Safety and Immunogenicity of a Recombinant Adenovirus Serotype 35-Vectored HIV-1 Vaccine in Adenovirus Serotype 5 Seronegative and Seropositive Individuals

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Abstract

Background: Recombinant adenovirus serotype 5 (rAd5)-vectored HIV-1 vaccines have not prevented HIV-1 infection or disease and pre-existing Ad5 neutralizing antibodies may limit the clinical utility of Ad5 vectors globally. Using a rAd vector serotype, such as Ad35, may circumvent these issues, but there are few data on the safety and immunogenicity of rAd35 directly compared to rAd5 following human vaccination.

Methods: HVTN 077 randomized 192 healthy, HIV-uninfected participants into one of four HIV-1 vaccine/placebo groups: rAd35/rAd5, DNA/rAd5, and DNA/rAd35 in Ad5-seronegative persons; and DNA/rAd35 in Ad5-seropositive persons. All vaccines encoded the HIV-1 Env A antigen. Antibody and T-cell responses were measured 4 weeks post boost immunization.

Results: All vaccines were generally well tolerated and similarly immunogenic. As compared to rAd5, rAd35 was equally potent in boosting HIV-1-specific humoral and cellular immunity and responses were not significantly attenuated in those with baseline Ad5 seropositivity. Like DNA, rAd35 efficiently primed rAd5 boosting. All vaccine regimens tested elicited cross-clade antibody responses, including Env V1/V2-specific IgG responses.

Conclusions: Vaccine antigen delivery by rAd35 is well-tolerated and immunogenic as a prime to rAd5 immunization and as a boost to prior DNA immunization with the homologous insert. Further development of rAd35-vectored prime-boost vaccine regimens is warranted.

Keywords: HIV Vaccine; Adenovirus 35; Adenovirus 5; DNA vaccine; Randomized clinical trial

Introduction

The development of a safe and effective preventive HIV vaccine remains an urgent public health priority in the setting of an estimated 2.1 million new infections globally [1]. Since the first preventive HIV vaccine candidate entered clinical testing in 1987, four distinct vaccine concepts including subunit protein, DNA, and viral vector vaccines have been evaluated in six completed efficacy trials [2-7]. Thus far, only one vaccine regimen, a canarypox vector encoding three HIV-1 genes with a gp120 subunit boost, demonstrated partial efficacy in a phase 2b study [6]. Subsequent findings indicated that non-neutralizing IgG antibody responses to HIV-1 Env V1/V2 were significantly correlated with decreased risk of infection in vaccines [8].

HIV vaccine candidates using recombinant adenovirus (rAd) vectors have been some of the most immunogenic [9]. The Step study (HIV Vaccine Trials Network [HVTN] 502) was the first efficacy trial to test an rAd5-vectored vaccine expressing HIV-1 clade B Gag, Pol and Nef; however the product failed to protect against infection or disease progression. Furthermore, this study suggested that pre-existing Ad5 neutralizing antibodies may have played a role in increased HIV susceptibility among vaccinees [4,10]. A multiclade/multigene DNA prime, rAd5 boost regimen encoding HIV-1 Gag, Pol, and Nef from
subjects, Materials and Methods

Vaccines

All vaccines tested in HVTN 077 were produced by the NIH VRC and GenVec, Inc. (Gaithersburg, Maryland). One prototype vaccine consisted of a single E1-deleted, replication-deficient group B rAd35 vector, constructed to express a truncated modified HIV-1 EnvA antigen. Both vaccines were formulated at a dose of 1 x 10^9 particle units and administered by needle and syringe intramuscularly.

The DNA-EnvA vaccine encodes for the clade A env gene and is one of the 6 plasmids included in HVTN 505 regimen [7]. The DNA vaccination was administered intramuscularly via the needle free injection device Biojector® 2000 (Tualitin, Oregon) at a dose of 4mg. The placebos for the adenovectors and DNA vaccines were final formulation buffer and phosphate-buffered saline (PBS), respectively.

