ABSTRACT

Most synapses in the central nervous system exhibit a prominent electron-opaque specialization of the postsynaptic plasma membrane called the postsynaptic density (PSD). We have developed a procedure for the isolation of PSDs which is based on their buoyant density and their insolubility in N-lauroyl sarcosinate. Treatment of synaptic membranes with this detergent solubilizes most plasma membranes and detaches PSDs from the plasma membrane so that they can be purified on a density gradient. Isolated PSDs appear structurally intact and exhibit those properties which characterize them in tissue. The isolated PSDs are of the size, shape, and electron opacity of those seen in tissue; they stain with both ethanolic phosphotungstic acid and bismuth iodide-uranyl lead and the fraction contains cyclic 3',5'-phosphodiesterase activity. Quantitative electron microscope analysis of the PSD fraction gives an estimated purity of better than 85%. Inasmuch as the PSD is associated primarily with dendritic excitatory synapses, our PSD fraction represents the distinctive plasma membrane specialization of this specific synaptic type in isolation.

Abbreviations used in this paper: BIUL, bismuth iodide-uranyl lead; E-PTA, ethanolic phosphotungstic acid; INT, p-iodonitrotetrazolium violet; KMnO₄-L, potassium permanganate lead; NLS, N-lauroyl sarcosinate; OsO₄-UL, osmium tetroxide-uranyl lead; PSD, postsynaptic density; SJC, synaptic junctional complex; SM, synaptic membrane.
describe a method for the isolation of PSDs from rat brain.

In the mammalian central nervous system PSDs, are characteristically associated with certain types of synapses. In cerebral cortex, for example, two morphologically distinct classes of synapses have been recognized based primarily on the characteristics of the PSD (Gray, 1959; Colonnier, 1968). One class (asymmetric or Gray type I) is characterized by a prominent PSD which extends 200–670 Å (van der Loos, 1964) into the cytoplasm and gives this synapse its asymmetric appearance (Gray, 1959; Colonnier, 1968). The other class (symmetric or Gray type II) is distinguished by the sparseness or absence of a PSD on the cytoplasmic side, giving these synapses a symmetrical appearance (Gray, 1959; Colonnier, 1968). In the cat visual cortex, asymmetric synapses are the major type and make up 80% of the total (Colonnier, 1968).

Synapses with a prominent PSD of the asymmetric type are usually excitatory (Gray, 1959, 1969; Walberg, 1968). In the cerebellum, asymmetric excitatory synapses include the parallel fiber-Purkinje cell synapses (Gray, 1961; Larramendi and Victor, 1967), mossy fiber-granule cell synapses (Gray, 1961), climbing fiber-Purkinje cell synapses (Chan-Palay and Palay, 1970), and climbing fiber-Golgi cell synapses (Chan-Palay and Palay, 1971); in the hippocampal formation, entorhinal-granule cell synapses (Naftad, 1967), mossy fiber-pyramidal cell synapses (Blackstad and Kjaerheim, 1961; Hamlyn, 1962); in the lateral geniculate, retinal-class I cell synapses (Guillery and Colonnier, 1970; Hendrickson, 1969); in the olfactory bulb, mitral cell-granule cell synapses (Rall et al., 1966; Price, 1968); and in the medial septal nucleus, pyramidal-septal cell synapses (Raisman, 1969; Raisman and Field, 1973). Most of the excitatory synapses are located on dendritic shafts or dendritic spines. On the other hand, inhibitory synapses are characterized by a thin sparse PSD and are usually located on neuronal soma (Chan-Palay and Palay, 1970; Gottlieb and Cowan, 1972). Thus, in general, prominent PSDs are a distinctive structural specialization associated with excitatory dendritic synapses.

The PSD exhibits a number of distinctive chemical and enzymatic properties as revealed by cytochemical methods. It is the site of a high cyclic 3',5'-nucleotide phosphodiesterase activity (Florendo et al., 1971). The PSDs, along with the other elements of the synaptic junction, selectively stain with ethanolic phosphotungstic acid (E-PTA) (Bloom and Aghajanian, 1966). Plasma membranes, even those immediately adjoining the synaptic complex, along with most other cellular structures (mitochondria, myelin, endoplasmic reticulum, etc.), fail to stain. Bismuth iodide-uranyl lead (BIUL) stains the specializations of the synaptic junction (PSD, cleft, and dense projections) and the extracellular coats of plasma membranes outside the synaptic region while it leaves relatively unstained the membrane itself and various intracellular organelles (Pfenninger, 1971).

