Crystal Structure of Hyaluronidase, a Major Allergen of Bee Venom

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aminoglycan of vertebrate extracellular spaces and is specifi- bound PH-20 protein [6], which is essential for fertilization, and cally degraded by a b**-1,4 glycosidase. Bee venom hyaluronidase the human lysosomal enzymes Hyal-1 [7] and Hyal-2 [8] that dases, which are involved in fertilization and the turnover of enzymes are Hya [6, 9] and PH-20, which also catalyses the** HA. On the basis of sequence similarity, mammalian enzymes transglycosylation reaction [10, 11]. PH-20 protein, in the acro**and Hya are assigned to glycosidase family 56 for which no somal membrane of the sperm, degrades the HA-rich matrix structure has been reported yet. in which cumulus cells surrounding the egg are embedded,**

Hya was determined at 1.6 A˚ resolution. The overall topology basis of sequence similarity mammalian and insect hyaluroniis composed of only seven strands. A long substrate binding hydrolases [13, 14]. groove extends across the C-terminal end of the barrel. Cocrys- Native Hya isolated from bee venom [9] is a single polypeptallization with a substrate analog revealed the presence of a tide composed of 350 residues. It is secreted as a basic glyco-HA tetramer bound to subsites 2**4 to** 2**1 and distortion of the protein with a carbohydrate content that accounts for 7% of**

an acid–base catalytic mechanism, in which Glu113 acts as otic (*Escherichia coli***) and eukaryotic (***Baculovirus***) hosts [15]. the proton donor and the** *N***-acetyl group of the substrate is Although native-like enzymatic activity and IgE binding capacthe nucleophile. The location of the catalytic residues shows ity were observed for the Baculovirus-expressed protein, markstriking similarity to bacterial chitinase which also operates via edly lower values were obtained with Hya expressed in** *E. coli***. a substrate-assisted mechanism. Hya is an endo-***N***-acetyl-D-hexosaminidase that specifically**

mals, insects, leeches, and bacteria [1, 2]. These enzymes hy- tures of other glycosyl hydrolases show usually at least two drolyse hyaluronic acid (HA), a high molecular weight nonsul- acidic residues in the active site. In lysozyme [16], chitinase fated linear glycosaminoglycan, which is composed of repeating [17], and cellobiohydrolase [18] these are aspartate and gluta-

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disaccharide units, D-glucuronic acid (GlcA), and *N***-acetyl glucosamine (GlcNAc). They also hydrolyse chondroitin sulfates and, to a small extent, dermatan sulfate. HA is found in almost *Division of Structural Biology all tissues and body fluids and is particularly abundant in the Biozentrum soft connective tissues of cartilage, synovial fluid, and the University of Basel vitreous humor of the eye. In the extracellular matrix HA is a CH-4056 Basel component of the ground substance that connects protein Switzerland filaments, collagen fibers, and the connective tissue cells. The †Polytech S.C.R.L. structural properties and biological role of HA are well studied 34012 Trieste [3, 4]. The structural role of HA as a stabilizer, lubricant, and Italy shock absorbant is based on its viscoelastic properties, which ‡Laboratory of Immunochemistry are dependent on the concentration and molecular weight of US Food and Drug Administration HA. The level of HA is markedly elevated during embryogene-Rockville, Maryland 20852 sis, wound healing, malignant transformation,, and whenever §CCLRC Daresbury Laboratory fast tissue turnover and repair is required (see recent review** Daresbury, Warrington, Cheshire WA4 4AD by R Stern and TB Csóka, on mammalian hyaluronidases on **United Kingdom the website http://www.glycoforum.gr.jp). In vertebrates, the Example 1** Iziegler Hospital Bern **the same of the set of the set of HA** is controlled by hyaluronidase, which is an **Medical Clinic endoglycosidase that acts jointly with two exoglycosidases, Postfach CH-3001 Bern** b**-glucuronidase, and** b**-***N***-acetyl glucosaminidase. Hyaluroni-Switzerland dase from bee venom (Hya) specifically degrades HA in the extracellular matrix of skin, thereby facilitating penetration of venom constituents into the body. Hya shares greater than 50% se-Summary quence identity with hyaluronidases from other hymenoptera [5] and, interestingly, also with several mammalian enzymes (as much Background: Hyaluronic acid (HA) is the most abundant glycos- as 30% identity). These include the sperm GPI-membrane-(Hya) shares 30% sequence identity with human hyaluroni- are involved in HA turnover (Figure 1). The best characterized enabling sperm–egg adhesion. It has been shown that PH-20 Results: The crystal structure of recombinant (Baculovirus) is an effective antigen for immunocontraception [12]. On the** resembles a classical (β/α ₈ TIM barrel except that the barrel dases have been classified as belonging to family 56 of glycosyl

