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# Affinity Labeling of Hepatitis C Virus Replicase with a Nucleotide Analog: Identification of binding site

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#### **Abstract**

We have used an ATP analog, 5'-[p-(fluorosulfonyl)benzoyl]adenosine (FSBA), to modify HCV replicase in order to identify ATP binding site in the enzyme. FSBA inactivates HCV replicase activity in a concentration dependent manner with a binding stoichiometry of 2 moles of FSBA per mole of enzyme. The enzyme activity is protected from FSBA in the presence of rNTP substrates or double stranded RNA template primers that do not support ATP as the incoming nucleotide but not in the presence of polyrU.rA<sub>26</sub>. The HPLC analysis of tryptic peptides of FSBA-modified enzyme revealed the presence of two distinct peptides eluted at 23 min and 36 min; these were absent in the control. Further we noted that both the peptides were protected from FSBA modification in the presence of Mg.ATP. The LC/MS/MS analysis of the affinity-labeled tryptic peptides purified from HPLC, identified two major modification sites at positions 382 (Tyr), and 491 (Lys) and a minor site at position 38 (Tyr). To validate the functional significance of Tyr38, Tyr382 and Lys491 in catalysis, we individually substituted these residues by alanine and examined their ability to catalyze RdRp activity. We found that both Y382A and K491A mutants were significantly affected in their ability to catalyze RdRp activity while Y38A remained unaffected. We further observed that both Y382A and K491A mutants were not affected in their ability to bind template primer but significantly affected in their ability to photo-crosslink ATP in the absence or presence of template primer.

HCV is currently the major cause of chronic liver disease, with approximately 3% of the world's population estimated to be infected with the virus <sup>(1)</sup>. Although some infected individuals are able to clear the virus without treatment, most infections persist if untreated, leading to chronic hepatitis C, which may further lead to liver cirrhosis and hepatocellular carcinoma <sup>(2)</sup>. Current therapy for HCV infection includes extended administration of a combination of ribavirin and pegylated interferon-α <sup>(3–5)</sup>. However, this combination produces sustained virological response of approximately 40–50% in genotype 1-infected individuals but does not show similar response in other genotypes <sup>(6,7)</sup>. Recently, the FDA approved use of the protease inhibitors, Boceprevir and Telaprevir, which in combination with ribavirin and pegylated interferon-α demonstrated a cure rate of 60–80% for genotype 1 patients <sup>(8–10)</sup> and thus offer a new treatment option for the patients who failed to respond conventional treatment <sup>(11)</sup>. However, these inhibitors have showed some serious side effects in clinical trials <sup>(10, 12)</sup>.

HCV is a (+) strand RNA virus having a genome size that is approximately 9.6 kb long and contains a single long open-reading frame encoding a polyprotein of approximately 3,000

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Sequences of synthetic oligomeric DNA and template primers, and fluorescence quenching data. This material is available free of charge via the Internet at http://pubs.acs.org.

amino acids. The structural proteins are located at the N-terminal portion, followed by nonstructural proteins. The genomic HCV (+) strand RNA is first copied into (–) strand HCV RNA, which is then used as the template to produce a large number of progeny (+) strand RNA. One of the nonstructural proteins is HCV replicase (NS5B), which contains RNA-dependent RNA polymerase activity (RdRp) and is responsible for replicating the viral RNA <sup>(13, 14)</sup>. NS5B is a 68-kDa protein that can carry out RNA synthesis on primed RNA template, as well as de novo synthesis of viral RNA in the absence of primer. However, the detailed mechanism of RdRp and de novo initiation steps remains unclear.

Many crystal structures of HCV NS5B have been reported (15-20). Like 3-dimensional structures of other polymerases, HCV replicase is folded into fingers, palm, and thumb domains resembling a right hand structure (16, 17). Unlike DNA polymerases with an open polymerase cleft, HCV NS5B is crystallized in closed conformation with a Δ1 finger loop anchored into the hydrophobic pocket on the thumb, thus closing the polymerase cleft (21). A similar closed conformation has been reported for other RNA replicases (22). The closed conformation assumes open conformation during the formation of active polymerase complexes (23). Crystal structure of HCV NS5B from the genotype 2a strain has been reported to exist in open and closed conformation, which is proposed to represent, respectively, inactive and active forms of the enzyme (20). Since both conformations lack bound RNA template or NTP substrate, it is yet to be determined whether binding of RNA template or substrate will induce similar structural change. HCV replicase in solution has been shown to exist in both monomeric and oligomeric forms (24–29). Using real-time ultracentrifugation analyses, it has been shown that HCV NS5B is predominantly monomeric in solution and can be induced into oligomeric form by short RNA templates <sup>(27)</sup>. It has been suggested that the oligomeric form of HCV NS5B is critical for catalyzing RdRp activity (24, 25, 28, 30, 31). Moreover, Bressanelli et al. suggested that the closed structure of NS5B represents a "preformed active site" in the absence of template (18).

Studies have proposed that HCVNS5B undergoes a transition to an open structure to accept the template RNA (20, 32–34). This contention is supported by experiments, in which a circular template has been shown to be used by HCV NS5B, suggesting that the "closed" polymerase cleft of the enzyme may open up to accommodate the circular template (23). Bressanelli et al. have co-crystallized NS5B with GTP, showing that GTP interacts with four residues in the thumb subdomain and two residues on the  $\Delta 1$  finger tips (18). Binding of GTP to the enzyme also been proposed to stabilized the interaction of  $\Delta 1$  finger tips and thumb domain, thus favoring monomeric conformation. The interacting sites on the thumb and fingertips are both on the surface of the enzyme molecule, located 30Å away from the polymerase active site residues Asp318, Asp319 and Asp220. Since NS5B may undergo significant conformational changes during the transition from initiation to elongation, it is expected to exist in multiple conformations. In this study, we used fluorosulfonyl benzoyl adenosine as an NTP analog for affinity labeling of HCV NS5B to map the NTP binding site(s) on the enzyme molecule. This analog has been used successfully to probe the nucleotide binding site(s) in polymerases and kinases (35-43). The structure and size of FSBA are such that it resembles ATP, and its reactive sulfonyl fluoride moiety may reside in the positions of  $\beta$  and  $\gamma$  phosphate of NTP (Fig. 1). The reactive sulfonyl fluoride group of FSBA reacts with the nucleophilic side chain of amino acid residues in the vicinity of the nucleotide binding domain (NBD) of the enzyme.

#### MATERIALS AND METHODS

IDA-Sepharose for immobilized metal affinity chromatography (IMAC) was obtained from Pharmacia. HPLC purified ultrapure NTPs were purchased from Roche Applied Sciences. FSBA and synthetic template-primers were from Sigma-Aldrich. Mass spectrometry grade

trypsin was obtained from Promega. The [ $^3$ H] labeled FSBA was prepared as described ( $^{(44)}$ ) and freshly dissolved in DMSO before use. A site-directed mutagenesis kit (QuickChange) was purchased from Stratagene. All reagents, purchased from Fisher, Millipore and Bio-Rad, were of the highest available purity. Fluorescence quenching was done on a Varian Cary Eclipse fluorescence spectrophotometer. GraphPad Prism 4.0 (GraphPad software) was applied for plotting the data using non-linear fit to the data using hyperbolic function. The pET21C-5B $\Delta$ 21 plasmid expressing His-tagged HCV NS5B with 21 amino acid deletion from the C terminal was a gift from Dr. Sergey Kochetkov ( $^{(45)}$ ).

