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Hyaluronan Synthase of Chlorella Virus PBCV-1

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Hyaluronan or hyaluronic acid (HA), a member of the glycosaminoglycan family that also includes heparin and chondroitin, is a linear polysaccharide composed of alternating β1,4-glucuronic acid (β1,4-GlcA) and β1,3-N-acetylg glucosamine (β1,3-GlcNAc) groups. Typically the full-length polymer chains are composed of $10^3$ to $10^4$ monosaccharides ($10^6$ to $10^7$ daltons). HA is an important structural element in the vitreous humor of eye, synovial fluid, and skin of vertebrates (1). Furthermore, HA interacts with proteins such as CD44, RHAMM, and fibrinogen, thereby influencing many natural processes such as angiogenesis, cancer, cell motility, wound healing, and cell adhesion (2). HA also constitutes the extracellular capsules of certain bacterial pathogens such as group A and C Streptococcus and Pasteurella multocida type A (3, 4). These capsules act as virulence factors that protect the microbes from phagocytosis and complement during infection (5, 6). Because HA, a component of the host tissues, is not normally immunogenic, the capsule serves as molecular camouflage (7).

HA synthases (HASs) are integral membrane proteins that polymerize the HA molecule using activated uridine.
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diphosphate (UDP)-sugar nucleotides as substrates. Amino acid sequences for some HASs have been deduced from gene sequencing (8); their sizes range from 419 to 588 residues. The vertebrate enzymes (DG42, HAS1, HAS2, and HAS3) and streptococcal HasA have several regions of sequence similarity. Recently, while sequencing the doublestranded DNA genome of virus PBCV-1 (Paramecium bursaria chlorella virus), we unexpectedly discovered an open reading frame (ORF), A98R (GenBank accession number U42580), encoding a 568-residue protein with similarity to the known HASs (28 to 33% amino acid identity in pairwise comparisons by FASTA) (Figure 1).

On the basis of its deduced amino acid sequence, the A98R gene product should be an integral membrane protein. To test this hypothesis, we produced recombinant A98R protein in Escherichia coli and assayed the membrane fraction for HAS activity (12, 13). UDP-GlcA and UDP-GlcNAc were incorporated into polysaccharide by the membrane fraction derived from cells containing the A98R ORF on a plasmid, pCVHAS, (average specific activity of 2.5 pmol of GlcA transferred per minute per microgram of protein), but not by samples from control cells (< 0.001 pmol of GlcA transferred per minute per microgram of protein). No activity was detected in the soluble fraction of cells transformed with pCVHAS. UDP-GlcA and UDP-GlcNAc were simultaneously required for polymerization. The activity was optimal in Hepes buffer at pH 7.2 in the presence of 15 mM MnCl$_2$, whereas no activity was detected if the metal ion was omitted. The ions Mg$^{2+}$ and Co$^{2+}$ were ~20% as effective as Mn$^{2+}$ at similar concentrations. The P. multocida HAS (14) has a similar metal requirement, but other HASs prefer Mg$^{2+}$.

We also tested the specificity of recombinant A98R for UDP-sugars (15). Only the two authentic HA precursors were incorporated into polysaccharide; neither UDP-galacturonic acid (UDP-GalA) nor UDP-N-acetylgalactosamine (UDP-GalNAc), the C4 epimers of UDP-GlcA or UDP-GlcNAc, respectively, were incorporated. Likewise, UDP-glucose (UDP-Glc) was not polymerized in place of either HA precursor. This strong substrate specificity

Figure 1. Sequence similarity of HASs. The Multalin program (26) was used to align the amino acid sequences of HASs Xenopus laevis DG42, human HAS2, PBCV-1 A98R, and Streptococcus pyogenes HasA (red, 90% consensus; green, 50% consensus, as calculated by Multalin) (8). In the consensus sequence, the symbols are: !, any one of I or V; $, any one of L or M; %, any one of F or Y; #, any one of N, D, E, or Q. Single-letter abbreviations for the amino acid residues are as follows: A, Ala; C, Cys; D, Asp; E, Glu; F, Phe; G, Gly; H, His; I, Ile; K, Lys; L, Leu; M, Met; N, Asn; P, Pro; Q, Gln; R, Arg; S, Ser; T, Thr; W, Trp, and Y, Tyr.

