Endogenous retroviral sequences are required for tissue-specific expression of a human salivary amylase gene

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The human salivary amylase genes are associated with two inserted elements, a γ-actin-processed pseudogene and an endogenous retroviral-like element. To test the contribution of these inserted elements to tissue specificity, 25 lines of transgenic mice carrying 10 amylase constructs were established. A 1-kb fragment of AMY1C (−1003 to +2) was found to be sufficient for parotid-specific expression of a human growth hormone reporter gene. The 1-kb fragment is entirely derived from inserted sequences. Deletion from −1003 to −826 resulted in reduced levels of transgene expression and loss of tissue specificity. The fragment −1003 to −327 was sufficient to transfer parotid specificity to the thymidine kinase promoter. The data demonstrate that the functional tissue-specific promoter of human AMY1C is derived from inserted sequences and that parotid expression can be conferred by sequences derived solely from the retrovirus. A role for retrotransposition in the evolution of gene regulation is indicated by these and other recent observations.

[Key Words: Human salivary amylase gene; inserted elements; endogenous retroviral sequences; tissue-specific expression; parotid-specific expression]

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The amylase genes provide an interesting model for analysis of the evolution of tissue-specific isozymes. All mammalian species produce amylase in the pancreas, but the only mammals that also produce salivary amylase are primates, rodents, and lagomorphs (for review, see Meisler and Gumucio 1986). We have investigated the origin of salivary amylase expression in the human genome.

Salivary and pancreatic amylase are encoded by distinct but closely related genes (Schibler et al. 1982; Nishide et al. 1986). The human genome contains three salivary and two pancreatic amylase genes (Gumucio et al. 1988; Samuelson et al. 1988; Groot et al. 1989, 1991). Our earlier studies indicated that these genes were derived from one ancestral gene copy during primate evolution (Samuelson et al. 1990). During the evolution of this gene family, insertion of a processed γ-actin pseudogene in the proximal promoter region of the ancestral amylase gene was followed by a retroviral insertion. The 5′-flanking regions of the salivary amylase genes contain both γ-actin and retroviral sequences (Fig. 1). The transcriptional orientation of the retrovirus is opposite that of the amylase gene. Insertion of the retroviral element is correlated with a switch from pancreatic to parotid expression and excision of the retrovirus with reversion to pancreatic expression (Samuelson et al. 1990). The exclusive association of the provirus with salivary amylase genes led us to propose that this inserted element may be responsible for salivary amylase expression in primates.

The amylase-associated retroviral-like elements belong to the family of human endogenous retroviruses designated 4-1, or HERV-E, which contains ~50 members (Rabson et al. 1983; Repaske et al. 1985; Larson et al. 1989). In the current study we tested the role of retroviral sequences in regulating expression of the human salivary amylase gene in transgenic mice.

Results

Transcription of AMY1C in the parotid gland of transgenic mice

Three genomic fragments containing the intact salivary amylase gene AMY1C with different amounts of 5′-flanking sequence were microinjected into fertilized mouse eggs as described in Materials and methods. The structures of the fragments are represented in Figure 2A. Transgenic mice were identified by polymerase chain reaction (PCR) of genomic DNA. RNA was isolated from...
Figure 1. Structure of the human salivary amylase gene AMY1C. Insertions of the γ-actin pseudogene (solid bar) and the retrovirus ERVA1C occurred ~40 million years ago (Samuelson et al. 1990). [●]Exon a and the NTE; the rest of the gene is not shown. The major start site for transcription is indicated by an arrow. Insertion of the retrovirus apparently activated a cryptic promoter within the γ-actin pseudogene.

