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Genetic Analysis of a Hydrophobic Domain of Coxsackie B3 Virus Protein 2B: a Moderate Degree of Hydrophobicity Is Required for a cis-Acting Function in Viral RNA Synthesis

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Received 5 July 1995/Accepted 8 September 1995

Coxsackie B virus protein 2B contains near its C terminus a hydrophobic domain with an amino acid composition that is characteristic for transmembrane regions. A molecular genetic approach was followed to define the role of this domain in virus reproduction and to study the structural and hydrophobic requirements of the domain. Nine substitution mutations were introduced in an infectious cDNA clone of coxsackie B3 virus. The effects of the mutations were studied in vivo by transfection of Buffalo green monkey cells with copy RNA transcripts. The results reported here suggest that a critical degree of hydrophobicity of the domain is essential for virus growth. The mutations S77M, C75M, I64S, and V66S, which caused a more pronounced increase or decrease in mean hydrophobicity, yielded viable viruses. The double mutations S77M/C75M and I64S/V66S, which caused a more pronounced increase or decrease in hydrophobicity, were nonviable. Negatively charged residues (mutations A71E, I73E, and A71E/I73E) abolished virus growth. The mutations had no effect on the synthesis and processing of the viral polyprotein. Replication and complementation were studied by using a subgenomic coxsackievirus replicon containing the luciferase gene in place of the capsid coding region. Analysis of luciferase accumulation demonstrated that the mutations cause primary defects in viral RNA synthesis that cannot be complemented by wild-type protein 2B provided in trans. The hydrophobic domain is predicted by computer analysis to form a multimeric transmembrane helix. The proposed interaction with the membrane and the implications of the mutations on this interaction are discussed.

The genus Enterovirus, a subgroup of the Picornaviridae, consists of the polioviruses, coxsackieviruses, echoviruses, and enterovirus serotypes 68 to 71. The structures, genomic organization, and molecular biology of these viruses are relatively well known from the extensive studies of poliovirus. Enteroviruses are positive-strand RNA viruses with a single-stranded genome of approximately 7,500 nucleotides (nt). A small virally encoded protein, VPg, is covalently attached at the 5' end of the RNA, and a genetically encoded poly(A) tail is present at the 3' end (see Fig. 2A). The viral genome acts as mRNA to direct the synthesis of a polyprotein. Translation occurs via a CAP-independent binding of ribosomes to the 5' nontranslated region (19, 32). The viral polyprotein is proteolytically processed by the virus-encoded proteases 2Apro, 3CP, and 3CDpro to functional cleavage intermediates and the individual proteins. The proteins encoded by the P1 region of the genome are capsid proteins. The nonstructural proteins encoded by the P2 and P3 regions are involved in viral RNA (vRNA) replication. For more details on the genomic structure of enteroviruses, we refer to a recent review of Wimmer et al. of the genetics of poliovirus (40). Replication of the enterovirus RNA genome starts with the synthesis of a complementary negative strand RNA in the cytoplasm of the host cell by the RNA-dependent RNA polymerase 3Dpol (3). Viral positive RNA strand synthesis takes place in replication complexes, which are specialized structures composed of partially double-stranded replicative intermediates plus the replication proteins (17). Genetic and biochemical studies have shown that the proteins encoded by the P3 region are involved in vRNA synthesis within these complexes (40). The replication complexes are closely associated with virus-induced membrane vesicles (6). Both the initiation of vRNA synthesis and the release of vRNA from the complexes are dependent on these vesicles (7, 37, 38). The inhibition of poliovirus RNA replication by Brefeldin A, an antibiotic that blocks membrane traffic between the endoplasmic reticulum (ER) and the Golgi apparatus and between the cis and trans Golgi apparatus, further indicates that intact secretory pathway traffic is required for vRNA replication (14, 18, 28).