#### Expression and purification of HCV NS5B and in-vitro mutagenesis

The pET21C-5BΔ21 expression clone of HCV NS5B was expressed in Escherichia coli Rosetta (DE3), as described <sup>(45)</sup>. In brief, the transformed cells were grown with vigorous shaking at 37°C in Luria Broth containing 100 mg/L ampicillin and 15 mg/L chloramphenicol to a density of 0.4 at 595 nm. NS5B expression was induced by the addition of 1 mM IPTG. The cells were further incubated at 25°C with vigorous shaking for 4 h. The cells were harvested, and lysed; His-tagged HCV NS5B was purified on an IMAC column. The eluted fractions containing small impurities were further purified on FPLC (Pharmacia) using a prepacked 1 ml Hi-Trap Heparin column (Pharmacia). In brief, the IMAC fractions were pooled and diluted 3-fold with buffer A (100 mM Tris HCl (pH 7.0), 5% glycerol, 0.5% triton X-100, and 1 mM β-mercaptoethanol) and loaded onto the Hi-Trap Heparin column. After washing the column with 2 column volumes of buffer A, the enzyme was eluted with a linear gradient (0% to 80%) of 1 M KCl in the same buffer in 20 min (1 ml/min). Eluted fractions showing more that 98% purity on SDS-PAGE (8%) were collected and stored at -80°C in a buffer containing 50 mM Tris HCl (pH 7.8), 1 mM DTT, 100 mM NaCl, and 50% glycerol. Site-directed mutagenesis was carried out using the QuickChange mutagenesis kit according to the manufacturer's protocol. Mutations were confirmed by sequencing at the UMDNJ Molecular Resource Facility.

#### Fluorescence quenching assay

The fluorescence-quenching (FQ)-based binding assay was done in a final volume of  $100~\mu l$  per well in a 96-well plate (Fluotrac 200). HCV NS5B (1  $\mu M$ ) was incubated in the absence and presence of FSBA in a buffer containing 50 mM Tris HCl (pH 7.8), 50 mM NaCl, 10% glycerol, 10% DMSO, and 0.5 mM MnCl $_2$ . The mixture was incubated at room temperature for 20 min. The fluorescence emission of enzyme control or enzyme-ligand complex was monitored at 300 to 420 nm with an excitation wavelength of 280 nm. The emission fluorescence intensity of HCV NS5B in the absence and presence of FSBA had the same fluorescence emission maxima, at 330 nm, suggesting that enzyme bound with FSBA did not shift the emission wavelength.

#### RdRp assay

This assay was done at 37°C in a in a total volume of 25  $\mu$ l containing 20 mM Tris HCl (pH 7.8), 40 mM NaCl, 40 mM sodium glutamate, 0.5 mM DTT, 0.01% BSA, 0.01% tween-20, 5% glycerol, 20 U/ml RNase Out, 25 $\mu$ M cold UTP, 1 $\mu$ Ci/assay <sup>3</sup>H UTP or  $\alpha$ <sup>32</sup>P UTP, 500 nM PolyrA.dT<sub>18</sub> and 0.5 mM MnCl<sub>2</sub>. The reaction was started by the addition of 588 nM HCV NS5B and terminated by the addition of 5% ice cold trichloroacetic acid. The terminated reaction mixture was kept on ice for 30 min, after which acid-insoluble material was filtered on glass fiber filters (GF/B). The filters were washed with 5% TCA, water, and ethanol. Filters were dried under infrared lamp and placed in vials containing 5 ml EcoLite scintillation fluid. Radioactivity was counted on a Packard 2200-CA Tri-Carb scintillation counter.

#### Inactivation of NS5B by FSBA

HCV NS5B ( $3\mu M$ ) was preincubated with varying concentrations of FSBA in a buffer containing 20 mM Tris HCl (pH 7.8), 40 mM NaCl, 40 mM sodium glutamate, 0.01% tween-20, 5% glycerol and 10% DMSO in a final volume of 100 $\mu$ l. An aliquot of the incubation mixture was withdrawn periodically and assayed for RdRp activity.

#### Stoichiometric of FSBA binding to NS5B

For determining the binding stoichiometry, we used [ $^3H$ ]-labeled FSBA. We incubated 4  $\mu$ M of NS5B in the presence of varying concentrations of [ $^3H$ ]-FSBA (0.25 mM-1 mM) as above in the incubation buffer in a total volume of 500  $\mu$ l. Aliquots were withdrawn periodically and spotted on Whatman chromatography paper strip #3 and subjected to ascending chromatography using methanol: chloroform (1:1) as the mobile phase. Free [ $^3H$ ]-FSBA was migrated near the solvent front whereas [ $^3H$ ]-SBA bound to NS5B remained at the origin. After one hour, the chromatography strip were removed, air dried and  $^3H$  radioactivity remained at the point of application was determined by liquid scintillation counting. A sample containing the standard reaction mixture minus the enzyme was also incubated similarly and the enzyme was added to this mixture just before to spotting on the chromatography paper strip. The  $^3H$  count obtained at the origin was used as the background.

# 5'-32P labeling of oligomeric primer or template primer

37-mer double stranded RNA looped template primer and  $dT_{18}$  primer were labeled at 5′ using [ $\alpha$ - $^{32}$ P]ATP and T4 polynucleotide kinase (Invitrogen) according to manufacturers protocol. The 5′ labeled oligomers were purified by illustra ProbeQuant G-50 micro columns.

#### Photocrosslinking of double stranded template primers to HCV NS5B

Two micro molar of HCV NS5B or its mutant derivatives were incubated on ice with 20 nM of 5'- $^{32}$ P labeled 37-mer double stranded looped RNA template primer (Chart S1) or poly rA.5'- $^{32}$ P-dT<sub>18</sub> (20K Cerenkov CPM) in a crosslinking buffer containing 20 mM Tris HCl (pH 7.8), 40 mM NaCl, 40 mM sodium glutamate, 2 mM DTT and 0.5 mM MnCl<sub>2</sub> in a final volume of 50  $\mu$ l. After 20 min incubation, samples were irradiated by UV light at 254 nm, with a dose rate of 3600 J/m² (Spectrolinker XL-1000) at a distance of 10.5 cm. The irradiated samples were mixed with gel loading dye and resolved by SDS-PAGE using 8% polyacrylamide gel. The crosslinked enzyme-TP complexes were visualized by phosphorImager and gels were stained using Coomassie blue staining.

# Photocrosslinking of [ $\alpha$ - $^{32}$ P]-ATP with HCV NS5B and its mutant derivatives in the binary and ternary complex

The wild type HCV NS5B or its mutant derivatives (7 $\mu$ g) was incubated with 20  $\mu$ Ci [ $\alpha$ - $^{32}$ P]-ATP in the crosslinking buffer in the absence or presence of 2  $\mu$ M of pre-annealed rU<sub>15</sub>.rA<sub>10</sub> template primer with 3′-deoxy terminated primer terminus. After 20 min incubation on ice, the mixture was UV irradiated as above and the crosslinked complexes were resolved by SDS PAGE and visualized by autoradiography.

# Nucleotidyl transferase activity of the enzyme crosslinked with double stranded RNA template primer

Two micro molar of HCV NS5B was incubated with 2  $\mu$ M of unlabeled 37-mer double stranded RNA template primer (Chart S1) by UV irradiation as described above. An aliquot of the crosslinked enzyme-TP binary complex was incubated with 10  $\mu$ Ci  $\alpha$ - $^{32}$ P UTP in a reaction mixture containing 20 mM Tris HCl (pH 7.8), 0.5 mM DTT and viable salt

concentration ranging from 50 mM to 1 M in a final volume of 50  $\mu$ l. The mixture was incubated at 37°C for 30 min and the incorporation of  $^{32}P$  labeled nucleotide onto the immobilized primer terminus was monitored by visualizing the labeled enzyme-TP covalent complex by resolving the complex on 8% SDS-PAGE followed by PhosphorImaging.