Tissues of transgenic animals and human amylase transcripts were assayed using riboprobe 1B-2 which contains the nontranslated exon (NTE) and the first coding exon (exon a) of AMY1C (Fig. 3). Human parotid RNA protects four major fragments from this probe (Fig. 3, lane 1). Parotid RNA from line 6713, carrying the N2 transgene, protected the same fragments (lanes 5–7), indicating that the human gene is transcribed accurately. Parotid RNA from nontransgenic mice does not protect this probe (Fig. 3, lane 11), demonstrating the species specificity of the assay. Another NTE probe was protected by parotid RNA from three independent lines carrying N2 and one line carrying N2–Apa, but not by four independent transgenic lines carrying the smallest fragment, N2–Bam (Fig. 4). All samples contained comparable levels of mouse amylase mRNA on Northern blots (data not shown). These results indicate that sequences required for expression in the parotid gland are located between the Apal site at −10 kb and the BamHI site at −826 bp.

Tissue specificity of AMY1C expression

The major site of transcription of these fragments was the parotid gland. In lines carrying the N2 construct, human amylase transcripts could not be detected in brain, submaxillary gland, stomach, intestine, pancreas, liver, spleen, kidney, muscle, or fat with the ribonu...
Using riboprobes complementary to the 5' and 3' long terminal repeats (LTRs), we were unable to detect retroviral transcripts in tissues of transgenic mice or in human parotid RNA (C.-N. Ting and M.H. Meisler, unpubl.).

Expression of human growth hormone fusion genes

To further localize sequences required for expression in the parotid gland, fragments containing the AMY1C promoter and various 5'-flanking sequences were ligated to a human growth hormone (hGH) reporter gene (Fig. 2B). The expression of each construct was tested in two or three independent transgenic lines, by use of a species-specific radioimmunomatrix assay for hGH. Constructs AGH1, AGH2, and AGH3, with 10, 4.2, and 1 kb of 5'-flanking DNA, respectively, were all expressed in the parotid gland (Table 1). Although the level of expression of each construct varied by as much as 20-fold in different independent lines, the range of expression for the three constructs was very similar. These observations indicate that the 1-kb amylase fragment in AGH3 con-

![Figure 3. Tissue distribution of transcripts from the N2 transgene. Total RNA was prepared from tissues of N2 transgenic line 6713 and nontransgenic C57BL/6J mice. The structure of the 1B-2 riboprobe used in the RNase protection assay is indicated (Samuelson et al. 1988). The 140- and 170-bp fragments are derived from the NTE, which is initiated at two major transcription start sites, the 217- and 205-bp products are derived from exon a (Samuelson et al. 1988). Transgenic and nontransgenic parotid RNA contained comparable amounts of intact mouse amylase mRNA, which was detected by Northern blotting and protection assays (not shown). (Lane 1) Human parotid; (lanes 2-7) transgenic; (lanes 8-11) C57BL/6J. (Par) Parotid; (pan) pancreas; (sub) submaxillary gland; (liv) liver.

Three lines carrying the construct N2-Bam did not produce amylase transcripts in any of the 10 tissues tested. In the fourth line, 9053, a low level of aberrant transcripts which lacked the NTE were detected in the parotid gland and several other tissues.
Table 1. Expression of hGH in transgenic mice carrying hGH constructs