Enterovirus proteins 2B, 2C, and 2BC seem to be involved in the functional and structural organization of the replication complexes. These proteins are rough-ER (rER) associated soon after their synthesis and are exclusively localized at the cytoplasmic surface of the ER-derived membrane vesicles that surround the viral replication complex (5, 6, 8). Electron microscopy and biochemical studies have shown that protein 2C or its precursor 2BC is involved in the attachment of the vRNA to the vesicular membranes (7, 9, 35). Furthermore, it has been shown that poliovirus protein 2C is an NTPase with both ATPase and GTPase activities (29, 35). The transient expression of poliovirus protein 2C or its precursor 2BC in eucaryotic cells resulted in the proliferation of cellular membranes and accumulation of membrane vesicles in a way similar to that seen during virus infection, suggesting that protein 2C belongs to the family of small GTPases involved in the regulation of membrane traffic (1, 10). Enterovirus protein 2B is a small hydrophobic protein which has profound effects on cellular membranes. Mutations mapping to the N terminus of poliovirus protein 2B exhibited primary defects in RNA synthesis (20, 27). These 2B mutants displayed a dosage-dependent dominance over wild-type virus, which led to the suggestion that protein 2B plays a structural role in the viral replication complexes, possibly by interacting with a limiting membrane at-
attachment site (20). The expression of both poliovirus and coxsackie B3 virus (CBV3) protein 2B in Escherichia coli resulted in a modified membrane permeability (26, 39a). Recently, it has been shown that the expression of poliovirus protein 2B in eucaryotic cells leads to permeabilization of the cell membrane and inhibition of secretory pathway function (13). These observations suggest that enterovirus protein 2B is a membrane-anchoring protein which may be involved in the accumulation of membrane vesicles for vRNA replication.

The regions of protein 2B that are involved in the interaction with the ER membrane are not yet known. Membrane anchors are typically composed of mainly hydrophobic amino acids and a significant portion of serine and threonine residues (15). About 18 amino acid (aa) residues are required to span the ER membrane (31). Analysis of the hydropathy of CBV3 protein 2B according to the method described by Kyte and Doolittle (24) demonstrates the existence of two hydrophobic domains (Fig. 1A). The hydrophobicity of these domains is well conserved in all enteroviruses, despite extensive differences in amino acid sequence among members of the CBV-like and poliovirus-like subgroups (Fig. 1B). The hydrophobic domain near the C terminus of CBV3 protein 2B is composed of a stretch of 18 aa (aa 63 to 80) which is devoid of charged residues. The amino acid composition of this hydrophobic domain is characteristic for transmembrane domains, since it contains 12 hydrophobic aa, 4 threonine residues, and 2 serine residues. The importance of this domain is reflected in the conservation of the hydrophobic features among all enteroviruses. The limited intratypic sequence heterogeneity found in five CBV1 isolates containing divergent 2A coding regions (41) further strengthens the importance of conservation in this region of protein 2B (Fig. 1B).
The results reported here indicate that a moderate degree of hydrophobicity of the domain is essential for the function of protein 2B. Mutations which caused a major increase or decrease in hydrophobicity as well as the introduction of negatively charged residues interfered with virus growth. These mutations caused primary defects in RNA synthesis that could not be complemented in trans by wild-type protein 2B. The proposed interaction of this hydrophobic domain with the membrane and the implications of the mutations on this interaction are discussed.

MATERIALS AND METHODS

Cells and viruses. Virus propagation and RNA transfections were performed with BGM cells. Plaque assays were performed with Vero cells. The cells were grown in minimal essential medium (MEM) (GIBCO) supplemented with 10% fetal bovine serum (GIBCO). After infection, cells were fed with MEM containing 3% fetal bovine serum. After transfection, MEM containing 10% fetal bovine serum was added to the cells. Virus yields were determined by endpoint titration. Serial 10-fold dilutions were tested in 96-well plates containing BGM cell monolayers (36). Viral titers were calculated according to the method described by Reed and Muench and were expressed in 50% tissue culture infective dose (TCID50) values (34).

Oligonucleotide-directed site-specific mutagenesis. Plasmid pCB3/T7 was kindly provided by R. Kandolf (University of Tübingen, Tübingen, Germany). This plasmid contains a full-length copy DNA of CBV3 (strain Nancy) cloned behind a T7 RNA polymerase promoter (22). In vitro mutagenesis was performed with the Altered Sites in vitro mutagenesis system (Promega) according to the recommendations of the manufacturer. For mutagenesis, the HindIII (nt 2080) to XhoI (nt 4947) fragment of pCB3/T7 was introduced in phagemid pALTER-1 to generate pALTCB3/2080-4947. Synthetic oligonucleotides were used to introduce site-specific mutations. The nucleotide sequence of the mutant pALTCB3 was verified by dideoxy chain termination sequence analysis of the plasmid DNA with oligonucleotide 5'-CTCACTAATGAGTTCCTGCTG (complementary to nt 417 to 434) with the Ampli Cycle sequencing kit according to the manufacturer's instructions. The plasmid DNA was linearized by digestion with Spel (nt 3837) and BglII (nt 3388) restriction enzymes. In the resulting plasmid (pCB3/T7-APal), nt 746 to 3279 of the PI capsid-coding region was replaced by the luciferase coding sequence, the PI capsid-coding region by the luciferase coding sequence, and the PI capsid-coding region was replaced by the luciferase coding sequence. Because of the cloning procedure, the luciferase product contains three additional amino acids (Gly-Gly-Arg), which correspond to the N-terminus of VP1 were retained to ensure a correct proteolytic processing by 2Apro.

