

B23/nucleophosmin is involved in regulation of adenovirus chromatin structure at late infection stages, but not in virus replication and transcription

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4 **RUNNING TITLE:** Regulation of adenovirus chromatin by B23/nucleophosmin

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1 **SUMMARY**

2 B23/nucleophosmin was identified *in vitro* as a stimulatory factor for
3 replication of adenovirus (Ad) DNA complexed with viral basic core proteins. In the
4 present study, we have studied the *in vivo* function of B23 in Ad life cycle. We found
5 that both the expression of a decoy mutant derived from Ad core protein V that tightly
6 associates with B23 and siRNA-mediated depletion of B23 impede the production of
7 progeny virions. However, B23 depletion did not significantly affect the replication
8 and transcription of the virus genome. Chromatin immunoprecipitation analyses
9 revealed that B23 depletion significantly increases the association of viral DNA with
10 viral core proteins and cellular histones. These results suggest that B23 is involved in
11 the regulation of association and/or dissociation of core proteins and cellular histones
12 with the virus genome. In addition, our results suggest that proper viral chromatin
13 assembly regulated in part by B23 is crucial for the maturation of infectious virus
14 particles.

15

1 INTRODUCTION

2 Adenovirus (Ad) is an icosahedral particle with linear double-stranded DNA of
3 approximately 36,000 base pairs in length. The linear DNA is covalently linked with
4 virally coded terminal proteins and condensed with viral basic proteins Mu, VII, and
5 V, thus forming a chromatin-like structure termed Ad core/Ad chromatin (Anderson
6 *et al.*, 1989; Black & Center, 1979; Chatterjee *et al.*, 1985). Protein VII, a 19 kDa
7 basic protein, is the major component of Ad core and most tightly associated with the
8 Ad genome (Sung *et al.*, 1983). Protein V associates loosely with Ad DNA and forms
9 an outer shell around Ad core to link it with the capsid through a dimer of polypeptide
10 VI (Brown *et al.*, 1975; Chatterjee *et al.*, 1985; Fedor & Daniell, 1983). The virus
11 genome is thought to be packed around the hexamer of core protein VII, and each unit
12 of viral DNA-VII hexamer complex is bridged by core protein V (Dery *et al.*, 1985;
13 Sung *et al.*, 1977).

14 Infecting virus particles are disassembled in the cytoplasm stepwise after
15 penetration through endocytosis, and Ad core enters into the nucleus through nuclear
16 pore complexes (Greber *et al.*, 1996; Martin-Fernandez *et al.*, 2004; Nakano *et al.*,
17 2000; Trotman *et al.*, 2001). During entry of the virus genome into the nucleus, core
18 protein V seems to be dissociated from viral chromatin. Thus, viral DNA associated
19 with core protein VII functions as a template for viral early gene transcription and
20 DNA replication in the infected cell nucleus (Chatterjee *et al.*, 1986; Haruki *et al.*,
21 2006; Xue *et al.*, 2005). However, it was reported that core proteins function as a
22 repressor for transcription and replication *in vitro* (Johnson *et al.*, 2004; Matsumoto *et*
23 *al.*, 1995; Nakanishi *et al.*, 1986). Therefore, it is suggested that core proteins are
24 either released or remodeled after entry into the host nucleus (Chen *et al.*, 2007;
25 Matsumoto *et al.*, 1993; Matsumoto *et al.*, 1995; Spector, 2007). Histones may

1 associate with incoming viral DNA (Sergeant *et al.*, 1979; Tate & Philipson, 1979).
2 Recently, it has been shown that not only protein VII but also cellular histones are
3 functional components of viral chromatin in early phases of infection (Komatsu *et al.*,
4 2011). During late stages of infection, the precursor of core protein VII (pre-VII) and
5 core protein V are synthesized at high levels concomitantly with viral DNA synthesis,
6 assembled onto newly replicated DNA, and incorporated into immature virions
7 (Daniell *et al.*, 1981). Newly replicated viral DNA may associate with cellular
8 histones (Dery *et al.*, 1985). However, mature Ad particles do not contain cellular
9 histones. It is still largely unknown how only the virus genome associated with viral
10 core proteins is selectively incorporated into virions. It is also unknown which
11 cellular factors are involved in this process.

12 Previously, we identified host factors termed Template Activating Factors
13 (TAF)-I, -II, and -III from uninfected HeLa cell extracts that remodel the Ad core
14 structure and stimulate replication and transcription from Ad core (Matsumoto *et al.*,
15 1993; Matsumoto *et al.*, 1995; Okuwaki *et al.*, 2001a). Recently, we have shown that
16 TAF-I remodels the Ad core structure by forming a ternary complex with Ad DNA-
17 core protein VII complexes and plays an important role in early stages of the Ad
18 infection cycle (Gyurcsik *et al.*, 2006; Haruki *et al.*, 2003; Haruki *et al.*, 2006;
19 Komatsu *et al.*, 2011). TAF-II is identical to nucleosome assembly protein-1 (NAP-
20 1), a structural and functional homologue of TAF-I (Kawase *et al.*, 1996; Nagata *et al.*,
21 1995). The major component of TAF-III was found to be B23/nucleophosmin
22 (Okuwaki *et al.*, 2001a).

23 B23/nucleophosmin is an abundant ubiquitously expressed cellular protein
24 that modulates diverse molecular functions such as ribosome biogenesis (Hingorani *et al.*
25 *et al.*, 2000; Savkur & Olson, 1998), centrosome duplication (Okuda *et al.*, 2000),

1 chromatin assembly/disassembly (Okuwaki *et al.*, 2005; Okuwaki *et al.*, 2001b), and
2 nucleo-cytoplasmic trafficking (Adachi *et al.*, 1993; Yu *et al.*, 2006). Two splicing
3 variants of B23, B23.1 and B23.2, which differ only in their C-terminal regions, are
4 expressed in a variety of growing cells. Both B23.1 and B23.2 contain highly acidic
5 domains, while the C-terminal region unique for B23.1 is essential for its RNA
6 binding activity. Recently, we have shown that B23 interacts with Ad core protein V,
7 VII, and pre-VII, and may have a role as chaperone in the assembly of core proteins
8 into Ad core (Samad *et al.*, 2007). However, an *in vivo* role(s) of B23 in the Ad life
9 cycle has not been clarified yet. Here, we developed a decoy molecule for the
10 interaction between B23 and core protein V based on the analysis of their interaction
11 domains. Furthermore, we studied the effect of siRNA-mediated knock down (KD)
12 of B23 on Ad proliferation. Perturbation of the B23 function either by over-
13 expression of the decoy molecule or KD was shown to impede the Ad proliferation
14 without significant inhibition of viral DNA replication or viral late gene expression.
15 However, ChIP experiments indicated that the association of both core proteins and
16 cellular histones with viral DNA was significantly increased upon B23 KD.
17 Altogether, the results suggest that B23 is required for maintenance of the proper
18 adenovirus chromatin structure.