| Transgene | Line    | parotid (hGH ng/g tissue wet weight) | heart | pancreas | spleen | liver |
|-----------|---------|-------------------------------------|-------|----------|--------|-------|
| None      | C57BL/6 | 9 ± 1 [7]                           | 5 ± 3 | 9 ± 6    | 14 ± 3 | 3 ± 1 |
| AGH1      | 13984   | 270 ± 22 [7]                        | 33 ± 9| 8 ± 2    | 38 ± 9 | 19 ± 3|
|           | 13987   | 1400 [1]                            |       | 9        |        | 7     |
| AGH2      | 19671   | 260 ± 10 [7]                        | 3 ± 1 | 6 ± 2    | 4 ± 3  | 5 ± 3 |
|           | 19683   | 530 ± 280 (2)                       | 4, 6  | 7, 7     | 2, 9   | 4, 8  |
|           | 14114   | 3900 ± 270 (4)                      | 5, 7  | 4, 5     | 3, 2   | 5, 2  |
| AGH3      | 16694   | 150 ± 10 [4]                        | 5 ± 1 | 14 ± 8   | 6 ± 3  | 7 ± 4 |
|           | 16704   | 3700 ± 100 (4)                      | 11 ± 4| 11 ± 5   | 12 ± 2 | 8 ± 2 |
| AGH4      | 864     | 8 ± 1 [7]                           | 4 ± 1 | 8 ± 2    | 4 ± 1  | 6 ± 3 |
|           | 866     | 25 ± 3 [4]                          | 11 ± 2| 50 ± 21* | 16 ± 5 | 7 ± 2 |
|           | 889     | 30 ± 4 [10]                         |       | 8 ± 3    |        | 4 ± 1 |
| AGH5      | 272     | 270 ± 20 [6]                        | 10 ± 1| 22 ± 5   | 45 ± 8 | 7 ± 1 |
|           | 269     | 380 ± 40 [7]                        | 11 ± 1| 27 ± 6   | 25 ± 10*| 11 ± 1|
|           | 266     | 2600 ± 250 (3)                      |       | 12 ± 2   |        | 5 ± 2 |
| ATK1      | 19021   | 700 ± 140 [4]                       | 12 ± 2| 8 ± 1    | 8 ± 2  | 7 ± 1 |
|           | 128     | 1300 ± 150 (5)                      | 10 ± 1| 20 ± 2   | 6 ± 1  | 40 ± 3|
| ATK2      | 19636   | 8 ± 1 [4]                           | 3 ± 1 | 126 ± 15 | 8 ± 2  | 2 ± 1 |
|           | 19635   | 9 ± 1 [4]                           | 7 ± 1 | 10 ± 1   | 4 ± 1  | 3 ± 1 |

Values are means ± s.e. [n] The number of animals assayed for parotid gland. For other tissues, n = 4-6. When less than three animals were available, the individual values are reported. Values marked with an asterisk (*) are influenced by a single individual with high activity.

Transfer of parotid specificity to a heterologous promoter

To determine whether the amylase promoter is required for parotid-specific expression, two constructs substituting the herpes virus thymidine kinase (tk) promoter were studied. In ATK1, the AMY1C fragment -1003 to -327 was ligated to the tk promoter (Fig. 2B). ATK1 was expressed specifically in the parotid gland of two independent lines, with a level of activity comparable to that of the constructs containing the AMY1C promoter (Table 1). The region -1003 to -327 thus appears to contain all of the parotid-specific enhancer activity of the larger constructs.

As a negative control, the 600-bp Apal–HindIII fragment [-10 to -9.4 kb] lacking enhancer activity was placed in a similar position [ATK2, Fig. 2B]. ATK2 was not expressed in the parotid gland or in other tissues [Table 1]. The lack of expression of this construct demonstrates that the growth hormone and tk sequences do not produce parotid-specific expression of ATK1.

Common sequence elements in the AMY1C enhancer

AMY1C sequence -1003 to -752 is compared with the corresponding region of the human endogenous retrovirus 4-1 in Figure 5. The high degree of sequence identity [85?] demonstrates that this region is derived from prival sequences. The sequence was also compared with seven other genes that are expressed in the parotid gland.
Parotid-specific amylase expression

AMY1C -1003 AAGCTT*GCT CAGGTGGAGT GGGCAAGTTG AAAAGACTTG
HER41  782 .C.....CA...AA...........C CS.....CA.

AMY1C -963 TCTTACTAAG TTTCAGATGT CTGGACTCCA AGTGCCAGTT
HER41  740 .G .... C ........ A ........... GG ....... T...

AMY1C -923 CTTTACTAAG TTTCAGATGT CTGGACTCCA AGTGCCAGTT
HER41  700 .G .... C ........ A ........... GG ....... T...

AMY1C -883 **********CAGGCTGGCC AGAGTCCCTG GCAGCCTGAG GGATGCCCCG
HER41  663 .A ............... CT . .****** ........ T.**

AMY1C -853 GCCTTACTCC ACAAGGCATG CC
HER41  601 ********* .... G .... G. ..

