Studies on the *Vibrio cholerae* mucinase complex. I. Enzymic activities associated with the complex

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Summary. Mucinase enzymes were isolated and partially purified from the culture fluid of *Vibrio cholerae* grown in proteose peptone-colostrum medium. The mucinase complex contained neuraminidase, endo-β-N-acetylhexosaminidase, nicotinamide-adenine-dinucleotidase and proteinases. Traces of phospholipase activity were detected but the complex lacked aldolase activity.

Introduction

The mucinase activity of *Vibrio cholerae* was originally shown by the desquamation of guinea-pig ileal segments after exposure to culture filtrates for 3 h at 37°C; this activity survived heating at 56°C for 30 min and was greater in the presence of Ca++ ions (Burnet and Stone, 1947a). Burnet and Stone (1947b) also demonstrated the activity of another enzyme (receptor destroying enzyme) that removed the influenza virus receptor from erythrocytes; this enzyme was sensitive to heating at 56°C for 30 min. It was identified subsequently as a sialidase (Heimer and Meyer, 1956) and finally as a neuraminidase (Gottschalk, 1957) which acted by removing the terminal neuraminic acid unit from a neuramin-lactose substrate by hydrolytic cleavage of the glycoside bond joining the keto-group of N-acetylneuraminic acid to D-galactose or galactosamine.

Intestinal sialomucin may provide a physical barrier to *V. cholerae* colonisation of the intestinal epithelium but, as reviewed by Arbuthnott and Smyth (1979) and Savage (1980), enzymes of the mucinase complex could hydrolyse the mucus, create random tracks and enable the organisms to reach the columnar epithelial surface.

In a series of studies on the development of an acellular cholera vaccine, the mucinase enzymes were isolated and characterised, in view of their potential use as immunogens.

Materials and methods

**Bacterium**

*Vibrio cholerae* NCTC 10732, the classical biovar, Inaba serovar was obtained from the National Collection of Type Cultures, Colindale, London.

**Medium for production of mucinase**

Bovine colostrum was obtained 24 h post-parturition at the Veterinary School experimental farm and kept at −20°C until required. The colostrum (1·5 L) was placed in dialysis tubing (Visking; diameter 7 cm) and dialysed against 7 L of distilled water, saturated with chloroform to prevent bacterial contamination, for 7 days. The distilled water containing dialysed substances was transferred to a 5-L pressure vessel (Millipore), aerated for 2 h with sterile filtered air to remove traces of chloroform and subsequently filtered through a 0·45-μm membrane in a Millipore filter holder (diameter 142 mm) under nitrogen pressure into a 15-L fermenter vessel containing meat peptone at a final concentration of 1% w/v.

Meat peptone (150 g; Gibco-Europe) was dissolved in 7·5 L of distilled water, placed in the fermenter vessel and sterilised by autoclaving at 121°C for 1 h. Glycerol (BDH, Poole, Dorset) was sterilised at 121°C for 30 min and 112 ml was added aseptically to the fermenter vessel. After the addition of the colostrum dialysate, the volume was adjusted to 14·75 L with sterile distilled water.

The large-scale batch fermenter was primed with a 250-ml inoculum of an overnight culture of *V. cholerae* NCTC 10732 grown in proteose peptone (Gibco-Europe) 2% w/v. The fermenter (L. H. Engineering Ltd, Stoke Poges, Bucks) was run at 37°C with an air-flow of 5 L/min, stirring at 300 rpm and with automatic dosing of 1% v/v Antifoam C Emulsion (Sigma Chemical Co., Poole, Dorset). After 24 h the organisms were killed *in situ* with merthiolate 0·01% (Sigma) or by increasing the temperature to 60°C for 2 h. The cells were removed from the culture fluid by centrifugation at 9000 g for 15 min.

**Partial purification of mucinase**

Methanol (1·5 L) cooled to −30°C was added in
300-ml volumes/h to 1 L of the culture supernate and left at 4 C for 48 h. The precipitate was harvested by centrifugation at 12000 g for 15 min. The pellets so obtained from the 15 L of culture supernate were resuspended in 100 ml of 0.05 M sodium acetate buffer, pH 5.5, and dialysed against the same buffer for 2-3 days at 4 C. The retentate was centrifuged at 12000 g for 15 min and the supernate, containing crude mucinase, was concentrated with polyethylene glycol 4000 (Carbowax, BDH) to a final volume of 50 ml.

