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# Arenavirus Z protein controls viral RNA synthesis by locking a polymerase–promoter complex

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**Arenaviruses form a noncytolytic infection in their rodent hosts, yet can elicit severe hemorrhagic disease in humans. How arenaviruses regulate gene expression remains unclear, and further understanding may provide insight into the dichotomy of these disparate infection processes. Here we reconstitute arenavirus RNA synthesis initiation and gene expression regulation in vitro using purified components and demonstrate a direct role of the viral Z protein in controlling RNA synthesis. Our data reveal that Z forms a species-specific complex with the viral polymerase (L) and inhibits RNA synthesis initiation by impairing L catalytic activity. This Z–L complex locks the viral polymerase in a promoter-bound, catalytically inactive state and may additionally ensure polymerase packaging during virion maturation. Z modulates host factors involved in cellular translation, proliferation, and antiviral signaling. Our data defines an additional role in governing viral RNA synthesis, revealing Z as the center of a network of host and viral connections that regulates viral gene expression.**

transcription factor | Machupo virus | matrix protein | negative-strand RNA virus | gene regulation

**T**ranscription initiation is a fundamental focal point of gene expression regulation in all orders of life. Modulation of RNA synthesis affords an important response to stress, alterations in growth conditions, and environmental cues. Although initiation of RNA synthesis is often indirectly controlled through gene promoters and accessory signaling elements, regulation also occurs through factors directly interacting with and altering the catalytic potential of the RNA synthesis machinery itself (1). Studies with phage T7 and the T7 lysozyme repressor system have provided an important paradigm for control of viral RNA transcription (2, 3), but these processes remain a mystery for many eukaryotic viruses and medically relevant pathogens. Gene expression regulation is particularly critical for negative-sense RNA viruses of the family *Arenaviridae*, which must respond to input from the host environment to maintain a noncytolytic infection in their rodent reservoir (4). In stark contrast to this persistent infection, arenavirus infection in humans can lead to severe hemorrhagic fever disease and many arenaviruses represent important emerging pathogens (5). Intriguingly, recent evidence indicates that the balance between persistent and acute viral infection can be altered by a single mutation within the viral polymerase (6). Mechanistic insight into how arenaviruses tune gene expression and control viral RNA synthesis is required to further our understanding of the response of arenavirus infection to various host and cellular environments.

Arenaviruses are the simplest of all hemorrhagic fever viruses, with only four viral proteins. The viral genome consists of two segments of single-stranded negative-sense RNA, where each segment encodes two proteins in an ambisense orientation. The small segment encodes for the viral glycoprotein, essential for entry into the host cell, and the nucleocapsid protein (NP), which coats the genomic RNA and protects it from degradation and detection by host antiviral sensors (7, 8). The large segment encodes for the large polymerase protein (L) as well as the smallest viral protein, Z. All of the enzymatic machinery required for viral RNA synthesis and mRNA transcription is contained within L, a 250-kDa multifunctional protein that is organized as a central ring-like RNA-dependent RNA-polymerase (RdRP) domain with three accessory appendages (9). As ectopic overexpression of

Z induces potent inhibition of replicon-based reporter protein expression (10, 11), Z has been hypothesized as an important regulator of L and arenaviral gene expression. However, Z is known to interact with several host factor binding partners, and it remains unclear whether Z directly regulates the viral RNA synthesis machinery or acts through alteration of an essential host process.

The Z gene is highly conserved across all arenavirus species and encodes a protein organized as an unstructured amino and carboxyl terminus bracketing a really interesting new gene (RING) domain that coordinates a pair of structural zinc ions (12). Z has been implicated in many facets of the arenavirus replication cycle and is the primary driving force of virion maturation (13). In addition to recruiting the cellular endosomal sorting complex required for transport (ESCRT) machinery to mediate virion budding (13, 14), Z has been demonstrated to interact with the translation initiation factor eIF4E (15) as well as retinoic acid inducible gene I (RIG-I), promyelocytic leukemia protein (PML), and proline-rich homeodomain protein (PRH) (16–18). Thus, Z is positioned at the center of a network of host and viral connections and may serve as a crucial viral sensor of the host cellular environment.