Figure 5. Sequence comparison of AMY1C and the human endogenous retrovirus HER 4-1. Dots represent identity with the AMY1C sequences. Asterisks mark deleted nucleotides. Three elements shared by other parotid-expressing genes are doubly underlined and numbered I, II, and III. The recognition sequence for restriction enzyme BamHI is underlined. The HER 4-1 sequence is from Repaske et al. (1985). The AMY1C sequence reported here differs at six nucleotides from the sequence reported by Emi et al. (1988).

Approximately 1 kb of 5'-flanking region from each gene was searched by computer to detect 10-bp segments with >85% sequence identity to this region of AMY1C. Three common elements were detected [Table 2]. The arrangement of these elements in the human AMY1C and proline-rich protein [PRP] genes is nearly identical, and the two unrelated genes differ at only 3 of 29 nucleotides within the elements. These elements do not appear to be related to known transcriptional regulatory sequences. Although functional information is not available for the other genes, the sequence similarity suggests that the elements may be important components of the amylase enhancer.

Parotid nuclear proteins bind the amylase enhancer

Because deletion of the fragment -1003 to -826 reduces hGH expression in the parotid gland, we tested the affinity of this fragment for nuclear proteins. Parotid nuclear extracts contain protein which binds specifically to this fragment [Fig. 6]. Binding was specifically competed by the same fragment [lanes 3–5] and by the subfragment -885 to -826 which contains conserved element I [data not shown]. Liver nuclear extracts do not contain binding activity [lane 9].

Discussion

We have demonstrated that sequences derived from the endogenous retroviral-like element ERVA1C are responsible for the tissue specificity of the adjacent AMY1C gene. The AMY1C sequence -1003 to +2, which functions as a parotid-specific enhancer/promoter in vivo, is composed of sequences derived entirely from retroviral and γ-actin inserts [Fig. 7]. This tissue-specific regulator is thus derived from the juxtaposition of two unrelated sequences. Transfer of parotid specificity to a heterologous promoter demonstrated the autonomy of enhancer-like elements within the region -1003 to -337. This is the first localization of parotid-specific regulatory elements of which we are aware.

The results confirm the earlier prediction that was based on comparison of the structures of the five human amylase genes. The three salivary amylase genes are each associated with an intact retroviral element. The two pancreatic amylase genes either lack retroviral sequences (AMY2B) or contain a solo LTR as a result of excision of the retrovirus (AMY2A) (Samuelson et al. 1990). The gene structures are consistent with a sequence of events in which insertion of the retrovirus converted a pancreatic amylase gene like AMY2B into a salivary-specific gene, and excision of the retrovirus resulted in a return to pancreas-specific expression of AMY2A. The combined structural and functional information provide strong evidence for the role of retroviral insertion in the evolution of tissue specificity of this gene family.

Comparison of the 5'-flanking region of AMY1C and

Table 2. Common elements in human and mouse salivary gland-specific genes

| Gene       | I            | II          | III         |
|------------|--------------|-------------|-------------|
| hAMY1C     | -849 TTTCCTACC | -802 AGAGTCCCTG | -784 TGAGGGATGC |
| hPRP       | -119 .A....... | -93 ........A. | -77 ........A. |
| mPsp       | -697* .G....... | -16 ..A...A. | -675* ........ATG |
| mPrp       | -332 .A.......T | -92* ....T....A | -104* ........T.A.G |