# Preparative scale modification of NS5B by [<sup>3</sup>H]-FSBA and tryptic digestion of modified protein

Modification of HCV NS5B with FSBA was carried out incubating 15 nmol of the enzyme with 1mM [³H]-FSBA or unlabeled FSBA at 37°C in the incubation mixture in a final volume of 1 ml, as described above. After 1 hr of incubation, 10 mM DTT was added to stop the reaction. For HPLC analysis, the modified protein was precipitated by adding TCA (10% final) and washed twice with 0.5 ml acetone to remove residual TCA. The protein pallet was first dissolved in 8M urea in 100 mM sodium bicarbonate buffer, pH 8.0, then diluted to 2M urea in the same buffer. Trypsinization was done at 37°C at a trypsin-to-HCV NS5B ratio of 1:50. The trypsinized protein was adjusted to 0.1% TFA and resolved on HPLC. HCV NS5B without FSBA treatment was processed and trypsinized in a manner similar to that for HPLC analysis and considered as the control for MALDI/MS analysis, the FSBA-modified enzyme and unmodified control enzyme were resolved on 8% SDS-PAGE. The gel was stained with 0.1% Coomassie brilliant blue R250 in 10% acetic acid, 50% methanol, and 40% H<sub>2</sub>O for 10 min. The gel was destained overnight in the same solution except the dye. The protein bands were excised and processed for MALDI/MS analysis.

### **HPLC** purification of tryptic peptides

Tryptic peptides were resolved on C18 reverse-phase HPLC column (Phenomenex Kinetex XB). The photodiode array detector was set to monitor the eluants at 214 and 254 nm. Elution was programmed by a linear gradient of solvent A: ( $H_2O$ , 0.1% TFA) into solvent B: (acetonitrile) over 10 min at 0–1% solvent B. Then a linear gradient was run from 1%–50% solvent B for 40 min and 50%–100% solvent B for 30 min with a flow rate of 1.5 ml/min. The sample collector was set to collect 1.5 ml fractions. An aliquot of each fraction was counted for  $^3H$  radioactivity in a liquid scintillation counter. The peptide peaks with significant radioactivity were further purified on HPLC C4 reverse-phase column (Delta Pak, 300 Å, 3.9 mm × 15 cm) and subjected to LC/MS/MS analysis.

#### MS analysis of HCV NS5B covalently modified with FSBA

For MALDI/MS analysis, the excised gel bands were dried in Speedvac and *in-situ* trypsin digested at 37 °C for 16 h. MS analyses were acquired using the MALDI-TOF/TOF technique on an Applied Biosystems 4800 MALDI-TOF/TOF mass spectrometer. Matrix solution contained 3.0 mg/ml α-cyano-4-hydroxycinnamic acid as a matrix. The spectra were internally calibrated using bradykinin and ACTH (18–39, human (CLIP) as an internal standard; protonated, monoisotopic masses of 1060.569 and 2465.199Da, respectively. Comparative MALDI analysis was done on a peptide fingerprint map (PFM) over the 600–4,500 Da mass range in reflector positive mode, summing 2,000 laser shots. The spectra of control and modified samples were compared to determine likely candidates for SBA-modified peptides. The complete protein sequence was entered into an in-house database to permit matching of the peptide. Modification of peptides by SBA (sulfonylbenzoyl adenosine; +433.08 Da) and the product of ester hydrolysis of FSBA, sulfonylbenzoic acid (SB; +183.983 Da), were entered as user-defined variable modifications.

LC-MS/MS analyses of HPLC-purified peptides were acquired on a Thermo Scientific LTQ Orbitrap equipped with a Waters Nanoacquity UPLC system using a Waters Symmetry C18 180  $\mu$ m  $\times$  20 mm trap column and a 1.7  $\mu$ m, 75  $\mu$ m  $\times$  250 mm nanoacquity UPLC column (35°C) for peptide separation. Trapping was done at 15  $\mu$ l/min in 99% buffer A (water,

0.1% formic acid) for 1 min. Peptide separation was done in a two-step gradient of buffer B (CH<sub>3</sub>CN, 0.075% formic acid) to buffer A at a flow rate of 300 nl/min. The elution was programmed with 5% buffer B, 50% buffer B at 50 min, and 85% buffer B at 51 min. MS was acquired in the Orbitrap using 1 microscan over the range of 400–2,000 m/z at 30,000 resolution and a maximum inject time of 900 ms, followed by six data-dependent MS/MS acquisitions in the ion trap. MS/MS spectra were searched in-house using the Mascot algorithm for uninterpreted MS/MS spectra after using the Mascot Distiller program to generate Mascot compatible files  $^{(46)}$ . The Mascot Distiller program combines sequential MS/MS scans from profile data that have the same precursor ion. A charge state of +2 and +3 are preferentially located with a signal-to-noise ratio of 1.2 or greater; a peak list is generated for database searching. The data was searched against a sequence- specific database using a peptide tolerance of  $\pm$  15 ppm, MS/MS fragment tolerance of  $\pm$  0.6 Da, and peptide charges of +2, +3, or +4. The modifications by SBA (+433.069 Da) and SB (+183.983 Da) were investigated.

#### Molecular docking

We used induced-fit docking (IFD) workflow of Schrodinger Suit (Schrodinger LLC, New York, NY) for this purpose. The crystal structure of NS5B bound to GTP (PDB entry 1GX5) at the allosteric site was used for docking FSBA to NS5B. The crystal structure was prepared using the protein preparation workflow in the Schrödinger Maestro modeling package version 9.2. The GTP, catalytic metals (Mn<sup>2+</sup>), and tri-phosphate moieties were removed from the crystal structure during protein preparation workflow. The structure of FSBA to be docked into the crystal structure was generated as follows. The initial structure of FSBA was obtained from PubChem (pubchem.ncbi.nlm.mih.gov) entry number 133129 in sdf format. The 3D conformations of FSBA from this structure were generated by the Epic and LigPrep ligand preparation protocols of the Schrodinger Suit incorporated in Maestro 9.2 molecular modeling environment. Two separate Induced-Fit docking runs were carried out. In first run, the docking was done using the centroid out at the centroid of K491 residues; in the second run, the centroid of Y382 atoms was considered. The box size was 50 ų in each case.

#### **RESULTS**

#### Binding affinity of FBSA to HCV NS5B

FSBA, a chemically reactive analog of ATP, has affinity for enzymes that use ATP as a substrate. Therefore, by measuring fluorescence quenching (FQ) of the enzyme in the presence and absence of FSBA, we tested whether FSBA can bind to HCV NS5B. The binding of NTPs to HCV NS5B significantly reduces the fluorescence emission intensities of the enzyme at 330 nm when excited at 280 nm  $^{(47,\,48)}$ . We therefore incubated a fixed concentration of HCV NS5B with varying concentrations of FSBA and measured the fluorescence emission between 300–420 nm. The emission fluorescence maximum was noted at 330 nm which was significantly quenched in the presence of FSBA (Fig. S1A). As shown, the increase in FQ was correlated with and proportional to the increase in FSBA concentration. At as low a level as 62  $\mu$ M of FBSA, the fluorescence emission was reduced by 12%, while at 1 mM concentration of the analog, 70% reduction in the emission intensity was noted (Fig. S1B). We could not determine the binding affinity of this analog since it is chemically reactive and forms covalent bond with amino acid residue at its binding site.

#### Inactivation of RdRp activity of HCB NS5B by FBSA

We noted that FSBA, being an ATP analog, rapidly binds to the enzyme and quenches its emission fluorescence, as does NTP substrate <sup>(48)</sup>. To examine the effect of FSBA on HCV NS5B activity, we measured the RdRp activity of the enzyme in the presence of increasing

concentrations of FSBA (Fig. 2A). We observed that FSBA binding to the enzyme inactivated its RdRp activity. Approximately 70% inactivation of RdRp activity was noted at 0.6 mM concentration of FSBA. At higher concentration of the analog (1 mM), the inactivation was only marginally increased probably due to its decreased solubility of the analog beyond 1 mM concentration. We then examined the inactivation pattern of the enzyme as a function of both time of incubation and FSBA concentration. We incubated HCV NS5B with 0.25 mM, 0.5 mM, and 1 mM FSBA at 4 °C and, at 5, 10, 20, 40, and 60 min, withdrew an aliquot of sample and assayed it for RdRp activity (Fig. 2B). We noted 15% inactivation of the RdRp activity of the enzyme when it was incubated with 0.25 mM FSBA for 20 min, but 20%–30% after 40 min of incubation. At 1 mM concentration of FSBA, enzyme activity was reduced approximately 70% within 10 min of incubation; 80% reduction occurred after 60 min of incubation.