Figure 2. Size exclusion chromatography of polymer product of recombinant A98R HAS. Membranes derived from E. coli cells transformed with pCVHAS were incubated with both radiolabeled HA precursors diluted to the same specific activity (27). After deproteinization and removal of unincorporated precursors, samples were injected onto a Sephacryl S-500HR size exclusion column, and the radioactivity in the fractions was measured ($^3$H, solid squares; $^{14}$C, solid circles). A duplicate sample was treated with HA lyase before deproteinization and chromatography ($^3$H, open squares; $^{14}$C, open circles); no polymer remains after digestion. Size standards: $V_o$, arrow, void volume, HA derived from recombinant streptococcal HasA (17 ml; ≥2 × 10$^7$ daltons) (13); crosshatched box, blue dextran (29 to 32 ml; average molecular size 2 × 10$^6$ daltons; Pharmacia); $V_t$, arrow, totally included volume, UDP-sugars (37 ml).
for UDP-GlcA and UDP-GlcNAc is a general feature of the HASs HasA (13) and DG42 (16).

The recombinant A98R enzyme synthesized a polysaccharide with an average molecular size of $3 \times 10^6$ to $6 \times 10^6$ daltons (Figure 2), which is smaller than that of the HA synthesized by recombinant HasA or DG42 in vitro ($10^7$ to $8 \times 10^7$ daltons, respectively) (13, 16). The polysaccharide was completely degraded by Streptomyces hyalurolyticus HA lyase, an enzyme that depolymerizes HA but not structurally related glycosaminoglycans such as heparin and chondroitin (17).

We examined PBCV-1-infected chlorella cells for A98R gene expression. A ~1700-nucleotide A98R transcript appeared about 15 min after infection and disappeared by 60 min after infection (18), indicating that A98R is an early gene. Consequently, we assayed membrane fractions from uninfected and PBCV-1-infected chlorella cells at 50 and 90 min after infection for HAS activity. Infected cells, but not uninfected cells, had activity (Table 1). Like the bacterially derived recombinant A98R enzyme, radioactive label incorporation from UDP-$^14$C-GlcA into polysaccharide depended on both Mn$^{2+}$ and UDPGlcNAc. This labeled product was also degraded by HA lyase. Disrupted PBCV-1 virions had no HAS activity.

PBCV-1-infected chlorella cells were analyzed for HA polysaccharide by means of a highly specific 125I-labeled HA-binding protein (19, 20). Extracts from cells at 50 and 90 min after infection contained substantial amounts of HA (0.7 and 1400 ng per microgram of protein, respectively), but not extracts from uninfected algae (< 0.04 ng per microgram of protein) or disrupted PBCV-1 virions (< 0.04 ng per microgram of dry weight). The labeled HA-binding protein was also detected with intact infected cells at 50 and 90 min after infection, but not with healthy cells (21). Therefore, a considerable portion of the newly synthesized HA polysaccharide was immobilized at the outer cell surface of the infected algae. The extracellular HA does not play any obvious role in the interaction between the virus and its algal host because neither plaque size nor plaque number was altered by including either testicular hyaluronidase (465 units/ml) or free HA polysaccharide (100 μg/ml) in the top agar of the PBCV-1 plaque assay (9).

Among chlorella viruses, HA biosynthesis during infection is not limited to the PBCV-1 prototype strain. Thirty-three independently isolated and plaque-purified viruses from the United States, South America, Asia, and Australia were tested for the presence of an A98R-like gene and for the ability to direct production of HA polysaccharide in Chlorella NC64A. Dot-blot hybridization analyses of the individual viral genomes with the PBCV-1 A98R probe indicated that 19 isolates (58%) had a similar gene; the algal host DNA did not cross-react with the probe (21). Chlorella cells infected with each of these 19 viruses produced cell surface HA as measured by interaction with the 125I-HA-binding protein (21).