2**1 sugar. the protein mass [9], and contains two disulfide bridges and four potential glycosylation sites (Asn–X–Thr, where X is any Conclusions: The structure of the complex strongly suggest amino acid). Recombinant Hya has been expressed in prokary-**

cleaves the b**-1,4 glycosidic bond between GlcNAc and GlcA Introduction of long HA chains. The end product of exhaustive hydrolysis is the tetrasaccharide GlcA–GlcNAc–GlcA–GlcNAc [10, 11]. The Hyaluronidases are widespread in nature, being found in mam- catalytic mechanism of Hya is unknown. However, the struc-**

Figure 1. Sequence Alignment of Bee Venom Hya with Human Hyaluronidases

The proteins are listed from top to bottom: lysosomal hyaluronidase Hyal-2 (residues 1–355), sperm PH-20 protein (1–353), lysosomal hyaluronidase Hyal-1 (1–349), and bee venom Hya (entire sequence, residues 1–350). Fully conserved residues are shown in red, partially conserved residues in yellow. Secondary structure elements for Hya are shown below the sequences: green arrows represent β strands, red cylinders α helices, and **orange cylinders 310 helices. Blue arrowheads denote the active-site residues Asp111 and Glu113, as well as Glu247 of bee venom Hya. The four cysteine residues forming two disulfide bridges, Cys22–Cys313 and Cys189– Cys201, are marked by black arrowheads. The sequence of mature Hya starts with Thr (TPDN…), as determined by protein sequencing [70] (D Hoffman, personal communication), that is, one residue earlier than reported by Kreil and coworkers [6], based on nucleotide sequencing.**

shown that Hya and phospholipase A2 are the major allergens studies combined with biochemical and immunochemical

mate, whereas two glutamate residues are found in xylanase because they can induce pathogenic reactions in the majority [19] and b**-amylase [20, 21]. Of these, one carboxylate group of patients allergic to bee venom (71% of patients had specific** functions as the acid/base catalyst and the other as the cata-

serum IgE to recombinant Hya and 78% to recombinant phos**lytic nucleophile [22]. The mechanism proceeds either via a pholipase A2 [26]). Nonallergic control individuals had no IgE double or a single nucleophilic substitution, resulting in reten- antibodies to either of the two recombinant enzymes whereas tion or inversion of the configuration of the anomeric carbon, 15% showed some IgE antibodies to whole bee venom, indicatrespectively. Some glycosidases that act on** *N***-acetylglucos- ing a superior diagnostic specificity of recombinant allergens amine-containing saccharides, such as chitinolitic enzymes, compared to whole bee venom. The allergic response is initilack the proteinaceous nucleophile and the** *N***-acetyl group of ated by allergens that cross-link the Fc-receptor-bound IgE the substrate acts as the nucleophilic base in a substrate- antibodies on the surface of mast cells. This is followed by assisted reaction (see [23] and references therein). The active- rupture of the mast cell membrane and the release of stored site residues of Hya have not yet been identified. However, mediators, such as histamine, which are responsible for the mutagenesis studies on the human sperm PH-20 protein sug- immediate hypersensitive reaction. A promising approach to gested Asp111, Glu113, and Glu247 as candidates [24]. inhibit the onset of allergy would be to prevent allergen binding The venom of honey bee contains proteins that can induce to IgE. Knowledge of the molecular interactions between the life-threatening allergic reactions in humans [25]. It has been allergen and the IgE antibodies may be obtained from structural**

Table 1. Diffraction Data and Phasing

^a Values in parentheses refer to the highest resolution shell.

b $R_{iso} = \sum ||F_{PH}| - |F_p|\sum |F_p|$, where F_p and F_{PH} are the structure factors of native and derivative crystals.
 c $R_{Cullis} = \langle \text{lack of closure} \rangle / \langle \text{isomorphous difference} \rangle$.

 h **Phasing power** = $\langle \text{F}_{\text{H}} / \text{lack}$ of closure).

Crystals of the monoclinic form are in space group P2, with unit-cell dimensions a = 39.9 Å, b = 90.2 Å, c = 49.1 Å, β = 92.4°; crystals of the triclinic form **are in space group P3221 with unit-cell dimensions a** 5 **71.2 A˚ , b** 5 **71.2 A˚ , c** 5 **152.1 A˚ . All data sets were collected from one crystal each on MAR-research image-plate detectors.**

mined in two different crystal forms and in complex with an HA is common to many hydrolases [28]. However, in Hya only oligomer, provides insight into the mode of substrate binding seven strands (β 1– β 7) form the barrel that is surrounded by α **and the catalytic mechanism. The structure of Hya will be used to helices A–J. There is a lack of closure between strands** b**1 and model the sequence-related mammalian enzymes and, hopefully,** b**2, which do not interact through hydrogen bonding, leaving** will lead to identification of the mechanism that triggers the a wide gap (7–8 Å) that could accommodate an additional **allergic response caused by the sting of the honey bee. strand (Figure 3). Loop 34–49 partly fills this gap with two**