To unravel the complex network of Z functions and host interactions, and to determine whether Z plays a direct role in regulating arenavirus gene expression, we reconstituted arenavirus RNA synthesis and regulation by Z in vitro using purified components. Here we use purified L from Machupo virus (MACV), the causative agent of Bolivian hemorrhagic fever, to establish an in vitro model of arenavirus RNA synthesis that maintains authentic promoter recruitment and RNA synthesis initiation while relying solely on recombinant L and short naked RNA templates. With this minimal system, we demonstrate that Z functions as a direct regulator of arenaviral RNA synthesis and forms a direct complex with L governed by species-specific protein interactions. Z controls gene expression by locking a promoter-bound polymerase complex in a catalytically inactive state. These findings offer mechanistic insight into Z mediation of L catalytic activity and uncover how a single viral protein coordinates both modulation of host processes and regulation of viral gene expression.

## Results

**Z Is a Direct Inhibitor of L Catalytic Activity.** To probe how arenaviruses control viral gene expression, we reconstituted Z regulation of arenavirus RNA synthesis in cells and in vitro with purified components. We designed a MACV S-segment replicon where the viral glycoprotein gene has been replaced with *gaussia* luciferase to quantitatively assess activity of the viral polymerase complex within transfected cells. Luciferase signal was only observed in the presence of MACV L, and a mutation to the predicted RdRP catalytic active site (SDD1328AAA) abolished reporter protein expression (Fig. 1A). Ectopic overexpression of MACV Z resulted in the inhibition of the MACV replicon-

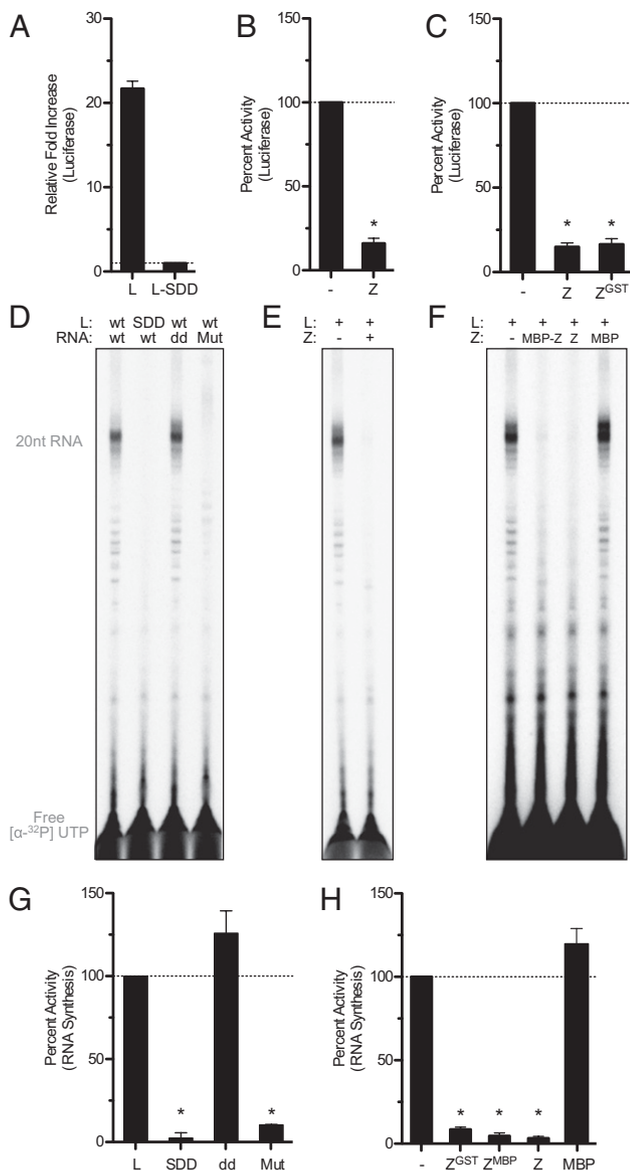
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**Fig. 1.** Z is a direct inhibitor of L catalytic activity. (A) Cellular-based replicon assay for MACV RNA synthesis where functional viral polymerase results in the production of gaussia luciferase. Luciferase activity was measured from the supernatant of cells expressing pMCM-Gluc replicon in the presence of L (L) or a catalytically inactivated mutant (L-SDD). Error bars represent the SD from the mean of at least three independent experiments. (B and C) Replicon assay as in A with cells transfected with the viral replicon and L alone (–) or with MACV Z (Z) or GST-tagged MACV Z (Z<sup>GST</sup>) plasmids. Luciferase values were normalized to cells expressing only the replicon plasmid and wild-type L. Error bars represent the SD as in A and conditions that are statistically significant from the wild-type L control are designated with the symbol \* ( $P < 0.001$  or for Z<sup>GST</sup>  $P = 0.0016$ ). (D) MACV in vitro RNA synthesis. Purified L (L: WT) or a catalytically inactivated mutant (L: SDD) was incubated with a wild-type template (template RNA: WT), a 3' terminal dideoxy template (template RNA: dd) or an RNA template with a mutated promoter binding site (template RNA: Mut) in the presence of GpC primer as described in the text. RNA products were labeled with [ $\alpha$ -<sup>32</sup>P]-UTP and separated by denaturing gel electrophoresis. (E and F) RNA products generated from RNA synthesis reactions supplemented with purified GST-tagged MACV Z (+), MBP-tagged MACV Z (MBP-Z), untagged MACV Z (Z), or free MBP (MBP) and analyzed as in D. (G and H) Total RNA synthesis was quantified with a PhosphorImager and graphed as values normalized to reactions containing wild-type L and RNA template. Error bars represent the SD from the mean of at least three independent experiments (\* $P < 0.001$ ). See also Figs. S1 and S2.