Surprisingly, the PBCV-1 genome also has additional genes, named A609L and A100R, that encode for a UDP-Glc dehydrogenase (UDP-Glc DH) and a glutamine:fructose-6-phosphate amidotransferase (GFAT), respectively. UDP-Glc DH converts UDP-Glc into UDP-GlcA, a required precursor for HA biosynthesis. GFAT converts fructose-6-phosphate into glucosamine-6-phosphate, an intermediate in the UDP-GlcNAc metabolic pathway. Both of these PBCV-1 genes, like the A98R HAS, are expressed early in infection and encode enzymatically active proteins (22); however, these three genes do not function as an operon. Although two of these genes, A98R and A100R, are near one another in the viral genome (bases 50,901 to 52,607 and 52,706 to 54,493, respectively), A609L is located ~240 kb away and is transcribed in the opposite orientation (bases 292,916 to 291,747). The presence of multiple enzymes in the HA biosynthesis pathway indicates that HA production must serve an important function in the life cycle of these chlorella viruses.

The details of the natural history of the phycodnaviruses are unknown. These viruses are ubiquitous in freshwater collected worldwide, and titers as high as $4 \times 10^6$ infectious viruses per milliliter of native water have been reported (23). The only known hosts for these viruses are chlorella-like green algae, which normally live as hereditary endosymbionts in some isolates of the ciliate, P. bursaria. In the symbiotic unit, algae are enclosed individually in perigal vacuoles and are surrounded by a host-derived membrane (24). The endosymbiotic chlorella are resistant to virus infection and are only infected when they are outside the paramaecium (9). We hypothesize that HA synthesis and its accumulation on the algal surface may block the uptake of virus-infected algae by the paramaecium. Alternatively, the chlorella viruses might have another host in nature (such as an aquatic animal); perhaps the virus is transmitted because this other host is attracted to or binds to the HA polysaccharide on virus-infected algae.

As depicted in Figure 1, HASs of Streptococcus, vertebrates, and PBCV-1 have many motifs of two to four residues that occur in the same relative order. These conserved motifs probably reflect domains crucial for HA biosynthesis. Regions of similarity between HASs and other enzymes that synthesize β-linked polysaccharides from UDP-sugar precursors are also being discovered as more glycosyltransferases are sequenced (25). The significance of these similar structural motifs will become more apparent as the three-dimensional structures of glycosyltransferases are determined.

The fact that Chlorella virus PBCV-1 encodes a functional glycosyltransferase that can synthesize HA is contrary to the general observation that viruses either (i) use host cell glycosyltransferases to create new carbohydrate structures, or (ii) accumulate host cell glycocjugates during virion maturation.

### Table 1. HAS activity of membranes derived from Chlorella cells infected with PBCV-1.

<table>
<thead>
<tr>
<th>Sample</th>
<th>UDP-GlcNAc</th>
<th>Mn$^{2+}$</th>
<th>HAS specific activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uninfected</td>
<td>+</td>
<td>+</td>
<td>≤ 6</td>
</tr>
<tr>
<td></td>
<td>–</td>
<td>–</td>
<td>≤ 6</td>
</tr>
<tr>
<td>50 min a.i.</td>
<td>+</td>
<td>+</td>
<td>42</td>
</tr>
<tr>
<td></td>
<td>–</td>
<td>–</td>
<td>≤ 6</td>
</tr>
<tr>
<td>90 min a.i.</td>
<td>–</td>
<td>–</td>
<td>170</td>
</tr>
<tr>
<td></td>
<td>+</td>
<td>–</td>
<td>≤ 6</td>
</tr>
<tr>
<td></td>
<td>–</td>
<td>+</td>
<td>≤ 6</td>
</tr>
</tbody>
</table>
Furthermore, HA has been generally regarded as restricted to animals and a few of their virulent bacterial pathogens. Though many plant carbohydrates have been characterized, to our knowledge, neither HA nor a related glycosyltransferase has been characterized, to our knowledge. Though many plant carbohydrates have been characterized, to our knowledge, neither HA nor a related glycosyltransferase has been characterized.