One kilobase of 5'-flanking sequence from each gene was compared with AMY1C nucleotides -1003 to -752. Three regions with >85% sequence identity were identified. (∗) Reverse orientation; [hPRP] human proline-rich protein gene [Kim and Maeda 1986]; [mPsp] mouse parotid secretory protein gene [Shaw and Schibler 1986]; [mPrp] mouse proline-rich protein gene [Ann and Carlson 1985].
are quite similar, there is no obvious advantage to dupli-
been strong positive selection for salivary amylase at
enzymatic activities of pancreatic and salivary amylases
some points during mammalian evolution. Because the
human AMY1 C, may be derived from a preexisting pan-
ance of rodents and primates. This is consistent with
rotid specificity has arisen independently in the mouse
creatic amylase gene. [Schibler et al. 1982] suggests that the mouse gene, like
creatic amylase genes within each species, compared
leading to complexity and diversity. The presence of
ths of copies of such elements in the mammalian
function for salivary amylase in facilitating oral micro-
bacterial colonization has been proposed [Scannapieco et al.
In the evolution of tissue-specific isozymes, three
mechanisms for duplication of coding sequence have
identified. The most common mechanism, as ob-
erved for the amylase genes, is gene duplication fol-
lowed by divergence of regulation of one gene copy. Sev-
eral testis-specific isozymes have been generated by a
different mechanism, that is, retroposition of processed
scripts with the new regulatory sequences provided
by the insertion site [Boer et al. 1987; McCarrey and
Thomas 1987; Dahl et al. 1990]. In the third case, a single
structural gene is regulated by alternative promoters
with different tissue specificity [Schibler and Sierra
1987]. In all of these situations, the mechanism of origin
of the new regulatory elements is of central importance
to understanding the evolution of diversity. For the hu-
man amylase genes, we have now demonstrated that ret-
roviral insertion contributed to altered tissue specificity.
It has been argued that changes in gene regulation may
be more important to the process of speciation than the
gradual accumulation of structural variation. How gen-
ral is the role of retroviral insertion in changing gene
expression? Two additional examples have recently been
described: insertion of an LTR-derived hormone-re-
sponse element upstream of a mouse complement-re-
lated gene [Stavenhagen and Robins 1988; Adler et al.
1991], and insertion of an LTR-derived nonspecific pro-
ponent upstream of the gonad-specific chicken aromatase
gene [Matsumine et al. 1991]. These examples suggest
that retroviral-like elements and other DNA inserts
could be important vectors of rapid qualitative changes
leading to complexity and diversity. The presence of
other salivary-specific genes identified three short ele-
ents that are present in a similar array in the proximal
promoter region of the human PRP gene. Deletion of
element I resulted in reduced parotid expression in vivo.
It will be of great interest to determine whether these
three elements are sufficient for parotid enhancer activ-
ity and whether they are shared by other salivary-specific
genes.

One surprising implication of these studies is that pa-
rotid specificity has arisen independently in the mouse
and human genomes, as the y-actin and proviral inser-
tions in the human gene were acquired after the diver-
gence of rodents and primates. This is consistent with
the closer sequence similarity of the salivary and pan-
creatic amylase genes within each species, compared
with the similarity of the orthologous genes [for review,
see Meisler and Gumucio 1986]. The fact that the mouse
salivary amylase gene is transcribed from an upstream
NTE that is not present in the pancreatic amylase gene
[Schibler et al. 1982] suggests that the mouse gene, like
human AMY1C, may be derived from a preexisting pan-
creatic amylase gene.
The independent, convergent evolution of salivary
amylase in human and mouse indicates that there has
been strong positive selection for salivary amylase at
some points during mammalian evolution. Because the
enzymatic activities of pancreatic and salivary amylases
are quite similar, there is no obvious advantage to dupli-
cation of the digestive activity per se in two different
organisms. One interesting hypothesis is that the sweet
taste of the sugars produced by the action of salivary
amylase in the mouth could aid in the recognition of
nutritious food sources. Another potentially selectable
function for salivary amylase in facilitating oral micro-
bial colonization has been proposed [Scannapieco et al.
1990].

![Figure 6](image-url)

Figure 6. Specific binding of parotid nuclear protein by the
AMY1C parotid enhancer. The HindIII–BamHI fragment
(−1003 to −826) was radiolabeled and incubated with 10 μg of
nuclear protein, as described in Materials and methods. The
nonspecific competitor was a 120-bp fragment from pBR322
[Howard et al. 1989]. [Lanes 1, 7] Probe alone; [lane 8] nuclear
extract from mouse liver; [lanes 2–6, 9] nuclear extract from
mouse parotid.