Study design and procedures

HVTN 077 was a randomized, double-blind, placebo-controlled phase 1b trial conducted at 11 clinical sites in the United States. The protocol was approved by the institutional review boards of all participating centers (Clinical Trials.gov registration NCT00801697). Between February of 2009 and January 2010, 192 adults aged 18-50 who reported low risk for infection and determined to be HIV-1-seronegative and healthy based on medical history, physical exam, and laboratory tests were enrolled after providing written informed consent. Eligible individuals who consented and enrolled were randomized to one of four treatment (T) groups (Table 1). Individuals randomized to treatment groups 2 (DNA/rAd5) or 3 (DNA/rAd35) were blinded to their assignment. For all groups, participants were blinded to assignment to vaccine or placebo. All participants were Ad35 neutralizing antibody (nAb) negative at baseline; for groups 1-3, participants were also Ad5 nAb negative. In group 4, participants were Ad5 nAb positive determined by nAb titers ≥ 18.

Safety evaluations included physical examinations and standard clinical chemistry and hematological tests. Local injection site (pain, tenderness, redness, erythema, and induration) and systemic (malaise, headache, fever, chills, myalgias, arthralgias, nausea, vomiting, and fatigue) reactogenicity symptoms were assessed for three days following each vaccination or until resolution. Adverse events were graded based on the HVTN Table for Grading Severity of Adverse Events (http://rsc.tech-res.com/Document/safetyandpharmacovigilance/Table_For_Grading_Severity_of_Adult_Pediatric_Adverse_Events.pdf). Several licensed diagnostic HIV ELISA assays (Abbott HIVAB HIV 1/2[rDNA], Abbott Architect HIV Ag/Ab Combo, BioRad Genetic System HIV 1/2 Plus O EIA, BioRad Genetic System HIV 1/2 rLAV, and BioRad Multispot HIV-1/HIV-2 Rapid Test) were performed on sera at all participants at the end of study (Day 364) to assess vaccine-induced seroreactivity.

Blood samples for assessment for primary immunogenicity were collected at days 28 (4 weeks after the single rAd35 priming injection in Group 1), 84 (4 weeks after the DNA priming series in Groups 2-4) and 196 (4 weeks after the boost vaccination in all groups).

Immune response assays

Humoral responses

Neutralizing Antibodies to Ad5 and Ad35: Baseline Ad5 neutralizing antibody titers were measured as previously described with titers ≥ 18 noted as positive [24]. Ad35 neutralizing antibody titers were measured by luciferase transgene detection [25], and titers ≥ 12 noted as positive.

HIV-Specific Binding Antibody Assays: Validated binding antibody multiplex assays [26] for measurement of vaccine elicited
HIV-1 Envelope-specific IgG to Group M Consensus (Con S gp140 CF1), Clade A (00MSA 4076 gp140), Clade B (B.con.env03 140 CF), and Clade C (C.con.env03 140 CF) were performed according to a pre-specified assay study plan following GCLP guidelines. Additional studies were performed for Env V1V2 reactive antibodies [8] utilizing scaffolds gp70 V1V2 VRC EnvA [27] and gp70 V1V2 (Case A2) [28]. HIV-1-specific IgG was detected from 1:50 serum dilution with biotin-conjugated mouse anti-human IgG (Southern Biotech, Birmingham, AL) (4 µg/ml), followed by washing and incubation with streptavidin-PE (BD Pharmingen). Mean fluorescent intensity (MFI) readouts were acquired on a Bio-Plex instrument (BioRad). Positive controls (purified HIV-1 positive immunoglobulin [HIVIG] and CH58 mAb [27] for the V1V2 assays) and negative controls (blank beads, HIV-1 negative sample, and baseline samples) were included to ensure specificity and for maintaining consistency and reproducibility between assays.

Positivity of antibody binding at Day 196 was defined by meeting all three conditions: (1) the MFI minus blank values are ≥ 2 antigen specific cutoff (based on the average + 3 standard deviations of 80 seronegative plasma samples), (2) the MFI minus blank values are greater than 3 times the baseline (Day 0) MFI minus blank values, and (3) the MFI values are greater than three times the baseline MFI values. For positive responses, binding magnitude was quantified by the net MFI concentration (subtracting the blank value) estimated using a 10-point standard curve (4P1 fit).