Taken together, these data show that PSDs are morphologically distinctive entities with special cytochemical properties. In the central nervous system they are generally associated with excitatory synapses on dendrites. Yet very little is known to indicate what their function may be.

The purification of PSD described in this paper now makes possible a direct chemical analysis of the PSD and may clarify its role in synaptic function. Our isolation method makes use of the selective solubilizing action of sodium N-lauroyl sarcosinate (NLS) which previous studies have shown is useful in the isolation of gap junctions from liver tissue (Goodenough and Stoeckenius, 1972; Evans and Gurd, 1972). The method, based on the solubilization of SMs with NLS, followed by density gradient centrifugation, yields a highly purified fraction of PSD.

MATERIALS AND METHODS

Subcellular Fractionation

A PSD fraction was prepared from a SM fraction that contained intact synaptic junctional complexes (SJCs). Most of the plasma membrane could be solubilized with sodium NLS and a fraction of PSDs obtained from the insoluble residue by density gradient centrifugation.

All subcellular fractions were prepared from the forebrains of Sprague-Dawley rats, 40–60 days of age (Simonsen Laboratory, Gilroy, Calif.). The procedure described below uses Beckman type 30, SW 25.1 and SW 25.2 rotors (Beckman Instruments, Inc., Spinco Div., Palo Alto, Calif.). It should be readily adaptable for other rotor combinations so long as the amount of starting material and solution volumes are adjusted proportionately.

Isolation of SMs: A crude SM fraction was prepared based on a minor modification of the method described by Cotman and Taylor (1972) which was
adapted from the procedure developed by Davis and Bloom (1970, 1973). Brains (20 g wet weight) were homogenized in 60 ml of 0.32 M sucrose, 1 mM MgCl₂, and the homogenate was diluted to 200 ml with 0.32 M sucrose before centrifugation. This material was pelleted (type 30 rotor, 1,000 g for 5 min), the supernate was saved, and the pellet was resuspended by homogenization in 0.32 M sucrose and pelleted again. The remaining pellet was discarded and the combined supernates were pelleted (type 30 rotor, 14,500 g for 10 min) to obtain a crude mitochondrial fraction. This material was resuspended by homogenization and subjected to osmotic shock by diluting with 0.05 mM CaCl₂, pH 7.0 to a final volume of 250 ml and was allowed to stand for 20 min at 0°C.

The osmotically shocked crude mitochondrial fraction was next incubated in a solution containing succinate and p-iodonitrotetrazolium violet (INT). When the INT is 0.05 mM in CaCl₂, pH 7.0) and 8-10 ml of suspension crude mitochondrial fraction. This material was fractionated by density gradient centrifugation consisted of 7 ml each of 0.8, 1.0, 1.2 M sucrose, 0.05 mM CaCl₂, 0.2 M Na phosphate buffer, pH 7.5 were added and the suspension was then incubated at 30°C for 25 min. (The INT was best dissolved in buffer before adding succinate, and the solution was freshly prepared for each experiment.) After incubation the material was centrifuged (type 30 rotor, 8,720 g for 7 min) and the supernate discarded. The pellet was resuspended by homogenization in 0.16 M sucrose containing 0.05 mM CaCl₂ diluted to 300 ml with the same solution and pelleted (type 30 rotor, 34,880 g for 15 min). The washing procedure was repeated once more, and then the final pellet was resuspended in 0.32 M sucrose, 0.05 mM CaCl₂ to a total volume of 40-50 ml. This material was fractionated by density gradient centrifugation. 5 ml of suspension were applied to each of three gradients designed for the SW 25.1 rotor which consisted of 7 ml each of 1.0, 1.4, and 2.2 M sucrose, all 0.05 mM in CaCl₂, pH 7.0 and the gradient was centrifuged at 63,600 g for 75 min. The material at each interface was collected, diluted 1:2 with 0.1 mM EDTA and pelleted (type 30 rotor, 78,500 g for 20 min). The final pellet was resuspended in 0.05 mM CaCl₂, pH 7.0. The yield of membrane was about 60 mg protein/20 g wet weight brain. This SM fraction served as the source for the preparation of the PSDs.