based reporter protein expression (Fig. 1B), and we observed equivalent levels of inhibition with a GST-tagged version of Z (Fig. 1C). Previous results have revealed a similar phenotype using overexpressed Z in other arenaviral replicon systems (10, 11, 19), yet it has been impossible to determine whether Z directly impacts the viral RNA synthesis machinery or functions through alteration of an essential host cellular process. We therefore established an in vitro system to examine a potential direct effect of Z on viral RNA synthesis.

The large size (~250 kDa) and flexible multidomain organization of arenaviral L proteins have limited biochemical characterization of viral RNA synthesis. Previously, we demonstrated purification of MACV L protein and formation of an L–RNA complex with the conserved 19-nt RNA promoter element (9). Using this recombinant L–RNA complex, we established an in vitro transcription system to study the initial events of viral RNA synthesis. The current model for initiation of arenavirus RNA synthesis is a “prime-and-realign” model, where the short dinucleotide primer (GpC) is first synthesized and realigned with the template RNA resulting in the insertion of an extra 5' guanosine (20). Consistent with this model, when recombinant L–RNA complexes were incubated with NTPs and [ $\alpha$ -<sup>32</sup>P]-UTP, we observed the formation of a predominant ~20-nt RNA product from the 19-nt RNA template as well as minor populations of ~21-nt RNA and shorter products likely resulting from premature termination (Fig. 1D). GpC dinucleotide primer was included in RNA synthesis reactions as it has previously been demonstrated to stimulate viral RNA synthesis (9, 20); but similar products are observed with and without exogenous GpC primer addition (discussed below).

Analogous to the cellular-based replicon assay, a mutation (SDD1328AAA) within the RdRP active site of purified L abolished all RNA synthesis in vitro (Fig. 1D). Purified viral polymerases have been shown to possess terminal transferase activity (21, 22). To demonstrate that the RNA synthesis observed in vitro was not dependent upon a terminal transferase, we used a template in which the 3' terminal guanosine is replaced by a dideoxy cytosine nucleotide (template RNA: dd). We previously demonstrated that the G1C RNA mutation does not impact L–RNA complex formation (9). In the presence of GpC dinucleotide primer, we observed similar RNA synthesis with this template, verifying that in vitro RNA synthesis is the result of de novo initiation and not an L-associated terminal transferase activity (Fig. 1D). In the absence of exogenous GpC primer, RNA synthesis with the G1C template resulted in slightly altered ratios of the predominant RNA products, which may indicate an effect of this mutation in normal prime-and-realign initiation events (discussed below). To further demonstrate that the products reflect de novo initiation of RNA synthesis, we examined whether they contained a reactive 5' triphosphate. We took advantage of the vaccinia virus guanylyltransferase, which can add a capping guanylate to free 5' triphosphate but not 5' monophosphate. The products of in vitro transcription were capped by the vaccinia virus guanylyltransferase (discussed below), demonstrating that they retained a 5' tri- or diphosphate RNA terminus, as observed previously with purified arenavirus genomic RNA (23). We next examined the products of RNA synthesis using an RNA template containing mutations to the high-affinity 3' RNA promoter binding site. We previously demonstrated that the absolutely conserved 3' C<sub>2</sub>G<sub>3</sub>U<sub>4</sub>G<sub>5</sub> motif is required for L–RNA complex formation (9) and, as further verification of the authenticity of this reconstituted RNA synthesis system, we observed that mutating this 3' binding site to C<sub>2</sub>C<sub>3</sub>G<sub>4</sub>G<sub>5</sub> (template RNA: Mut) eliminated in vitro RNA synthesis activity (Fig. 1D).