### Binding stoichiometry of FBSA to HCV NS5B

We noted that FSBA being an ATP analog inactivates the RdRp activity of NS5B and expected to bind with enzyme in an irreversible manner. Covalent attachment of the [<sup>3</sup>H]-labeled FSBA to the enzyme was monitored as a function of time and concentration of FSBA (Fig. 2C). The covalent attachment of FSBA to the enzyme was found to vary with time and was dependent on the FSBA concentration. At 1 mM concentration of FSBA, the binding stoichiometry was approximately 2 moles per mole of enzyme after 90 min incubation.

#### Protection of HCV NS5B from FSBA in the presence of rNTPs and different templateprimers

To determine whether inactivation of HCV NS5B by FSBA can be protected in the presence of nucleotides, or template primer, we preincubated HCV NS5B in the presence of individual NTPs (ATP, GTP, CTP, and UTP), or with template-primer and then supplemented with 0.5 mM FSBA. The reaction mixture was incubated at 4°C for 1 h and an aliquot of the incubation mixture was assayed for RdRp activity (Table 1). We observed that in the presence of rNTP, poly rA.dT<sub>18</sub>, 37-mer looped RNA or poly rA.rU<sub>12</sub> template primer, the RdRp activity of the enzyme is protected from FSBA, but similar protection was not observed in the presence of poly rU.rA<sub>26</sub>. The protection of enzyme activity in the presence of poly rA.dT<sub>18</sub>, 37-mer looped RNA and poly rA.rU<sub>12</sub> template primers can be explained since incoming nucleotide with these template primers is UTP; therefore binding of the ATP analog is expected to be excluded. In the presence of poly rU.rA<sub>26</sub>, enzyme activity was not protected from FSBA because incoming nucleotide with this template primer is ATP.

We further observed that the protection of the enzyme activity from FSBA in the presence of TP is significantly reduced at 150 mM salt concentration but salt had no effect on the protection in the presence of rNTP substrate. This may be possible that binding of double stranded nucleic acid substrate to HCV NS5B requires low ionic strength. To examine this possibility we incubated the enzyme with 5′-3²P labeled 37-mer double stranded looped RNA template primer as well as with ploy rA.5′-3²P-dT<sub>18</sub> in the presence of increasing salt concentrations. The incubation mixture was UV irradiated and the resulting 3²P labeled E-TP complexes were resolved by SDS-PAGE. The results shown in Fig. 3 indicate that HCV NS5B efficiently binds to heteromeric 37-mer dsRNA looped TP (Fig. 3A) and homopolymeric double stranded nucleic acid substrate (Fig. 3B) at 50 mM salt concentration. The formation of enzyme-TP binary complex was reduced to 50% and 75% at 100 mM and 150 mM salt, respectively, and nearly abolished at or above 200 mM salt concentration. These results suggest that template primer binding to the enzyme is drastically reduced even at physiological salt concentration and therefore offers no

protection to the enzyme from FSBA mediated inactivation. This could be due to low electropositive potential in the template primer binding track of the enzyme. Lohman et al., have reported 80% reduction in RdRp activity of HCV NS5B at 100 mM KCl and 50% reduction at 150 mM NaCl concentration. This reduction in the activity could be due to low binding of double stranded nucleic acid substrates used in their assay system <sup>(49)</sup>.

We further noted that high salt concentrations have no effect on the protection of enzyme activity in the presence of rNTP substrates. This could be due to hydrophobic nature of rNTP binding pocket which may be insensitive to high salt concentration. To examine this hypothesis, we incubated HCV NS5B with unlabeled 37-mer looped RNA TP and UV irradiated to covalently crosslink with the enzyme. An aliquot of the enzyme-TP binary complex was incubated with first incoming nucleotide, α<sup>32</sup>P UTP, at different salt concentrations ranging from 50 mM to 1 M in the RdRp reaction buffer. The reaction mixture was incubated at 37°C for 30 min and then resolved by SDS-PAGE. The results shown in Fig. 4 indicate that enzyme-TP binary complex was able to incorporate a single nucleotide on to the immobilized primer terminus of the double stranded TP even at 1 M salt concentration suggesting salt resistant hydrophobic nature of the nucleotide binding pocket.

#### Comparison of tryptic peptides and LC/MS/MS analysis of affinity-labeled peptides

To identify the binding site of FSBA, we did tryptic peptide mapping of the [3H]-FSBAmodified enzyme, using HPLC. We used unmodified enzyme as the control. The tryptic digests of the modified and unmodified enzyme were separated by HPLC using a C<sub>18</sub> reverse-phase column. The eluants were monitored at 220 nm and 254 nm to detect the concurrence of SBA-labeled peptides; the peptide labeled with SBA moiety was expected to give relatively high absorbance at 254 nM. Since HCV NS5B was modified by [3H]-FSBA, we also monitored the eluants for radioactivity. The results shown in Fig. S2B indicate the absorbance of tryptic peptides monitored at 254 nm to detect SBA linked peptides. As shown, three distinct radioactive peaks were eluting at Rt 20.3, 23.6, and 36.8 min. The radioactive peak eluting at Rt 20.3 was unreacted free FSBA since free FSBA also eluted at the same position. The radioactive peaks eluted at 23 min and 36 min were found to be associated with peptides (Fig. S2B). Both of these peptides were conspicuously absent in the control without modification (Fig. S2A). We also analyzed tryptic peptides of the enzyme modified by FSBA in the presence of ATP (Fig. S2C). We noted that both of the radioactive peptide peaks eluting at 23 and 36 min were significantly reduced, suggesting that in the presence of ATP the modification of enzyme by FSBA is significantly protected. We further purified these peptides and subjected them to LC/MS/MS analysis to identify the peptides and site of modification. The LC/MS/MS analysis showed that the peptide eluted at 23 min spanned residues 33–43 in the primary sequence of HCV NS5B, with Y38 as the modification site. The peptide peak eluted at 36 min was found to contain two peptides spanning residues 381-386 and 491-498, with Y382 and K491 as the respective modification sites (Table 2). A representative MS/MS fragmentation pattern of peptide eluting at 36 min is shown in Fig. 5.

#### Characterization of the FSBA labeling site using MALDI-TOF/TOF

To confirm the FSBA modification site in HCV NS5B, we used highly sophisticated MALDI-TOF/TOF analysis of the tryptic digest of FSBA-modified HCV NS5B. Both FSBA-modified enzyme and unmodified control were subjected to SDS PAGE. The protein bands were excised from gels and processed for *in situ* trypsin digestion. The digested samples were then analyzed on MALDI-TOF/TOF to acquire the MS data. The MS data of the FSBA-modified sample was compared with the unmodified control (Fig. 6). Microscopic analysis of the MS spectra showed two major peptides with masses of 998.43, 1311.63, and a minor peptide of 1,500.60 Dalton in the FSBA-modified sample which were absent in the

control (Fig. 6). Further MS analysis of two major peptides showed that the 998.43-Da peptide corresponded to residues 381–386 and contained Y382 as modification site while 1311.63 Da peptide spans residues 491-498 with K491 as the site of modification. The minor peptide of 1500.60 Da spans residue 33-43 with Y38 as the modification site (Table 2). The peptide mass of 1311.63 Da represents an additional mass of 433.08 Da due to sulfonylbenzoyl adenosine (SBA) attached with the peptide, while the peptides having masses of 998.43 Da and 1,500.60 Da represent additional mass of 183.98 Da due to the attachment of sulfonylbenzoyl moiety (SB) of SBA <sup>(43)</sup>. This might be due to ester hydrolysis of the SBA moiety during sample preparation, in which only the SB moiety remains intact, with the modified residue showing a mass increase of 183.98 Da. In MALDI-MS experiments, we found the coverage of 89% of the primary sequence of HCV NS5B obtained by MS analysis (Fig. 7).