**Gel filtration of crude enzyme concentrate**

The most suitable grade of Sephadex for the partial purification of the crude mucinase was G100. The beads were suspended in 0.05 M sodium acetate buffer at pH 5.5 and poured into an 80 x 1.5 cm column. The crude concentrate (2 ml) was filtered through the column with the same buffer as eluate, at a flow rate of 0.8 ml/min. The absorption value at 280 nm and the Lowry protein value were determined for each 2-ml column fraction. Thiobarbituric acid and methoxyphenol-neuraminate assays were also done on each fraction. The fractions within the peak of enzyme activity were pooled, concentrated with Carbowax to 2 ml and stored at -20 C. The neuraminidase-containing preparations were designated G100/N.

**Estimation of protein**

The protein content of the partially purified mucinase complex was determined by a Lowry assay with bovine serum albumin as standard (Lowry et al., 1951) or by a modification of the microburet method of Levin and Brauer (1951) with ovalbumin as the standard. In the latter test, mucinase samples (1 ml) were mixed with 0.3 ml of 3 N NaOH and heated at 100 C for 5 min. The mixtures were cooled to room temperature and 0.2 ml of CuSO,·5H2O 2.5% w/v was added. After 5 min, the mixtures were centrifuged at 1200 g and the absorption values at 555 nm of the supernates were recorded.

**Determination of neuraminidase activity**

(a) **Thiobarbituric acid (TBA) assay** (Warren, 1959). Bovine submaxillary mucin (20 mg) was dissolved in 5 ml of 0.05 M sodium acetate buffer, pH 5.5, containing CaCl2 0.1% w/v and NaCl 1% w/v. The *V. cholerae* neuraminidase as the G100-N fraction (0.1 ml) was added to 0.4 ml of the solution of bovine submaxillary mucin. The volume was adjusted to 1 ml with buffer and the mixture was incubated at 37 C for 30 min. Subsequently, 1 ml of phosphotungstic acid (5 g in 100 ml of 2 N HCl) was added and the mixtures were centrifuged at 1200 g for 10 min. A 0.5-ml volume of each supernate was removed and added to 0.1 ml of periodate solution (4.27 g sodium periodate in 38.3 ml distilled water and 61.7 ml phosphoric acid). After 20 min at room temperature, 1 ml of arsenite solution (Na2SO3 17.75 g in 250 ml of distilled water with 5 ml of 5 N H2SO4 and 25 g of sodium arsenite) was added and the mixtures were shaken until unreacted iodides were removed, as indicated by loss of yellow colour. Thiobarbituric acid solution (3 ml of solution containing 4.5 g of twice-crystallised thiobarbituric acid in 750 ml of 0.5 M Na2SO3) was added. Tubes containing the reaction mixtures were placed in a heating-block at 100 C for 15 min before cooling to room temperature. Re-distilled cyclohexanone (4.6 ml) was added to each tube, the mixtures were shaken and centrifuged at 1200 g for 3 min and finally the absorption value at 549 nm of the organic phase was recorded.

The amount of neuraminic acid (NANA) released from the mucin was determined by reference to a standard curve of neuraminic acid. Positive controls were *V. cholerae* neuraminidase (Koch Light) and *Clostridium perfringens* neuraminidase (Sigma). The activity of enzyme preparations was calculated on the basis that 1.0 International Unit (IU) liberates 309.3 μg of NANA/min.

(b) **Methoxyphenol-neuraminate (MPN) assay of Palese et al. (1973).** The substrate in this assay, kindly provided by the Research Resources Branch, NIAID, Bethesda, MD, USA, was 2-(3' methoxy-phenol)-N-acetyl-α-neuraminic acid; the stock substrate solution contained 50 mg in 11.82 ml of 0.1 M sodium phosphate buffer, pH 5.9. Neuraminidase-containing G100/N fractions (0.1 ml) were mixed with 0.1-ml volumes of 0.1 M sodium phosphate buffer, pH 5.9, in the presence of 2 mM CaCl2 and warmed at 37 C for 3 min before the addition of 0.1 ml of the stock substrate solution. The mixtures were incubated at 37 C, and the reaction was stopped after 30 min by the addition of 25 μl of disodium ethylene diamine tetraacetic acid (BDH). The mixtures were kept in an ice-bath for 30 min and 0.2 ml of freshly prepared and filtered diazonium salt of 4-amino-2,5-dimethoxy-4' nitroazobenzene (Koch-Light Laboratories; diazonium salt ADNB 6.0 mg in 1 ml of 0.4 M sodium phosphate buffer, pH 7.0). After 30 min, 4 ml of 0.5 N NaOH was added to dissolve the precipitate, the tubes were centrifuged at 1200 g for 3 min and the absorption values of the supernates at 580 nm were recorded. The amount of methoxyphenol liberated was determined from a standard curve; 1 IU of neuraminidase liberates 121-14 μg of methoxyphenol/min. A conversion factor of 0.401 was used to calculate the amount of NANA (in μg) released from the substrate.