Two crystal forms of Hya, monoclinic and trigonal, were used between strands 3 and 4 (helices C and D) and 4 and 5 (helices for the structure determination. Two heavy-atom derivatives E and F) and two β hairpins comprising residues 63–75 and **were found for the monoclinic form (Table 1). The resulting 269–279 at the C-terminal end of** b**2 and** b**6, respectively. Hya multiple isomorphous replacement (MIR) map at 2.7 A˚ resolu- contains two disulfide bridges (Figure 3b). The first, Cys189– tion was rather poor, but allowed the building of a partial model. Cys201, stabilizes the base of a long loop at the C-terminal For phase improvement multi-crystal averaging was employed. For this, the experimental electron density corresponding to** one molecule was transferred into a large P1 cell employing a **mask based on the partial model. The derived structure factors were used for molecular replacement to solve the trigonal crystal** form. An outstanding solution with an R factor of 42% was found. Cyclic electron-density averaging between the two crys**tal forms to 2.7 Å** resolution resulted in a markedly improved map. Finally, after tracing most of the chain in the monoclinic form, the model was subjected to conventional refinement us-**No. of ligand molecules ⁵³ ing the high-resolution synchrotron data set to 1.6 A˚ resolution** (Table 2). The final model consists of residues 10–330. The conformations of the nine N-terminal and the 20 C-terminal **residues (plus the His₆ tail) are not defined by electron density.** In addition, density is missing for loop segments comprising **residues 65-71, 193, 194, and 255-259. The model shows good stereochemistry and has a free R factor of 22.7% (Table 2). An) example of the final 2F_o–F_c map is shown in Figure 2.**

Overall Fold and Structure Comparison
Hya is a globular single-domain protein with approximate core
dimensions 52 × 44 × 39 Å. It comprises ten α helices, 11 β
dimensions 52 × 44 × 39 Å. It comprises ten α helice

characterization. Such information might provide the basis for strands, six 3₁₀ helices (one comprising seven residues and the **the development of allergen-specific forms of therapy. others three residues each; Figure 3). The overall fold (Figures 3 The X-ray structure of hyaluronidase from bee venom, deter- and 4) resembles that of a classical (**b**/**a**)8 TIM barrel [27], which polar mainchain atoms of Leu39 engaged in parallel** β sheet **Results and Discussion hydrogen-bonding interactions with the mainchain atoms of Val13 and Trp15 of** b**1. Other unusual features, that deviate Structure Determination from the regular (** β $/\alpha$)₈ barrel, are the presence of two α helices

Figure 2. Representative Section of the Final Electron Density Map

Close-up stereo view of the segment 111– 116, including the proton donor Glu113, overlaid onto the final SigmaA-weighted 2F_o-F_c **map [71] contoured at 1**s**. Figures 2, 3, and 5–7 were produced using the program DINO (A. Philippsen, http://www.biozentrum.unibas. ch/**z**xray/dino).**

end of b**4, whereas the Cys22–Cys313 bridge joins distant malian hyaluronidases as shown in Figure 5b. Grooves of simisecondary structure elements, a 310 helix near the N terminus lar shape have been found in several endoglycosidases where** and α helix I near the C terminus. The Hya sequence contains they form the binding site for long polysaccharide substrates **four potential glycosylation motifs (Asn–X–Thr) at asparagine [16, 19, 31]. In Hya, a number of conserved residues (Asp111, residues 4, 83, 191, and 231. Asn191 and Asn231 can be ruled Glu113, Arg116, Arg244, Tyr55, Trp123, Tyr184, Tyr190, Tyr227, out as glycosylation sites because they are buried in the pro- Tyr265, Trp301, and Ser225) line the groove (Figure 6). A large tein. The N terminus including Asn4 is not defined by electron number of aromatic residues are observed, reminiscent of density, whereas weak electron density extends from Asn83, other sugar binding proteins. The two conserved arginines** corresponding to two sugar residues. **observed at the opposite walls of the binding groove**, probably

