RELATIONSHIP OF A TRYSIN-LIKE ENZYME IN RABBIT SPERMATOZOA TO CAPACITATION

L. J. D. ZANEVELD, P. N. SRIVASTAVA AND W. L. WILLIAMS

Department of Biochemistry, University of Georgia, Athens, Georgia 30601, U.S.A.

(Received 28th April 1969, revised 26th June 1969)

Summary. The existence of a trysin-like enzyme (TLE) in acrosomes of epididymal spermatozoa was confirmed and was further demonstrated to be present in acrosomes of ejaculated and capacitated spermatozoa. TLE rapidly removes the zona pellucida of the ovum. Extracts of acrosomes of ejaculated spermatozoa contain an inhibitor that is separated from the TLE by purification of the TLE.

The inhibitor of TLE is also present in seminal plasma. This enzyme–enzyme inhibitor relationship appears analogous to the corona-removing enzyme–decapacitation factor relationship and part of capacitation very likely involves removal of the inhibitor from TLE and decapacitation factor from the corona-removing enzyme. TLE is inhibited by soybean tryspin inhibitor and less effectively by mercaptoethanol.

Yamane (1935a, b) showed that canine sperm extracts obtained by toluene treatment had the ability to remove the external layers of ova and contained a tryspin-like enzyme (TLE). Buruiana (1956) also reported the presence of tryspin activity in spermatozoa of several species.

Hartree & Srivastava (1965) and Srivastava, Adams & Hartree (1965) described an acrosomal enzyme preparation from ram spermatozoa that removed the corona and zona pellucida from rabbit ova. Using essentially the same technique, Zaneveld, McRorie & Williams (1968) confirmed these results using rabbit sperm acrosomes and showed that removal of the corona by acrosomal extracts was inhibited by decapacitation factor (DF) preparations.

Stambaugh & Buckley (1968) extracted a TLE from epididymal sperm acrosomes that removed the zona pellucida from rabbit ova. Zaneveld, McRorie & Williams (1969) did not find a TLE in the acrosomal extracts of ejaculated spermatozoa employing the method of Hartree & Srivastava (1965). On further fractionation of these extracts by DEAE-column chromatography, TLE activity was found in one of the eluted protein peaks. This peak also possessed the ability to remove the zona pellucida from rabbit ova.

The purposes of the present experiments were to explain the absence of TLE activity from crude acrosome extracts of ejaculated spermatozoa in view of the demonstration of TLE in epididymal spermatozoa (Stambaugh & Buckley, 1968) and to determine the effect of capacitation on TLE activity of spermatozoa.
Epididymal, ejaculated and capacitated spermatozoa (the last obtained by flushing the uteri of does 12 hr after mating) were washed and treated with 0·15% Hyamine 2389 (Rohm and Haas Co.). The solution was subjected to centrifugation, precipitation with 80% alcohol, dialysis and lyophilization. Each extract was assayed using benzoyl arginine ethyl ester (BAEE) as substrate according to the method of Schwert & Takenake (1955). One unit of TLE was defined as that amount causing a change of 0·001 optical density/min. Under these conditions, 1 µg pancreatic trypsin (Worthington Biochem. Co.) has an activity of 20 units. Ejaculated sperm extracts were further purified by DEAE-column chromatography (Zaneveld et al., 1969).

Experiments on ova were performed by incubating 0·3 ml of the test solution with four ova from which the cumulus had been removed by treatment with hyaluronidase.

As shown in Table 1, only one of several crude extracts of ejaculated spermatozoa showed TLE activity and this activity was very low. After fractionation and apparent removal of an inhibitor, TLE activity was present. Epididymal sperm extracts always showed high TLE activity, and the TLE activity of crude extracts of the acrosomes of capacitated spermatozoa was as high as partially purified material from ejaculated spermatozoa. Five units of TLE activity (epididymal spermatozoa) was completely inhibited by 200 µg soybean trypsin inhibitor. Mercaptoethanol (10 mm) gave 50% inhibition of TLE activity. To demonstrate the presence of a trypsin inhibitor, boiled and centrifuged rabbit seminal plasma (0·1 ml = 2·5 µg protein) was added to 18 units of TLE from epididymal spermatozoa. This resulted in 85% inhibition of activity. Boiling was necessary to destroy the enzyme activity in seminal plasma. Crystalline serum albumin (2·5 µg), as a non-specific control, caused only an 8% decrease in TLE activity. Different extracts of epididymal spermatozoa, as a source of TLE activity, were inhibited by various samples of heated seminal plasma. One mg of crude DF (ultracentrifuged pellet) caused only 4% reduction in activity of pancreatic trypsin (20 units). Purified DF (Robertson, Pinsker, Caster & Williams, 1969), at 100 µg, did not inhibit TLE activity (15 units). Alcohol-precipitated, dialysed and lyophilized acrosomal extracts of ejaculated spermatozoa did not remove the zona pellucida from rabbit ova.

Table 1

<table>
<thead>
<tr>
<th>Activity of Various Acrosomal Extracts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sperm extract</td>
</tr>
<tr>
<td>----------------</td>
</tr>
<tr>
<td>Epididymal*</td>
</tr>
<tr>
<td>Ejaculated (crude)*</td>
</tr>
<tr>
<td>Ejaculated (purified)†</td>
</tr>
<tr>
<td>Capacitated*</td>
</tr>
</tbody>
</table>

* Alcohol precipitated, dialysed and lyophilized.
† After purification on DEAE column chromatography.
‡ Only one sample possessed 10 units of TLE activity, several others were not active.
Enzyme involved in ovum penetration

whereas further purification by DEAE-column chromatography yielded a fraction that readily removed the zona. Zona removal, like TLE activity, was inhibited by soybean trypsin inhibitor (200 μg) and mercaptoethanol (75 mm).

Since TLE activity is high in epididymal spermatozoa, absent in ejaculated spermatozoa and again present in capacitated spermatozoa, it is postulated that during contact with seminal plasma the TLE inhibitor is added to spermatozoa. As capacitation takes place, this inhibitor is removed. Although capacitated spermatozoa are frequently contaminated with white blood cells, these contain no trypsin-like enzymes (Zeya & Spitznagel, 1969). Previously, Haendle, Fritz, Trautschold & Werle (1965) and Fritz, Trautschold, Haendle & Werle (1968) demonstrated a heat-stable polypeptide trypsin inhibitor in the seminal vesicles of mice, guinea-pigs and bulls. The existence, after contact with seminal plasma, of a TLE-inhibitor complex is analogous to the complex between corona-removing enzyme (CRE) and DF described by Zaneveld et al. (1968). It is probable that both TLE inhibitor and DF are removed during capacitation enabling the CRE and TLE to act in sequence on the corona cell layer and the zona pellucida to effect penetration of the ovum. DF did not inhibit TLE or pancreatic trypsin, indicating that the inhibitors are specific for each penetration enzyme.

Characterization of the acrosomal enzyme as trypsin-like is justified because of hydrolysis of benzoyl arginine ethyl ester at pH 8 and inhibition by soybean trypsin inhibitor.

REFERENCES