Analysis of viral protein synthesis in vivo. BGM monolayer cells (2 X 10^5) infected with virus at a MOI of 2 to 4 were infected with virus at a MOI of 2 to 4. Medium was collected at 0.5, 2, and 4 h pi and analyzed by sodium dodecyl sulfate (SDS) (25). The samples were resolved on a 12.5% polyacrylamide gel containing sodium dodecyl sulfate (SDS) (25). The proteins were transferred to polyvinylidene difluoride (PVDF) membranes and detected with an antibody to VP1. Analysis of viral protein synthesis in vivo. BGM monolayer cells (2 X 10^5) infected with virus at a MOI of 2 to 4 were infected with virus at a MOI of 2 to 4. Medium was collected at 0.5, 2, and 4 h pi and analyzed by sodium dodecyl sulfate (SDS) (25). The samples were resolved on a 12.5% polyacrylamide gel containing sodium dodecyl sulfate (SDS) (25). The proteins were transferred to polyvinylidene difluoride (PVDF) membranes and detected with an antibody to VP1. Analysis of viral protein synthesis in vivo. BGM monolayer cells (2 X 10^5) infected with virus at a MOI of 2 to 4 were infected with virus at a MOI of 2 to 4. Medium was collected at 0.5, 2, and 4 h pi and analyzed by sodium dodecyl sulfate (SDS) (25). The samples were resolved on a 12.5% polyacrylamide gel containing sodium dodecyl sulfate (SDS) (25). The proteins were transferred to polyvinylidene difluoride (PVDF) membranes and detected with an antibody to VP1.
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were incubated at 33°C, and four were incubated at 36°C. Cells infected at either 33 or 36°C after transfection with copy RNAs posttransfection. No CPE was observed up to 5 days posttransfection. The result of the effect of the mutations on virus growth was summarized in Fig. 2B.

**RESULTS**

Introduction of mutations in a hydrophobic domain of CBV3 protein 2B. Nine CBV3 mutants containing amino acid substitutions in a hydrophobic 18-residue domain of protein 2B were constructed by site-directed mutagenesis. The mutations were verified by sequence analysis and introduced in the infectious cDNA clone pCB3/T7. The integrity of the resulting plasmids was checked by restriction enzyme analysis. Figure 2 shows the genotypes of the mutants and the predicted effects of the mutations on hydrophobicity. On the basis of the effects of the mutations on the mean hydrophobicity and net charge of this region, the pCB3/T7 2B mutants can be divided into three groups as follows. (i) In mutants 2B-S77M, 2B-C75M, and 2B-S77M/C75M, serine 77 (TCG) and cysteine 75 (TGT) were replaced by more-hydrophobic methionine (ATG) residues. (ii) In mutants 2B-I64S, 2B-V66S, and 2B-I64S/V66S, the mean hydrophobicity is decreased by substitution of hydrophobic residues isoleucine 64 (ATC) and valine 66 (GTG) by polar serine residues (codons AGT and TCG, respectively), which contain aliphatic hydroxyl side chains. (iii) In mutants 2B-A71E, 2B-I73E, and 2B-A71E/I73E, the mean hydrophobicity is decreased by the replacement of alanine 71 (GCC) and isoleucine 73 (ATC) by negatively charged glutamic acid (GAG) residues.

**Effect of the mutations on virus growth.** The effect of the mutations on virus viability was studied by transfection of BGM cells with copy RNA transcripts. For each mutant, eight transfections were performed. Four transfected cell cultures were subjected to infection at either 33 or 36°C after transfection with copy RNAs of mutant 2B-S77M/C75M, 2B-I64S/V66S, 2B-A71E, 2B-I73E, and 2B-A71E/I73E. These cell cultures were subjected to three cycles of freezing and thawing, and 250 μl was subsequently passaged to fresh BGM cell monolayers. CPE was observed after passage of one of the cell cultures transfected with copy RNA of mutant 2B-S77M/C75M (see the following paragraph), but not after passage of cells transfected with copy RNAs of any of the other mutants.