A search for proteins with a similar fold was performed using contacts and is located far from the active site. the Dali [32] server. The top scoring proteins were all glycosidases with a regular (b**/**a**)8 barrel fold (Table 3), with the excep- Structure of the Enzyme–Hyaluronic Acid Tetramer Complex tion of the seven-stranded barrel of thermophilic endocellulase Hya was cocrystallized in the presence of a HA-hexamer (GlcA– E2** [31]. The observed similarity is mainly confined to the β GlcNAc)₃, modified with a fluoresceinamine group at C1 of the barrel and the surrounding α helices, with the β strands in reducing sugar (see the E **barrel and the surrounding** α helices, with the β strands in reducing sugar (see the Experimental Procedures section). The register (i.e., the first strand of Hya being structurally equivalent structure was solved to 2.6 **register (i.e., the first strand of Hya being structurally equivalent structure was solved to 2.65 A˚ resolution in the trigonal form with the first strand of the corresponding protein, etc.). The (Tables 1 and 2). Electron density accounting only for a (GlcA– seven-stranded** β barrel of endocellulase with evenly spaced GlcNAc)₂ fragment was observed in the substrate binding
strands, however, does not fit the 'open' Hya barrel closely. aroove (Figure 7), with the reducing en **Significant similarity in the active-site region (see below) was putative proton donor Glu113 (see below). Thus, the tetrasaconly observed for bacterial chitinase A [17], which showed a charide most probably represents a cleavage product that is** somewhat lower overall score $(Z = 9.2)$, but had similar loca-
bound to subsites -4 to -1 , according to the nomenclature **tions for the catalytic residues and for several conserved aro- for sugar binding subsites in glycosyl hydrolases [33] (–n is a**

is \sim 30 Å \times 10 Å and would be large enough to accommodate successive removal of GlcA–GlcNAc dimers from the nonre**is the smallest oligomer that can be hydrolysed [10, 11]. Most (three months). residues in the groove are conserved between insect and mam- Although the saccharide in subsite** 2**4 appears loosely bound**

Another seven-stranded 'open' barrel has been observed guide the substrate into the active site through electrostatic in the structure of quinolinic acid phosphoribosyltransferase interactions with the carboxylic groups of hyaluronic acid. (QAPRTase) [29]. Here, the gap that occurs between strands Many of the conserved active-site residues are located in the 2 and 3 is partially occupied by the substrate. In Hya there long loops adjacent to b **strands 3 and 4 comprising residues are no hydrogen bonds between strands 1 and 2 whereas in 111–124 and 184–201, respectively (Figure 6). The hydrophilic QAPRTase the analogous strands interact through a single loop 65–71 (KDPNGNV) joining the** b **strands of the hairpin hydrogen bond. Seven-stranded** b**/**a **barrels with equally spaced 62–74, located above the active site, is not visible in the elecstrands have been reported for cellobiohydrolase II [30] and tron-density map of the monoclinic crystal form. It is, however, the related thermophilic endocellulase E2 [31]. fully defined in the trigonal form, where it is stabilized by crystal**

groove (Figure 7), with the reducing end oriented close to the **matic residues in the binding cleft (data not shown). subsite at the nonreducing end and** 1**n at the reducing end,** with cleavage taking place between sugars in the -1 and $+1$ **The Substrate Binding Groove subsites). The bound tetrasaccharide is formed by the removal The dominant feature of the Hya protein surface is a large of a disaccharide from the reducing end of the fluorosceinated groove that extends perpendicularly to the barrel axis, approxi- hexasaccharide. In contrast, Takagaki et al. [11] reported that mately from strand 1 to strands 4 and 5 (Figure 5). Loops a HA hexamer labeled at the reducing end with 2-aminopyridine following** b **strands 2, 3, and 4 form one wall of the groove and was not digested by sperm PH-20 protein following 3–4 hr those following 1, 5, 6, and 7 the other. The size of the groove of incubation, whereas higher oligomers were hydrolyzed by a hexasaccharide. This is consistent with solution binding stud- ducing end. Degradation observed in our study might be due ies on the testicular enzyme which showed that (GlcA–GlcNAc)**³ to the long time required for crystallization of the complex

180–184 (strand 4), 220–226 (strand 5), 261–268 (strand 6), and 296–302 shown) but make no further protein contacts. The average dis- (strand 7). The substrate binding groove is located at the C-terminal end of tance between the four carboxyl oxygens of the two carboxylates the β barrel (top). Asp111 and Glu113 are shown as stick models. is 3.8 Å, considerably shorter than the 5.5 Å found in retaining
(b) View along the barrel axis, that is, after rotation by 90° around a horizontal ally

to the protein, the *N***-acetyl group of sugar residue** −3 and the and 111O δ 1−113O ϵ 2−113C δ are 117[°] and 118[°], respectively). **carboxylate group of** 2**2 are hydrogen bonded to the hydroxyl For steric reasons the two carboxylates cannot approach the groups of nonconserved Ser304 and Ser303, respectively (Fig- anomeric carbon of the substrate from opposite sides, as obure 7). In addition, the carboxylate of sugar residue** 2**2 interacts served in many retaining glycosidases. The close interaction sugar ring against Tyr55. The** *N***-acetyl group of GlcNAc in the clinic and trigonal crystal forms grown at pH 4.2 and 6.5, re-**2**1 subsite is deeply buried in a pocket with its carbonyl and spectively, as well as in the liganded structure at pH 6.5. The** amide groups hydrogen bonded to the sidechains of Tyr227 close proximity of the two COO- groups should result in ele**and Asp111, respectively. The chair conformation is observed vated pK values and protonation of one of the sidechain carfor the saccharides bound in subsites** 2**2,** 2**3, and** 2**4, whereas boxylates. Short hydrogen bonds between carboxylates have GlcNAc in subsite** 2**1 lacks density for the C6–O6 sidechain and either buried in the protein ('low-barrier hydrogen bonds', deexhibits weak density for C5 and O1. This may indicate partial scribed by Cleland and Kreevoy [39]), or exposed to the solvent**

