The LTR, v-src, LTR Provirus Generated in the Mammalian Genome by src mRNA Reverse Transcription and Integration

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Different types of altered proviruses of Rous sarcoma virus (RSV) have been detected in mammalian tumor cell lines. We cloned and sequenced one of these altered proviruses with the structure LTR, v-src, LTR. The presence of an intact viral splice junction, as well as duplications of the chromosomal sequence GCGGGG flanking the two 2-base-pair-deleted LTRs, demonstrated reverse transcription and normal retroviral integration of src mRNA in mammalian cells. In addition, a 1-nucleotide deletion 2 bases upstream from the AAUAAA polyadenylation signal is suspected to be responsible for the absence of a poly(A) track in the src mRNA present in virions of rescued viruses.

The simplified transcriptionally and translationally active LTR, v-src, LTR provirus has been detected in the H-19 hamster sarcoma cell line transformed by the Prague strain of Rous sarcoma virus (RSV) subgroup C (PR-RSV-C) rescued from XC cells (provirus II) as reported previously (10, 20, 35, 39). It was demonstrated that this cryptic provirus is functional, because if properly complemented (by fusion with chicken fibroblasts preinfected with Rous-associated virus 1), transforming viruses can be rescued from H-19 cells which encapsidate RNA having the character of nonpolyadenylated src mRNA (37). According to restriction enzyme analysis and S1 mapping, the proviral structures of most of the rescued viruses, as well as of the original H-19 provirus, corresponded to that of reverse-transcribed src mRNA (8, 37). The aim of this work was to elucidate the structure of the LTR, v-src, LTR provirus integrated in the hamster genome to understand its genesis.

For cloning, we used the finding that the whole LTR, v-src, LTR provirus, together with flanking cellular sequences, is contained within a 5.9-kilobase-pair HindIII fragment and is excised by EcoRI as a 2.6-kilobase-pair fragment (37, 38).

HindIII fragments 5.4 to 6.0 kilobase pairs long were cut out of agarose and electropheluted, and the purified DNA was ligated with AIN1149 DNA opened at the HindIII site (18, 21). After molecular cloning, both of the expected HindIII and EcoRI fragments were observed (data not shown). The molecular clone obtained was designated λH19cIC2.1.

The integrity of λH19cIC2.1 was verified in cotransfection experiments in which 10 µg of each of λH19cIC2.1 and λ11411 DNAs containing a complete MAV-1 insert (25) was used to treat chicken fibroblasts (9). Transforming virus was detected 12 days after transfection (7 × 10⁴ focus-forming units/ml) and reached a titer of 6.5 × 10⁵ focus-forming units/ml after 4 weeks of culture. The relatively low titer of this transforming virus has already been noticed in virus rescue experiments (8). This indicates that in spite of the presence of the main structures required for packaging in src mRNA (see below), some additional cis-acting and packaging signals of the gag region are absent (1, 2, 14, 27, 33). Moreover, a lack of appropriate structure and the small size of src mRNA might be responsible for decreased binding to the encapsidation proteins (5), and the efficient packaging of viral genomic RNA might compete with that of src mRNA (7).

λH19cIC2.1 DNA was digested with HindIII or EcoRI and restriction enzyme Accl, Apal, BalI, BamHI, BglII, BglIII, BstEI, EcoRV, Eco47III, HincII, HpaI, KpmI, Narl, Ncol, NdeI, PstI, PvuI, PvuII, SacI, SmaI, SphI, SstI, Stul, XbaI, or Xhol and hybridized to the 3²P-labeled src11 probe (the PvuII fragment spanning base pairs [bp] 8098 to 8671 of PR-RSV-C) or a long terminal repeat (LTR) 2 probe (the HaeIII fragment spanning bp 8992 to 9335 of PR-RSV-C) or probe pATV-8, which detects all RSV genes (13). In this way, a restriction enzyme cleavage site map of λH19cIC2.1 was obtained (Fig. 1). This map corresponds to reverse-transcribed src mRNA after expected RSV genomic RNA splicing.

To determine the precise structure of λH19cIC2.1 relative to PR-RSV-C, we examined the nucleotide sequence of the provirus restriction fragments that seemed likely to contain important features. The viral leader, including the primer-binding site for Trp tRNA from which synthesis of the minus-strand DNA initiates during reverse transcription, was found to be intact (data not shown). As expected, a deletion was found to have occurred precisely between the splice donor site, which is located 18 bases downstream from the beginning of the gag gene sequence, and the acceptor site for the subgenomic src mRNA, which is located 75 bases upstream from the v-src initiation codon (Fig. 2).