![Figure 7](image-url)

Figure 7. The functional parotid-specific promoter of AMY1C
is derived entirely from two inserted elements. Nucleotides
−1003 to −237 are derived from the retrovirus. Nucleotides
−236 to +2 are derived from the 3′-untranslated region of a
y-actin-processed pseudogene. The first codon of the retroviral
insertion is located at nucleotides −1245 to −1247. The com-
plete sequence of this region is available [Fig. 5]. Emi et al. 1988;
Samuelson et al. 1988; Samuelson et al. 1990]. [H] HindIII; [X]
XbaI; [B] BamHI; [S] StuI.
genomes, some of which are capable of transposition [Dombroski et al. 1991; Evans and Palmiter 1991], is consistent with more widespread effects than are currently appreciated.

The human salivary amylase enhancer described here has potential applications in basic and applied research. Direction of oncogene expression to the parotid gland may be used to develop cultured cell lines that retain some of the differentiated characteristics of salivary acinar cells. Such lines are not currently available and would be quite useful in salivary research. Treatments for oral disease may be tested by expression of potentially therapeutic proteins in transgenic animals. Finally, isolation of valuable proteins from saliva of transgenic animals could be practical in some situations.

Materials and methods

Transgenic lines

DNA fragments were electroeluted from agarose gels and prepared for microinjection as described previously [Osborn et al. 1987]. Fertilized mouse eggs were obtained from matings of C57BL/6 × C3H/HeJ mice, except for constructs AGH3 and ATK1, which utilized C57BL/6 × SJL/J mice. Transgenic founders were crossed to inbred strain C57BL/6J, and 25 independent transgenic lines were established. Tissues for analysis were obtained from heterozygous transgenic individuals from generations N1, N2, and N3. The transgene copy numbers in the transgenic lines were estimated by Southern blotting of genomic DNA with transgene-specific probes and comparison with standards as described previously [Jones et al. 1989]. Most lines contained between one and five copies of the transgene, with a higher copy number in lines 9257 (20-25 copies), 13987 (10-20 copies), and 889 (15-20 copies). Mice carrying the hGH fusion genes were within the normal range of body weight.

Construction of hGH fusion genes

To generate the constructs in Figure 2, a 2.1-kb BamHI–EcoRI fragment containing the hGH structural gene was isolated from pOGH [Nichols Institute, Los Angeles, CA] and subcloned into the vector pSP72 [Promega]. The 12-kb DNA fragment extending from the XhoI site (-12 kb) to the XbaI site (+2 bp) of AMY1C was inserted upstream of the hGH gene in pSP72 to generate the cloning intermediate pXGHX. Constructs AGH1, AGH2, AGH3, and AGH4 were isolated from pXGHX by digestion with EcoRI and Apal, XbaI, HindIII, or BamHI, respectively. pAGH5 was constructed by digestion of pXGHX with HindIII and religation. The AGH5 fragment was isolated after digestion of pAGH5 with Apal and EcoRI. The 2.9-kb fragment ATK1 was isolated by HindIII and EcoRI digestion of a derivative of pAGH5 in which the AMY1C promoter (-325 to +2) was replaced with herpes simplex virus tk (-110 to +20) [McKnight 1982]. pATK2 was generated from pAGH5 by replacing AMY1C (-1003 to +2) with tk (-159 to +56). The ATK2 fragment was isolated from pATK2 by digestion with Apal and EcoRI.

Identification of transgenic individuals by PCR of genomic DNA

Transgenic individuals were identified by PCR of genomic DNA isolated from the tail. The AMY1C primers HTA-C (5'-

CACCAT TGG GTT CTG GGC TCA GTA TTC-3') and HTA-N (5'-CGG TCA CAT ACA AGA GCA ATA TCA ACC CAT-3') amplify exon a of AMY1C from +584 to +704 [Gumucio et al. 1988] and do not amplify mouse amylase. Reactions contained 1 µg of genomic DNA and 2 units of Taq DNA polymerase in 10 mm Tris-HCl [pH 8.3], 50 mm KCl, 1.5 mm MgCl2, 0.01% BSA, with 0.2 µm dNTPs and 0.5 µm concentration of each primer. Reactions were subjected to 24 cycles alternating between 94°C (75 sec) and 72°C (3 min). The 110-bp product was detected on 6% polyacrylamide gels stained with ethidium bromide. PCR primers for the hGH gene amplify a 330-bp fragment (+410 to +729) [DeNoto et al. 1981]: hGH-C (5'-CCA CAA ATT CCC TTA TCC AGG CTT TTT GAC-3') and hGH-N (5'-TAC TTC TGT TCC TTT GGG ATA TAG CTC TCT-3'). Amplification of hGH was carried out with the solutions described above for 29 cycles at 94°C (30 sec), 60°C (90 sec), and 72°C (2 min). The products were analyzed on polyacrylamide gels as described above.