#### Mutational analysis of the identified residues

Since all three identified sites are protected from modification from FSBA in the presence of ATP, it is likely that they are part of the NTP channel or NTP binding pocket of the enzyme. We therefore carried out site-directed mutagenesis to generate Y38A, Y382A, and K491A mutants of the enzyme. We purified the mutant enzymes and examined their RdRp activity. As shown in Fig. 8A, Y382A and K491A had severely impaired polymerase activity, while Y38A remained unaffected. These results indicate that Y382 and K491 are important residues for catalysis, while Y38 is dispensable. K491 is located in the NTP channel and substitution of the Lys side chain with Ala may perturb the NTP channel  $^{(50)}$ . The Y382 is located on the  $\beta$ 16 sheet at the interface of finger and thumb, very close to the suggested primer grip region corresponding to  $\beta$ 14- $\beta$ 15  $^{(15)}$ .

#### Effect of Y382A and K491A mutations on the binding of template-primer and ATP substrate

The reduction in RdRp activity of Y382A and K491A mutants could be due to defect in either template primer binding step or at the nucleotide binding step. In order to determine the template primer binding ability of the mutant enzymes, we used a double stranded 5'-32P labeled 37-mer self annealing RNA template primer (Chart S1) to photo-crosslink with the wild type and its mutant derivatives. The results shown in Fig. 8B indicated no significant change in the template primer binding ability of both Y382A and K491A mutant enzymes as compared to the wild type enzyme. We then examined their ability to crosslink ATP substrate in the binary (enzyme-ATP) and ternary (enzyme-TP-ATP) complexes. The template primer that we used had 3' deoxy terminated rA<sub>10</sub> primer annealed with rU15 template (rU<sub>15</sub>/rA<sub>10-3'deoxy</sub>) which supports ATP as the incoming nucleotide. The results shown in Fig. 8C indicated that both the mutant enzymes were significantly affected in their ability to crosslink ATP in the binary and ternary complexes. We further found that extent of reduction in the ATP crosslinking ability of K491A mutant was similar either in the binary and ternary complexes (lanes 3 and 6) whereas reduction in the crosslinking was more pronounced with Y382A mutant in the ternary complex. The Y382 and K491 may be part of multiple non-covalent interactions/contact points between the ATP and the enzyme molecule. The reduction in the extent of crosslinking of ATP to Y382A and K491A mutants may be due to removal of one of the multiple contact points with ATP in the nucleotide binding pocket.

#### Molecular docking of FSBA at the nucleotide binding site of HCV NS5B

The most favorable poses (i.e., the least energy-docking conformer) for two Induced-Fit docking (IFD) workflows were used for analyses of the binding of FSBA to NS5B (Fig. 9A, C). The first IFD complex obtained after docking of FSBA at the centroid of K491 is shown in Fig. 9A, displaying specific interactions of FSBA with neighboring amino acid residues. The polar (H-bond) interactions are shown in cyan-dotted lines, while the van der Waals

interactions are shown as orange double-dotted lines; the positions of active-site residues (D318, D319, and D220) are included as references. The nitrogen of epsilon amino group of K491 side chain is within interacting distance of 3.4 Å. From the fluorosulfonyl group of FSBA. The other important residues that interact in this modeled complex are F145, K155, S367, R386, D387, and T390. Bressanelli et al. showed that R158, S367, R386, and T390 interact with one of the tri-phosphates in the crystal structure of NS5B <sup>(18)</sup>. The position of some of these interacting residues can be clearly seen in the surface view of the putative NTP channel (Fig. 9B). These residues have been proposed to be part of priming site <sup>(18)</sup>. The initiation site residues R48, K51, K151, K155 and R158 have been suggested to be the part of NTP-tunnel <sup>(18)</sup>. In our complex, residues K155 interact with FSBA. Therefore, this complex can be considered to represent a combination of priming-site and initiation-site complexes described previously.

The second IFD complex, shown in Fig. 9C, was obtained by docking FSBA near Y382. In this complex, the FSBA appears to be stabilized by polar (H-bond), van der Waals, and hydrophobic interactions. The distance between the oxygen atom of Y382 and the fluorosulfonyl group of FSBA is 3.7 Å. Amino acid residues H467, S470, D375, K379, and K209 form H-bond interactions with various moieties of FSBA. G198, E202, and V205 interact through van der Waals interactions. The ring moieties of FSBA are stabilized by hydrophobic interactions with Y383, Y382, and V381. The Y382 located on  $\beta16$  in the thumb domain is in the vicinity of the primer grip region ( $\beta14-\beta15$ ) on the palm domain. A surface view of this region is shown in Fig. 9D. A crystal structure of the ternary complex showing interaction of NTP with these residues is yet to be elucidated.

#### DISCUSSION

Affinity labeling of enzyme with substrate analogs is frequently used to identify the substrate binding site on the enzyme molecule. FSBA is a unique ATP analog that has been used to probe the ATP-binding region in ATP-using enzymes (35-43). We have used this analog, as well as its deoxy form (FSBdA), to covalently modify interacting amino acid residues in the polymerase cleft of E. coli DNA pol I (36). In our present study, we used FSBA to modify HCV NS5B to identify the nucleotide interacting regions on the enzyme molecule. The electrophillic sulfonyfluoride moiety of FSBA may readily react and form covalent linkage with Ser, Cys, Lys, Tyr, and His residues located within the region interacting with the incoming nucleotide (44, 51, 52). The position of the fluorosulfonybenzoyl moiety of the FSBA molecule is similar to that of the triphosphate moiety of ATP; therefore, that moiety is expected to bind and react with amino acid residues in the vicinity. The reactive sulfonyl fluoride group of enzyme-bound FSBA may form stable covalent adduct with the amino acid residue in the binding site, which could be identified by tryptic peptide profiling followed by mass spectrometry analysis. The relevance of using FSBA to probe the ATP binding site is derived from the fact that it's binding to the HCV replicase is protected in the presence of polyrA.rU<sub>12</sub> but not in the presence of polyrU.rA<sub>26</sub> that supports ATP as the incoming nucleotide. We first confirmed that FSBA binds to HCV NS5B in a concentration dependent manner with a binding stoichiometry of approximately 2 moles of FSBA per mole of enzyme. We further observed that binding of FSBA to HCV NS5B irreversibly inactivates the RdRp activity of the enzyme. Inactivation of the enzyme was readily protected in the presence of NTP, suggesting that reactive sites may be located within the region interacting with NTP substrate. The crystal structure of HCV NS5B soaked with rCTP, rGTP, and rATP in the presence of Mn<sup>2+</sup> displayed density only for the triphosphate moiety of the nucleotide bound to two metal ions (18). This indicated that all the NTPs, specifically their triphosphate moieties share the same binding site on the enzyme molecule. Since the reactive fluorosulfonyl group of FSBA is positioned similar to triphosphate moiety of NTP, its reactivity with the enzyme is protected in the presence of

any of the four NTPs. In contrast, in the presence of template primer, the binding of individual NTPs is guided by the template base and thus offering protection to the enzyme in the presence of template primers that do not support ATP as the incoming nucleotide.

