Cloning and sequence analysis of the cDNA for the precursor of the beta subunit of ovine luteinizing hormone

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Clones containing cDNA for the beta-subunit of ovine luteinizing hormone (LHbeta) were isolated from an ovine pituitary pBR322 cDNA library using a rat LHbeta cDNA as a probe. The complete DNA sequence for the LHbeta subunit, presented in Figure 1, was obtained by sequencing three overlapping recombinant inserts using both the chemical cleavage (1) and chain terminator (2) methods. The 533 base-pair cDNA included the entire coding region for a 141 amino acid LHbeta subunit precursor preceded by a short 5' non coding region as for other mammals (3—5), and followed by a 103 nucleotide long 3' region. Comparison of the amino acid sequence deduced from the open reading frame of the cDNA with that previously established by direct polypeptide sequencing (7) implies a 20 amino acid leader peptide, and a 121 amino acid mature subunit in which we noted four conservative substitutions and a two amino acid extension to the carboxyl terminus giving the ovine LHbeta a size identical to that of the other mammals studied (3—6). In addition, residues 106 and 111 were identified as Gin and Asp, respectively.

Comparison of either the entire cDNA or the coding region of the ovine LHbeta with that of the corresponding bovine sequence (5) gave an overall homology of 98%. Interestingly, although it has long been thought that ovine and bovine LHbeta subunits share complete homology in their primary structures, DNA sequencing of the different clones has revealed two amino acid differences. In addition to the 98% homology with bovine LHbeta, the amino acid sequence of the ovine LHbeta determined in the present study displays an overall homology of 84% with porcine (6), 81% with rat (4), 74% with equine (8), 69% with human (3) and 43% with the chicken (9) or eel (10) sequences.

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REFERENCES


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