To determine whether Z directly inhibits the viral RNA synthesis machinery, we purified recombinant GST-tagged MACV Z (Fig. S1). Supplementing RNA synthesis reactions with purified GST-Z resulted in potent inhibition of L catalytic activity (Fig. 1E). To verify this inhibition was not related to GST or the presence of a large solubility tag, we purified Z tagged with maltose binding protein (MBP) and removed the tag using tobacco etch virus protease (Fig. S1). Purified untagged Z, as well





end. We observed that the GpC primer is incorporated into elongated RNA synthesis products (Fig. 4C), confirming that GpC can be used to prime arenavirus RNA synthesis. As with unprimed reactions, Z directly inhibited all detectable [ $\alpha$ - $^{32}$ P]-GpC primed polymerase activity (Fig. 4C and E), demonstrating that L is incapable of initiating de novo or primed RNA synthesis in the presence of Z. As an additional method to confirm that smaller products are not generated in the presence of Z, we labeled all de novo initiated transcripts after RNA synthesis through subsequent capping of the 5' end with vaccinia virus guanylttransferase and [ $\alpha$ - $^{32}$ P]-GTP (Fig. 4D and E). Together these results demonstrate the formation of a Z-L-RNA complex and define the mechanism of Z regulation as inhibition of the initial catalytic steps of RNA synthesis following promoter engagement.

## Discussion

Here we use purified components to reconstitute individual steps of arenavirus RNA synthesis initiation and demonstrate that the viral Z protein inhibits RNA synthesis by locking the polymerase in a catalytically inactive promoter-bound state. Z-L complex formation provides a direct link between viral gene expression, host cellular modulation, and virion packaging, revealing the multifarious activities of Z as an important network that regulates the dynamics of arenaviral infection.

**Mechanism of Z Inhibition.** Arenavirus genomic replication requires L and an appropriate NP-encapsidated RNA template (27, 28). Although NP is not essential for the catalytic activity of L itself (Fig. 1), it is likely that NP is critical for activities not recapitulated by the initial steps of RNA synthesis initiation in vitro, including integrity of the full-length genome and possibly reduction of RNA secondary structure (29, 30). Here we demonstrate that Z is a direct modulator of the viral RNA synthesis machinery. In the cell, overexpression of Z results in inhibition of replicon-based reporter protein expression (Fig. 14), and Z similarly inhibits RNA synthesis reactions using purified components in vitro (Fig. 1E and F). Z has been identified as a binding partner to both host and viral proteins (15–18, 31–34). However, by establishing an in vitro model system of arenavirus RNA synthesis (Fig. 1D), we demonstrate that host interactions are not required for Z to inhibit L catalytic activity (Fig. 1).

Z inhibits L catalytic function, but interaction with Z does not restrict recruitment of L to the viral RNA promoter (Fig. 44). Our data reveal that Z interacts directly with L (Fig. 3 and Fig. S3) and the promoter-bound L-RNA complex (Fig. 44) and that Z alone does not interact with viral RNA (Fig. 44). In the presence of Z, L maintains interaction with the high-affinity 3' RNA promoter binding site (Fig. 44) (9), but Z-L-RNA complex formation results in an inhibited polymerase incapable of RNA synthesis (Figs. 1 and 4 and Fig. S5). This block in an early step of RNA synthesis initiation prevents both de novo and GpC-primed transcription (Figs. 1 and 4), indicating that the polymerase is frozen in a catalytically inactive conformation or occluded complex. In our in vitro RNA synthesis system, L never becomes resistant to Z inhibition (Fig. S2), but determining the effect of Z on an actively elongating complex will require further analysis of single polymerase molecules.