Two acidic residues constitute the catalytic site in most glyco- hevamine [36], and chitinase A1 from *Bacillus circulans* **[43]. sidases, one acting as the proton donor and the other as the It was proposed that interactions with ordered water molecules**

nucleophile. The only carboxylic residues present in the Hya binding groove are the strictly conserved Asp111 and Glu113 residues (Figure 1) which protrude from the C-terminal end of β 3 **(Figure 3). Glu113 appears well positioned to act as the proton donor with a distance of 2.6 A˚ between its sidechain carboxylate and the glycosidic oxygen (Figure 7a). In contrast, the carboxylate oxygens of Asp111 are separated by more than 5 A˚ from the glycosidic oxygen. Indeed, functional analysis of respective mutants in the related human sperm PH-20 protein indicated that Glu113 is more important for catalysis than Asp111. Replacement of Glu113 by glutamine and Asp111 by asparagine in PH-20 (equivalent to Glu113 and Asp111 in Hya) resulted in a protein with no detectable activity and 3% activity, respectively [24]. This, together with the crystal structure, suggests direct involvement of these residues in catalysis. Interestingly, the glutamic acid was also identified as the proton donor in the structurally equivalent carboxylic pairs in endo-**b**-***N***-acetylglycosaminidase H (endo H) [34, 35] (Asp130–Glu132), hevamine [36] (Asp125–Glu127), and chitinase A from** *Serratia marcescens* **[23] (Asp313–Glu315). The impaired function of the three other PH-20 mutants (Glu249** → **Gln, Arg252**→**Thr, Arg176**→**Gly), investigated in the same study, is probably caused by structural perturbations. Glu249 and Arg252 correspond to Glu247 and Arg250 in Hya (Figure 1), where they form an intrahelical (helix G) salt bridge that is partly buried by the adjacent helix E. Although this salt bridge is too far away from the putative active site for direct involvement in catalysis (Figure 6), its disruption may destabilize the entire structure. The other mutated residue, Arg176 (the equivalent of Arg176 in PH-20), is also not part of the binding groove, but is buried between helices B and D with its guanidinium moiety bound to mainchain carbonyl groups of both helices. It is interesting to note that nonconservative mutation of Glu268 to lysine in Hyal-1, which corresponds to Glu247 in Hya, was identified in a patient deficient in serum hyaluronidase. This disorder is known as muco-**

Figure 3. Ribbon Representation of the Crystal Structure of Hya
 α Helices are colored in red, 3₁₀ helices in orange, and β strands in green.

(a) Side view of the barrel formed by β strands 1–7 (green arrows) **(b) View along the barrel axis, that is, after rotation by 90**⁸ **around a horizontal glycosidases [38]. Atoms Glu113 O**d**2 and Asp111 O**d**1 are axis. The two disulfide bridges are in yellow. separated from each other by 2.5 A˚ and have the right geometry for hydrogen-bond formation (angles 111C**g**–111O**d**1–113O**e**2** between the two carboxylic acids was observed in the monobeen observed in the active sites of diverse enzymes and are **disorder due to fewer stabilizing interactions after cleavage. as for the periplasmic glucose/galactose binding protein [40] and, more relevant, in the glycosidases cellobiohydrolase II Catalytic Groups and Reaction Mechanism [30, 41], endo-**b**-***N***-acetylglycosaminidase F1 [42] (endo F1),**

Figure 4. Stereo Ca **Trace of the Hya Molecule**

The view is rotated by 908 **around an axis perpendicular to the paper with respect to the view in Figure 3a. Every tenth residue is highlighted with a black circle and every twentieth residue is labeled.**

stabilize the short hydrogen bond [44]. Accordingly, substrate 2.9 A˚ (Figure 7b), which compares well with the corresponding

mine [23], where the corresponding residues are Asp539, nonhydrolysable cello-oligosaccharide [41]. Tyr669, Trp616, and Trp639 and Asp125, Tyr183, and Tyr6, In conclusion, Hya is a retaining b**-glycosyl hydrolase that respectively. In Hya, the tight interaction of the N-acetyl group catalyzes an acid/base reaction mechanism with Glu113 acting with the surrounding protein groups requires deformation of as the proton donor and the** *N***-acetyl carbonyl group of the** the sugar ring -1 from the ⁴C₁ chair to the ^{4,1}C boat conforma- substrate acting as the nucleophilic base. Assignment of the **tion (Figure 7b). The carbonyl group of the** *N***-acetyl moiety conserved Glu113 as catalytic residue is in agreement with points toward C1 of the** 2**1 sugar ring with a O7–C1 distance of mutagenesis studies on the sequence-related human sperm**