19

1 **RESULTS**

2 **Domains of core protein V required for its interaction with B23**

3 Recently, we have shown that B23 interacts with Ad core proteins V and VII
4 (Samad *et al.*, 2007). However, the function of B23 in Ad proliferation has not yet
5 been clarified. To gain additional insight into the *in vivo* function of B23, we have
6 designed a decoy molecule based on the analyses of interaction domains between B23
7 and core protein V. First, we determined the domain of core protein V required for
8 the interaction with B23. Core protein V contains lysine- and arginine-rich basic
9 clusters in its N- and C- terminal regions. We have postulated that core protein V
10 interacts with B23 through these basic clusters, as it has been shown that the acidic
11 region of B23 is essential for its function (Okuwaki *et al.*, 2001a). To test this
12 hypothesis, we constructed a series of deletion mutants as shown in Fig. 1A. GFP-
13 and Flag-tagged core protein V mutants were co-expressed with HA-tagged B23.1 in
14 293T cells, and immunoprecipitation assays were carried out with anti-Flag antibody.
15 HA-B23.1 was co-immunoprecipitated with full length core protein V (Fig. 1B, lane
16 10). The mutants, V(1-313) and V(44-369), lacking the C- and N-terminal regions,
17 respectively, similarly bound to HA-B23.1 (Fig. 1B, lanes 11 and 12). However, the
18 mutants, V(44-313) and V(79-313), lacking both N- and C-terminal basic clusters
19 showed virtually no ability to interact with HA-B23.1 (lanes 13 and 14). In contrast,
20 the N- and C-terminal fragments, V(1-78) and V(314-369), respectively, efficiently
21 co-precipitated HA-B23.1 (lanes 15 and 16). These results indicate that both N- and
22 C-terminal regions are involved in the interaction between B23 and protein V, and
23 these fragments are good candidates for decoy molecules for its interaction with B23.

24

25 **Inhibition of infective virus production by B23 decoy molecule**

1 We hypothesized that over-expression of these domains would interfere with
2 the function of B23 in Ad proliferation if B23 is involved in Ad life cycle. To test
3 this, HeLa cells were transfected with either GFP-empty vector or vectors for the
4 expression of GFP-V, GFP-V(1-78), GFP-V(79-313), and GFP-V(314-369), and
5 super-infected with HAdV5 at 20 hours after transfection. At 24 hpi, progeny virus
6 particles were collected, and the infectivity titer was examined as described in
7 Methods (Supplemental Fig. 1 and Fig. 1). Results demonstrate that the over-
8 expression of GFP-core protein V, GFP-V(1-78), and GFP-V(314-369), but neither
9 GFP alone nor GFP-V(79-313), inhibited the production of infectious virus particles.
10 These results suggest that the mutant proteins that tightly associate with B23 inhibit
11 the infectious virus production. It was further demonstrated that the co-expression of
12 exogenous B23 with GFP-V(1-78) (Fig. 1D) rescued the negative effect of GFP-V(1-
13 78). These results support the idea that V(1-78) functions as a sort of decoy for the
14 interaction between B23 and core protein V, and the impairment of this interaction
15 decreases the progeny virus production level.

16

17 **B23 knock-down (KD) inhibits the production of infective viral particle**

18 To further show that B23 is involved in Ad proliferation, we decreased the
19 cellular B23 level using siRNA specific for B23.1. Although both B23.1 and B23.2
20 have been suggested to be differentially involved in Ad replication (Hindley *et al.*,
21 2007), we focused on B23.1 as it is concentrated in the nucleoli in which core protein
22 V is located at the late stage of infection (Matthews, 2001), while B23.2 is distributed
23 throughout the nuclei. In addition, the depletion of B23.1 alone efficiently decreased
24 the nucleolar function of B23 (Murano *et al.*, 2008). Treatment of HeLa cells with
25 B23.1 siRNA decreased the cellular B23.1 protein level but not β -actin, whereas

1 control siRNA had no effect (Fig. 2A). The expression level of nucleolar proteins
2 such as nucleolin and fibrillarin were found to be unchanged upon B23 knock down
3 (KD) (Fig. 2B). To examine the effect of B23 KD on Ad proliferation, control
4 siRNA- and B23 siRNA-treated HeLa cells were infected with HAdV5. We first
5 examined the effect of siRNA treatment on the localization of viral proteins (Fig. 2C).
6 In control siRNA-treated cells, DBP was concentrated in nuclear foci and core protein
7 VII distributed throughout nuclei at 24 hpi. As previously reported (Hindley *et al.*,
8 2007; Matthews, 2001), the nucleolar localization of B23 was slightly suppressed
9 upon Ad infection, and B23 was partially co-localized with core protein VII but not
10 with DBP. We also demonstrated that the localization patterns of DBP and core
11 protein VII were not significantly affected by B23.1 siRNA treatment. At 24 hpi, the
12 supernatant fraction containing progeny virus particles were collected, cleared by low
13 speed centrifugation, and then examined for the infectious titer (Fig.2D). The
14 production of infectious progeny virus particles from B23 KD cells were decreased to
15 approximately 50 to 60% of that from control siRNA-treated cells (Fig. 2D, lanes 1
16 and 2). Even under the decreased B23.1 level, Ad virus production was increased
17 until 48 hpi, although the amounts of produced infective virus during 24-36 hpi were
18 lower than those of control siRNA-treated cells (Fig. 2C and supplemental Fig. 2).
19 This result suggests that B23.1 is not essential but plays a crucial role in Ad virus
20 production, and/or other cellular factor(s) could also be involved (see Discussion).
21 Next, we investigated whether the effect of B23 KD on Ad virus production could be
22 rescued by B23 over-expression. At 36 hours after introduction of control or B23
23 siRNA, cells were transfected with either empty vector or vector encoding HA-B23.1.
24 Cells were then infected with Ad at 24 hours after transfection of plasmid DNA, and
25 the production of progeny virus particles was examined. The expression level of

1 exogenous HA-B23.1 is shown in Fig. 2F. Over-expression of B23.1 in control cells
2 slightly inhibited the infectious progeny virus production (Fig. 2D), although this
3 result was not statistically significant. Importantly, the exogenous expression of
4 B23.1 counteracted the negative effect of B23 siRNA-mediated KD on the progeny
5 virus production. These results support the idea that B23.1 plays an important role in
6 the production of infectious virus particles.

7

8 **B23 KD has no significant effect on viral DNA replication and late gene** 9 **expression**

10 Given that B23 KD decreased the production of infectious virus particles, it was
11 highly expected that this inhibition might be due to the interference with viral DNA
12 replication. To test this notion, control or B23 siRNA-treated HeLa cells were
13 infected with Ad and the amplification of viral DNA at 12, 18, and 24 hpi was
14 examined by quantitative PCR using a primer set specific for Ad DNA (Fig. 3A).
15 Because the amount of Ad DNA was increased as a function of incubation periods
16 after infection and the amplification of DNA is strongly inhibited by hydroxyl urea
17 (HU), it is confirmed that the PCR products detected under the condition employed
18 here correspond to the amounts of the viral DNA. Surprisingly, no significant
19 decrease in the amount of viral DNA upon B23 KD was observed (Fig. 3A). We also
20 examined the effect of B23 KD on the late gene expression. The expression level of
21 the late gene was examined by western blotting with anti-pVII and anti-V antibodies
22 (Fig. 3B) and RT-PCR with primer sets for mRNAs of the major late promoter and
23 pVII (Figs. 3C and D). Consistent with the fact that the late gene transcription
24 depends on viral DNA replication, the expression level of late genes was strongly
25 inhibited by the presence of HU. We did not find any significant decrease in both

1 mRNA and core proteins expression levels upon B23 KD (Figs. 3B-D). These results
2 indicated that B23 plays a crucial role(s) in progeny virus production at a step(s) later
3 than virus genome DNA replication and mRNA synthesis. Since Ad genome
4 replication completely depends on viral early gene products, we could exclude the
5 possibility that B23 is involved in the early gene transcription.