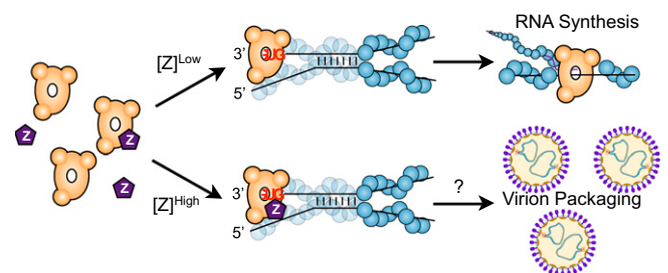
As we observe here with Z regulation of L RNA synthesis initiation, modulating gene expression through control of RNA transcription initiation is a common theme found in all orders of life. The eukaryotic transcription factor Maf1 directly interacts with RNA polymerase III to repress RNA synthesis, but unlike the Z-L-RNA complex observed in vitro (Fig. 44), Maf1 prevents promoter engagement and closed complex formation (35). Similarly, RNA synthesis initiation in phage T7 is controlled by interaction with T7 lysozyme that causes noncatalytic inhibition of T7 RNA polymerase resulting in short abortive transcript production (36). In contrast, Z prevents all detectable RNA synthesis, suggesting a direct impairment to the catalytic activity of L (Figs. 1 and 4 and Fig. S5). This situation mirrors repression of RNA synthesis initiation maintained by the bacterial transcription factor Gre-family homolog 1, which disrupts catalytic

activity by physically occluding the NTP channel and locking the RNA polymerase in an inactive ratcheted state (37). Although the exact molecular mechanism of Z inhibition will require further biochemical and structural analysis, our data are consistent with a model where the catalytic potential of L is physically compromised within the Z-L complex.

**Regulation of Arenavirus Gene Expression.** Z interacts with several host factors and has been implicated in modulating a variety of host processes including mRNA translation, cell proliferation, and antiviral signaling (15–18). Our data now reveal a direct role of Z in regulating viral gene expression, placing Z at a crossroads that links coordination of viral RNA synthesis with pathways essential for cellular viability. These results confirm a direct feedback loop where increased viral gene expression and higher levels of Z will act to suppress RNA synthesis and balance the infection process (10) (Figs. 2 and 5).

Z is highly conserved across all arenavirus species, although only Z from MACV and other New World arenaviruses has been demonstrated to interact with RIG-I (16). In contrast, we now demonstrate that MACV Z interacts with eIF4E (Fig. S4B), indicating that at least some of the binding partners of Z are shared between both New and Old World arenaviruses. However, interaction with eIF4E was not sufficient to regulate Z-induced inhibition of L in vitro (Fig. S4). JUNV Z inhibition of MACV RNA synthesis (Fig. 24), and previous reports of cross-regulation by Z of LASV and LCMV (19), support that the Z-L interface is sufficiently conserved such that closely related arenaviruses share similar binding and regulatory interactions within their polymerase machinery. However, we demonstrate both in cells and in vitro that LCMV Z is unable to interact with or inhibit MACV L (Figs. 2–44). These results show that Z inhibition of arenaviral RNA synthesis is species specific and unique to the distinct geographic and phylogenetic arenavirus clades.

**Polymerase Packaging.** As the viral genome itself is not an infectious unit, all reverse transcribing, double-stranded, and negative-sense RNA viruses must package their polymerase machinery to generate an infectious particle. Whereas many viruses solve this problem by including their polymerase protein as an integral structural component essential for virion formation, negative-sense RNA viruses do not appear to invoke this strategy. In addition to complex formation with L, Z interacts with the cellular ESCRT machinery and is the major driving force of viral budding and virion formation (13). Importantly, inhibition by Z does not disrupt the recruitment of L to the genomic promoter (Fig. 44), and the resulting Z-L-RNA complex may thus serve as a critical intermediate to ensure genomic RNA is packaged along with a polymerase poised to reinitiate viral RNA synthesis in the newly infected cell (Fig. 5). Although the individual mechanisms remain undetermined, inhibition of viral RNA gene expression by overexpression of the matrix, or matrix-like, protein has been observed for influenza A virus and in a multitude of nonsegmented negative-sense RNA viral systems (38–41). It is therefore possible



