furthermore, assigning CRPAs to each patient is a new method of translating genotype to phenotype.
suggest that the protective effect of CCR5 polymorphisms may largely be limited to progression to AIDS in HIV+ individuals, the precise mechanism underlying the role of CCR5 levels in HIV disease remains unclear.

In this study, we took a different approach to understanding the potential role of CCR5 polymorphisms in clinical manifestations of HIV infection. We genotyped the CCR5 promoter region of HIV+ patients and determined the haplotypes for each allele. We then assessed the activity of each CCR5 promoter haplotype using an in vitro gene reporter assay. After normalizing the promoter activities to the ancestral HHA haplotype, we assigned each haplotype a numerical value termed Relative Promoter Activity (RPA). We then calculated the Combined Relative Promoter Activity (CRPA) for each patient by adding the RPAs of each allele. As a result, we could convert the genotypic information into a numerical value that could be used to correlate with various clinical manifestations in HIV patients. We find that CRPA correlates with CD4 counts, AIDS phenotype and CD4 T cell apoptosis in HIV patients. In normal individuals CRPA correlates with CCR5+ cells in peripheral blood confirming a direct relationship with CCR5 expression. This indicates that CCR5 promoter polymorphisms alter CCR5 expression levels that influence CD4 apoptosis and consequently CD4 decline in HIV patients. To the best of our knowledge, this is the first study demonstrating a new approach of assigning numerical scores (CRPA) based on CCR5 haplotype that also correlates with HIV disease progression/pathogenesis.

Results

**CCR5 promoter polymorphisms in HIV infected population.** CCR5 promoter polymorphisms have been associated with HIV disease progression2–4. Seven SNPs in the promoter region have been identified in the human population that collectively make up the haplotypes HHA through HHG (Supplementary Table 1). A 32 bp deletion in the CCR5 gene is also prominent in the human population and differentiates the HHG1 and HHG2 haplotypes18. We determined the SNPs in the CCR5 promoter region as well as the CCR5 Δ32 genotype and the allele frequency of each haplotype in our HIV infected and control population (Table 1). The most frequent haplotypes in HIV infected individuals were HHC (42%) and HHE (28%) which collectively made up 70% of the total haplotypes. Our data is much in line with other studies showing that in terms of HIV infection, no particular SNP or haplotype was associated with acquisition of HIV. In fact, we found 3/50 (6%) of the patients in our study were CCR5Δ32 heterozygous which corresponds with the frequency of the mutation in Hispanic population19. We did not find CCR5Δ32 homozygous individuals in our HIV+ group, which is consistent with resistance of this population to HIV infection6,8. Overall, our data indicate that our study population shows similar distribution of genotypes as seen in other Hispanic populations and CCR5 promoter polymorphism in our study population does not protect against acquisition of HIV infection.

**Individual CCR5 promoter SNPs are associated with CD4 levels in HIV infected patients.** Multiple studies have reported CCR5 promoter polymorphisms to be associated with disease progression and CD4 levels in patients6,20–22. We determined whether specific SNPs correlated with CD4 levels in HIV infected individuals. We studied 6 SNPs: 58755A, 58934G, 59029A, 59353C, 59402A and 59653C. Frequency of SNPs in the HIV+ and normal population is listed in Supplementary Table 2. Interestingly, we found that 58934GG (p = 0.02), 59029AA (p = 0.04), 59353CC (p = 0.04) and 59402AA (p = 0.03) were significantly associated with lower CD4 counts in our patient population (Fig. 1A–F). Closser analysis revealed that these SNPs collectively make up the non HHC group (Supplementary Table 1). Thus, our preliminary data analysis revealed that certain CCR5 promoter SNPs may play a role in regulating HIV disease progression as evident from the CD4 counts.

**HHC haplotype is associated with higher CD4 counts in HIV infected patients.** Previous studies have reported a protective effect of HHC haplotype on the decline in CD4 counts in Caucasian population6. Our individual SNP analysis also suggested that HHC haplotype may be associated with higher CD4 levels (Fig. 1). We hence divided our population into HHC group (N = 33), where at least one allele was HHC, or non HHC group (N = 17), where neither allele was HHC (Supplementary Table 3). We then asked whether HHC versus non HHC haplotype affected the immunopathological features of HIV infection, including CD4 counts, immune activation, CD4 apoptosis and viremia. Interestingly, we found that CD4 counts in the non HHC group were lower than HHC group (p = 0.0174) (Fig. 2A), and this effect was more dramatic in viremic patients (viremic defined as ≥100 virus copies/ml) (p = 0.0014) (Fig. 2B) (Supplementary Table 4). Analysis of CD4:CD8 ratios, another

| Population (frequency) | HIV+ (N = 50) | Normal (N = 28) |
|------------------------|--------------|----------------|
| HHA                    | 4 (0.040)    | 3 (0.054)      |
| HHB                    | 0 (0.000)    | 0 (0.000)      |
| HHC                    | 42 (0.420)   | 19 (0.339)     |
| HHD                    | 2 (0.020)    | 0 (0.000)      |
| HHE                    | 28 (0.280)   | 14 (0.250)     |
| HHE                    | 11 (0.110)   | 14 (0.250)     |
| HHE                    | 10 (0.10)    | 4 (0.07)       |
| HHG2                   | 3 (0.03)     | 2 (0.035)      |