6

7 **B23 regulates the amounts of core proteins and cellular histones on the Ad**
8 **genome**

9 Our biochemical data suggested that the Ad core proteins forms aggregates with
10 viral DNA when mixed directly, and B23 dissociates the aggregation between DNA
11 and core proteins (Samad *et al.*, 2007). In addition, we demonstrated that B23, as a
12 histone chaperone, regulates the histone density around the rRNA gene region in
13 uninfected cells (Hisaoka *et al.*, 2010). Therefore, it is possible that B23 KD affects
14 the virus genome chromatin structure in infected cells. To test this possibility, we
15 examined whether B23 is associated with the virus genome in infected cells. HeLa
16 cells were infected with Ad, and at 20 hpi cells were cross-linked with formaldehyde
17 and then sonicated to release chromatin. The average size of DNA purified from
18 chromatin fragments was <1 kb (data not shown). The extracts were subjected to
19 immunoprecipitation with antibodies against core proteins V, VII, or B23. We found
20 that B23 associates with the virus genome (the VA gene region) as do core proteins V
21 and VII (Fig. 4A). We examined the association of B23 with the virus genome using
22 primer sets as shown in the bottom panel of Fig. 4B. Next, we assessed the amounts
23 of core proteins and cellular histones on the virus genome with ChIP assays using
24 cells treated with control and B23 siRNAs. HeLa cells treated with siRNAs were
25 infected with HAdV5 at an MOI of 10. At 20 hpi, cells were subjected to ChIP assays

1 as described above. Five different primer sets as shown in Fig. 4B were used to
2 examine the amounts of core proteins and histones on the Ad genome. In B23 KD
3 cells, the association of both core proteins V and VII with viral DNA were found to
4 be increased in all regions tested (Figs. 4C and D). We also found that the association
5 of histone H3 along the virus genome is increased (Fig. 4E).

6 We also examined whether the amounts of core proteins and histone H3 on the
7 virus genome increased by B23.1 KD were counteracted by exogenously expressed
8 B23.1. HeLa cells treated with control or B23.1 siRNA were transiently transfected
9 with empty and HA-B23.1 expression vectors, and then infected with Ad. At 24
10 hours after Ad infection, ChIP assays were carried out as shown in Figure 4 using
11 primer sets specific for VA gene region (Figure 5). Western blotting analyses
12 demonstrated that the amount of B23.1 was decreased efficiently by siRNA treatment,
13 and that was recovered by transient expression of exogenous HA-tagged B23.1
14 (Figure 5A, lanes 5 and 6). Consistent with the data in Figure 4, the amounts of core
15 proteins V, VII, and histone H3 on the virus genome was increased by B23.1 KD
16 (Figure 5B-D). However, those were decreased upon over-expression of HA-B23.1.
17 Interestingly, even in control siRNA-treated cells, HA-B23.1 over-expression
18 decreased the association level of core protein V and histone H3. Taken together,
19 these results suggest that B23 is involved in the regulation of viral chromatin
20 formation in infected cells by restricting the access of core proteins and cellular
21 histones.

22

1 **DISCUSSION**

2 In this paper, we have studied the *in vivo* function of B23 in Ad life cycle.
3 Based on previous reports, it was expected that B23 may be involved in Ad DNA
4 replication (Hindley *et al.*, 2007; Okuwaki *et al.*, 2001a). However, we could not
5 detect any significant decrease in the amount of viral DNA (Fig. 3A) as well as the
6 level of both transcription and translation of core proteins (Figs. 3B-D) upon B23.1
7 KD. Therefore, it is possible that B23.1 is not involved in Ad DNA replication in
8 infected cells under the conditions employed here or the loss of B23.1 may be
9 compensated alternatively. In this sense, it is noted that not only B23 but also other
10 histone chaperones are identified as factors for Ad DNA replication (Kawase *et al.*,
11 1996; Matsumoto *et al.*, 1993; Okuwaki *et al.*, 2001a). In addition, we could not
12 exclude the possibility that B23.2 remained in B23.1 KD cells plays a compensatory
13 role in Ad DNA replication. Nevertheless, the data presented here demonstrate that
14 B23.1 KD did not significantly affect DNA replication and transcription.

15 We have shown that the decrease of B23 reduces the production of progeny
16 virions and increases the association level of viral core proteins and cellular histones
17 on the progeny virus genome DNA. Based on these observations, we would propose
18 that (1) B23 is involved in the adenovirus infection cycle at a step later than late gene
19 expression, and (2) proper virus chromatin assembly is required for the Ad virion
20 maturation. The precise mechanism of how B23 regulates viral chromatin and is
21 involved in the final maturation step of infective virus particles remains unclear. It is
22 also possible that in addition to B23, other cellular factor(s) is involved in these
23 processes. TAF-I is a candidate of such additional factors, because we have reported
24 that TAF-I is also associated with pre-VII in the late phases of infection (Gyurcsik *et*
25 *al.*, 2006). It is suspected that the viral DNA associated with appropriate amounts of

1 basic proteins is important for encapsidation. In fact, the Ad genome DNA is
2 condensed into a core structure only by viral basic proteins within virions, although
3 viral DNA is associated with histones throughout the infection cycle (Dery *et al.*,
4 1985; Levy & Noll, 1981). Thus, it is likely that those virus genomes associated with
5 cellular histones are restricted and eliminated for encapsidation. For efficient
6 encapsidation to occur, cellular histones must be replaced with viral basic proteins
7 through an unknown pathway.

8 Newly replicated DNA is associated with histones, and this viral DNA-histone
9 complex might be important for ongoing replication and transcription throughout the
10 infection cycle. At later stages of infection, the synthesis of cellular DNA and
11 histones is inhibited with the concomitant accumulation of a large pool of viral basic
12 proteins. The virus genomic DNA associated with viral core proteins might become
13 prominent through direct interaction of ongoing replicated DNA with viral basic
14 proteins and/or replacement of histones on the replicated DNA with viral basic
15 proteins. Based on these observations, B23 may be involved in the final
16 encapsidation step either by replacing histones with core proteins or by restricting the
17 access of excessive amounts of viral basic proteins/histones to viral DNA. These
18 models are in agreement with earlier assumption (Dery *et al.*, 1985; Komatsu *et al.*,
19 2011).