**Fig. 5.** Model of arenavirus RNA synthesis regulation by Z. Low concentrations of Z permit ongoing RNA synthesis, whereas high concentrations of Z result in an inhibited Z-L-RNA complex bound to the viral promoter. As described in the text, the Z-L-RNA complex may serve as an important intermediate ensuring L is packaged into mature virions.

that the Z–L–RNA complex and Z–L inhibition mechanism identified here may serve as an important paradigm for the link between transcription regulation and polymerase packaging of negative-sense RNA viruses.

## Methods

**In Vitro RNA Synthesis.** Arenavirus RNA synthesis was reconstituted using recombinant MACV L and a 19-nt template corresponding to the conserved 3' terminus of the viral genome. L–RNA complexes were established as described previously (9). Briefly, 0.3  $\mu$ g of purified L (final reaction concentration of 0.12  $\mu$ M) was incubated with 2  $\mu$ M of RNA in transcription buffer (50 mM Tris pH 7, 40 mM NaCl, 5 mM MnCl<sub>2</sub>, 10 mM KCl, 1 mM DTT, and 0.1 mg/mL BSA (New England Biolabs) at 25 °C for 30 min. After initial incubation, 0.5  $\mu$ L of [ $\alpha$ -<sup>32</sup>P]-UTP (~10  $\mu$ Ci), and cold NTPs (1 mM ATP/CTP/GTP and 25  $\mu$ M UTP) were added and reactions were incubated at 30 °C for 2 h in a final volume of 10  $\mu$ L. Reactions included 40  $\mu$ M GpC primer (Dharmacon) as indicated in the text. Alternatively, reactions were labeled with [ $\alpha$ -<sup>32</sup>P]-GTP and cold NTPs (1 mM ATP/CTP/UTP and 100  $\mu$ M GTP), or with 1 mM cold NTPs and 5' [<sup>32</sup>P]-GpC primer labeled with [ $\gamma$ -<sup>32</sup>P]-ATP and T4 polynucleotidyl kinase (New England Biolabs). Where indicated, RNA synthesis reactions were supplemented with recombinant Z or MBP (2  $\mu$ M final concentration) during L–RNA complex assembly. After 2 h incubation, reactions were terminated with 10  $\mu$ L of stop solution (95% deionized formamide, 20 mM EDTA) and incubation at 95–100 °C for 2.5 min. Reactions were analyzed by denaturing gel electrophoresis on a 36 cm long, 20% polyacrylamide-urea (7 M) sequencing gel (with 0.5 $\times$  TBE). RNA products were visualized using

a PhosphorImager, sized according to chemically synthesized RNA markers, and quantified with ImageQuant (Amersham). Statistical significance was calculated using an unpaired, two-tailed *t* test.

**GST Pull-Down and L–RNA Gel Shift Assays.** Recombinant L (5  $\mu$ g) and/or eIF4E (8  $\mu$ g) was incubated with GST-tagged Z or GST alone (3  $\mu$ g) in binding buffer (PBS with 250 mM KCl and 0.2% Nonidet P-40) (12) in a total volume of 50  $\mu$ L at room temperature for 30 min. Reaction volumes were raised to 500  $\mu$ L with binding buffer and 15  $\mu$ L of packed glutathione sepharose resin (GE Healthcare) and incubated at room temperature for 1 h while rotating. Glutathione resin and bound proteins were gently pelleted by centrifugation at 400  $\times$  *g* for 30 s and washed three times with wash buffer (PBS with 500 mM KCl and 1.0% Nonidet P-40). Proteins were eluted by boiling in 25  $\mu$ L of SDS loading buffer, and 10  $\mu$ L was analyzed by 12% denaturing SDS/PAGE and staining with colloidal Coomassie (Sigma). L–RNA complex formation and the electrophoretic shift assay were performed as described previously (9), and reactions were supplemented with 0.35  $\mu$ g GST-tagged MACV Z or LCMV Z as indicated.

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