**Protease activity**

The digestion of casein with trypsin as standard (Kunitz, 1947) was the method used. The partially purified *V. cholerae* mucinase G100/N (1 ml) was added to 1 ml of casein solution (1 g of casein, Hammarsten grade, BDH, dissolved in 0.1 M sodium phosphate buffer, pH 7-6, to a final volume of 100 ml) and incubated at 37 C for 30 min. The reaction was stopped by adding 3 ml of trichloroacetic acid 5% w/v and incubating at 45 C for 15 min. Undigested casein was removed by centrifugation at 1200 g for 10 min and the absorption
values of the supernates at 280 nm were recorded. Protease activity was determined from the standard plot for trypsin.

**Phospholipase C activity (Boehringer, Mannheim GmbH, 1983)**

The lecithin substrate used in this assay was prepared by a modification of the method of Pangborn (1951) and was described in detail by Stewart-Tull et al. (1978). The partially purified *V. cholerae* fraction (20 μl) was added to 1 ml of 0-1 M triethanolamine buffer, pH 7-5, containing 20 mM CaCl₂, 1 ml of freshly prepared lecithin suspension (20 mg/ml in distilled water), 50 μl of lipase (ex *Rhizopus arrhizus* 10 mg/ml in 3-2 M ammonium sulphate, EC. 3.1.1.3) and 0-5 ml of distilled water. After 15 min at 37°C the reaction was stopped with 0-3 ml of 1-5 M trichloroacetic acid. The mixtures were heated at 100°C for 2 min, cooled and centrifuged at 1200 g for 5 min. The supernate (0-25 ml) was pipetted into a cuvette with 2-5 ml of 0-3 M triethanolamine buffer, pH 7-6, containing 4 mM magnesium sulphate, 0-15 ml of a solution containing 32 mM ATP, 6 mM NADH and 45 mM PEP, 0-01 ml of lactate dehydrogenase 5-0 mg/ml in 3-2 M ammonium sulphate and 0-01 ml of pyruvate kinase 2 mg/ml in 3-2 M ammonium sulphate. The reagents were mixed in the cuvette and the absorption value at 340 nm was recorded, A₁. Finally, 0-04 ml of glyceral kinase (1 mg/ml in 3-2 M ammonium sulphate) was added and after 10 min the absorption value at 340 nm was recorded, A₂. The relative phospholipase C activity ΔA was equal to A₁ - A₂ and was quantified by reference to a standard plot of values obtained with *Bacillus cereus* phospholipase C (Boehringer, EC 3.1.4.3, 2 mg/ml in 3-2 M ammonium sulphate). One IU of phospholipase C hydrolyses 1 μmole of lecithin/min.

**Endoglycosidase activity (Dubois et al., 1956; Chien et al., 1975)**

*Streptomyces griseus* endoglycosidase (Seikagaku Kogyo Co. Ltd, Tokyo, Japan; 0-1 unit of freeze-dried enzyme was dissolved in 100 μl of distilled water) was compared with the *V. cholerae* partially purified mucinase against ovalbumin substrate (Sigma grade V; 2 mg in 0-2 ml of 0-05 M sodium acetate buffer, pH 5-5, which contained sodium dodecyl sulphate 0-5% w/v).

The G100/N enzyme sample (20 μl) or the *S. griseus* endoglycosidase (20 μl) of stock solution 2 μl in 100 μl of 0-1 M NaCl containing bovine serum albumin 0-1% w/v) was added to 0-5 ml of ovalbumin substrate and incubated for 1 h at 37°C. An equal volume of cold trichloroacetic acid 12-5% w/v was added to stop the reaction and after 15 min at 0°C the mixtures were centrifuged for 30 min at 35 000 g. Samples (0-2 ml) of the clear supernate were assayed for neutral sugar by the phenol sulphuric acid method; 50 μl of phenol 80% w/v and 0-5 ml 36 N H₂SO₄ (Analar) were added and after 30 min at room temperature the absorption values at 490 nm were recorded. The endoglycosidase activity of the G100/N fraction was measured by the release of neutral sugar relative to glucose. The units of endoglycosidase activity were calculated from a standard curve of glucose released from ovalbumin substrate against different concentrations of commercial enzyme.