20

1 **METHODS**

2 **Cell culture and viruses**

3 HeLa cells were maintained in minimal essential medium (MEM; Nissui)
4 supplemented with 10% fetal bovine serum (FBS) at 37°C. 293T cells were cultured
5 at 37°C in Dulbecco's modified Eagle medium (DMEM; Nissui) containing 10% FBS.
6 Human adenovirus type 5 (HAdV5) used in this study was amplified and purified as
7 previously described (Haruki *et al.*, 2006).

8

9 **Plasmid construction and transfection**

10 Construction of plasmids for a series of protein V mutants is described in
11 Supplemental Methods. pCHA-B23.1 was prepared as described (Okuwaki *et al.*,
12 2002). Transient transfection of each plasmid was performed by the calcium
13 phosphate precipitation method into 293T cells and by Gene-Juice (Novagen) into
14 HeLa cells.

15

16 **Antibodies**

17 The antibodies used in this study were as follows: mouse monoclonal antibody for
18 B23 that recognizes endogenous B23.1 was purchased from Invitrogen. Polyclonal
19 antibody that recognizes both B23.1 and B23.2 was generated in rabbits using
20 B23 Δ C2 expressed in *E. coli* (Okuwaki *et al.*, 2001a) as an antigen. Rabbit anti-core
21 protein V and mouse anti-DBP antibodies were obtained from Dr. W.C. Russel as
22 generous donation. Rat polyclonal anti-pre-VII (Haruki *et al.*, 2003), mouse
23 monoclonal antibody for Flag-tag, rabbit monoclonal antibody for hemagglutinin
24 (HA)-tag, rabbit polyclonal antibody for histone H3, and mouse monoclonal antibody
25 for β -actin were described elsewhere (Murano *et al.*, 2008).

1

2 **Immunoprecipitation assays**

3 293T cells transiently transfected with plasmids where indicated were lysed in 1 ml of
4 IP buffer (50 mM Tris-HCl, pH7.9, 1 mM PMSF, 0.1% Triton X-100, and 1 mg/ml
5 bovine serum albumin) containing 150 mM NaCl on ice for 10 min followed by
6 extensive sonication. Cell extracts recovered by centrifugation were mixed with anti-
7 Flag antibody, and incubated at 4°C for 3 hours. Then, protein A sepharose beads (10
8 µl of resin; GE health care) were added and further incubated for 1 hour with gentle
9 agitation. The beads were washed three times with 0.5 ml of IP buffer.
10 Immunoprecipitated proteins were eluted by an SDS sample buffer, boiled, separated
11 on a 12.5% SDS-PAGE, and transferred to a polyvinylidene difluoride (PVDF)
12 membrane (Millipore). The membrane was subjected to western blotting analysis
13 using anti-HA antibody.

14

15 **Decoy assays**

16 HeLa cells were transfected with either GFP-empty vector or the vector for the
17 expression of GFP-V(1-78). At 20 hours post transfection, cells were super-infected
18 with HAd5 at an MOI of 10. At 24 hpi, culture supernatant (5 µl) clarified by low
19 speed centrifugation was used for infection of HeLa cells seeded on coverslips in 12
20 well-plates (1×10^5 cells/well) to determine the virus titer. After incubation for 1 hour,
21 cells were supplemented with MEM containing 2% FBS and maintained at 37°C in a
22 5% CO₂ environment for additional 15 hours. Cells on cover slips were collected,
23 washed with PBS (-), fixed with 3% paraformaldehyde for 10 min at room
24 temperature, and stained with anti-DBP antibody. Cells were also counter-stained
25 with 4',6-diamidino-2-phenylindole (DAPI) and the infectious titers (% of infected

1 cells) was determined by counting DBP positive cells out of DAPI positive cells.
2 This assay was carried out in doublet, and error bars indicate SD.

3

4 **B23 knock down by siRNA**

5 B23 Stealth RNAi (NPM1-HSS143154) and Stealth RNAi negative control (catalog
6 no. 12935-200) were purchased from Invitrogen and introduced into HeLa cells with
7 LipofectamineTM RNAiMAX (Invitrogen) according to the manufacturer's protocol.
8 At 24 hours post transfection, the medium was replaced, and cells were harvested
9 after 60 hours post siRNA transfection. Total HeLa cell lysates were prepared, and
10 proteins were separated through 10% SDS-PAGE and detected by western blotting.
11 To examine the effect of B23 KD on the virus production, control and B23 siRNA-
12 treated HeLa cells at 60 hours post transfection were infected with HAdV5 at an MOI
13 of 10. After 24 hours, the culture medium was recovered and examined for virus titer
14 as described above.

15

16 **Immunofluorescence assay**

17 Indirect immunofluorescence assays were carried out essentially as described
18 previously (Haruki *et al.*, 2006). Briefly, cells grown on cover slips (15 mm;
19 Matsunami) were fixed with 4% paraformaldehyde in PBS for 10 min at room
20 temperature (RT) and then treated with 0.5% NP-40 in PBS for 5 min at RT. After
21 blocking with 5% nonfat milk in TBS-T, samples were subjected to
22 immunofluorescence analyses using antibodies described above. Localization of the
23 protein was visualized with the secondary antibodies (anti-rabbit IgG conjugated with
24 AlexaFluor 488, anti-mouse IgG conjugated with AlexaFluor 568, and anti-rat IgG
25 conjugated with AlexaFluor 568; Invitrogen). DNA was visualized by staining with

1 TO-PRO-3 iodide (Invitrogen). Labeled cells were observed with confocal laser
2 scanning microscopy (LSM5 Exciter; Carl Zeiss) using argon laser (488 nm) and
3 He/Ne laser (546 and 633 nm) lines.

4

5 **Quantitative PCR**

6 Control and B23 siRNA-treated HeLa cells were infected with HAdV5 at an MOI of
7 10. At 12, 18, and 24 hours post infection, cells (1×10^5) were collected and
8 suspended in lysis buffer (20 mM Tris pH7.9, 100 mM NaCl, 5 mM EDTA, and 0.5%
9 SDS), and total DNA was purified with proteinase K treatment at 50°C for overnight
10 followed by phenol-chloroform extraction and ethanol precipitation. The amount of
11 DNA was then examined by quantitative (q) PCR with a primer set specific for the Ad
12 VA gene region (see below). Total RNA was purified from infected cells (1×10^5)
13 using RNeasy mini kit (QIAGEN), and the purified RNA was treated with DNase I
14 according to the manufacturer's protocol. The concentration of RNA in each sample
15 was determined by using NanoDrop (Thermo Scientific). cDNA was synthesized
16 from total RNA (1 μ g) using ReverTraAce (Toyobo) and oligo-dT as primer
17 according to the manufacturer's protocol. qPCR with FastStart SYBER Green Master
18 Mix (Roche) and Thermal Cycler Dice Real Time System (Takara) was performed
19 using synthesized cDNA as a template with primer sets specific for the mRNA from
20 major late promoter (MLP), 5'-ACTCTCTTCCGCATCGCTGT-3' and 5'-
21 GTGACTGGTTAGACGCCT-TTCT-3', and β -actin gene, 5'-
22 ATGGGTCAGAAGGATTCCTATGT-3' and 5'-GGTCATCTTCTCGCGGTT-3'.