distances of 3.0 A˚ for chitobiase and 2.8 A˚ binding will disorder the solvent network and facilitate the for hevamine. At uptake of a proton by the substrate. **the substrate of the carbonyl** this close distance, the partial negative charge of the carbonyl **As in Hya, there is no obvious proteinaceous nucleophile in oxygen is well positioned to stabilize the positive charge of several other retaining** b**-1,4-glycosidases which act on GlcNAc the anomeric carbon, as postulated for the oxocarbonium ion in the** 2**1 position, such as hevamine, chitinase, and chitobiase intermediate. This distortion, induced by binding into the active (see [23] and references therein). In these enzymes, the nucleo- site, facilitates the reaction by bringing the glycosidic bond into philic function is performed by the carbonyl oxygen of the a near equatorial position and moving the glycosidic oxygen substrate's** *N***-acetyl group at C2 [23, 36]. The prerequisite for nearer to Glu113, the proton donor [46]. Clearly, Hya belongs this substrate-assisted reaction is the accurate positioning of to the group of retaining enzymes, because the anomeric C1 is** the *N*-acetyl sidechain of the substrate with respect to the found in a β configuration in the enzyme–HA-tetramer complex. **enzyme. The reaction can then proceed either via a positively Distortion of the pyranoside ring bound at site** 2**1 was first charged oxocarbonium ion intermediate or an oxozolinium ion observed for planar oxocarbonium ion intermediate in lysowith a covalent bond between the** *N***-acetyl carbonyl oxygen zyme [47]. Half-chair conformations were observed in the enand the C1 atom [23]. zyme–product complexes of hen egg white lysozyme [48, 49] In Hya, as well as in the chitinolytic enzymes, the position and bacteriophage T4 lysozyme [50]. Twisted boat conformaof the** *N***-acetyl sidechain is tightly defined by two hydrogen- tions, in which C1, C2, C3, C5, and O5 atoms are nearly coplabonding interactions: one between its amide NH and Asp111, nar, were found in chitobiase complexed with uncleaved dichiand the other between its carbonyl oxygen and the hydroxyl tobiose in the active site [45] and in the crystal structure of the group of the conserved Tyr227. Furthermore, the methyl group retaining** b**-endoglucanase I in complex with a nonhydrolysable of the** *N***-acetyl sidechain forms hydrophobic interactions with substrate analog [51]. Distortion of the glycosyl unit in the** 2**1 Tyr184 and Trp301 (Figure 7a). Similar interactions have been** site from the usual chair to the ²S_o conformation has recently **observed in the liganded structures of chitobiase [45] and heva- been reported for cellobiohydrolase (Cel6A) in complex with**

^a As defined in the program DALI [32]. Several entries between 1tml and 1ctn have been omitted for convenience.

The 1.6 A˚ denotes residues that are identical in all four sequences shown in Figure 1; resolution crystal structure of hyaluronidase, a

venom hyaluronidase with human hyaluronidases, belonging to the Hya epitope(s). Susceptible individuals respond to expofamily 56 glycosidases, indicates conservation of the active- sure with Hya by producing IgE antibodies which, following site residues, suggesting a similar catalytic mechanism for the reexposure, bind to the allergen and thus induce an anaphylac-

occasionally fatal, systemic IgE-mediated anaphylactic reac- to have a strong impact on the treatment of Hya-induced altions in human [25]. Only a few three-dimensional structures lergy. Rationally designed ligands that can compete with IgE of allergens are known to date, four from plants and five from antibodies for the allergen binding site could prevent the alleranimals (see [52] and references therein) [53]. These structures gen-induced release of mediators from effector cells. Alternaare different from the (b**/**a**)7 barrel fold of Hya. Mammalian tively, mutant proteins that bind weakly to IgE antibodies but lipocalins [52, 53] show an antiparallel** β barrel fold as does can still interact with allergen-specific T cells could be used **the plant allergen Phl p 2 from timothy grass [54]. Profilin for immunotherapy to suppress IgE sensitization to Hya, with-**

allergens [55, 56] have a large β sheet with flanking helices, and **the major mite allergen Der f 2 [57] resembles immunoglobulin. Thus, no unique structural feature responsible for the allergenicity can be identified. However, Rouvinen et al. [52] pointed out that all these proteins are monomers (except for the dimeric major horse allergen [53]) of similar dimensions. This also applies to Hya.**