**SDS-PAGE**

The method used for sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) was based on that of Laemmli (1970) as modified by Ames (1974). The separating gel (11% w/v) contained N,N' bis-acrylamide (acrylamide 30-0 g; N,N'methylene bis-acrylamide 0-8 g; water to 100 ml) 36-7 ml; 1 μ Tris/HC1 buffer, pH 8-8, 37-5 ml; SDS 20% w/v, 0-5 ml; N,N,N',-tetramethylethylenediamine (TEMED; Sigma), 25-0 μl; ammonium persulphate 0-8% w/v, 10-0 ml; and water, 15-3 ml. The stacking gel (5%) contained 16-7 ml of the N,N'- bis-acrylamide solution; 0-5 μ Tris/HCl buffer, pH 6-8, 25-0 ml; SDS 20% w/v, 0-5 ml; TEMED 25 μl; ammonium persulphate 0-8% w/v, 10 ml; and water, 47-8 ml.

Each sample was mixed with an equal volume of solubilising buffer (sodium dodecyl sulphate 20% w/v, 20 ml; 0-5 μ Tris/HCl buffer, pH 6-8, 25 ml; β-mercaptoethanol, 10 ml; glycerol, 20 ml; bromophenol blue 0-1% w/v, 2 ml; water, 23 ml) and heated at 100°C for 5 min. Samples of 75 μl volume were run for approximately 3 h at 15 mA in a Shandon analytical slab PAGE outfit with a Tris/glycine/SDS buffer (3-0 g, 14-4 g, 0-1 g respectively in 100 ml of water). Protein bands were visualised by staining for 1 h with Coomassie brilliant blue (Sigma) 1-25 g in 454 ml of methanol plus 46 ml of glacial acetic acid (Analar; BDH). Gels were destained in methanol, 50 ml: acetic acid, 75 ml; water, 875 ml.

**ADP-ribosylating activity**

ADP-ribosylation was measured by the indirect method of Tait and Nassau (1984). Each assay was done with a mixture (total volume 100 μl) containing 200 mM potassium phosphate (KH₂PO₄–K₂HPO₄), pH 6-5, 20 mM dithiothreitol, 0-2 mM NAD (Grade 1, Boehringer-Mannheim) containing 2 × 10⁵ cpmp [4-3H]-NAD (1-01 mCi/mmol; Amersham International), 20 μM 10-(p-methoxyphenyl methyl decany1)-guanidine; kindly provided by Glaxo Group Research Ltd, Greenford, Middlesex, and 10-0 μg of cholera toxin (List Biological Laboratories, CA, USA) or the experimental sample to be assayed. The mixture was incubated for 30 min at 37°C; after this time, the reaction was stopped by dilution with 1 ml of distilled water.

[3H]-Nicotinamide was separated from the unreacted [4-3H]-NAD by ion-exchange chromatography on 0-5 × 1-0 columns of QAE-25 Sphadex. The diluted reaction mixture was loaded on to the column and the [3H]-nicotinamide was eluted with 3 ml distilled water into scintillation vials containing 15 ml of emulsifying
Fig. 1. Elution profile of *V. cholerae* methanolic precipitate of culture fluid separated on Sephadex G100 measured by absorption at 280 nm. The maximum neuraminidase activity associated with the mucinase complex was detected in the interpeak fractions between peaks 1 and 2 by both the MPN-assay (curve B, ○—○) and the thiobarbituric acid assay (curve C, ●—●). The Lowry protein content of the interpeak fractions is shown in curve A (■—■).

Scintillant (Optiphase Safe, LKB). The unreacted NAD remained on the column and was eluted with 5 ml of 1 M HCl, followed by 10 ml of water.

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0.78 international milliunits (ImU) of neuraminidase/mg of protein. The minor peak contained 5.4 ImU of neuraminidase/mg of protein. However, it became apparent that there was a peak of enzyme activity between these two peaks which contained 9.3 ImU of neuraminidase/mg of protein. This result was consistently obtained and therefore the inter-peak fractions were routinely collected as the partially purified mucinase. The neuraminidase peak fractions from six column separations were pooled to provide a sufficient supply of mucinase, designated as G100/N.