Sequence analysis of the 2B coding region of the viable mutant viruses showed that the original mutations introduced by site-specific mutagenesis were retained in the vRNA and that no other amino acid replacements had occurred. Viral growth characteristics were examined by plaque assays and single-cycle growth analysis. Viruses vCB3-2B-S77M, -I64S, and -V66S exhibited wild-type growth. Virus vCB3-2B-C75M exhibited a small-plaque phenotype (Fig. 3). In single-cycle infections, this virus produced 10 and 20% of the wild-type virus yield at 6 and 8 h postinfection, respectively (data not shown). All mutant viruses showed similar reproductive capacities at 33, 36, and 39°C, which indicates that none of the mutations caused temperature-sensitive growth defects. The results of the transfections and the characterization of the viable mutants are summarized in Fig. 2B.

**Isolation of two revertant viruses.** Passage of cells transfected with copy RNA of mutant 2B-S77M/C75M revealed virus growth in one of the cultures grown at 33°C. Sequence analysis demonstrated that these viruses still contained the methionine at position 77 but a reversion of the introduced methionine (AUG) at position 75 to a valine (GUG) residue. This partial revertant virus, vCB3-2B-S77M/C75M, exhibited a small-plaque phenotype at all temperatures (data not shown). In single-cycle infections, this virus produced 3 and 5% of the wild-type virus yield at 6 and 8 h postinfection, respectively (data not shown). Amplification and sequence analysis of the 2B coding region of virus from the primary transfected culture showed that the reversion had already occurred in the transfected cells.

The plaque assay plates of mutant vCB3-2B-S77M/C75M contained predominantly small plaques; however, larger plaques were also observed (Fig. 3). Three large plaques and three...
small plaques were picked from the plates, and the viruses were grown on BGM cells. Sequence analysis demonstrated that the viruses isolated from the small plaques had retained the introduced methionine (AUG) residue. However, the viruses isolated from the three large plaques contained a valine (GUG) residue at amino acid position 75 instead of the introduced methionine and were named vCB3-2B-C75V. No other mutations in the 2B coding sequence were observed. This revertant, vCB3-2B-C75V, exhibited a wild-type plaque size at all temperatures.

To investigate if the reversions in the mutants vCB3-2B-C75V and vCB3-2B-S77M/C-75V were solely responsible for the observed phenotypes, we introduced these mutations into the wild-type pCB3/T7 clone. For this purpose, the 519-bp PCR products (nt 3733 to 4251) that were used for the sequence analysis of the 2B coding region (nt 3746 to 4041) were cut with Spe1 (nt 3837) and BssHII (nt 4238), purified, and cloned in pCB3/T7. nt 4042 to 4238 were sequenced to verify that no additional mutations had occurred in this region. BGM cells were transfected with copy RNA generated from the resulting plasmid clones 2B-C75V and 2B-S77M/C75V. Sequence analysis of the 2B coding regions of the resulting viruses demonstrated that the introduced mutations were retained in the vRNA. These viruses exhibited the same plaque phenotype that the revertants obtained after transfection of cells with copy RNA of mutants 2B-C75M and 2B-S77M/C75M, respectively, exhibited, which indicates that the reversions were sufficient to confer the observed phenotypes (data not shown).

Analysis of vRNA replication with subgenomic replicon pCB3/T7-LUC. The observation that the capsid-coding P1 region could be deleted from poliovirus RNA without any effect on the efficiency of genome replication (21) has led to the construction of chimeric subgenomic poliovirus replicons in which the P1 region was replaced by reporter genes (2, 33). Andino et al. (2) have described a subgenomic poliovirus replicon (RLuc-31) in which the P1 region was replaced by the luciferase gene and have shown that the activity of the reporter protein following transfection correlated with the levels of accumulated RNA (2).

To study vRNA replication of the 2B mutants, chimeric subgenomic replicon pCB3/T7-LUC was constructed (Fig. 4A). In this replicon, the P1 region of pCB3/T7 is substituted by the firefly luciferase gene. The suitability of this replicon to study vRNA replication was examined by transfection of BGM cells with copy RNA of pCB3/T7-LUC or the replication-defective control pCB3/T7-LUC-Δ3D. This plasmid contains a large in-frame deletion in the 3Dpol region by deletion of the BbrPL fragment (nt 6117 to 7152). At several times posttransfection, the cells were lysed and the levels of luciferase activity were measured. Figure 4B demonstrates that the kinetics of luciferase production by pCB3/T7-LUC copy RNA can be divided into three phases, which are similar to those described for RLuc-31 (2). In phase I (1 to 3 h posttransfection), luciferase activity increased as a result of translation of the input RNA. Phase II (3 to 5 h posttransfection) is characterized by a steady state in luciferase activity. Phase III (5 to 12 h posttransfection) demonstrated a second increase (10- to 50-fold) in luciferase activity as a result of the translation of newly synthesized chimeric RNA strands. In contrast, phase III of RLuc-31 is already detectable from the 3rd h posttransfection, and replication results in a 100- to 500-fold increase in luciferase activity (2). This difference is probably due to the low infectivity of pCB3/T7-derived copy RNA, which is only 14 to 30 PFU/μg of copy RNA (22). Upon transfection of pCB3/T7-LUC RNA, luciferase activity is generated in all transfected cells, but not in a few of these cells will an infectious cycle be initiated. The exponential increase in luciferase activity as a result of vRNA replication will, therefore, be less evident and cannot be detected before 6 h posttransfection. However, the absence of phase III in cells transfected with copy RNA of pCB3/T7-LUC-Δ3D (Fig. 4B) demonstrates the suitability of replicon pCB3/T7-LUC to study vRNA replication.