**T Cell Response:** Peripheral blood mononuclear cells (PBMC) were isolated and cryopreserved from whole blood within 8 hours of phlebotomy using standard procedures [29]. A 10-color intracellular cytokine staining (ICS) assay was performed on cryopreserved PBMC as previously described [30-32]. For the detection of HIV-specific T cells, thawed PBMC were rested overnight and then stimulated for 6 hours with overlapping HIV-1 15-mer peptide pools matched to the vaccine insert (VRC EnvA). Positivity was established at p<10^-3 using a Fisher’s exact test comparing stimulated and unstimulated samples.

**Statistical analysis**

All data from enrolled participants who received at least one vaccination were analyzed. Five study groups were evaluated for immunogenicity: the four vaccinated groups individually plus the placebo group. HIV-1 specific IgG binding antibody and T-cell responses were evaluated at baseline (Day 0, IgG only) and at the primary immunogenicity timepoint, Day 196 (one month after the final injection). Rates of HIV-1 Env-specific antibodies and positive CD4+ and CD8+ T-cell responses (for cells expressing IFN-γ and/or IL-2 as measured by ICS) were estimated for each study group and timepoint. Lachenbruch’s test was used for comparing primary immunogenicity endpoints between study groups [33]. Response rates were compared between groups using Fisher’s exact tests. Magnitudes of responses among positive responders were compared between study groups using Wilcoxon rank sum tests. All statistical tests were 2-sided. Primary and secondary analyses comparing immunogenicity endpoints between vaccinated groups were considered statistically significant if p ≤ 0.033, chosen to control the overall type I error rate at 0.10 and correcting for three pairwise comparisons. All other analyses used p ≤ 0.05 to judge statistical significance. All descriptive and inferential statistical analyses were performed using SAS and/or R statistical software.

**Results**

**Participant accrual, demographic data, and vaccine safety**

Of the 736 individuals who underwent screening procedures, 8% were Hispanic, and 16% Non-Hispanic Black. All participants received their initial vaccination and 98% received the second vaccination; of those assigned to 4 injections, 94% received the third, and 90% all four vaccinations. No significant differences were observed in vaccine completion rates between treatment groups. The primary reasons for study discontinuation included loss to follow-up or participant relocation (n= 9), incarceration (n =1), refusal (n =2) or other reasons (n=2); none were due to adverse experiences or reactogenicity related to vaccination.

Overall, each of the vaccine components was well tolerated. Pain or tenderness at the injection site was reported most commonly (88% of study participants). Differences between groups were detected; as seen in Figure 1, those receiving the rAd35/rAd5 heterologous adenovector regimens were less likely to report local reactions compared to the DNA prime, adenovector boost regimens (p<0.001). Maximum systemic symptoms were less commonly reported by those with pre-existing neutralizing antibodies to Ad5 (T4, Figure 1). There were 22 adverse events that were at least probably or definitely attributed to the vaccine, and most were local injection site reactions that were characterized as mild or moderate in severity. One case of transient mild leukopenia deemed probably related to vaccination was observed after receiving DNA in a T4 participant. Four expedited adverse events were reported in the trial including rectal bleeding, post traumatic lower extremity and right intraorbital ethmoid fractures, bipolar disorder, and gallstone pancreatitis; none were attributed to product. No significant differences in laboratory parameters were noted among groups. Overall, 60% had evidence of vaccine-induced HIV seroreactivity at the end of study using several commercially available HIV test kits. Rates were highest among rAd35/rAd5 vaccinees (81.8%) and lowest among participants who received DNA/rAd5 (45.7%).

**HIV-1-specific antibody responses**

Each of the vaccine regimens induced high frequency and magnitude cross-clade binding antibody responses (Figure 2). The antibody response rates were 100% for all treatment groups recognizing the consensus M gp140 (data not shown) and 97-100% recognizing the clade A Env antigen. For the clade B antigen, responses were also detected in greater than 92% of individuals. For the clade C antigen, response frequencies were highest for rAd35/rAd5 (96%) compared to the DNA prime/adenovector boost groups (76-78% for groups 2-4), however at lower magnitude in comparison to the DNA/rAd5 group (p=0.02).

Based on evidence that IgG binding antibodies to V1/V2 were correlated with reduced risk of HIV infection in the RV144 vaccine...