23

24 **ChIP assays**

1 ChIP assays were carried out according to the manual of ChIP assay kit (Millipore)
2 with anti-core protein V, pVII, B23, and histone H3 antibodies. The amount of
3 immunoprecipitated DNA was determined by qPCR as described above. The reaction
4 condition was described previously (Komatsu *et al.*, 2011). The following primer sets
5 were used; 5'-GGGTCAAAGTTGGCGTTTTA-3' and 5'-
6 CAAAATGGCTAGGAGGTGGA-3' for the E1a promoter region, 5'-
7 GCGGTCCTCCTCGTATAGAA-3' and 5'-CCCACCCCCTTTTATAGCC-3' for the
8 ML promoter region, 5'-GCTGGAGCAAAACCCAAATA-3' and 5'-
9 TATCTTGCGGG-CGTAAACT-3' for the VA region, 5'-
10 GTGTAGACACTTAAGCTCGCCTT-3' and CTTCAAAGTGCCTGACCAAGT-3'
11 for the E2A (DBP) region, and 5'-TGGCGTGGTCAAAGTCTACA-3' and 5'-
12 GATTTTTACAATGGCCGGACT-3' for the E4 ORF region.

13

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7

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- 34
35

1 **FIGURE LEGENDS**

2 **FIG. 1. N- and C-terminal regions of core protein V are required for the**

3 **interaction with B23.** (A) Schematic diagram of wild type and mutant protein V.

4 GFP-Flag-tag was fused at the N-termini of core protein V and its mutant proteins.

5 The basic clusters of core protein V are indicated by filled boxes. (B)

6 Immunoprecipitation of Flag-core protein V. 293T cells were transiently co-

7 transfected with HA-B23.1 and either GFP-Flag-V or its mutants. After

8 immunoprecipitation without (-) or with (+) anti-Flag antibody, proteins in input

9 extracts (lanes 1-8) and precipitated proteins (lanes 9-16) were separated through a

10 12.5% SDS-PAGE and detected by western blotting with anti-HA and anti-Flag (top

11 and bottom panels, respectively) antibodies. Positions of GFP-Flag-tagged protein V

12 and its mutants are indicated by arrow heads at the left side of each lane. (C)

13 Inhibition of virion production by a decoy molecule. HeLa cells were transfected

14 with either GFP-empty vector or a vector expressing GFP-Flag-tagged V(1-78)

15 mutant. At 20 hours after transfection, cells were infected with HAd5. At 24 hpi,

16 viruses in the culture fluid were collected and examined for virus titer. In the bottom

17 panel, expression of exogenous proteins as well as β -actin was confirmed by western

18 blotting. (D) Rescue experiments. Cells expressing either GFP-Flag (lane 1), GFP-

19 Flag-V(1-78) with pCHA empty vector (lane 2), or GFP-Flag-V(1-78) with pCHA-

20 B23.1 vector (lane 3). At 24 hour after transfection, cells were super-infected with

21 Ad and virus production was examined by the procedure as described in (C).

22 Exogenously expressed proteins were detected by western blotting as shown at the

23 bottom of the panel.

24

1 **FIG. 2. B23 is involved in Ad proliferation.** (A) Knockdown (KD) of B23.1.
2 HeLa cells were transfected with control siRNA (lane 4) or B23 siRNA (lane 5), and
3 the expression level of B23 was examined by western blotting with anti-B23 antibody.
4 HeLa cell lysates from 5×10^3 , 1.5×10^4 , and 5×10^4 cells for lanes 1, 2, and 3,
5 respectively, were loaded on the same gel and used as standards. β -actin is shown as
6 a loading control. (B) Expression of nucleolar proteins. The expression level of
7 indicated proteins was determined by western blotting using HeLa cells treated with
8 control siRNA or B23.1 siRNA (lanes 1 and 2, respectively) as in (A). (C)
9 Localization of viral proteins in B23.1 siRNA-treated cells. HeLa cells treated with
10 control- or B23.1-siRNA as indicated at the top of the panels were super-infected with
11 Ad and subjected to indirect immunofluorescence analyses at 24 hpi. B23 and DBP
12 (top panels) or B23 and core protein VII (bottom panels) were simultaneously stained
13 and visualized. DNA was counter-stained with To-Pro-3. (D) Inhibition of infectious
14 virus production by KD of B23. HeLa cells treated with either control- or B23.1-
15 siRNA were infected with HAd5. At 24 hpi, virus titers in the culture medium were
16 examined as described in Methods (lanes 1 and 2). At 36 hrs after siRNA transfection,
17 pCHA empty vector (lanes 3 and 4) or pCHA-B23.1 (lanes 5 and 6) were transfected
18 and incubated for 24 hrs. Then, cells were super-infected with HAd5, and the virus
19 infectivity was determined at 24 hpi as described above. Experiments were carried
20 out in triplicate, and error bars indicate standard deviations (SD). Statistical *P* values
21 are indicated at the top of the graph. (E) Expression level of endogenous and
22 exogenous B23. Lysates prepared as indicated in (D) were analyzed by western
23 blotting with anti-B23.1 and anti β -actin antibodies (top and bottom panels,
24 respectively).
25

1 **FIG. 3. Effect of B23 KD on Ad DNA replication and late gene expression.** (A)
2 Effect of B23.1 KD on Ad DNA replication. Control or B23.1 siRNA-treated HeLa
3 cells (black and white bars, respectively) were infected with Ad and incubated
4 without (-) or with (+) 2 mM hydroxyl urea (HU) as indicated at the bottom of the
5 panel. At 12, 18, and 24 hpi, DNA was purified from infected cells, and the amount
6 of viral DNA was examined by quantitative PCR using a primer set specific for the
7 VA region of the Ad genome. Genomic DNA purified from HeLa cells infected with
8 Ad was used as standards for the amount of the Ad genome in infected cells, and the
9 relative amounts of Ad DNA was normalized by that of β -actin gene. (B) Expression
10 level of late gene products. Lysates were prepared from infected HeLa cells as
11 described in (A) at 24 hpi. Proteins were separated on SDS-PAGE, and analyzed by
12 western blotting with anti- β -actin, -B23, -core protein V, and -pVII antibodies. (C, D)
13 Expression level of Ad late genes. HeLa cells treated with siRNA as described in A
14 were infected with Ad. Total RNA was prepared at 18 hpi, and the expression level
15 of MLP mRNA (C) and pVII mRNA (D) was determined by quantitative RT-PCR
16 using specific primer sets as described in Methods. PCR reactions were performed in
17 triplicate, and error bars indicate SD. Three independent experiments showed similar
18 results.