However, protection offered by poly rU.rA<sub>26</sub> template primer was significantly reduced at 150 mM salt concentration suggesting the double stranded template-primer binding to the enzyme may be highly sensitive to salt concentration. It has earlier been proposed that the wild type HCV replicase cannot accommodate double stranded nucleic acid substrate due to close interaction between fingertips and thumb as well as due to protruding beta flap (amino acid residues from 443–454) from the thumb domain into the catalytic cleft (53). We noted that binding of double stranded RNA substrate to the enzyme requires low salt concentration. HCV NS5B efficiently binds to the dsRNA substrates (poly rA.dT<sub>18</sub> and 37mer RNA TP) at 50 mM salt concentration. At or above 100 mM salt concentration, the formation of Enzyme-TP binary complex is drastically reduced (Fig. 3). This could be the reason for reported 80% reduction of RdRp activity with poly rC-oligo dG at 100 mM KCl and 50% reduction at 150 mM NaCl concentrations <sup>(49)</sup>. It is possible that at higher salt concentration, the hydrophobic interaction between finger tips and thumb may be strong enough to keep the cleft in closed conformation and thus not accessible for binding to dsRNA substrate. In contrast, at lower salt concentration, the reduced hydrophobic interaction between fingertips and thumb may allow the polymerase cleft to breath and open up to accommodate dsRNA substrate. Another possibility of salt sensitivity could be low electropositive potential of the RNA binding track responsible for dsRNA binding.

Most interestingly, the addition of a single nucleotide on to the immobilized primer terminus of the crosslinked template primer is highly resistant to high salt concentration. As illustrated in Fig. 4, the addition of  $\alpha^{32}$ P-UTP onto the crosslinked 37-mer dsRNA TP could be seen even at 1 M salt concentration. This observation indicates that binding of dsRNA template primer to HCV NS5B was specifically in the primer extension mode and therefore protected FSBA mediated modification of the enzyme. Since both poly rA.rU<sub>12</sub> and poly rA.dT<sub>18</sub> template primers allow UTP as the incoming nucleotide, the binding of ATP analog is expected to be excluded. In contrast, poly rU.rA<sub>26</sub> did not protect the enzyme from FSBA because incoming nucleotide with this template primer is ATP.

HPLC analysis of the tryptic digest of HCV NS5B modified with FSBA in the absence and presence of ATP demonstrated the presence of three distinct affinity-labeled peptides, which were protected from modification in the presence of ATP. LC/MS/MS analysis of the HPLC-purified affinity-labeled peptide identified the peptides spanning residues 33-43, 381-386, and 491-498, each containing a unique site of modification at Tyr38, Tyr382, and Lys491, respectively. However the MALDI-MS analysis of the tryptic digest of FSBA-modified HCV NS5B, detected two major peptides and a minor peptide that were absent in the tryptic digest of unmodified control. Further MALDI-TOF/TOF analysis of these peptides identified Tyr382 and Lys491 as the major sites and Tyr38 as the minor site of modification.

Since Y38 is located on the surface of the  $\Delta 1\text{-}\alpha A\text{-}\beta 2$  finger loop which closes the polymerase cleft, we believe that the finger loop containing Y38 may undergo repositioning during FSBA entry in to the polymerase cleft and thus transiently exposing Y38 to FSBA. Bressanelli et al. have shown that soaking the HCV NS5B crystal with GTP identified six amino acid residues on the surface that make direct water-mediated contact with the nucleoside moiety of the bound nucleotide <sup>(18)</sup>. The position of Tyr38 on the  $\Delta 1\text{-}\alpha A\text{-}\beta 2$  loop and its interaction with FSBA suggest that it may be part of the surface binding site for the nucleotide and thus may not be an essential residue for the RdRp activity of the enzyme. This was confirmed by our site-directed mutagenesis studies in which Tyr side chain at

position 38 was substituted with Ala; the resulting Y38A mutant was found to display wild-type activity. The Lys491 that we have identified as the reactive site of FSBA is located on the aS helix in close proximity to the active site cavity of the enzyme. Deval et al., have proposed that Lys491 is positioned in the NTP channel <sup>(50)</sup>; its interaction with NTP analog, FSBA, is in agreement with this contention. Lys491 seems to be an important but dispensable residue in the NTP channel since K491A mutant retained 25% of the wild-type activity.

FSBA's major interaction was seen with Tyr382, as judged by MALDI-MS analysis of tryptic peptides of the modified HCV NS5B. This residue is located on  $\beta16$  in the thumb, in the vicinity of the putative primer grip  $\beta14–15$  loop in motif E on the palm domain. In the structure of UTP-bound HCV NS5B, Ser367 in the motif E on the  $\beta14$ - $\beta15$  hairpin and Arg386 at the base of the  $\beta16$ - $\alpha$ O helix loop have been shown to hydrogen bond with  $\beta$ -phosphate; Thr390, on the  $\alpha$ O helix, hydrogen bonds with the  $\gamma$  phosphate of UTP  $^{(18)}$ . Both Arg386 and Tyr382 share the highly conserved motif YYLTRD, which is essential for RdRp activity of the enzyme. Ala substitution at position 386 inactivates RdRp activity of the enzyme, since substitution of Tyr to Ala at this position severely impaired nucleotide binding and the polymerase function of the enzyme (Fig. 8). The alignment of 3-dimensional structure of RNA-dependent RNA polymerases from other RNA viruses, including Norwalk virus, bovine viral diarrhea virus, foot-and-mouth-disease virus, coxsackie virus, and poliovirus showed that structurally, both Tyr38 and Tyr382 are fairly conserved, while Lys491 is only moderately conserved (Fig. 10).

The induced-fit molecular docking of FSBA at the nucleotide binding domain shows that beside Lys491, which is within interacting distance from FSBA, other residues, including Phe145, Arg158, Lys155, Ser367, Arg386, Asp387, and Thr390, also interact in this modeled complex (Fig. 9B). It has been proposed that among these, residues Arg158, Ser367, Arg386, and Thr390 are part of the primer-site (P-site) and interact with the triphosphate moiety of NTP <sup>(18)</sup>. The second induced fit docking near Tyr382 shows His467, Ser470, Asp375, Lys379, and Lys209 forming H-bond interactions with FSBA, while Tyr383, Tyr382 and Val381 appear to stabilize the ring moieties of FSBA by hydrophobic interactions (Fig. 9C) and thus may represent nucleotide binding site in HCV NS5B.

## **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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#### Abbreviations used

**HCV** Hepatitis C virus

SDS-PAGE sodium dodecyl sulfate polyacrylamide gel electrophoresis

**DTT** dithiothreitol

**TP** template primer

dsRNA double stranded RNA

Poly (rA).(dT)<sub>18</sub> polyriboadenylic acid annealed with (oligodeoxythymidylic acid)18

**RdRp** RNA-dependant RNA polymerase **FSBA** 5'-p-fluorosulfonylbenzoyladenosine

**SBA** sulfonylbenzoyladenosine

**IMAC** immobilized metal affinity chromatography

Ni-IDA nickel-iminodiacetic acid

FPLC fast protein liquid chromatography
HPLC high pressure liquid chromatography

LC MS/MS Liquid chromatography coupled with MS/MS

MALDI-TOF matrix assisted laser desorption ionization time-of-flight mass

spectroscopy

BSA Bovine serum albumin

NBD Nucleotide binding domain

**IFD** Induced-Fit docking

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**Figure 1.**(A) Chemical structure of FSBA. (B) Chemical structure of ATP. (C) Reaction scheme of tyrosine modification by FSBA is shown as a representative of FSBA linked to peptide. SBA and SB add masses of 433.08 Da and 183.98 Da, respectively.