Table 1. Frequency of different CCR5 promoter haplotypes in the HIV+ population and normal controls. The HHG2 haplotype includes CCR5Δ32 genotype.
predictor of HIV pathogenesis, did not show significant differences between HHC and non HHC groups in either all HIV+ (Fig. 2C) or viremic (Fig. 2D) HIV patients. Similarly, we also looked at CD4 apoptosis (defined as active caspase + cells) between the groups and found that non HHC group showed modestly higher CD4 apoptosis (Fig. 2E) that trended but was not statistically significant (p = 0.0681) in the HIV+ population. However, within the viremic group, there was a stronger trend towards higher apoptosis in non HHC group compared to HHC (p = 0.0527) (Fig. 2F). Immune activation, defined by the presence of CD8+CD38+HLADR+ cells is a classical immunopathological feature of HIV infection and correlates strongly with CD4 decline 23–25. We hence looked at immune activation in these groups and found no significant difference in immune activation in CD8+ cells in the entire HIV+ group (p = 0.181) (Fig. 2G), or viremic patients (p = 0.1097) (Fig. 2H). With regards to log viremia, no significant differences were observed between the HHC and non HHC group in either the HIV+ (Fig. 2I) or viremic patients (Fig. 2J). Based on these data we can conclude that non HHC haplotype is associated with lower CD4 counts, which in part may be related to higher CD4 apoptosis in this group. The lack of difference in viremia in the two groups also indicates that in our study the effect of CCR5 polymorphism is likely to be independent of viremia.

**CCR5 promoter haplotypes have different transcriptional activity.** CCR5 promoter polymorphisms potentially modify gene expression and consequently affect cell surface expression of CCR5 3, 26. As CCR5 is the major co-receptor for HIV and interactions between CCR5 and Env glycoprotein of HIV are believed to be fundamental to HIV infection, replication and HIV pathogenesis. Thus alterations in CCR5 levels are functionally significant to HIV disease. We have previously demonstrated that the levels of CCR5 on the cell surface can in fact alter bystander apoptosis mediated by HIV Env glycoprotein 27. Hence, we next determined the promoter activity of each of the seven CCR5 promoter haplotypes (HHA-HHG) using a luciferase based gene reporter assay (Fig. 3A). We then normalized the activity of each promoter haplotype to the ancestral HHA haplotype. Based on these data, each haplotype was assigned a numerical value or the Relative Promoter Activity (RPA) (Fig. 3B). We found that the haplotypes could be divided into two groups. The low group included HHA, HBH, HHC and HHD while the high group comprised of HHE, HHF and HHG. Interestingly, the high group showed almost 30% higher RPA in our assay than HHA while HHC haplotype showed a low RPA of 96% (Fig. 3B). We assigned HHG2, which is in complete linkage disequilibrium with CCR5Δ32, a RPA value of 0 as the 32 bp deletion in CCR5 ORF results in a truncated protein that is not expressed on the cell surface 10. Using this approach we could now assign each patient a Combined Relative Promoter Activity (CRPA) score based on the sum of the RPA of each allele. The CRPA of the common allele pairs in our study is shown in Supplementary Table 3. The CRPA score now provides us the opportunity of assigning a numerical value to the CCR5 genotype that can be correlated with different immunopathological parameters in HIV infection. Interestingly, we found that all the three CCR5Δ32 individuals in our HIV infected population had a high RPA allele (HHG1, HHE or HHF) as the second allele (Supplementary Table 3). This is interesting, as in the context of CCR5Δ32, the protective effect in heterozygous populations against HIV acquisition may be related to the RPA of the second allele, as has...
been reported by Hladik et al. Similar luciferase based gene reporter assays were also conducted after cloning a smaller version of the CCR5 promoter as described by Mummidi et al. Data with smaller CCR5 promoter yielded similar Relative Promoter Activity (Supplementary Figure 1).