19

20 **FIG. 4. Effect of B23 KD on the Ad viral chromatin structure.** (A) ChIP assays
21 were carried out with extracts prepared from Ad-infected HeLa cells at 20 hpi. DNA
22 immunoprecipitated without (lane 2) or with anti-pVII, anti-V, and anti-B23 (lanes 3-
23 5) antibodies was examined by PCR using a primer set specific for the Ad VA gene.
24 (B) B23 associates with entire Ad genome in infected cells. Immunoprecipitated
25 DNA with anti-B23 as described in (A) was subjected to quantitative PCR using

1 primer sets specific for Ad genome. Positions of primer sets used are schematically
2 represented at the bottom of the panel. Arrows indicate the positions and direction of
3 the transcription of each gene. DNA extracted from input extracts was used as
4 standards to quantify the amount of DNA immunoprecipitated with anti-B23 antibody.
5 (C-E) ChIP assay of Ad-infected HeLa cells treated with siRNAs. Control or B23.1
6 siRNA-treated HeLa cells were infected with Ad, and ChIP assays were carried out
7 with anti-core protein V, anti-pVII, anti-histone H3 antibodies (C-E, respectively).
8 Immunoprecipitated DNA was quantitatively examined by q-PCR using primer sets
9 shown in (B). Black and white bars in graphs indicate the results obtained from
10 extracts prepared from control and B23.1 siRNA-treated HeLa cells, respectively.
11 The amounts of immunoprecipitated DNA were quantitatively analyzed compared
12 with those of DNAs extracted from input extracts. For B-E, PCR reaction was carried
13 out in triplicate, and error bars indicate SD. Two independent experiments showed
14 similar results.

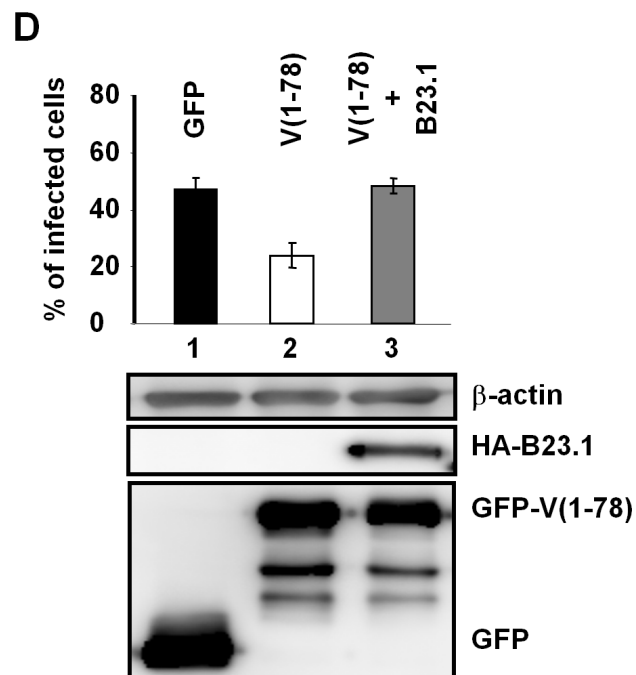
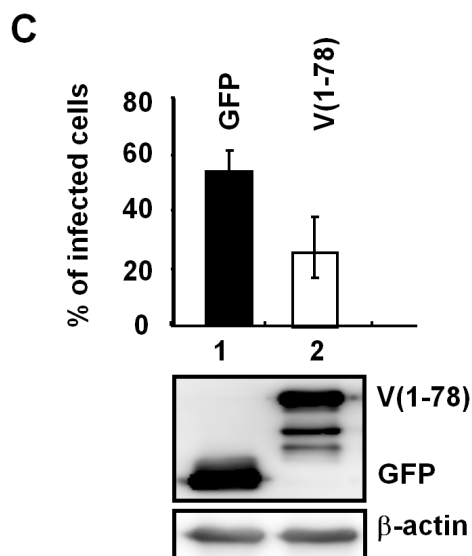
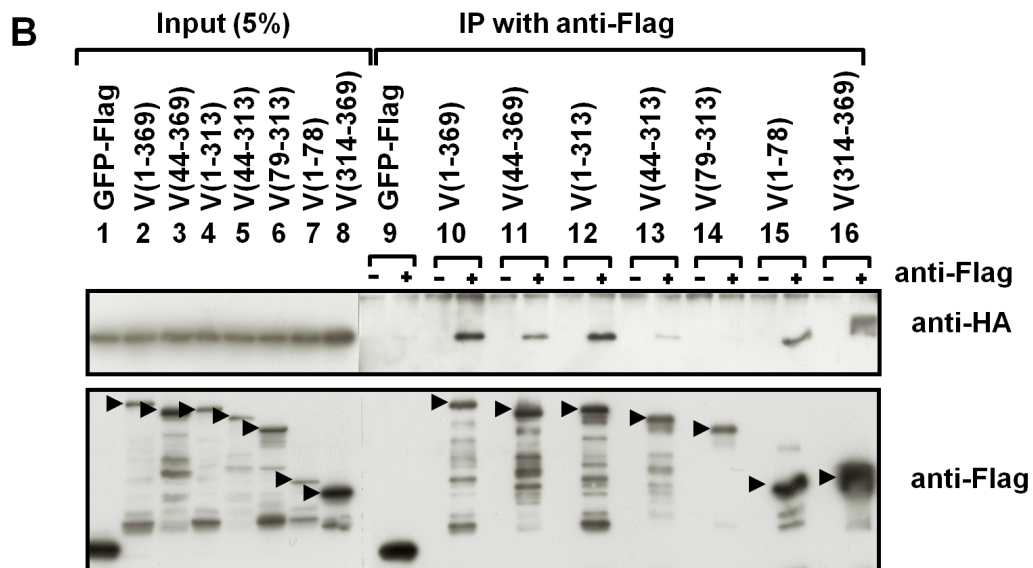
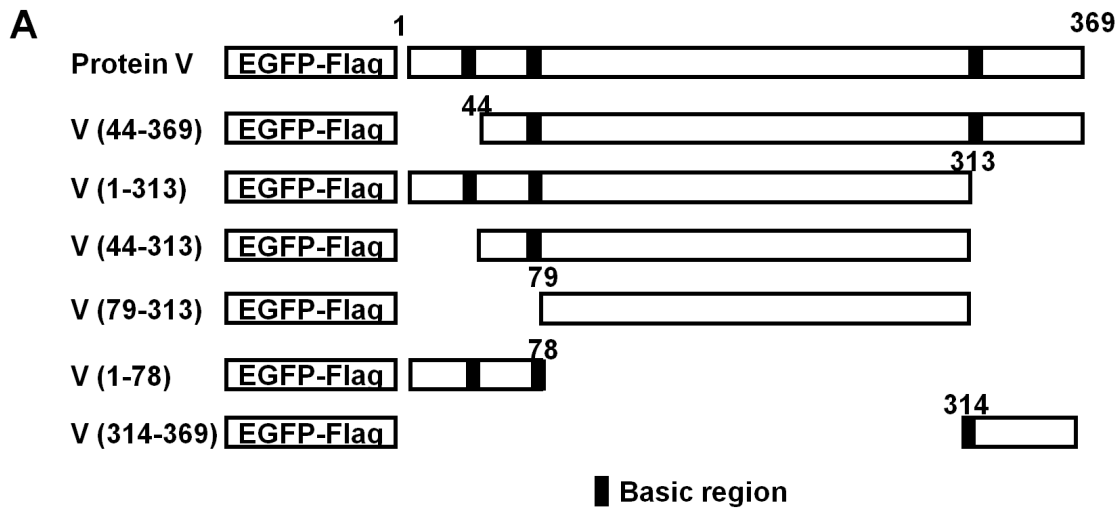
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16 **FIG. 5. Over-expression of exogenous B23.1 counteracts the effect of B23.1 KD**
17 **on viral chromatin structure.** (A) The expression levels of endogenous and
18 exogenous B23.1. HeLa cells were treated with control- or B23.1-siRNA (lanes 1, 3,
19 and 5 or 2, 4, and 6) without or with transfection of pCHA (lanes 3 and 4) or pCHA-
20 B23.1 (lanes 5 and 6), and then infected with Ad. The expression level of B23 and β -
21 actin was examined by western blotting analyses. (B-D) ChIP assays. HeLa cells
22 prepared as described in (A) were subjected to ChIP assays with anti-core protein V
23 (B), -core protein VII (C), or histone H3 (D) antibody. Immunoprecipitated DNA
24 was quantitatively examined by q-PCR using primer sets specific for the Ad VA gene
25 region. Black and white bars in graphs indicate the results obtained from extracts