**Growth-defective 2B mutants display defects in vRNA synthesis.** vRNA synthesis of the nine originally constructed mutants was examined after introduction of the SpeI-to-BssHII fragments in replicon pCB3/T7-LUC. BGM cells were transfected with copy RNA of pCB3/T7-LUC replicons containing the 2B mutations and grown at 36°C, and the luciferase activities were determined at several times posttransfection. Replicons carrying the nonviable 2B mutations (S77M/C75M, I64S/V66S, A71E, I73E, and A71E/I73E) displayed defects in RNA replication (Fig. 4C). These replicons exhibited phases I and II of the luciferase production, which are the result of translation of the input RNA, but not the phase III increase that reflects translation of newly synthesized RNA strands. Replicons carrying the viable mutations S77M, I64S, and V66S, which produced viruses with wild-type growth characteristics, displayed the same kinetics of luciferase production that wild-type pCB3/T7-LUC did (Fig. 4D). Mutation C75M, which yielded virus with a small-plaque phenotype, caused a severe reduction in RNA synthesis. The absence of an increase in luciferase activity above the translation level, which might have been expected from a low level of replication, is most likely due to the low level of infectivity of the copy RNA transcripts.

Figures 4C and D show that the luciferase activity generated by the replication-defective replicons remained constant up to 10 h posttransfection. Because luciferase is an unstable enzyme in vivo (2), the constant level must represent an equilibrium between translation of the input RNA and degradation of luciferase. The constant level of luciferase activity suggests that none of the 2B mutations affected the translation ability or stability of the chimeric input RNA.

**In vitro protein synthesis and processing of RNAs containing mutations in protein 2B.** In vitro translation reactions were performed to investigate whether the mutations affect translation and processing of the polyprotein. Figure 5 shows that the patterns of [35S]serylabeled proteins obtained with the 2B mutant plasmids were similar to that produced by wild-type plasmid, which indicates that none of the mutations caused defects in synthesis and processing of the viral polyprotein. The positions of proteins 2BC and 2C were identified by in vitro
FIG. 4. Analysis of vRNA replication of the 2B mutants with chimeric luciferase replicon pCB3/T7-LUC. (A) Nucleotide and amino acid sequences of construct pCB3/T7-LUC at the N and C termini of the introduced luciferase gene. The last 5 aa are conserved in this construct to ensure a correct cleavage by protease 2A. Underlined are the NotI and HpaI restriction sites used for the introduction of the luciferase gene. (B to D) Luciferase accumulation after transfection of BGM cells with copy RNAs of wild-type pCB3/T7-LUC and pCB3/T7-LUC-Δ3D (B), pCB3/T7-LUC containing the lethal 2B mutations (C), and pCB3/T7-LUC containing the viable 2B mutations (D). Luciferase activities were measured as described in Materials and Methods and are expressed on a logarithmic scale.

translation of copy RNAs carrying the 2BC or 2C coding sequence behind the encephalomyocarditis virus 5' nontranslated region (data not shown). In vitro-translated protein 2B (± 11 kDa) could not be visualized because of the presence of only two cysteines in protein 2B and the interference of the excess of globin in the reticulocyte lysate with a correct migration of proteins with molecular masses of less than 15 kDa. However, the production of protein 2C at levels equal to that produced by wild-type RNA demonstrates that none of the mutants displayed processing defects at the 2B-2C junction.