This arrangement demonstrates that src mRNA was used as the template for genesis of the LTR, v-src, LTR provirus. With a packaging mutant deleted in the leader sequence, high-frequency (50%) generation of proviral structures that contain the splice junction of src mRNA was recently reported in chicken fibroblasts (15, 22). The published size (2.3 kilobase pairs) indicates that the proviruses retained the deleted leader. In contrast, our provirus was found in only 1 of 24 mammalian tumors tested (J. Pichrtová, Ph.D. thesis, Institute of Molecular Genetics, Czechoslovak Academy of Sciences, Prague, Czechoslovakia, 1988) but retained all structures of src mRNA (see below). Therefore, it most likely represents an infrequent event of unaltered src mRNA packaging and reverse transcription. In fact, the src mRNA is equipped with the leader sequence containing the main packaging signals (14-16, 27, 32) and with one copy of the
FIG. 1. Restriction enzyme cleavage site map of the proviral part of the molecular clone λH19eC2.1. (A) Proposed structure of the LTR, v-src, LTR genome corresponding to reverse-transcribed src mRNA after regular retroviral splicing in which the splice donor site (sd) at position 397 would be exactly joined to the splice acceptor site (sa) for subgenomic src mRNA at position 7,054. Restriction enzyme cleavage sites are numbered according to Schwartz et al. (26), and each number designates the first nucleotide of the recognition sequence. (B) Actual restriction map obtained by Southern blot restriction analysis which satisfactorily fits with the proposed structure (vertical lines), except for one PstI site at position 8,050 (indicated by a broken line). Wavy lines represent flanking hamster chromosomal DNA. Abbreviations: D, expected duplications of hamster chromosomal DNA after integration; PBS, primer-binding site; L, viral leader; src', noncoding region of viral src gene; DR, direct repeat; E, inverted repeat element; PPT, polypurine tract. The LTRs are boxed.

direct repeat to the right of v-src (see below) which is implicated also in RNA encapsidation (30). The src mRNA probably was transmitted to the hamster cell by the chicken-generated RSV particles responsible for H-19 cell transformation (38, 39). Because of general nonpermissiveness of mammalian cells to RSV (36), it is difficult to imagine that a src mRNA produced in RSV-infected hamster cells can be transmitted to other cells. Packaging and reverse transcription of subgenomic viral mRNAs have been reported for the env mRNA (17, 31) and repeatedly observed in the myc mRNA of the MH2 virus, which harbors two onecogenes (3, 19, 24).

Furthermore, the rest of the noncoding sequences at the 3' end of the cloned LTR, v-src, LTR structure, e.g., the inverted repeat element, one copy of the direct repeat sequence localized downstream from v-src, and especially the polypurine tract from which plus-strand DNA synthesis initiates during reverse transcription (29, 40), were also found without changes (Fig. 2). The same is true for both LTRs of the LTR, v-src, LTR provirus, which were fully conserved after reverse transcription and integration, including the imperfect inverted repeat at both outer ends of the LTRs, with one exception discussed below (data not shown). Duplications of the chromosomal sequence GCGGGG are found flanking the two 2-bp-deleted LTRs (Fig. 3).

Retroviruses are supposed to produce equally long duplications of host cell DNA regardless of the host cell (42).

FIG. 2. Nucleotide sequence analysis of the splice junction site of the LTR, v-src, LTR genome. The deletion was found to have occurred between the splice donor site (sd) at position 397 and the acceptor site (sa) for subgenomic src mRNA at position 7,054. The only change near this splice junction is at position 7,059, where G (asterisk) substitutes for A according to the predicted sequence (26). Sequencing was performed with Sequenase in the M13 mp18, M13 mp19, pUC18, and pUC19 cloning system by standard methods (18). Abbreviations: L, viral leader; DR, direct repeat; E, inverted repeat element; PPT, polypurine tract.

FIG. 3. Model of integration of src mRNA into the hamster genome via normal retroviral integration based on actual sequence data (see the text). (A) Nucleotide sequence of the deduced integration site of the circular double-LTR integration precursor of reverse-transcribed src mRNA. LTRs are in open boxes. (B) Deduced target site of hamster chromosomal DNA (hatched box). (C) Sequence of the left edge of the LTR, v-src, LTR genome. (D) Sequence of the right edge of the LTR, v-src, LTR genome.
RSV-related viruses were shown to produce hexanucleotide duplications during integration (11, 12). In accord with this, we have observed a hexanucleotide repeat, GCGGGG. This leads us to suggest a model of integration of src mRNA into the hamster genome via normal avian retroviral reverse transcription and integration (for reviews of retroviral DNA integration, see references 4, 23, and 28).

All of the sequences compared with the nucleotide sequence of PR-RSV-C (26) were highly conserved. Only 1 bp changes were noted (in 1,595 bases examined, two 1-bp changes and one 1-bp deletion were detected). One of the two 1-bp changes found was an A-to-G transition at position 7,059, 6 bases downstream from the src mRNA splice acceptor site (Fig. 2). The same change has been correlated with the increase of src mRNA splicing efficiency (34). Such elevated src mRNA synthesis might have created a situation favorable for its packaging in virions required for generation of the LTR, v-src, LTR provirus in H-19 cells. A deletion of 1 bp, AT, 2 bp upstream from the polyadenylation signal, AATAAA, was found in the U3 region of the 3' LTR but not in the 5' LTR. Because of this location, the deletion should be maintained in virions rescued from H-19 cells and might influence the pattern of the viral RNA polyadenylation signal (6). In agreement with this assumption are our previous findings that the src mRNA present in virions of viruses rescued from H-19 cells is nonpolyadenylated, in contrast to the polyadenylated src mRNA isolated from transformed cells (37).

Here we document for the first time that unaltered src mRNA may be reverse transcribed and integrated in the mammalian genome in a way characteristic of the whole provirus. These results confirm and extend previous conclusions based on restriction analysis and S1 mapping (8, 38). The data presented thus provide additional support, at the molecular level, for Temin's original idea suggesting transmission of an oncogene via an RNA transcript and its integration into the cell genome (41).

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