Using the above methods we assigned each patient a numerical value or CRPA score for CCR5 promoter activity and then asked whether CRPA of each patient correlated with CD4 counts and other immunopathological parameters. Based on CRPA score, our population could be divided into two groups of CRPA > 229 or CRPA < 229. This classification was based on the observation that > 229 did not contain any HHC haplotype (Supplementary Table 3). Once again, we found that CRPA < 229 was associated with higher CD4 counts (p = 0.0082) (Fig. 4A), which was more dramatic in viremic patients (p < 0.0001) (Fig. 4B). We also looked at CD4:CD8 ratios and found a similar trend towards higher CD4:CD8 ratio in RPA < 229 group (Fig. 4C), which was significant (p = 0.0423) in the viremic group (Fig. 4D). With regards to immune activation, there was no significant difference in patients with CRPA > 229 or CRPA < 229 in either all HIV+ patients or the viremic patients group (Fig. 4E and F), although RPA > 229 had slightly higher immune activation. However, with regards to CD4 apoptosis, there was a trend towards higher apoptosis in the CRPA > 229 group in all HIV+ patients (p = 0.0797) (Fig. 4G) but not in viremic patients (p = 0.1091) (Fig. 4H). Closer look at the viremic group revealed that one of the samples in the CRPA < 229 group was from an HHG2 patient, who had detectable viremia (8810 copies/ml) and also showed high apoptosis in CD4 cells. This patient may be harboring a dual or X4 tropic virus and when excluded from the analysis, we saw significant difference (p = 0.0351) in CD4 apoptosis between CRPA < 229 no HHG2 vs CRPA > 229 (Fig. 4I). There was no significant difference in log viremia between CRPA > 229 vs CRPA < 229 in all HIV+ (Fig. 4I) or in viremic patients (Fig. 4K).

**Combined Relative promoter activity of both alleles in HIV patients as a new parameter for disease progression.** Using the above methods we assigned each patient a numerical value or CRPA score for CCR5 promoter activity and then asked whether CRPA of each patient correlated with CD4 counts and other immunopathological parameters. Based on CRPA score, our population could be divided into two groups of CRPA > 229 or CRPA < 229. This classification was based on the observation that > 229 did not contain any HHC haplotype (Supplementary Table 3). Once again, we found that CRPA < 229 was associated with higher CD4 counts (p = 0.0082) (Fig. 4A), which was more dramatic in viremic patients (p < 0.0001) (Fig. 4B). We also looked at CD4:CD8 ratios and found a similar trend towards higher CD4:CD8 ratio in RPA < 229 group (Fig. 4C), which was significant (p = 0.0423) in the viremic group (Fig. 4D). With regards to immune activation, there was no significant difference in patients with CRPA > 229 or CRPA < 229 in either all HIV+ patients or the viremic patients group (Fig. 4E and F), although RPA > 229 had slightly higher immune activation. However, with regards to CD4 apoptosis, there was a trend towards higher apoptosis in the CRPA > 229 group in all HIV+ patients (p = 0.0797) (Fig. 4G) but not in viremic patients (p = 0.1091) (Fig. 4H). Closer look at the viremic group revealed that one of the samples in the CRPA < 229 group was from an HHG2 patient, who had detectable viremia (8810 copies/ml) and also showed high apoptosis in CD4 cells. This patient may be harboring a dual or X4 tropic virus and when excluded from the analysis, we saw significant difference (p = 0.0351) in CD4 apoptosis between CRPA < 229 no HHG2 vs CRPA > 229 (Fig. 4I). There was no significant difference in log viremia between CRPA > 229 vs CRPA < 229 in all HIV+ (Fig. 4I) or in viremic patients (Fig. 4K).

**CRPA correlates with CD4 decline.** Correlation analysis and regression models are difficult with data divided into groups. Assigning CRPA scores allows us to not only conduct regression analysis but also account for both alleles in the patients. We found a strong correlation between CD4 counts and CCR5 CRPA in all HIV patients (p = 0.0042) (Fig. 5A), as well as in viremic patients (p = 0.0038) (Fig. 5B). CCR5 CRPA also correlated...
with CD4:CD8 ratios (Fig. 5C), being significant in viremic patients \( (p = 0.0414) \) (Fig. 5D). Finally, CRPA was significantly higher \( (p = 0.0008) \) in patients with AIDS, defined as CD4 counts less than 200/μl of blood (Fig. 5E). Linear regression models showed that the CRPA based classification correlated better with CD4 counts compared to HHC vs non HHC classification both before and after adjusting for viremia (Table 2). Taken together, these findings suggest that the CRPA based approach correlates with HIV disease progression and could be used as a parameter to predict disease outcomes.