Analysis of the viral protein synthesis in vivo. The rates of viral protein synthesis in cells infected with small-plaque viruses vCB3-2B-C75M and vCB3-2B-S77M/C75V were compared with that in wild-type virus-infected cells to examine whether the defects in RNA synthesis are a consequence of an impaired level of viral protein synthesis in vivo. BGM cells were infected with either vCB3-2B-C75M, vCB3-2B-S77M/C75V, or wild-type virus at a MOI of 50, and proteins were labeled with [35S]methionine at different times postinfection. Figure 5 shows that the shutoff of host cell translation after infection with these two mutant viruses was similar to the shutoff induced by wild-type virus. At 4 and 5 h postinfection, the level of viral protein synthesis in the mutant virus-infected cells was slightly below the level in wild-type-infected cells. At 6 h postinfection, the rates of viral protein synthesis in the mutant-infected and wild-type-infected cells were similar, which indicates that it is unlikely that the mutations cause primary defects in the rate of synthesis of the viral polyprotein.

In the previous paragraphs, it has been shown that mutations in protein 2B cause defects in vRNA synthesis. The slightly decreased level of viral protein synthesis in cells infected with vCB3-2B-C75M and vCB3-2B-S77M/C75V at 4 and 5 h postinfection is probably a reflection of the defects in RNA synthesis.

FIG. 5. In vitro translation of RNA derived from wild-type pCB3/T7 and the 2B mutant plasmids. SstI linearized plasmid DNA (0.5 μg) was incubated for 3 h in TNT reticulocyte lysate, a coupled transcription-translation system, supplemented with HeLa initiation factors. The [35S]cysteine-labeled translation products were analyzed on a SDS-12.5% polyacrylamide minigel. An extract from wild-type virus-infected cells, labeled with [35S]methionine for 1 h at 4 h postinfection, was used as marker.
and, as a consequence, the reduced amount of available viral positive RNA strands. The reduction in vRNA synthesis is probably also responsible for the delay in CPE and, as result, for the prolonged synthesis of viral proteins up to 7 h postinfection, when viral protein synthesis in wild-type virus-infected cells is already severely decreased.

Defects in vRNA synthesis caused by mutations in a hydrophobic domain of protein 2B cannot be rescued in trans. To study whether the defects in vRNA synthesis could be rescued in vivo by wild-type protein 2B provided in trans, genetic complementation experiments were performed. We studied complementation of the defects caused by mutation C75M, which yields small-plaque viruses, and the nonviable mutations S77M/C75M, I64S/V66S, A71E, I73E, and A71E/I73E. To assay for complementation, BGM cells were transfected with copy RNAs of the mutant pCB3/T7-LUC replicons, were incubated at 36°C, and at 90 min posttransfection were either mock or virus infected with wild-type virus at a MOI of 50. At 4 and 8 h posttransfection, the cells were lysed and the luciferase activity was measured. Wild-type pCB3/T7-LUC copy RNA transcripts were tested to exclude the possibility that the replication of wild-type virus interfered with the luciferase accumulation. The results shown in Table 1 demonstrate that virus growth did not adversely affect replication and translation of the pCB3/T7-LUC RNA strands. Complementation tests with the mutant pCB3/T7-LUC replicons showed similar levels of luciferase activity at 8 h posttransfection in mock- and virus-infected cells (Table 1). The luciferase activities measured at 8 h posttransfection in both mock- and virus-infected cells were comparable to the activities measured at 4 h posttransfection, a time point at which the level of luciferase activity reflects the translation of the input copy RNA. The absence of major increases in luciferase activity in the virus-infected cells suggests that the defects in RNA synthesis caused by mutations in a hydrophobic domain of protein 2B could not be efficiently complemented by the function of wild-type protein 2B provided in trans. However, the possibility that low levels of complementation have occurred cannot be excluded. Low levels of replication are difficult to detect because of the low infectivity of the copy RNA transcripts (see above) and the small variations in transfection efficiency and luciferase assays that were observed.