These findings were supported by SDS-PAGE analyses of (a) the methanolic precipitate containing crude mucinase complex, (b) pooled peak fractions from the Sephadex G100 column and (c) the inter-peak, pooled fractions from the G100 column that showed maximum neuraminidase activity. The result shown in fig. 2 indicated that there were three major bands shared in these preparations, together with several minor bands. Partial purification by Sephadex filtration removed a large amount of low-molecular-weight material, probably medium constituents. It was apparent that the mucinase consisted of a mixture of proteins with or without enzymic activity. Further attempts at purification by ion-exchange chromatography produced fractions with much lower neuraminidase activity and were not pursued.

The protein content of the Sephadex G100/N preparation varied from batch to batch when determined either by the Lowry or by the microbiuret method. The range was 37–562 μg of protein/ml by the former method and 32.8–319 μg/ml by the latter method. As shown in the table there was reasonable agreement between the values obtained by the two methods.

**Enzymic activities detected in the mucinase complex**

**Neuraminidase.** The level of neuraminidase activity in the partially purified preparations G100/N₁–N₄ from the culture fluid of *V. cholerae* grown in the colostrum-proteose peptone medium were in the range 30–55 ImU/mg of protein, as determined from the release of nanomoles of N-acetyl-neuraminic acid from bovine submaxillary mucin detected in the TBA assay (table). The comparative range of neuraminidase levels was 81–740 ImU/mg of protein when the more specific MPN assay was used.

**Endoglycosidase.** The G100/N₁–N₄ preparations contained levels of endoglycosidase which varied from 0.62 enzyme units (EU)/mg protein for G100/N₁ to 10.88 EU/mg of protein for G100/N₄ (table).

**Proteinase.** All the G100/N preparations with the exception of G100/N₄ contained proteinase equivalent to micrograms of trypsin; the values were in the range 37.0–56.5 (table). In addition, the methanolic precipitate of the culture fluid decreased the ADP-ribosylating activity of pure cholera enterotoxin after incubation for 30 min at 37°C. The activity ratio of treated cholera toxin: untreated toxin, was 0.64 for the methanolic precipitate and 0.68 for G100/N₂ mucinase complex.

**Phospholipase C.** The levels of phospholipase C were calculated relative to the activity of *B. cereus* enzyme; the calculated values were not significant and were in the range 0.0014–0.0141U (table).

**Nicotinamide-adenine-dinucleotidase (NADase)**
**Fig. 3.** The structure of bovine sub-maxillary mucin with points of action by *V. cholerae* mucinase complex.

**Table.** The protein composition and enzymic activities of the partially purified mucinase complex

<table>
<thead>
<tr>
<th>Preparation</th>
<th>Amount of protein (µg/ml) by</th>
<th>Amount of enzyme/mg of microbiuret protein</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Lowry (as bovine serum albumin)</td>
<td>Microbiuret (as ovalbumin)</td>
</tr>
<tr>
<td>G100 N1</td>
<td>562</td>
<td>319</td>
</tr>
<tr>
<td>G100 N2</td>
<td>134</td>
<td>108</td>
</tr>
<tr>
<td>G100 N3</td>
<td>37</td>
<td>32-8</td>
</tr>
<tr>
<td>G100 N4</td>
<td>167</td>
<td>136</td>
</tr>
</tbody>
</table>

*One ImU of neuraminidase liberates 0-31 µg NANA/min; TBA = thiobarbituric acid assay; MPN = methoxyphenol neuraminate assay.
†Calculated relative to the release of glucose from substrate by commercial endoglycosidase.
‡One IU of phospholipase C hydrolyses 1-0 µmole of lecithin (804-2 µg)/min.
activity. Neither the crude methanolic precipitate nor the G100/N fractions contained cholera enterotoxin in an indirect ELISA assay (Svennerholm and Holmgren, 1978). However, both the methanolic precipitate and the G100/N preparations showed ADP-ribosylating activity ratios, of 2-14 and 2-00 respectively; this was most likely to be NADase activity which cannot be distinguished from ADP-ribosylation in this particular assay.