CRPA correlates with number of CD4+ CCR5+ T cells in the peripheral blood. As promoter activity is likely to affect protein expression, we next looked at the potential correlation between the CRPA methodology and cell surface expression of CCR5 in CD4+ T cells. Other groups have demonstrated that CCR5 promoter polymorphism associates with the number of CCR5+ CD4 T cells in the peripheral blood \(^6, 26\). Using samples from healthy controls, we found that in fact CCR5 CRPA of the individuals correlates significantly \( (p = 0.0215) \) with percent CCR5+ CD4 T cells in the peripheral blood (Fig. 6A). Also, after dividing the populations in to CRPA > 229 and CRPA < 229 groupings, we found that there was a significant difference \( (p = 0.0246) \) in CCR5+ cells between the groups (Fig. 6B). These data are in line with results of others who have shown a correlation between CCR5 polymorphisms and CCR5 expression \(^3, 12, 28\). However, we show here that assigning a CRPA value to each sample can be an alternate method for correlating polymorphisms/genotype with different phenotypes.

R5 virus infected viremic patients show correlation between CCR5 CRPA and CD4 apoptosis and CD4 decline. We next asked whether we could get additional information regarding the mechanism behind the correlation between CCR5 polymorphisms and HIV pathogenesis. CCR5 is the major co-receptor for HIV and is directly involved in binding to the viral Env glycoprotein. Env-CCR5 interactions are fundamental to both viral entry and pathogenesis. In fact, the role of CCR5 binding by Env in HIV mediated bystander apoptosis has been demonstrated \(^27, 29\). We previously cloned full length Env glycoproteins from 11 of the high viremic patients (>10,000 copies/ml) in our cohort and found that all of these Envs were CCR5 tropic \(^24\). We hence asked whether in individuals with high viremia, harboring R5 tropic virus, if there was a correlation between CCR5 CRPA and CD4 decline and/or CD4 apoptosis. As shown in Fig. 7A there was a significant correlation \( (p = 0.0128) \) between CD4 counts and CCR5 CRPA and a strong trend between CCR5 CRPA and CD4:CD8 ratios (Fig. 7B). Moreover, a significant correlation \( (p = 0.0368) \) was found between CD4 apoptosis and CCR5 CRPA (Fig. 7C) consistent with data above. These findings suggest that the correlation between CCR5 CRPA and CD4 counts could in part be mediated by the susceptibility of cells to Env mediated bystander apoptosis.

Discussion

The correlation between CCR5 promoter polymorphisms in HIV disease progression has been extensively studied and a strong relationship between certain CCR5 promoter haplotypes and disease progression has been demonstrated \(^3, 6, 15, 21, 30\). While these correlations have often been associated with CCR5 levels, the mechanism behind the role of CCR5 promoter polymorphisms and HIV disease progression remains less certain.
Gonzalez et al. demonstrated that HHE haplotype is associated with accelerated progression to AIDS in Caucasian population, while HHC may be associated with delayed progression. Although HHC and HHE are two of the most common haplotypes, as was also confirmed in our study, there are other haplotypes that need to be investigated. With the frequency of other haplotypes being low in many cohorts it becomes difficult to undertake these studies based on genotypic classification alone. Also, it is imperative that allele pairs be accounted for in these studies. Since CCR5 promoter polymorphisms may influence CCR5 protein levels, and thus affect disease progression, we investigated whether promoter activity could be used as a surrogate for CCR5 levels and consequently disease progression in HIV patients.

We established the RPA of each haplotype using an in vitro gene reporter assay. After normalizing to HHA, the ancestral haplotype, we assigned numerical values to each promoter haplotype. Similarly, each patient could be assigned a CRPA based on the sum of the RPAs of the 2 alleles of each individual. This method converts genotypic information into a numerical score based on aggregate promoter activity of the allele pair. Using this approach we could now stratify our population based on functional CRPA scores, and not just on allele combinations. With this method we found that HHE, HHF and HHG had similar promoter activity that was about 130% of the ancestral HHA haplotype, and that HHC activity was slightly lower than HHA at 96%. We were able to stratify our population into two groups of patients, those with CRPA > 229 or CRPA < 229. Notably, patients with HHE/HHE, HHE/HHF or HHE/HHG belonged to the high CRPA group and those with HHC and other low activity haplotypes like HHA, HHB or HHD were in the low group. Using this strategy, we found a correlation between CD4 counts and CRPA values, with the cutoff between CRPA > 229 vs CRPA < 229.

Interestingly, we found that the differences between low and high CRPA groups were more significant when considering viremic patients (≥100 virus copies/ml of viral RNA) compared to the total population. As CCR5 is the major co-receptor for HIV and the viral Env glycoprotein directly binds to it, it is not surprising that the effects are more relevant in the presence of active virus replication/viremic patients. At the same time, we also found that differences between CRPA groups showed a strong trend in terms of CD4 apoptosis. We and others...