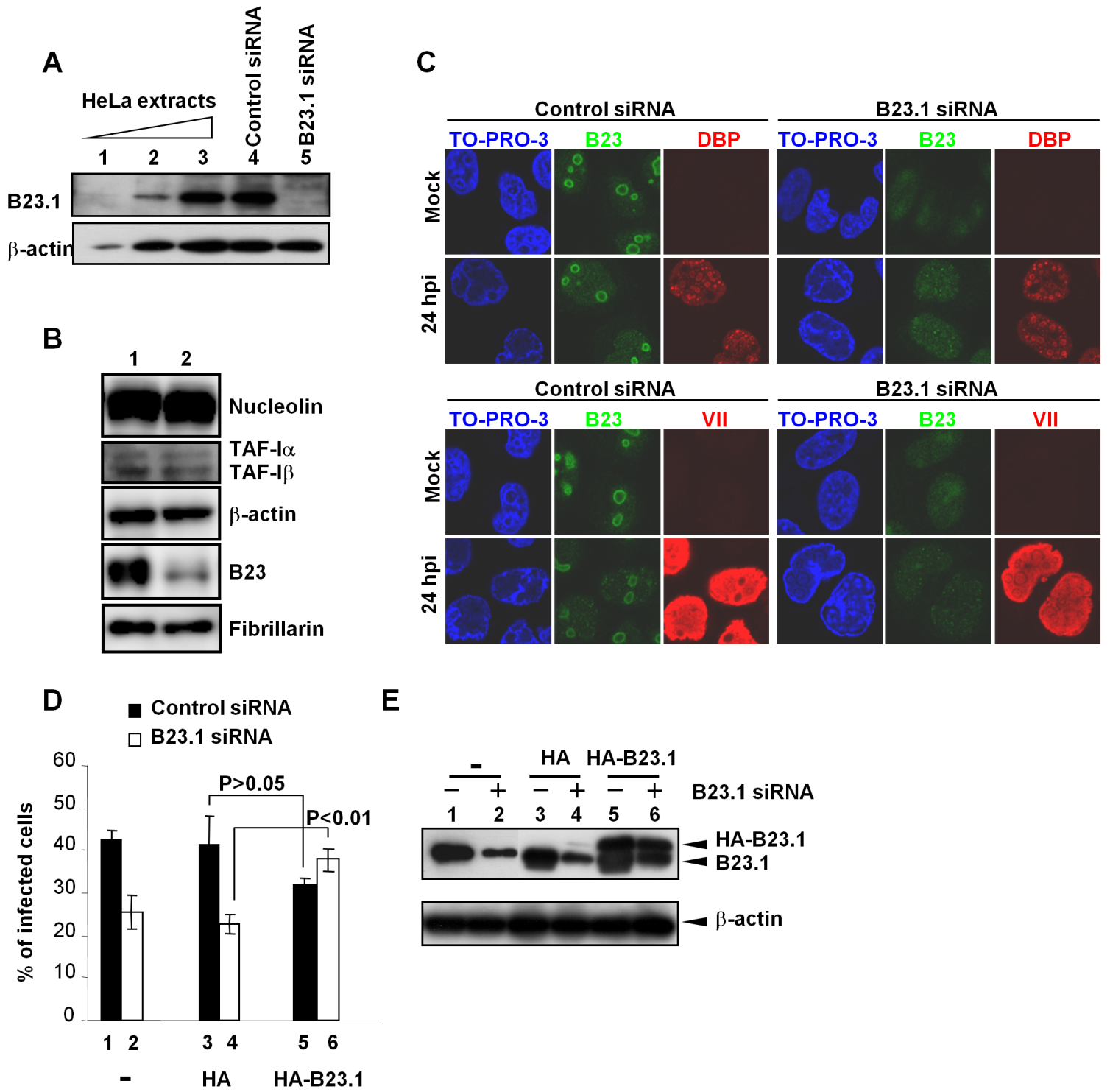
1 prepared from control and B23.1 siRNA-treated HeLa cells, respectively. The
2 amount of immunoprecipitated DNA was quantitatively analyzed compared with
3 those of DNAs extracted from input extracts. PCR reactions were carried out in
4 triplicate, and error bars indicate SD.