Figure 1: Safety assessment, showing maximum local reactogenicity A) and systemic reactogenicity, B) by treatment group. Subjects in control (C) groups 1-4 received group phosphate buffered saline. Subjects in treatment groups 1-3 (T1-T3) were Ad5 seronegative at baseline and received: T1- recombinant Ad35 (rAd35) prime and rAd5 boost; T2- three DNA priming injections boosted by rAd5; and T3- three DNA priming injections boosted by rAd35. Subjects in T4 were Ad5 seropositive at baseline and received three DNA priming injections boosted by rAd35. P values for comparisons of local and systemic reactogenicity were determined using the Kruskal-Wallis test.
Figure 2: Binding antibody net responses to Clades A (OOMSA 4076 gp140), B (B.con.env03 140CF), and C (C.con.env03 140CF) isolates 4 weeks after the boost vaccination as measured by median fluorescence intensity (MFI)-Blank where 'Blank' is a sample specific background measure. Responders are shown in red circles and non-responders in blue triangles. Box plots display the distribution of positive responses for the vaccinees for each antigen. P-values are derived from Lachenbruch’s test comparing rAd5 and rAd35 boosted groups in Ad5 seronegative individuals and the Ad35-boosted group in Ad5 seropositive individuals.
efficacy trial [8], we explored whether the EnvA constructs tested in HVTN 077 elicited these responses. Among positive responders, all treatment groups elicited gp70V1/V2 responses using the V1/V2 scaffold tested in the RV144 study (Case A2, Figure 3A) and there were no significant differences by group in the magnitude or frequency of the responses. In addition, we looked at the matching clade A V1/V2 sequence in the HVTN 077 vaccine regimen (clade A gp70V1/V2). Although the frequency of response did not differ substantially by group, as seen in Figure 3B, the response magnitude for binding antibodies to the vaccine-matched clade A gp70V1/V2 was significantly higher among the DNA/rAd5 group compared to the rAd35/rAd5 group, (p=0.005).

HIV-1--specific T-cell responses

As seen in Figure 4, HIV-1-specific T cells producing IFN-γ and/or IL-2 in response to vaccine insert-matched peptides were detected readily in each of the treatment groups. With regard to the CD4+ T-cell responses, the highest post-boost response rates were seen among Ad5 seronegative individuals receiving DNA/rAd35 (25/36, 69.4%) and lowest among those receiving rAd35/rAd5 (9/24, 37.5%); responses did not differ significantly across groups. Vaccine-induced CD8+ T-cell responses were elicited most frequently among Ad5 seronegative individuals who received the DNA/rAd5 regimen (32/42, 76.2%) and least among Ad5 seropositive who received the DNA/rAd35 regimen (10/25, 40.0%); responses did not differ significantly across groups. Overall, we found that among Ad5 seronegative subjects, 15%, 39%, and 32% of rAd35/rAd5, DNA/rAd5, and DNA/rAd5 recipients, respectively, and 14% of Ad5 seropositive participants receiving the DNA/rAd5 regimen developed both CD4+ and CD8+ T-cell responses. The HIV-specific CD4+ or CD8+ T-cell response magnitudes for positive responders after boosting were similar across groups.

In addition, we assessed expression of TNF-α and Granzyme B (GzB) in response to stimulation with insert-matched peptides. As shown in Figure 5, priming with DNA followed by either rAd35 or rAd5 led to the induction of significantly more polyfunctional CD4+ T cells than vaccination with rAd35/rAd5 in Ad5 seronegative subjects (p=0.0005 for three and p=0.007 for four functions, respectively, comparing the combined DNA/rAd5 and DNA/rAd35 groups with rAd35/rAd5 for Ad5 seronegative subjects). Interestingly, the patterns of combined expression of these functional markers did not differ significantly across groups.
Figure 4. HIV-specific CD4+ and CD8+ T-cell responses: The percentage of CD4+ (panel A) and CD8+ (panel B) T cells producing γ-interferon (IFN-γ) and/or interleukin-2 (IL-2) in response to EnvA matched peptide pools 4 weeks after the priming immunization(s) and 4 weeks after the boost as measured by intracellular cytokine staining. Responders are shown in red circles and non-responders in blue triangles. Boxplots show the distribution of the magnitude of response in positive responders only. The box indicates the median and interquartile range (IQR); whiskers extend to the furthest point within 1.5 times the IQR from the upper or lower quartile. Numbers at the top of each panel show the number of responders / number with an assay result and the percent with positive response. P-values are derived from Lachenbruch’s test comparing rAd5 and rAd35 boosted groups in Ad5 seronegative individuals and the Ad35-boosted group in Ad5 seropositive individuals. Data from samples with high background cytokine secretion was filtered, leading to differences in the number of samples with available data for CD4+ and CD8+ T cells.
higher than the 17.5% rate of response elicited by the HVTN 505 regimen tested in the RV144 trial (64%, 95% CI 58% to 70%) [8] and to the rate observed for the partially efficacious ALVAC/gp120 vaccine V1/V2 IgG responses (52% to 66% across the four groups) was similar and Nef was unable to prevent HIV acquisition or reduce viral load in the setting of pre-existing Ad5 nAbs, there were no significant differences between rAd35 and rAd5 vaccine efficacy trial [8].