Figure 4. Combined CCR5 relative promoter activity (CRPA) of both alleles in HIV patients correlates with CD4 counts. Each patient was assigned a numerical CRPA score based on the RPA of both alleles. The patient population was then divided into two groups of CRPA > 229 or CRPA < 229. CD4 counts in CRPA > 229 vs CRPA < 229 in (A) all HIV+ patients (B) viremic patients. CD4:CD8 ratio in CRPA > 229 vs CRPA < 229 in (C) all HIV+ patients (D) viremic patients. CD8 immune activation in CRPA > 229 vs CRPA < 229 in (E) all HIV+ patients (F) viremic patients. CD4 apoptosis in CRPA > 229 vs CRPA < 229 in (G) all HIV+ patients (H) viremic patients. (I) CD4 apoptosis in CRPA > 229 vs CRPA < 229 no HHG2 (after removing one data point from an HHG2 patient). Log viremia in CRPA > 229 vs CRPA < 229 in (J) all HIV+ or (K) viremic patients. One tailed student’s t-test was used for statistical analysis.
have shown that CD4 apoptosis correlates with CD4 decline in HIV patients. Also, binding of HIV Env to CCR5 is required for bystander apoptosis induction. We have also shown that levels of CCR5 on cell surface determines bystander apoptosis of cells via HIV Env, with higher CCR5 expression associated with increased bystander apoptosis in vitro. The higher apoptosis in the CRPA > 229 group is consistent with this idea. Furthermore, within the group of high viremic patients where we were successful in cloning full length CCR5 tropic Envs, we found a strong correlation between CCR5 CRPA and CD4 apoptosis. Taken together, these data suggest that higher CCR5 promoter activity and consequently higher CCR5 expression may be related to higher CD4 apoptosis mediated by primary R5 Envs (Fig. 8).

While bystander apoptosis is one the factors in HIV pathogenesis, it is clear from recent studies that HIV pathogenesis is multifactorial. In a model presented in our previous study, we proposed a role for the Env glycoprotein phenotype, viremia and immune activation in collectively mediating CD4 bystander apoptosis. Based on the findings from this study, inclusion of CCR5 promoter polymorphisms in this model is justified (Fig. 8). This is because levels of CCR5 can affect other HIV pathogenesis regulating factors as well as be altered by them. We know that increased cell surface CCR5 levels would likely support higher virus replication, as is clear from the lower viremia in CCR5Δ32 heterozygous individuals. The strong association between viremia and immune activation in HIV infections suggests that higher virus replication in CCR5 WT individuals may be related to higher immune activation. A trend towards higher immune activation in our samples is indicative of the complexity of HIV pathogenesis.

**Figure 5.** Correlation of CCR5 CRPA with CD4 decline. Correlation between CD4 counts and CCR5 CRPA in (A) all HIV+ or (B) viremic patients. Correlation of CCR5 CRPA with CD4:CD8 ratios in (C) all HIV+ or (D) viremic patients. (E) CRPA in patients with CD4 counts less than or more that 200/μl of blood. Spearman’s correlation with linear regression was used for correlation determination. One tailed student’s t-test was used for comparative statistics.
of this phenomenon. Recently, Gornalusse et al. have shown that immune activation can directly affect CCR5 promoter activity and alter cell surface CCR5 expression\(^3\). Finally, levels of CCR5 can affect HIV-mediated CD4 loss as seen in vitro by us\(^2\) as well as in SCID-hu mice studies by Scoggins et al.\(^4\). Interestingly, a role of anti-CCR5 antibodies in mediating CCR5 downregulation has been demonstrated in some long term non progressors (LTNP)\(^4\), further supporting a role for CCR5 levels in disease progression. Hence, a new and more extensive model of HIV pathogenesis can now be presented that incorporates CCR5 levels regulated via CCR5 gene and promoter polymorphisms (Fig. 8).

Overall, we have quantified the activity of different CCR5 promoter haplotypes, and used this information to assign each individual a CRPA score based on their allele pair. Our results yielded strong correlation between CRPA and CD4 levels in HIV+ patients. The correlation of CD4 T cell apoptosis with CCR5 polymorphism is also shown for the first time in our study. Hence our study provides a mechanistic explanation to the relationship between CCR5 polymorphisms and CD4 loss/disease progression. Use of the CRPA strategy for determining disease outcome in HIV patients needs to be further validated in longitudinal studies as well as in larger sample sizes. We anticipate that this methodology can be used for prognosis and for improved design of anti-HIV therapeutic regimens.