**DISCUSSION**

Enterovirus protein 2B is a small hydrophobic protein that is localized at the outer surfaces of the virus-induced membrane vesicles on which vRNA synthesis takes place. The nature of the interaction of protein 2B with the membrane is yet unknown. Examination of the amino acid sequence of protein 2B of CVB3 revealed an 18-aa hydrophobic domain (aa 63 to 80) with a composition that is characteristic of integral transmembrane regions. In this study, we examined the effects of amino acid substitutions in this hydrophobic domain of CVB3 protein 2B on virus growth, vRNA replication, and viral polyprotein processing to gain more insight into the role of this domain in the function of protein 2B. The mutations that were introduced can be divided into three groups on the basis of their effects on hydrophobicity and net charge of the domain (Fig. 2). In the first group, serine 77 and cysteine 75 were replaced by more hydrophobic methionine residues. Mutation S77M yielded virus with wild-type growth characteristics. Mutation C75M yielded mutant virus with a small-plaque phenotype because of a reduction in vRNA synthesis. Large-plaque revertants of this mutant were isolated and found to contain a reversion of methionine 75 to a valine residue (vCB3-2B-C75V). The double mutation S77M/C75M caused a defect in vRNA synthesis and was nonviable. However, the isolation of small-plaque revertant virus vCB3-2B-S77M/C75V demonstrates that mutation S77M/C75M did not completely abolish vRNA replication. In the second group, hydrophobic amino acids with aliphatic side chains (isoleucine 64 and valine 66) were replaced with polar serine residues with aliphatic hydroxyl side chains. Mutations I64S and V66S produced mutant viruses with wild-type plaque phenotypes. However, the double mutation I64S/V66S disturbed vRNA synthesis and was nonviable. In the third group, alanine 71 and valine 73 were replaced with negatively charged glutamic acid residues. Mutations A71E, I73E, and A71E/I73E disturbed vRNA replication and were all nonviable. RNA synthesis seemed to be completely abrogated, since no revertant viruses could be isolated. In vitro translation reactions and analysis of the viral protein synthesis in infected cells demonstrated that none of the mutations interfered with the synthesis or processing of the viral polyprotein, indicating that the mutations in the hydrophobic domain primarily affected vRNA synthesis.

The results presented here suggest that the hydrophobic character of the domain is crucial for the function of protein 2B in vRNA replication. Furthermore, the data suggest that for efficient functioning of the domain, the hydrophobicity (<H>)

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**TABLE 1. Complementation tests between pCB3/T7-LUC copy RNAs containing the 2B mutations and wild-type virus.**

<table>
<thead>
<tr>
<th>Mutation or wild type</th>
<th>4 h pt</th>
<th>8 h pt (mock infected)</th>
<th>8 h pt (virus infected)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C75M</td>
<td>142</td>
<td>244 (1.7)</td>
<td>376 (2.6)</td>
</tr>
<tr>
<td>S77M/C75M</td>
<td>106</td>
<td>181 (1.7)</td>
<td>137 (1.3)</td>
</tr>
<tr>
<td>I64S/V66S</td>
<td>354</td>
<td>212 (0.6)</td>
<td>283 (0.8)</td>
</tr>
<tr>
<td>A71E</td>
<td>185</td>
<td>326 (1.8)</td>
<td>152 (0.8)</td>
</tr>
<tr>
<td>I73E</td>
<td>198</td>
<td>95 (0.5)</td>
<td>93 (0.5)</td>
</tr>
<tr>
<td>A71E/I73E</td>
<td>373</td>
<td>405 (1.1)</td>
<td>370 (1.0)</td>
</tr>
<tr>
<td>Wild type</td>
<td>251</td>
<td>3,491 (13.9)</td>
<td>3,050 (12.1)</td>
</tr>
</tbody>
</table>

*BGM cells were transfected with 0.5 µg of RNA of pCB3/T7-LUC carrying the 2B mutations and were either mock or virus infected with wild-type virus at a MOI of 50 at 90 min posttransfection (pt). At 4 and 8 h posttransfection, the cells were lysed and the luciferase activities were determined.

*Values in parentheses are the ratios of luciferase activities to those measured at 4 h posttransfection.*
MUTATIONAL ANALYSIS OF COXSACKIEVIRUS B3 PROTEIN 2B

The expression of poliovirus proteins required for vRNA synthesis. Alterations in the hydrophobic domain may interfere with the interaction of protein 2B with the membrane, which may disturb the interaction of protein 2B with the membrane. The mean hydrophobicity of these segments is probably too low for the integration as a monomeric transmembrane helix but may be sufficient to form a multimeric transmembrane helix. Classification of these segments on the hydrophobic moment plot also suggests that they form multimeric transmembrane helices, i.e., protein segments that are cooperatively associated within the lipid bilayer and stabilized by either an intermolecular (e.g., in the form of multimers) or an intramolecular linkage with other transmembrane α-helices.