Antigenicity of Hya is fully determined by the structure of its epitopes, the areas of the protein surface that are recognized by specific antibodies. The structure of the Hya surface, which is now known in full detail, contains an unusually high number of charged residues which are evenly distributed on the surface (Figure 5a). Of all surface residues, 41% comprise aspartate, glutamate, lysine or arginine residues. This is considerably higher than the average value of 27% [58]. Although some of these residues may now be subjected to mutagenesis, the identification of the Hya epitope(s) must await further studies. The full characterization of the antigen–antibody recognition sites will require the elucidation of the structure of complexes between Hya and Fab fragments (from various mAb against Hya), because epitopes are generally composed of a discontinuous array of amino acids [59]. Site-directed mutagenesis of epitope residues may lead to mutant variants with low IgE binding activity (hypoallergens) [60] which might be used in immunotherapy without the risk of side effects. An example of this approach is a six-point mutant of Bet v 1, the major allergen of birch pollen, which exhibits extremely low reactivity with serum IgE from birch pollen-allergic patients [60].

Biological Implications

Hyaluronidases are a group of hydrolytic enzymes which degrade hyaluronic acid (HA), a high molecular weight polysaccharide found in virtually all mammalian tissues and body fluids. HA is a ubiquitous component of the vertebrate extracellular matrix where it fills the space between cells and fibers and acts as a lubricant as well as a barrier to penetration by foreign particles. HA levels are markedly increased during embryogenesis, inflammation, malignant transformation, wound healing, Figure 5. The Molecular Surface of Hya Viewed as in Figure 3b and whenever fast tissue turnover and remodeling is required. (a) The surface is color coded according to residue type with red, blue, and

yellow representing acidic, basic and aromatic residues, respectively. The

(GIcA-GIcNAC₂) tetrame bound to the substante binding groove and

yellow denotes similar residues. major allergen from bee venom, gives insights into the mode of substrate binding and the substrate-assisted catalytic mechanism. It also represents the first step toward the elucidation of PH-20 protein. Moreover, a multiple sequence alignment of bee a Hya–antibody complex structure which is needed to identify mammalian enzymes. tic reaction. Nonallergic individuals (e.g., beekeepers) produce mainly IgG4 antibodies associated with an immunoprotective Allergenicity role. Knowledge of the structure of the Hya epitopes will pro-Hya is a major allergen of bee venom that can induce serious, vide a basis for understanding its allergenicity and is expected

Figure 6. The Substrate Binding Groove Stereoscopic close-up view of the groove ex-

tending across the C-terminal end of the b **barrel (green arrows). Bold labels indicate conserved residues (Figure 1).**

out risk of side effects. This T cell targeted therapeutic ap- Crystallization and Data Collection

uronidases involved in HA turnover, the malfunctioning of contained one molecule in the asymmetric unit, resulting in a solvent content which is related to various diseases. For example, the synovial of 54% (Vm = 2.7 Å³/Da). Monoclinic crystals were obtained by mixing the
fluid of joints in patients suffering from osteoarthritis and rheu. protein (1:1) w fluid of joints in patients suffering from osteoarthritis and rheu-
matoid arthritis has increased hyaluronidase activity that re-
sults in a decrease of HA size and concentration. Interestingly,
sults in a decrease of HA **increased expression of a hyaluronidase similar to PH-20 has Prior to data collection, crystals were transferred into stabilizing solutions: been found in tumor cells [61]. There, the enzyme is used to 33% PEG 8000, 0.1 M Na-cacodylate (pH 6.5), 0.2 M ammonium sulfate for break down the basement membrane and facilitates tumor** the trigonal crystals and 12% PEG 6000, 1.0 M NaCl, 5 mM Na-acetate (pH
invasion angiogonosis, and motostasis. With the knowledge 4.4) for the monoclinic crystals. D invasion, angiogenesis, and metastasis. With the knowledge
of the Hya structure, modeling of the human enzymes should
now be feasible. These structures will enable the rational de-
sign of specific inhibitors which could b **sign of specific inhibitors which could be used to combat hyal- resolution data collection, monoclinic crystals were flashed-cooled to 120K in**

Recombinant enzymatically active Hya, containing a C-terminal His₆ tag, CCP4 package [63]. was produced by *Baculovirus*-infected insect cells and purified by Ni²⁺**chelate chromatography, as described previously [15]. For crystallization experiments the protein was dialyzed against 5 mM sodium acetate buffer Structure Determination and Refinement of the Monoclinic Form (pH 5.4) and concentrated to 8–10 mg ml**2**1. The structure was solved by MIR at 2.7 A˚ using phase information from two**