**Methods**

**Ethical Statement.** The study was reviewed and approved by the Texas Tech University Health Sciences Center, El Paso, Institutional Review Board (IRB) and all experiments were performed in accordance with relevant guidelines and regulations. The study design was cross-sectional and the study number recorded as IRB# E12092, approval date 07/31/2012. All participants were provided with written and oral information about the study. Written informed consent of all study participants in accordance with the institutional policy was documented. In accordance with institutional IRB guidelines, all participants were identified by coded numbers to assure anonymity and all patient records kept confidential.
Patient population. Fifty HIV-infected individuals and 28 healthy controls were recruited from the outpatient HIV clinic at the Texas Tech University Health Sciences Center at El Paso. The mean age of the HIV+ patient population was $37.9 \pm 11.9$. The HIV group comprised of 9 females (18.0%) and 41 males (82.0%). The study was cross sectional consisting of patients at different stages of the disease. Viremic patients were defined as those having $\geq 100$ copies of viral RNA. An age matched healthy population control group ($n = 28$) was also recruited from the same geographical location and comprised of 9 females (32%) and 19 males (68%). The mean age of the healthy control group was $34.5 \pm 10.12$. Further details of patient population including CD4 counts, viral load and HAART status have been described previously.\(^{24}\)

Sample collection and storage. Each patient provided a 20 ml blood sample that was separated into plasma and cellular components using Ficoll based separation. Genomic DNA was extracted from whole blood samples using the QIAamp DNA Blood Mini kit (Qiagen). All components from the sample including plasma, cells and DNA were aliquoted and stored at $-70^\circ C$ till further analysis.

CCR5 gene and promoter polymorphisms. Genomic DNA isolated from whole blood was used for PCR amplification of CCR5 promoter region or for CCR5$\Delta 32$ genotype determination. CCR5$\Delta 32$ genotype was determined by PCR amplification using primers spanning the 32 bp deletion as described by Kaur et al.\(^{42}\). Individuals with the $\Delta 32$ allele showed a 32 bp smaller PCR fragment compared to wild type. CCR5 promoter polymorphisms were determined by PCR amplification of CCR5 promoter region from genomic DNA using Phusion high fidelity PCR kit (New England Biosciences) followed by Sanger’s sequencing. Sequences were analyzed using the DNA STAR software and polymorphisms identified based on sequence peaks.

Generation of CCR5 promoter haplotypes. CCR5 promoter regions from HHC or HHG homozygous individuals were PCR amplified using specific primers spanning the region $-3139$ to $-1317$ (CCR5 long promoter) or $-2761$ to $-1814$ (CCR5 short promoter) based on numbering by Mummidi et al.\(^{5}\) and cloned into
TA cloning kit (Invitrogen). Subsequently, the promoter fragment was digested with Pkn-1 and Xho-1 and sub-cloned into pGL-3 basic vector (Promega) upstream of the luciferase gene. Site directed mutagenesis with Quick Change Site Directed Mutagenesis Kit (Stratagene) was used to generate the haplotypes HHA, HHB, HHD, HHE and HHH.

**CCR5 promoter activity determination.** CCR5 promoter activity was determined in 293T cells by transfecting with plasmid DNA (0.3 μg/well of a 96-well plate) from each CCR5 gene promoter haplotype (HHA, HHB, HHC, HHD, HHE, HHH and HHHG) using the TurboFect™ Transfection Reagent (Fisher Scientific). Promoter activity was determined 48 h later by measuring luciferase activity in cell lysates using Brite Lite Substrate (Perkin Elmer). Relative promoter activity (RPA) of each haplotype was calculated after normalizing to HHA, the ancestral allele. Each HIV patient was given a CRPA score based on their two alleles.

**Immunostaining.** Staining of cells for different immune markers has been described previously. Briefly, lymphocytes isolated from the blood samples obtained from HIV-infected or normal patients were stained for cell surface markers using specific antibodies (Supplementary Figure 2). The immune activation panel consisted of antibodies CD3- Cy7, CD4-Tx Red, CD8-APC (Beckman Coulter) along with immune activation markers CD38 PE and HLA-DR FITC (BD Biosciences). The apoptosis panel comprised of the following antibodies CD3-Cy7, CD4-Tx Red, CD8-APC (Beckman Coulter), CCR5 PE (BD Biosciences) along with apoptosis marker CaspACE FITC-VAD-FMK (Promega). Stained cells were washed and fixed using Cytofix reagent (Beckman coulter) and acquired on a 10 color Beckman Coulter Gallios flow cytometer. At least 20,000 events for each sample were acquired. Data was analyzed using FlowJo software (Tree Star). Cells were first gated on CD3+ population and immune activation/apoptosis on CD4+ and CD8+ T cell subsets determined.