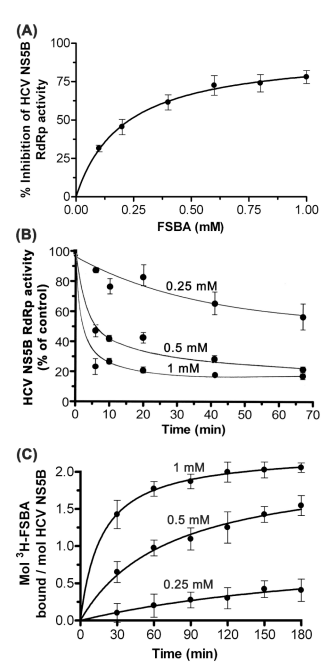


Figure 2. Inactivation of RdRp activity of HCV NS5B by FSBA

(A) RdRp activity assay of HCV NS5B was done in the presence of increasing concentrations of FSBA. The reaction was carried out for 60 min and percent of inactivation of RdRp activity was determined with respect to control. (B) HCV NS5B was pre-incubated with indicated concentration of FSBA at 4 °C. At indicted time points the samples were withdrawn from the incubation mixture and assayed for RdRp activity. (C) Binding stoichiometry of FSBA to NS5B as a function of FSBA concentration and time of incubation. Four micro molar NS5B was incubated at 30°C in the presence of indicated amount of [<sup>3</sup>H]-FSBA in a final volume of 500 µl. At indicated time point, aliquots were withdrawn and subjected to ascending paper chromatography followed by quantification of the radio labeled protein using liquid scintillation counting as described in the Materials and

Methods. The GraphPad PRIZM software was applied to plot the data using nonlinear fit to the data using hyperbolic function equation (Y=Bmax1\*X/(Kd1+X)) where Bmax is the maximal inactivation and Kd is the concentration of ligand required to reach half maximal inactivation or binding of ligand to the enzyme.

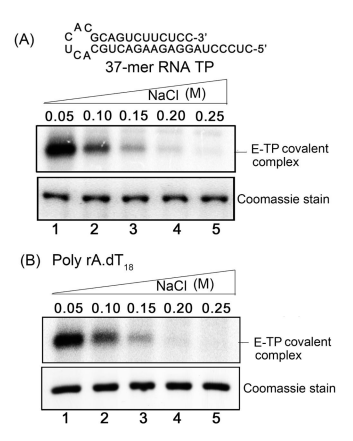


Figure 3. Effect of salt concentration on the binding of double stranded RNA TP to HCV NS5B HCV NS5B was incubated with either  $5^{\prime}$ <sup>32</sup>P labeled 37-mer self-annealing RNA template primer (A) or with poly rA.5 $^{\prime}$ <sup>32</sup>P-dT<sub>18</sub> (B) in the presence of indicated salt concentration on ice for 20 min and then UV irradiated as described in the Materials and Methods. The enzyme-TP binary covalent complexes were resolved by SDS-PAGE. The E-TP covalent complex was visualized by autoradiography and protein band was stained by Coomassie blue.

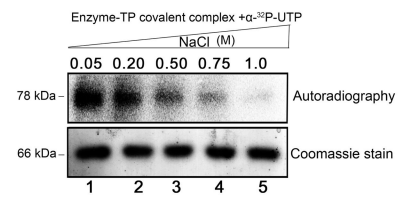
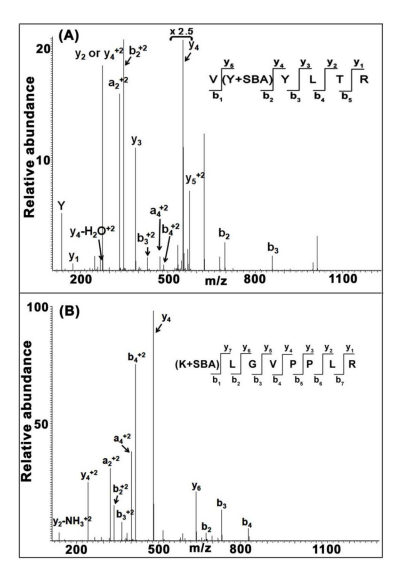


Figure 4. Effect of salt concentration on the incorporation of a single nucleotide onto the immobilized primer terminus of Enzyme-TP covalent complex  ${\bf E}$ 

The unlabeled 37-mer self-annealing RNA TP was covalently crosslinked with the enzyme by UV irradiation. An aliquot of the E-TP covalent complex was incubated with  $\alpha$ - $^{32}$ P UTP in the presence of indicated salt concentration as described in the Materials and Methods. After 30 min incubation at 37°C, the reaction mixture was subjected to 8% SDS-PAGE. The incorporation of a single nucleotide on to the immobilized primer terminus of E-TP covalent complex was visualized by autoradiography and protein band by Coomassie blue staining.



**Figure 5. Tandem mass spectrometry of FSBA modified peptides**(A) The representative MS/MS spectrum of triply charged ion at m/z 1246.50 shows the peptide <sup>381</sup>VYYLTR<sup>386</sup> modified by SBA at Tyr382. (B) The MS/MS spectrum of the triply charged ion at m/z 1311.63 shows the peptide <sup>491</sup>KLGVPPLR<sup>498</sup> modified by SBA at Lys491. Fragmentation pattern of the peptides are shown in the inset.

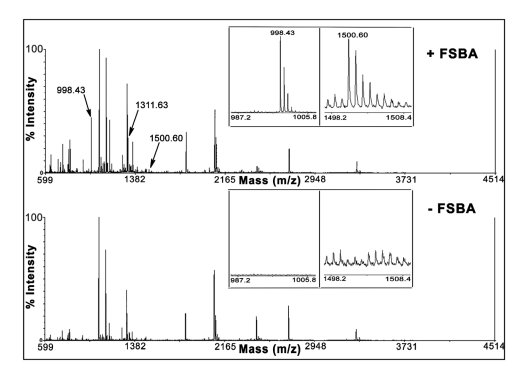


Figure 6. Identification of the FSBA binding sites using MALDI-TOF MS HCV NS5B (5  $\mu$ M) was incubated in the presence and absence of 1 mM FSBA at 37 °C for 1 h in a total volume of 100  $\mu$ l. After labeling, samples were resolved on 8% SDS-PAGE. The corresponding protein bands were visualized by mild staining. The protein bands were excised and in-gel trypsin digested and analyzed by MALDI-TOF. Comparison of full-scan MS data indicates the modification of FSBA-treated sample at m/z 998.43, 1311.63, and 1500.60 (indicated by arrow). The inset shows the zoomed modification at m/z 998.43 and 1550.60 as a representative of all three modifications identified. "+ and – FSBA" denotes for NS5B treated in presence and absence of FSBA.

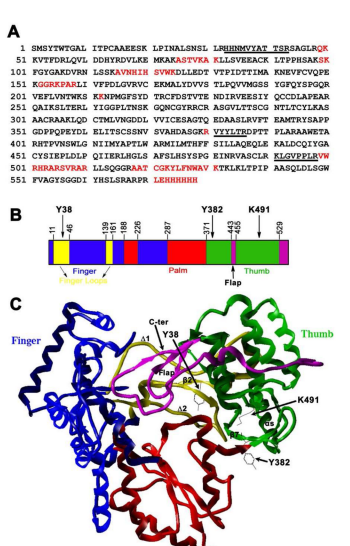


Figure 7. MS coverage and structural representation of NS5B showing FSBA modification sites (A) MS analysis identified peptides that had coverage of 89% of the sequence of HCV NS5B (bold). The sequence(s) that could not be detected in MS are shown in red. The peptides modified by FSBA and identified by MALDI-MS are underlined. (B) and (C) *Blue, red,* and *green* highlight the finger, palm, and thumb subdomains, respectively; *Yellow* represents loops interconnecting the fingers and thumb subdomains; *purple,* represents loops interconnecting the thumb subdomain. (B) and (C) respectively, the linear and three-dimensional representation of NS5B. The identified modification at Y38, Y382, and K491 are shown by the arrow. Y38 is located on the  $\beta$ 2 finger loop connecting the thumb and finger domains; Y382 is located on the  $\beta$ 7 strand at the interface of thumb and palm. K491 is positioned on the  $\alpha$ 5 helix in the thumb domain.