Cross-clade binding antibodies as well as V1/V2-specific IgG antibodies, to substantially dampen immunogenicity [23]. Finally, we found that responses from these serologically distinct adenoviruses are unable to induce EnvA-specific antibody and T-cell responses. In addition, we found that as a boost in DNA-primed subjects, there were no significant differences between rAd35 and rAd5 in the ability to induce EnvA-specific antibody and T-cell responses. We also observed that in the setting of pre-existing Ad5 nAbs, there was no significant reduction of HIV-1 specific antibody or cellular responses to an Ad35-based regimen when given once as a boost. This confirms findings from preclinical models that cross-reactive immune responses from these serologically distinct adenoviruses are unable to substantially dampen immunogenicity [23]. Finally, we found that each of the vaccine regimens tested in this trial was capable of inducing cross-clade binding antibodies as well as V1/V2-specific IgG antibodies, which were correlated with reduced risk of HIV infection in the RV144 vaccine efficacy trial [8].

The HVTN 505 Phase IIb efficacy trial showed that a DNA/rAd5 prime-boost regimen encoding EnvA, EnvB, EnvC, clade B Gag, Pol, and Nef was unable to prevent HIV acquisition or reduce viral load [7]. Of note, in HVTN 077, we found the proportion of vaccinees with V1/V2 IgG responses (52% to 66% across the four groups) was similar to the rate observed for the partially efficacious ALVAC/gp120 vaccine regimen tested in the RV144 trial (64%, 95% CI 58% to 70%) [8] and higher than the 17.5% rate of response elicited by the HVTN 505 regimen [7]. In addition, the V1/V2 IgG responses were similar between groups 2 and 3 (DNA/rAd5 and DNA/Ad35 in Ad5 seronegative subjects), and group 4 (DNA/rAd35 in Ad5 seropositive subjects). This is in contrast to findings from the phase 2 study of the VRC multiclade DNA/rAd5 regimen [12], where V1/V2 IgG responses were significantly lower in Ad5 seropositive vaccines compared to Ad5 seronegative individuals (G. Tomaras, personal communication). These findings suggest that presentation of the EnvA antigen alone may produce a more favorable antibody response to the V1/V2 region than presentation of multiple Env proteins with additional competing antigens. It is important to note that we do not know if the V1/V2 IgG correlate of risk translates into a correlate of protection, and, if so, whether it is a mechanistic or non-mechanistic correlate [35] or whether IgG V1/V2 responses will be a correlate of HIV-1 risk or protection for vaccines in other populations that differ from the community-based sample evaluated in Thailand [6]. However, studies in non-human primate models suggest that envelope binding antibodies, V2-specific antibodies, and the avidity to which anti-Env antibodies bind to native trimer [18,36-38] can correlate with protection. Therefore, future HIV-1 vaccines designed to elicit Env binding responses and V2-specific IgG antibodies, should consider these observations from preclinical studies and recent clinical trials.