**Statistical analysis.** Data were analyzed using GraphPad Prism (GraphPad Software, Inc, La Jolla, CA) and SAS 9.3 (SAS Institute, Inc., Cary, North Carolina). Differences between groups were assessed using the two-sample t-test; either one sided or two sided based on the hypothesis. All p values were considered significant at p < 0.05. Spearman's correlation with linear regression was used for all correlation determination using the GraphPad Prism Software. The association of CCR5 haplotypes (HHC and non-HHC) and CRPA groups (CRPA < 229 and CRPA >= 229) with CD4 counts, CD8 immune activation, CD4 apoptosis and CD4:CD8 ratio was determined using linear regression models, unadjusted and adjusted for viremia.

**References**

1. McDermott, D. H. et al. CCR5 promoter polymorphism and HIV-1 disease progression. Multicenter AIDS Cohort Study (MACS). *Lancet* 352, 866–870 (1998).
2. Martin, M. P. et al. Genetic acceleration of AIDS progression by a promoter variant of CCR5. *Science* 282, 1907–1911 (1998).
3. Hladik, F. et al. Combined effect of CCR5-Delta32 heterozygosity and the CCR5 promoter polymorphism — 2459 A/G on CCR5 expression and resistance to human immunodeficiency virus type 1 transmission. *Journal of virology* 79, 11677–11684, doi:10.1128/JVI.79.18.11677-11684.2005 (2005).
4. Catano, G. et al. Concordance of CCR5 genotypes that influence cell-mediated immunity and HIV-1 disease progression rates. *The Journal of infectious diseases* 203, 263–272, doi:10.1093/infdis/jiq023 (2011).
5. Mummidi, S. et al. Evolution of human and non-human primate CC chemokine receptor 5 gene and mRNA. Potential roles for haplotype and mRNA diversity, differential haplotype-specific transcriptional activity, and altered transcription factor binding to polymorphic nucleotides in the pathogenesis of HIV-1 and simian immunodeficiency virus. *The Journal of biological chemistry* 275, 18946–18961, doi:10.1074/jbc.M000169200 (2000).
6. Gonzalez, E. et al. Race-specific HIV-1 disease-modifying effects associated with CCR5 haplotypes. *Proceedings of the National Academy of Sciences of the United States of America* 96, 12004–12009 (1999).
7. Malhotra, R. et al. Association of chemokine receptor gene (CCR2-CCR5) haplotypes with acquisition and control of HIV-1 infection in Zambians. *Retrovirology* 8, 22, doi:10.1186/1742-4690-8-22 (2011).
8. Liu, R. et al. Homozygous defect in HIV-1 coreceptor accounts for resistance of some multiply-exposed individuals to HIV-1 infection. *Cell* 86, 367–377 (1996).
9. Samson, M. et al. Resistance to HIV-1 infection in caucasian individuals bearing mutant alleles of the CCR-5 chemokine receptor gene. *Nature* 382, 722–725, doi:10.1038/382722a0 (1996).
10. Dean, M. et al. Genetic restriction of HIV-1 infection and progression to AIDS by a deletion allele of the CRK5 structural gene. *Hemophilia Growth and Development Study, Multicenter AIDS Cohort Study, Multicenter Hemophilia Cohort Study, San Francisco City Cohort, ALIVE Study*. *Science* 273, 1856–1862 (1996).
11. Huang, Y. et al. The role of a mutant CCR5 allele in HIV-1 transmission and disease progression. *Nature medicine* 2, 1240–1243 (1996).
12. de Roda Husman, A. M., Blaak, H., Brouwer, M. & Schuitemaker, H. CC chemokine receptor 5 cell-surface expression in relation to CC chemokine receptor 5 genotype and the clinical course of HIV-1 infection. *J Immunol* 163, 4597–4603 (1999).
13. Wu, L. et al. CCR5 levels and expression pattern correlate with infectability by macrophage-tropic HIV-1, in vitro. *The Journal of experimental medicine* 185, 1681–1691 (1997).
14. de Roda Husman, A. M. et al. Association between CCR5 genotype and the clinical course of HIV-1 infection. *Annals of internal medicine* 127, 882–890 (1997).
15. Mummidi, S. et al. Genealogy of the CCR5 locus and chemokine system gene variants associated with altered rates of HIV-1 disease progression. *Nature medicine* 4, 786–793 (1998).
16. Liu, H. et al. Polymorphisms in CCR5 chemokine receptor gene in Japanese. *International journal of immunogenetics* 34, 325–335, doi:10.1111/j.1744-313X.2007.00694.x (2007).
17. Kulkarni, H. et al. Role of CCL3L1-CCR5 genotypes in the epidemic spread of HIV-1 and population vaccine efficacy. *PloS one* 3, e3671, doi:10.1371/journal.pone.0003671 (2008).
18. Burns, J. C. et al. Genetic variations in the receptor-ligand pair CCR5 and CCL3L1 are important determinants of susceptibility to Kawasaki disease. *The Journal of infectious diseases* 192, 344–349, doi:10.1086/430953 (2005).
19. Salas-Alarcon, J. C., Mellerio, J. E., Ashton, G. H. & McGrath, J. A. Frequency of the CCR5 gene 32-basepair deletion in Hispanic Mexicans. *Clinical and experimental dermatology* 24, 127–129 (1999).
20. Anastassopoulou, C. G. & Kostridis, L. G. The impact of human allelic variation on HIV-1 disease. *Current HIV research* 1, 185–203 (2003).
21. Mangano, A. et al. Concordance between the CC chemokine receptor 5 genetic determinants that alter risks of transmission and disease progression in children exposed perinatally to human immunodeficiency virus. *The Journal of infectious diseases* **183**, 1574–1585, doi:10.1086/320705 (2001).