Isolation of Other Fractions: A mitochondrial fraction and a myelin membrane fraction were obtained from the same gradient used to isolate SM. The material banding at the 0.32-0.8 M sucrose interface was taken as a myelin membrane fraction. A mitochondrial fraction was isolated as the pellet on a modified gradient which included a layer of 1.4 M sucrose underlying the 1.2 M layer in the usual gradient. The SJC fraction was isolated from the SM fraction using Triton X-100 as previously described (Cotman and Taylor, 1972). Triton X-100 was added at a ratio of 2 µl/mg of membrane protein and solubilization was carried out at 4°C. SJCs were then obtained by pelleting through a layer of 1.0 M sucrose. Mitochondrial, myelin membrane, and SJC fractions were treated with 3.9% NLS in 10 mM Bicine, pH 7.5. NLS was recrystallized once from 90% ethanol before use. In the case of solubilization with 0.5% NLS, 0.66 ml of NLS solution was added per milligram of SM protein (final concentration about 0.4%) and incubated for 6-10 min at 4°C before loading the suspension onto the gradient. With 3.9% NLS, 20 ml of the NLS solution were added to 5-8 ml of the membrane suspension containing 40-80 mg of SM protein, so that the final detergent concentration was approximately 3%. The mixture was incubated at 4°C for 10 min before application to the gradient. After solubilization, 8-9 ml of either suspension were applied to a discontinuous sucrose gradient for the SW 25.1 rotor which consisted of 7 ml each of 1.0, 1.4, and 2.2 M sucrose, all 0.05 mM in CaCl₂, pH 7.0 and the gradient was centrifuged at 63,600 g for 75 min. The material at each interface was collected, diluted 1:2 with 0.1 mM EDTA and pelleted (type 30 rotor, 78,500 g for 20 min). The PSD sedimented primarily at the 1.4-2.2 M sucrose interface, while remaining insoluble membrane fragments banded primarily on top of 1.0 M sucrose. About 0.5 mg of PSD protein was obtained from 20 g of rat forebrain.

In some experiments, to minimize handling when the material was to be used for electron microscope analysis, 2.2 M sucrose was sometimes omitted from the gradient and the PSD fraction was obtained as a pellet.

Isolation of PSDs: In order to isolate a PSD fraction from the SM fraction, the latter was treated with sodium NLS and the suspension resolved on a sucrose density gradient into PSD, membrane, and soluble fractions. We used NLS as either a 0.5% (wt/vol) or 3.9% (wt/vol) solution in 10 mM Bicine, pH 7.5. NLS was recrystallized once from 90% ethanol before use. In the case of solubilization with 0.5% NLS, 0.66 ml of NLS solution was added per milligram of SM protein (final concentration about 0.4%) and incubated for 6-10 min at 4°C before loading the suspension onto the gradient. With 3.9% NLS, 20 ml of the NLS solution were added to 5-8 ml of the membrane suspension containing 40-80 mg of SM protein, so that the final detergent concentration was approximately 3%. The mixture was incubated at 4°C for 10 min before application to the gradient. After solubilization, 8-9 ml of either suspension were applied to a discontinuous sucrose gradient for the SW 25.1 rotor which consisted of 7 ml each of 1.0, 1.4, and 2.2 M sucrose, all 0.05 mM in CaCl₂, pH 7.0 and the gradient was centrifuged at 63,600 g for 75 min. The material at each interface was collected, diluted 1:2 with 0.1 mM EDTA and pelleted (type 30 rotor, 78,500 g for 20 min). The PSD sedimented primarily at the 1.4-2.2 M sucrose interface, while remaining insoluble membrane fragments banded primarily on top of 1.0 M sucrose. About 0.5 mg of PSD protein was obtained from 20 g of rat forebrain.