Discussion

Growth of *V. cholerae* in the proteose peptone-bovine colostrum medium stimulated good yields of the mucinase complex, containing on average 12.42 IU of neuraminidase per L. After Sephadex G100 partial purification, this yield increased to 43.61 IU/L. As found by other groups interested in the exoenzymes of *V. cholerae* (Schick and Zilg, 1977; Schneider and Parker, 1978 and 1982), it was apparent that a high degree of purification had to be sacrificed in order to maintain high yields of mucinase.

The main difficulty encountered with the use of the colostrum dialysate was the apparently variable induction of enzymes. As shown in the table, the relative amounts of each enzyme in the mucinase complex varied from one preparation to another; this indicated that there was no definite relationship between the enzymes in the complex. However, it was significant that in three of the four G100/N preparations up to 0.5% w/w was proteinase(s). In further studies, to be published separately, these proteinases caused autodigestion of the mucinase complex and it is possible that in this study some of the variable quantitative results were due to this phenomenon. Chien et al. (1975) detected endo-β-N-acetylglucosaminidase and proteinase in commercial neuraminidase. The G100/N preparations contained considerable quantities of an endo-N-acetylhexosaminidase which could act on the β-glycosidic link in the bovine mucin between N-acetylglucosamine and the β-carboxyl group of aspartic or the γ-carboxyl group of glutamic acid (fig. 3).

Although not described in detail, the preparations were assayed for the presence of N-acetylneuraminic acid aldolase; no activity was found in any of the G100/N preparations. Ada et al. (1961) warned that this enzyme would interfere with an assay in which neuraminidase activity was measured by the liberation of N-acetyle neuraminic acid (NANA) because the aldolase would degrade this to N-acetyl-mannosamine. The difference between the Warren thiobarbituric acid assay and the methoxyphenol-neuraminate assay values for the release of NANA may reflect the availability of neuraminidase-sensitive residues. The submaxillary mucin has some five hundred carbohydrate groups per molecule and it produces a highly viscous solution (Montgomery, 1970) whereas methoxyphenol-neuraminate produces a non-viscous solution. Similar problems with the substrate were examined by Fraser and Smith (1975) who showed that a soluble glycoprotein fraction from pooled human plasma was sensitive to *C. perfringens* neuraminidase; the α1, α2 acid glycoprotein has an intermediate number of carbohydrate groups per molecule (Montgomery, 1973). One unit of NANA/mg of the glycoprotein was released compared with 0.58 unit/mg with bovine submaxillary mucin (Fraser and Smith, 1975). It does seem worthwhile to propose that in studies of the neuraminidase action on viscous mucin, one of the more sensitive assays containing soluble substrate should be included as a control.

As mentioned above, because of the reported difficulties in separating these *V. cholerae* exoenzymes there is a tendency to group the activities under the term mucinase complex (Burnet and Stone, 1947a; Burnet, 1948, 1949; Jensen, 1953; Singh and Ahuja, 1953; Freter, 1955; Kusama and Craig, 1970; Schneider and Parker, 1982). In the small intestine it is reasonable to propose that these exoenzymes in the complex interact in the breakdown of mucus and adhesion of the organism to the epithelial surface (Schrank and Verwey, 1976; Reed and Williams, 1978; Freter et al., 1978, 1981). Schneider and Parker (1982) proposed several non-exclusive mechanisms for the mucinase. Recently, Ramphal and Pyle (1983) noted that the source of neuraminidase and the chemical nature of the host mucus indicated specificity for certain sialic-acid glycoproteins. The associated enzymic activities in the mucinase complex may be partly responsible for this phenomenon. However, we would agree with Schneider and Parker (1982) that the mucinase complex facilitates *V. cholerae* penetration of the mucus barrier and, in addition, it may attack the glycosidic link between N-acetyl-galactosamine and β-galactose in the monosialoganglioside Gm1 and increase the receptivity for *V. cholerae* enterotoxin molecules. The precise role of the neuraminidase is still unsolved. Gascoyne and van Heyningen (1979) and Ackerman et al. (1980) attributed the conversion of membrane gangliosides to enterotoxin-binding gangliosides to the action of neuraminidase, but Holmgren (1981) did not endorse this. However, Vertiev et al. (1981) restated the
problem of solving this controversy because experimental and commercial preparations of *V. cholerae* neuraminidase are contaminated with other enzymes.

Further studies are being pursued in this laboratory to characterise the individual enzyme components of the mucinase complex and their role in *V. cholerae* pathogenesis.

REFERENCES