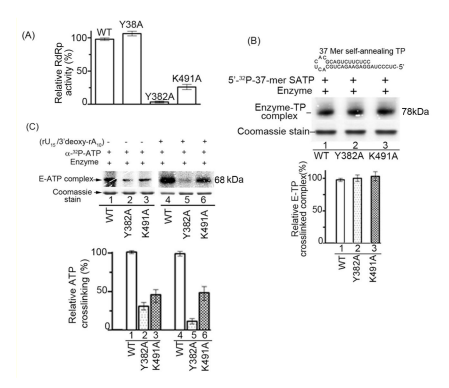


Figure 8. (A) Effect of alanine substitution at position 38, 382 and 491 on the RdRp activity of HCV NS5B  $\,$ 

The RdRp activity of the WT and its mutant derivatives (1  $\mu$ M) were determined and expressed as percent relative activity with respect to the WT enzyme. (B) Photocrosslinking of  $^{32}$ P-labeled RNA template primer to the wild type enzyme and its mutant derivatives. The 5'-[ $^{32}$ P]-labeled 37-mer self-annealing RNA template primer was used as the template primer to crosslink with the WT enzyme and its mutant derivatives. The  $^{32}$ P labeled 37-mer TP (200 nM) was incubated on ice with 1  $\mu$ M of enzyme in a final volume of 50  $\mu$ l. The mixture was UV irradiated and the crosslinked E-TP covalent complexes were resolved by SDS PAGE and visualized by PhosphorImaging. (C) UV mediated Photocrosslinking of 5'-  $\alpha$ - $^{32}$ P-ATP to the wild type enzyme and its mutant derivatives in the absence and presence of template primer. The wild type enzyme and its mutant derivatives were photo-crosslinked with [ $\alpha$ - $^{32}$ P]-ATP in the absence of template primer (lanes 1–3) or in the presence of rU<sub>15</sub>,rA<sub>10</sub> template primer with 3'-deoxy terminated primer terminus (lanes 4–6). The crosslinked complex was resolved on SDS PAGE and visualized by PhosphorImaging.

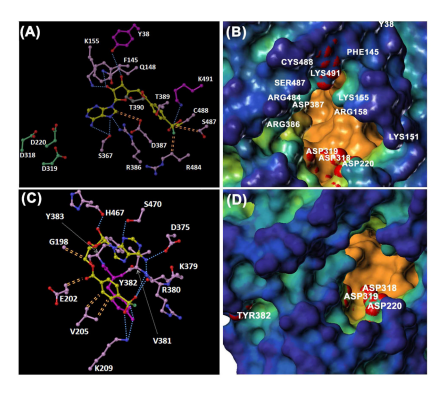
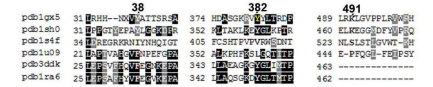


Figure 9. Molecular docking of FSBA at the nucleotide binding site in HCV NS5B
The induced-fit docking of FBSA with NS5B was done using workflow of a Schrodinger
modeling package version 9.2 (Schrodinger LLC., New York, NY). The crystal structure of
GTP-bound HCV NS5B (PDB entry 1GX5; (13)) was used for docking of FSBA near
LYS491 (A) as well as TYR382 (C). The oxygen of hydroxyl group of TYR382 and
nitrogen of epsilon amino group of LYS491 are, respectively, 3.7Å and 3.4Å away from the
fluorosulfonyl group of FSBA. The van der Waals interactions of FSBA with the
neighboring residues are shown as orange double-dotted lines; polar (H-bonding)
interactions are shown as cyan dotted lines. FSBA and target residues are shown as yellow
and magenta, respectively. Surface representation of the NTP tunnel showing the position of
LYS491 (B) and TYR382 (D). The NS5B residue (protein data bank code 1NB7), which
was devoid of RNA and metal, was prepared in Sybyl8.1 with MOLCAD probe radius 1.4 Å
(type: cavity depth). The residues LYS491 and TYR382 and the catalytic residues (ASP220,
ASP318, and ASP319) are shown in red.



 ${\bf Figure~10.~The~structural~alignment~of~RNA-dependent~RNA~polymerases~of~RNA~viruses~based~on~conserved~structural~elements}$ 

The PDB codes are: 1gx5, HCV NS5B; 1sh0, Norwalk virus (Triclinic); 1s4f, bovine viral diarrhea virus (BVDV); 1u09, Foot-and-Mouth Disease virus (FMDV); 3ddk, Coxsackie virus; and 1ra6, Polio virus.

Table 1

Effect of various additions on the FSBA-mediated inhibition or inactivation of RdRp activity of HCV NS5B<sup>a</sup>.

Additions	RdRp activity (% of control)		
Additions	50 mM NaCl	150 mM NaCl	
0.5 mM FSBA	33 ± 4	35 ± 8	
+ 1mM ATP	83 ± 7	82 ± 6	
+ 1mM GTP	87 ± 8	86 ± 8	
+ 1mM CTP	84 ± 8	82 ± 5	
+ 1mM UTP	90 ± 9	87 ± 8	
+ 10 μg Poly rA.rU <sub>12</sub>	70 ± 9	56 ± 5	
+ 10 µg poly rA.dT <sub>18</sub>	83 ± 6	50 ± 3	
+37-mer looped RNA	90 ± 8	56 ± 5	
+ 10 μg poly rU.rA <sub>26</sub>	35 ± 5	46 ± 4	

<sup>&</sup>lt;sup>a</sup>Reaction mixture was incubated in 100 µl final volume containing 10 µM HCV NS5B with the indicated additions in absence or presence of 0.5 mM FSBA. After 1 h incubation at 4°C, an aliquot of the incubation mixture was withdrawn and assayed for RdRp activity. The percent of HCV NS5B RdRp activity is expressed with respect to the control. Data represents the mean values (±SD) obtained from two independent experiments. The sequence of 37-mer double stranded RNA looped template primer is shown in Fig. 3.

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Table 2

Peptide modification by FSBA as determined by LC/MS/MS and MALDI-TOF/TOF.

Comments	Tyr382 is located on the $\beta 7$ strand at the interface of thumb and palm		Lys491 is located on the aS helix in the thumb domain	whereas Tyr38 is located on the $\beta2$ finger loop connecting thumb and finger domain
Target residue	Tyr382		Lys491	Tyr38
ATP prevents cross-linking $^{\mathcal{C}}$	yes		yes	yes
NS5B residue with modification	381-VYYLTR-386 SB	381-VYYLTR-386 SBA	491-KLGVPPLR-498 SBA	33-HHNMVYATTSR-43 SB
(z/m)	998. 43 <sup>6</sup>	1246.50 <sup>a</sup>	1311.634.6	1500.60 <i>a,b</i>
Peptide	1	2	3	4

 $^a_{
m HPLC}$  purified peptides were analyzed by LC/MS/MS.

The FSBA- modified HCV NS5B was resolved over 8% SDS-PAGE and subjected to in-situ trypsin digestion, followed by MALDI-TOF/TOF analyses. Peptide modified at a particular position by SBA or SB is shown as identified by MS fragmentation pattern. The molecular masses of SBA and BA were calculated as 433.08 Da and 183.98 Da, respectively. <sup>c</sup>HCV NS5B was incubated with 1 mM ATP before the addition of FSBA to the labeling reaction. The protein was TCA precipitated and digested by trypsin, followed by HPLC analyses. Comparisons of tryptic peptide profiles of the samples in the presence and absence of ATP were acquired where "yes" indicates the protection of HCV NS5B from FSBA-mediated modification (see Fig. S2 for details).