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12. The A98R ORF was cloned after amplification of genomic viral DNA
    with 22 cycles of polymerase chain reaction (PCR) with Taq polymerase
    [F. M. Ausubel et al., in Short Protocols in Molecular Biology
    (Wiley, New York, ed. 3, 1995). The oligonucleotide primers in bases
    S'-gacagctgATGGTTTACACCTATATG-3'; antisense S'-gagggcgg-CTCACACTGAGATTCAAT-3';
    Great American Gene Company] contained Nco I or Sal I restriction sites
    (underlined, respectively) flanking the ORF encoding 588 amino acids
    (uppercase letters). Codons 4 and 5 were altered to optimize bacterial
    expression. The PCR product was purified, digested with Nco I and Sal I, and ligated into
    a modified version of the plasmid pET-
    BC [B. A. Moffatt and F. W. Studier, J. Mol. Biol. 189, 113 (1986)] (it has an extra Sal I site
    in the polylinker) cleaved with Nco I and partially digested with Sal I. This construct
    placed the A98R ORF under the control of a T7 phage promoter. The resulting plasmid,
    pCVHAS, was transformed into the expression host, E. coli BL21 (DE3). The A98R protein
    was expressed by induction with 1 mM isopropylthigalactoside. After 3 to 5 hours
    of further growth, the membrane fraction was isolated [1]. Control membrane
    preparations were made from cultures with the same vector containing an irrelevant gene (a
    protein kinase). Total protein was measured according to M. M. Bradford
    [Anal. Biochem. 72, 248 (1976)]. The paper chromatography method was used to assay for HAS activity
    [13].
15. Assays with E. coli-derived A98R were incubated with 120 μM UDP-[14C]GlcA or with 150 μM UDP-[3H]GlcNAc. The reactions also contained one unlabelled sugar nucleotide (an authentic precursor
    for HAS catalysis) and 75 μM of the other nucleotide, or with 150 μM UDP-[3H]GlcNAc, present was
    detected if an unnatural UDP-sugar was substituted for UDP-GlcA and UDP-GlcNAc, or if only a
    single precursor was present.
18. M. V. Graves, D. Landstein, J. L. Van Etten, unpublished results.
20. A competitive radiometric assay based on a [125I]-labeled HA-binding protein (19)
    (Phar-macia Hab TA) was used to measure the amount of HA in disrupted (freeze-thawed)
    virus particles or the cultures of NC64A cells. The cells were disrupted by vigorous
    agitation with glass beads (1-mm beads, agitated for 3 min four times; Biospec Mini-
    Beadbeater-8). The cell lysate was clarified by centrifugation (15,000
    ×g, 5 min) before the deproteinization (Microcon 1000, ultracentrifugation)
    step. This treatment degraded the radioactive polymer to small oligosaccharides
    (tetramers and hexamers) that were removed by ultrafiltration before gel
    filtration chromatography.
27. Membranes (860 μg of protein) were incubated with 120 μM UDP-[14C]GlcA (0.36
    μCi) and 840 μM UDP-[3H]GlcNAc (2.6 μCi) in 300 μl of 50 mM Heps, pH 7.2, with 15 mM
    MnCl2, for 3 hours at 37°C. EDTA (18 mM final concentration) was then added
    to stop the HAS activity. Half of the reaction was deproteinized by treatment with 0.5%
    SDS (w/v) and Pronase (final concentration of 200 μg/ml, 5 hours at 37°C, Boehringer
    Mannheim). Unincorporated precursors and other small molecules were removed
    by ultrafiltration (Microcon I0, 105 daltons cutoff; Amicon). Half of this semipurified
    sample was injected onto a Sephacryl S-500HR column (1 cm by 50 cm; Pharmacia)
    equilibrated in 0.2 M NaCl, 5 mM tris, pH 8 (0.5 ml/min, 1-ml fractions). To verify that the
    identity of the labeled polysaccharide was HA, we treated the other half of the
    original reaction with HA lyase (30 units at 37°C overnight; Sigma) before the depro
    tinization step. This treatment degraded the radioactive polymer to small oligosaccharides
    (tetramers and hexamers) that were removed by ultrafiltration before gel
    filtration chromatography.
28. Two cultures of NC64A cells (0.9 liter; 1.9 × 1010 cells) were infected with PBCV-1
    (multiplicity of infection of 5) and incubated for 50 or 90 min after infection. Another
    culture served as an uninfected control. The cells were harvested, and the mem
    brane fraction (yield ~3 mg of protein) was prepared as described [P. L. DeAngelis
    and A. M. Achyuthan, J. Biol. Chem. 271, 23657 (1996)], except that 1mM mercaptoethanol
    was substituted for diethiothreitol. The paper chromatography method was used to assay
    for HAS activity [13].
29. We thank A. M. Achyuthan, G. M. Air, M. K.
    Brakke, R. D. Cummings, L. C. Lane, M. Nelson, and P. H. Weigel for helpful discussions.
    R. A. Steinberg provided the plasmids and host strain for T7 expression system. Sup
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