(Aldrich) solution in 30% AcOH/DMSO (3 ml) and fluoresceinamine (130 mg) ment of the heavy-atom parameters and calculation of initial phases were was added (Aldrich). After heating the mixture at 90°C for 3 hr, the reaction performed with the program MLPHARE [63] yielding a rather poor figure of **was stopped by adding an equal volume of methanol. The product was merit of 0.36. Additional sites were identified by difference Fourier techprecipitated from acetone and filtered. The solid material was recrystallized niques. Final refinement of the heavy-atom parameters and phase calculaceinated oligomer, was concentrated and further purified by gel-filtration 2.7 A˚ . Three binding sites for each of two derivatives were identified. Phase chromatography (Sephadex G-10 column). The total yield of fluoresceinated improvement was achieved by solvent flattening using the anticipated sololigomer was 48%. The identity and purity of the final compound was deter- vent content of 47% (SOLOMON [65]). The resulting map at 2.7 A˚ resolution mined by TLC (RP-18, eluent acetonitrile-water 8:2) and NMR analysis was of sufficient quality to reveal** a **helices and** b **strands. A partial polyala- (Bruker AC200 200 MHz spectrometer). nine model was built into regions of well-defined density with the aid of the**

proach will hopefully result in the induction of allergen-specific $\frac{1}{2}$ Two crystal forms suitable for X-ray analysis were obtained by the hanging-
immune tolerance.
Hya is closely related to the human PH-20 sperm pr **latter solution. Crystals (0.3** \times 0.5 \times 0.2 mm) grew within 6–8 months and unit and a solvent content of 47% (Vm = 2.3 \AA^3 /Da).

uronidase-related disorders. a cryosolvent containing stabilizing solution plus 10% ethylene glycol. X-ray data were collected (native-2, Table 1) at the synchrotron radiation source at EMBL/DESY, Hamburg. High-resolution data of native trigonal crystals Experimental Procedures (Table 1) were collected at room temperature at the Daresbury (UK) synchrotron radiation source. Image data were indexed and integrated using Purification Purification MOSFLM [62]. The data were reduced and scaled using programs of the

heavy-atom derivatives: Pb(CH3COO)2 (3 mM, two weeks soak) and PtCl4 (1 mM, 24 hr soak). Derivative and native monoclinic data (native-1, Table 1) Synthesis of the Fluoresceinated Oligomer sets were scaled with program SCALEIT. The major lead and platinum sites The HA hexamer (150 mg) was dissolved in a 1 M sodium cyanoborohydride were easily identified using isomorphous difference Patterson maps. Refine from acetonitrile and vacuum dried. Filtered supernatant, containing fluores- tion was done with the program SHARP [64] using data between 15 and

Figure 7. Stereoviews of the HA Tetramer GlcA-GlcNAc-GlcA-GlcNAc Bound to Hya

SigmaA-weighted simulated annealing [72] Fo–Fc electron-density map (contour level 3s**) has been calculated with the ligand omitted. (a) The atomic model of the tetrasaccharide and of all protein residues within a distance of 5 A˚ to the sugar are shown. The reducing** end of the tetrasaccharide (sugar residue -1) **is at the top, close to Glu113. The view is similar to that in Figure 6.**

(b) Close-up view of the distorted sugar ring of the GlcNAc residue in position -1. The view is rotated by about 90° around the verti**cal axis with respect to the view in (a).**

BONES options in O [66]. The electron-density map, however, was not of inated HA hexamer, using the hanging-drop method. The protein was mixed sufficient quality to complete the model. The structure was eventually solved (1:1) with reservoir solution containing 25% PEG-4000, 0.2 M ammonium by multicrystal averaging (DMMULTI [67]) of the phased P21 and the un- sulfate, 0.1 M Na-acetate (pH 4.6) and incubated over the reservoir. The phased P3221 crystal form, each having one monomer per asymmetric unit. complex structure was solved by molecular replacement with the final mono-The rotation/translation relationship between the molecules of the two crys- clinic model. tal forms was determined by molecular replacement (AMoRe [68]), using the MIR density of the monoclinic form. Acknowledgments

Initial refinement of the monoclinic form was carried out by REFMAC [63] using data set native-1. After manual adjustment of the model as indicated The technical assistance of Y. Rasser is greatly acknowledged. We thank ued using the high-resolution data set to 1.7 A˚ (native-2). For this, iterative We are grateful to the synchrotron staff at SRS in Daresbury, UK, and DESY, to 1.7 A˚ resolution. At 2 A˚ resolution water molecules were added to the National Foundation grant No. 31-50507 to Z. M.-H. and T. S. model using the program ARP. After each refinement cycle, the model was checked against $2F_o-F_c$ maps and manually adjusted to improve geometry **Received: June 7, 2000 and fit to the electron density. The model quality was checked with PRO- Revised: July 31, 2000 CHECK [69] which showed that 91% residues are within the most favored Accepted: August 3, 2000 regions of the Ramachandran plot and only one residue, Ala124, exhibits a disallowed conformation. References**

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Protein Data Bank Accession Codes

Coordinates and structure factors have been deposited in the Protein Data Bank with accession codes 1FCQ (native monoclinic data), 1FCU (native trigonal data) and 1FCV (trigonal form in complex with the HA tetramer).