5

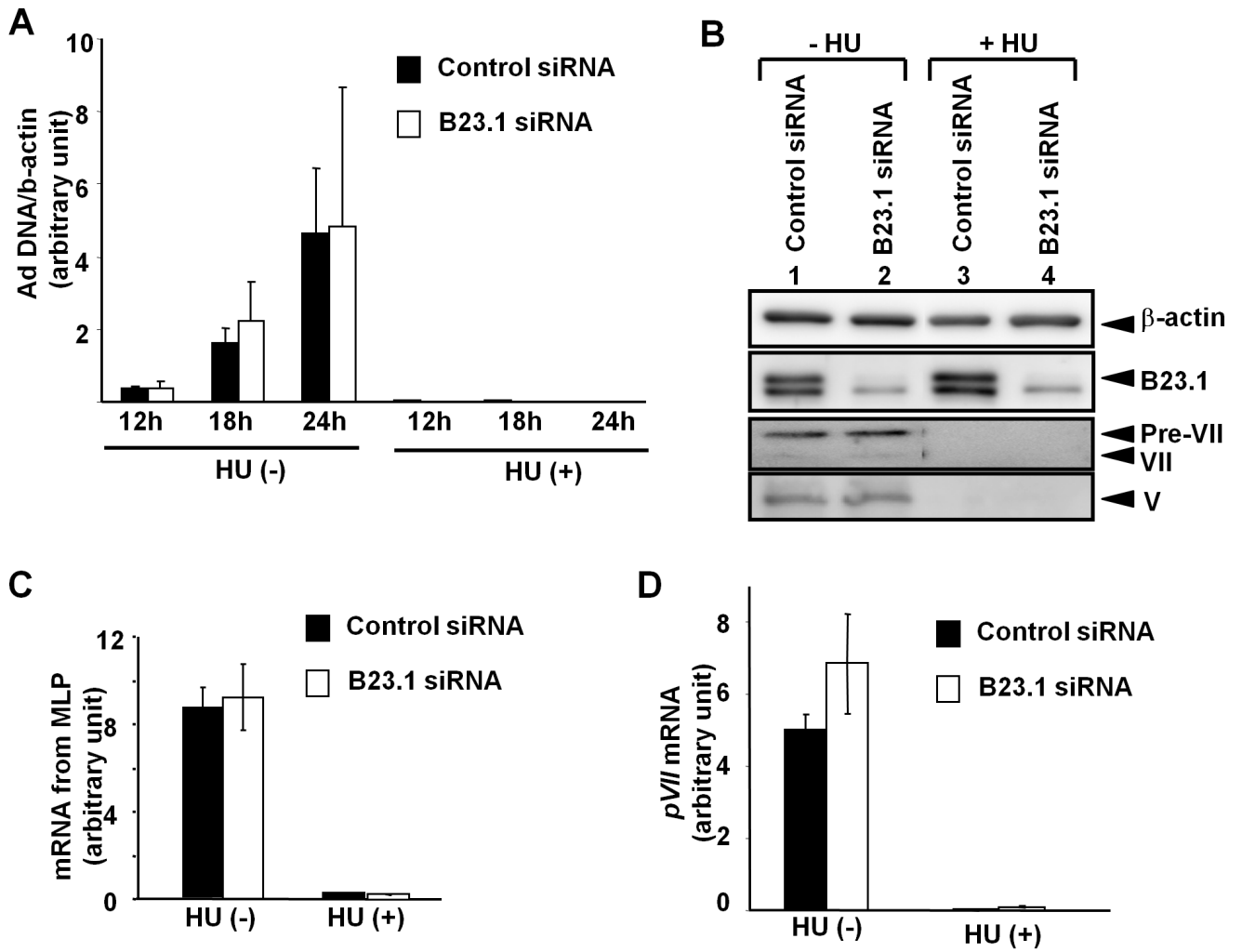
Samad Fig. 1



Samad Fig. 2

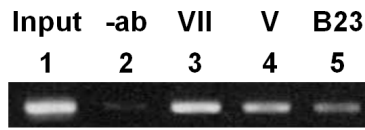


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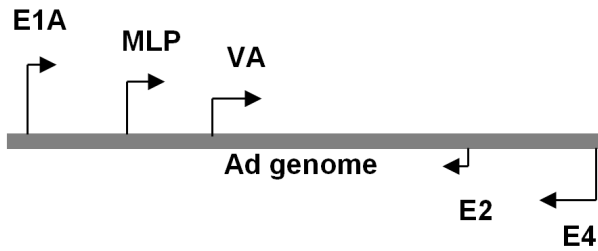
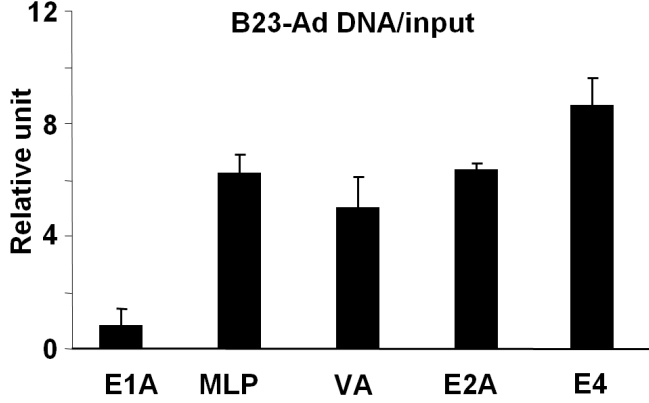


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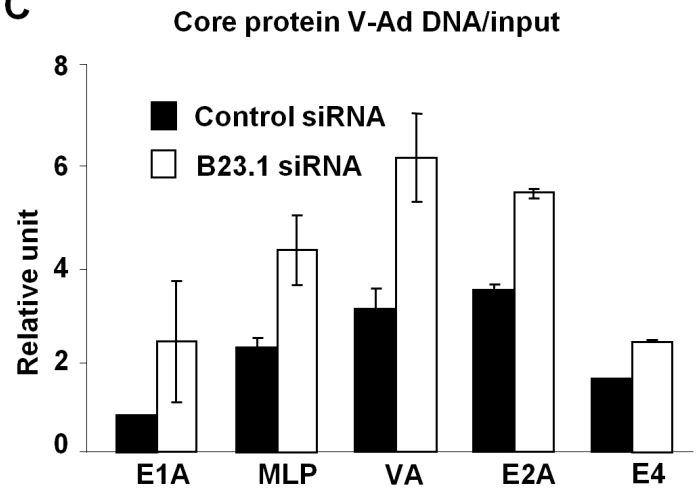
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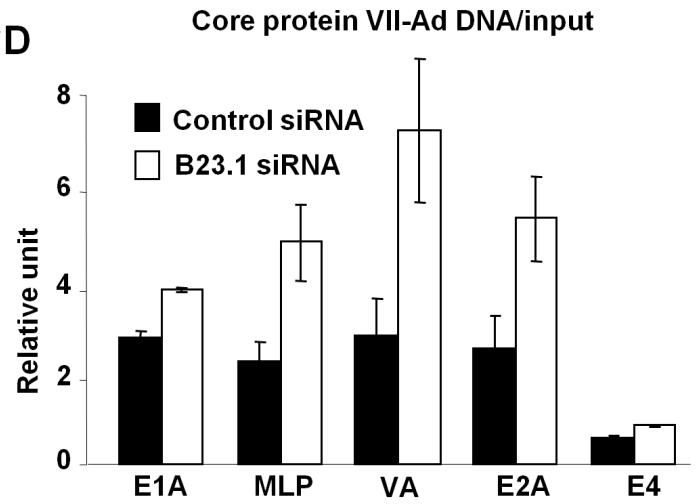
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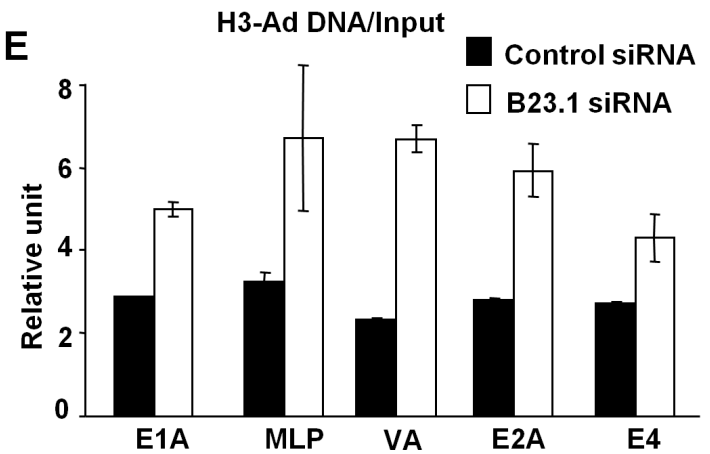
C



D



E



Samad Fig. 5