22. Tang, J. et al. Distribution of chemokine receptor CCR2 and CCR5 genotypes and their relative contribution to human immunodeficiency virus type 1 (HIV-1) seroconversion, early HIV-1 RNA concentration in plasma, and later disease progression. *Journal of virology* **76**, 662–672 (2002).

23. Hunt, P. et al. Relationship between T cell activation and CD4+ T cell count in HIV-seropositive individuals with undetectable plasma HIV RNA levels in the absence of therapy. *The Journal of infectious diseases* **197**, 126–133, doi:10.1086/524143 (2008).

24. Joshi, A. et al. HIV-1 Env Glycoprotein Phenotype along with Immune Activation Determines CD4+ T Cell Loss in HIV Patients. *J Immunol* **196**, 1678–1679, doi:10.1089/jimmunol.1901388 (2016).

25. Liu, Z. et al. Elevated CD38 antigen expression on CD8+ T cells is a stronger marker for the risk of chronic HIV disease progression to AIDS and death in the Multicenter AIDS Cohort Study than CD4+ cell count, soluble immune activation markers, or combinations of HLA-DR and CD38 expression. *Journal of acquired immune deficiency syndromes and human retrovirology: official publication of the International Retrovirology Association* **16**, 83–92 (1997).

26. Shieh, B. et al. Influence of nucleotide polymorphisms in the CCR2 gene and the CCR5 promoter on the expression of cell surface CCR5 and CXCR4. *International Immunology* **12**, 1311–1318 (2000).

27. Joshi, A., Nyakeriga, A. M., Ravi, R. & Garg, H. HIV ENV glycoprotein-mediated bystander apoptosis depends on expression of the CCR5 co-receptor at the cell surface and ENV fusogenic activity. *The Journal of biological chemistry* **286**, 36404–36413, doi:10.1074/jbc.M111.281659 (2011).

28. Salkowitz, J. R. et al. CCR5 promoter polymorphism determines macrophage CCR5 density and magnitude of HIV-1 propagation in vitro. *Clin Immunol* **108**, 234–240 (2003).

29. Tsao, L. C., Guo, H., Jeffrey, J., Hoxie, J. A. & Su, L. CCR5 interaction with HIV-1 Env contributes to Env-induced depletion of CD4+ T cells in vitro and in vivo. *Virology* **366**, 22, doi:10.1016/j.virol.2007.02.022 (2007).

30. Gornalusse, G. G. et al. Epigenetic mechanisms, T-cell activation, and CCR5 genetics interact to regulate T-cell expression of CCR5, the major HIV-1 coreceptor. *Proceedings of the National Academy of Sciences of the United States of America* **112**, 14658–14663, doi:10.1073/pnas.1514876112 (2015).

31. Ketas, T. J. et al. Cell surface expression of CCR5 and other host factors influence the inhibition of HIV-1 infection of human lymphocytes by CCR5 ligands. *Virology* **364**, 281–290, doi:10.1016/j.virol.2007.02.022 (2007).

32. Gornalusse, G. G. et al. Polymorphism in the CCR5 gene promoter and HIV-1 infection in North Indians. *Human immunology* **68**, 1–34, doi:10.1016/j.humimm.2007.01.016 (2007).

Acknowledgements

The authors would like to thank the patients who agreed to participate in the study and provided valuable samples. The authors also thank Peter Rotwein for critical review of the manuscript. This work was supported by National Institutes of Health R15 Grant AI116240-01A1 (to H.G.) and the Texas Tech University’s intramural seed grant program.

Author Contributions

A.J. and H.G. conceived and performed experiments, acquired and analyzed data and wrote the manuscript. E.B.P., M.S., B.B., R.P. and C.H. performed experiments and analyzed data. O.A. and D.M. helped in patient recruitment and provided critical input. J.G. performed data analysis and wrote the manuscript.

Additional Information

Supplementary information accompanies this paper at doi:10.1038/s41598-017-00192-x

Competing Interests: The authors declare that they have no competing interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/

© The Author( s) 2017