Ribonuclease protection assay

RNA was isolated from various tissues by homogenization in guanidine thiocyanate, followed by centrifugation through cesium chloride as described by Samuelson et al. [1988]. RNA concentrations were determined by OD260 and quality was assessed by examination of the 28S and 18S rRNA bands after electrophoresis through agarose and staining with ethidium bromide. Human-specific riboprobes 1B-2 and Act-2 were used to detect the presence of the human amylase transcripts by ribonuclease protection assay as described previously [Samuelson et al. 1988]. Single-stranded and uniformly labeled riboprobes were generated by use of [α-32P]UTP [800 Ci/mM, Amersham] according to the procedure recommended by Promega Biotec. The protected products were detected by autoradiography after electrophoresis in 6% polyacrylamide gels with 8 M urea. Parotid RNA samples were prepared from pooled glands of six individuals.

Radioimmunomatrix assay of hGH

Tissue samples from transgenic mice (15–60 mg wet weight) were homogenized in 1 ml of 0.85% NaCl with a Polytron homogenizer for 5 sec and centrifuged at 4°C for 5 min. Supernatant hGH was measured with a solid-phase two-site radioimmunomatrix assay kit by use of 125I-labeled and biotin-coupled anti-hGH antibodies [Nichols Institute]. Samples were counted in a Beckman Gamma Counter 5500. Tissue homogenates were diluted to levels within the linear range of the assay, which was 1–50 ng/ml (~500–25,000 cpm).

Saliva was collected from mice 5 min after intraperitoneal injection of 0.1 ml of pilocarpine nitrate (Sigma) (6 mg/ml). Fifty microliters of saliva was diluted with 200 µl of 0.85% NaCl for radioimmunomatrix assay.

Sequence analysis

Sequencing reactions were carried out by use of the Sequenase kit [U.S. Biochemical, v. 2.0]. The 252-bp HindIII–SpII fragment containing nucleotides -1003 to -752 of AMY1C was subcloned into pSP72, and both strands were sequenced by using the T7 and SP6 primers. Sequencing products were resolved on a 6% polyacrylamide gel with 8 M urea. Sequence alignments of AMY1C and other genes were performed on an IBM PC computer with the aid of Pustell DNA analysis software [International Biotechnology, Inc.]
Isolation of nuclei and gel retardation assay

Parotid glands were dissected from 60 mice and stored frozen at −70°C. Nuclei were prepared from frozen tissue by the method of Blobel and Potter [1966], with modifications. Frozen tissues were pulverized in a precooled mortar and pestle and homogenized in buffer A with 0.32 M sucrose and centrifuged at low speed for 30 sec to remove debris and unbroken cells. The supernatant was carefully layered on an equal volume of buffer A containing 0.88 M sucrose and centrifuged at 800g for 5 min at 4°C. The nuclear pellet was collected, and protein was extracted by the method of Dignam et al. [1983]. Dialysis and quantitation of protein were carried out as described by Howard et al. [1989]. Isolated DNA fragments were radiolabeled with [α-32P]dCTP by use of the Klenow fragment of DNA polymerase (Boehringer Mannheim). Gel retardation assays were performed as described by Dignam et al. [1983]. Samples were run on 4% polyacrylamide gels (acrylamide/bisacrylamide, 25:1) at 25 mA for 2 hr in a buffer containing 45 mM Tris (pH 8.0), 45 mM borate, and 1 mM EDTA. Gels were dried and visualized by autoradiography.

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