At present, there is no biochemical evidence for the prediction that the hydrophobic domain forms an integral transmembrane helix that is associated with other transmembrane helices. However, the suggestion that the hydrophobic domain forms a multimeric rather than a monomeric transmembrane helix is supported by several observations made in this study. (i) The mean hydrophobicity values of the mutants with wild-type phenotypes varied between 0.44 and 0.57. These values are too low for monomeric transmembrane proteins to efficiently penetrate the membrane and maintain an energetically stable configuration. (ii) The substitution of cysteine 75 with a hydrophobic methionine residue interfered with vRNA synthesis, which makes it unlikely that this residue is involved in a hydrophobic interaction with the lipid bilayer. (iii) The mutations which caused the greatest increase in the hydrophobicity of the domain were either lethal for virus growth (S77M/ C75V <H> = 0.59) or resulted in a small-plaque virus, whereas a revertant containing a revertant vCB3-2B-C75V makes it unlikely that the mutation C75M is restricted by an upper limit. (iv) The suggestion that the hydrophobic domain forms an integral transmembrane helix, a multimeric transmembrane helix, or a surface-seeking helix with the aid of the hydrophobic moment plot, on which the hydrophobic moment of the helix is plotted as a function of its hydrophobicity. We have identified candidate membrane-associated helices in enterovirus protein 2B using 18- to 21-residue windows run over the entire protein. Candidate helices must have a mean <H> value of at least 0.42. Furthermore, there must be either one highly hydrophobic helix with an <H> value of at least 0.68 or one or more moderately hydrophobic helices, which may be provided either intramolecularly, with a summation value of at least 1.10. Candidate helices are then classified as either a monomeric transmembrane helix, a multimeric transmembrane helix, or a surface-seeking helix with the aid of the hydrophobic moment plot, on which the hydrophobic moment of the helix is plotted as a function of its hydrophobicity (15). We have identified candidate membrane-associated helices in enterovirus protein 2B containing 18- to 21-residue windows, since it has been reported that 18 aa are appropriate for spanning the ER membrane (31). In members of the CBV-like subgroup, the most hydrophobic segment is formed by aa 63 to 80 (<H> = 0.52 to 0.60). In members of the poliovirus-like subgroup, the segment with the highest <H> value is formed by residues 64 to 81 (<H> = 0.49 to 0.55). The mean hydrophobicity of these segments is probably too low for the integration as a monomeric transmembrane helix but may be sufficient to form a multimeric transmembrane helix. Classification of these segments on the hydrophobic moment plot also suggests that they form multimeric transmembrane helices, i.e., protein segments that are cooperatively associated with the lipid bilayer and stabilized by either an intermolecular (e.g., in the form of multimers) or an intramolecular linkage with other transmembrane α-helices.

A complementation analysis was performed to examine whether the defects in vRNA synthesis caused by the mutations in the hydrophobic domain could be rescued by wild-type proteins provided in trans. Mutations in the N-terminal part of poliovirus protein 2B have been reported to cause noncomplementable defects in vRNA synthesis (4, 12, 20, 27). However, complementation experiments with protein 3A mutants have shown that even within one protein, cis-acting and trans-acting functional regions can be discriminated (4, 16). We found that none of the mutations in the hydrophobic domain that caused phenotypes in vRNA synthesis could be efficiently complemented by wild-type protein 2B. Both the N-terminal region and the hydrophobic domain of protein 2B appear to be...
volved in a cis-action function in vRNA replication. It is not clear whether the effects that are observed upon expression of poliovirus protein 2B (i.e., permeabilization of the plasma membrane and inhibition of protein secretion) (13) reflect different functions of protein 2B or whether one of these effec-
tives is the consequence of the other. If permeabilizing the plasma membrane were the primary function of the hydropho-
dic domain and if the inhibition of protein secretion were only a side effect of this function, then a certain degree of trans-
 complementation should have been expected. The lack of complementability does not exclude the possibility that the hydrophobic domain is involved in permeabilization of the plasma membrane but suggests that the hydrophobic domain is also involved in another function of protein 2B in vRNA syn-
thesis.

Several explanations may account for the cis-action function of protein 2B. First, protein 2B may interact directly with the ER membrane to function in the accumulation or structural organization of virus-induced membranous structures around the same positive RNA strand from which it is synthesized. This model would ensure a tight linkage between transla-
tion and replication and prevent RNAs other than viral posi-
tive RNA strands from being replicated by 3Dpol, which lacks template specificity (20). A second possibility is that wild-type 2B proteins cannot reach the mutant templates. Replication of the vRNA takes places in close conjunction with membranes. This membranous environment may interfere with the func-
tional interchange of mutant and wild-type protein 2B. A third possibility is that the mutant 2B proteins interact with a limiting target and that wild-type protein 2B, which is provided later, cannot efficiently compete with them. This possibility was raised by Johnson and Sarnow (20), who found that poliovirus 2B mutants exhibited a dosage-dependent dominance over wild-type protein 2B. The interaction of the mutant CBV3 2B proteins with this limiting target may interfere with the action of wild-type protein 2B and thereby account for the lack of complementation.

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