Discussion

In this clinical study of a prototype rAd35 vectorized HIV-1 vaccine, we found that it was well tolerated with a similar safety profile to that of rAd5-based experimental vaccines [12,34]. We sought to explore how the immunogenicity of the heterologous rAd35/rAd5 regimen compared to a DNA/rAd5 regimen, and found that rAd35/rAd5 elicited similar frequency and magnitude of HIV-1–specific antibody responses, and slightly (although not statistically significant) lower T cell responses. In addition, we found that as a boost in DNA-primed subjects, there were no significant differences between rAd35 and rAd5 in the ability to induce EnvA-specific antibody and T-cell responses. We also observed that in the setting of pre-existing Ad5 nAbs, there was no significant reduction of HIV-1 specific antibody or cellular responses to an Ad35-based regimen when given once as a boost. This confirms findings from preclinical models that cross-reactive immune responses from these serologically distinct adenoviruses are unable to substantially dampen immunogenicity [23]. Finally, we found that each of the vaccine regimens tested in this trial was capable of inducing cross-clade binding antibodies as well as V1/V2-specific IgG antibodies, which were correlated with reduced risk of HIV infection in the RV144 vaccine efficacy trial [8].

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Figure 5. Vaccine-induced HIV-specific CD4+ and CD8+ T cells expressing multiple functional markers: The proportion of HIV-specific CD4+ (left) or CD8+ (right) T cells with one, two, three or four functions measured by expression of Granzyme B, IFN-g, IL-2, or TNFα is shown for positive responders from Figure 4. Boxplots show the distribution of responses; the box indicates the median and IQR, whiskers extend to the furthest point within 1.5 times the IQR from the upper or lower quartile.

Whether multi-dose DNA priming for an adenovector boost confers an immunologic advantage over heterologous adenovector regimens is relevant given the desire for less complex vaccine regimens with fewer required immunizations. Compared to rAd35/rAd5, we found that the DNA/rAd5 regimen elicited higher magnitude binding antibody responses to some of the antigens tested (e.g., the EnvA clade-matched V1/V2 expressed by the vaccine and EnvC) but not to others (e.g., the Clade A Env). And while there was no overall difference in the HIV-specific CD4+ or CD8+ T-cell responses elicited by these regimens, DNA priming may generate more polyfunctional responses than the heterologous adenovector prime-boost regimen, a desirable feature of vaccines designed to elicit cellular immunity [39,40]. Several studies in the field will provide further insights into the relative immunogenicity of prime-boost regimens combining different adenovectors such as

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Ad35 and Ad26 encoding an EnvA antigen (IAV1 B003-IPCAVD004- HVTN091, NCT01215149) and DNA administered by electroporation with a multi-antigen rAd35 construct (IAVIB004; NCT01496899). Furthermore, to optimize immunogenicity, trials should carefully consider the administration interval between adenovector prime and boost. Our study delivered rAd35 and rAd5 6 months apart, eliciting higher magnitude HIV-specific T-cell responses compared to rAd35/ rAd5 given only 3 months apart in VRC’012 (NCT00479999) [22].

Conclusion

In this phase 1b study, we have demonstrated that rAd35 is well tolerated and immunogenic as a boost, is as potent as rAd5 in DNA primed individuals. In addition, the humoral and cellular responses elicited by rAd35 boosting are better preserved in the setting of pre-existing Ad5 seropositivity than responses to rAd5 boosting, suggesting that rAd35 is a reasonable choice for an alternative adenoviral vaccine vector to diminish the impact of antivector immunity. Therefore, as additional safety data emerge from studies exploring alternative adenoviral vectors in different global contexts, rAd35 should be considered for use as a vaccine delivery vector, particularly as effective antigen designs become available. This is particularly relevant when CD8+ T cell-mediated immunity is desirable in addition to antibody-based immunity, in subjects already primed with the antigens expressed by the rAd35 vector, and in settings with a high prevalence of pre-existing immunity to Ad5, such as in sub-Saharan Africa.

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Author contributions

Designed the study: JF, PAB, PG, NF, SD, TW, BG*, MJM. Oversaw study conduct and managed participants at study sites: JF, PAB, LB, BK, MS, KM, PG, NR, SK, MK, MJM. Medical monitoring and study oversight: JF, PAB, ES*, CM, BG*. Oversaw performing immunogenicity assays and result interpretation: JF, NF, SD, SR, GT, JM, BG*. Analyzed the data: JF, NK, PG, NF, GT, BG*, MJM. Wrote the manuscript: JF, PG, NF, GT, BG, MJM.

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