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Previously Cotman and Taylor (1972) referred to this fraction as the synaptic complex fraction. The notation is changed here to conform to that formally adopted by Davis and Bloom (1973) and to be more consistent with the nomenclature used by electron microscopists (Peters et al., 1970). This also avoids the possibility of confusion with synapticosomal fractions which have also been referred to as synaptic complex fractions by some (Korneguth et al., 1969).
Electron Microscopy

In order to assess the composition of fractions, samples were prepared for quantitative electron microscope analysis by the method of Cotman and Flansburg (1970). Samples were pelleted in small capsules in order to form pellets which were flat and sufficiently thin so that they could be completely examined in a single cross section (Cotman and Flansburg, 1970). Samples were handled as previously described, except for PSD fractions. PSD fractions adhered to the walls of these tubes, so it was necessary to prepare the micropellet by placing a small volume (10–20 μl, containing roughly 20–40 μg of protein) directly in the tip of the capsule. Optimal structural preservation of PSD fractions required omission of EDTA and a minimum of mechanical shear. After centrifugation of the capsules, pellets were fixed with 4% glutaraldehyde in Caulfield buffer (Caulfield, 1957), postfixed with osmium tetroxide in Caulfield buffer, block stained with uranyl acetate (Kellenberger et al., 1958), and embedded in Epon-Araldite. Sections were cut on an LKB III ultramicrotome and stained with lead citrate for 5 min (Venable and Coggeshall, 1965). In some cases samples were stained with E-PTA, based on the method of Bloom and Aghajanian (1966) as previously described (Cotman et al., 1971), or with BIUL as described by Pfenninger (1971). Sections from some E-PTA specimens were counterstained with 1% potassium permanganate (Glaue, 1965) for 1–2 min, followed by lead citrate. For quantitative analysis of fractions, micrographs were taken with a Zeiss microscope (EM 9S) at × 8,000 magnification and examined in prints at × 24,000 or 32,000. Micrographs for detailed structural analysis were made with a Siemens IA electron microscope.

The composition of fractions was quantitatively evaluated from a series of micrographs taken at uniform intervals through the thickness of the disklike pellet. Electron micrographs were taken at intervals of about 20 μm thick. Within each micrograph a 2 × 2 μm square was selected at random and all particles within it were classified, counted, and their cross-sectional areas measured. Areas were estimated from measurements of length and width for roughly rectangular objects or by using an overlying grid for objects of irregular shape. For membrane vesicles the diameter was measured and the cross-sectional area of membrane computed assuming a membrane thickness of 100 Å.

Biochemical Assays

Protein was determined by the method of Lowry et al. (1951). The method was slightly modified for fractions which contained formazan since these gave erroneously high readings. In most cases the interference was corrected for by reading the sample vs. a blank which contained sample, but to which the Folin reagent was not added. In a few instances protein was solubilized in 1 N NaOH at 37°C for 1 h and the formazan removed by centrifugation before assay of the supernate.

Cytochrome oxidase (cytochrome c oxidoreductase, E.C. 1.9.3.1), Na+,K+-ATPase (Na+,K+-activated ATP phosphohydrolase, E.C. 3.6.14), acid phosphatase (orthophosphoric monoester phosphohydrolase, E.C. 3.1.3.2), and 5'-nucleotidase (E.C. 3.1.3.5) were assayed by methods which have been described elsewhere (Cotman and Matthews, 1971). Cyclic 3',5'-phosphodiesterase was assayed by the method of Breckenridge and Johnson (1969). Total phosphorus was assayed by the method of Ames (1966).

Reagents

N-Lauroyl sarcosinate, sodium salt was obtained from K and K Laboratories, Inc. (Plainview, N. Y.). It was recrystallized once from 90% ethanol before use. Cytochrome c, p-iodonitrotetrazolium violet, and all substrates for enzyme assays were purchased from Sigma Chemical Co. (St. Louis, Mo.). Glutaraldehyde (electron microscope grade) and osmium tetroxide were obtained from Polysciences, Inc. (Warrington, Pa.). Other chemicals were of analytical grade and were purchased from standard suppliers. All solutions were prepared in glass-distilled water.

RESULTS

Isolation of the PSD Fraction

The strategy we developed to isolate a PSD fraction involved the preparation of an SM fraction containing synaptic junctions, incubation of the SM fraction with NLS, and separation of the PSD, which remained insoluble, from other residual contaminants by density gradient centrifugation.

The SM fraction which served as the source of PSD contained plasma membranes of various types, some opaque bodies, and a few synaptic junctions with prominent PSDs (Fig. 1, arrows). In this fraction the length of the PSD varied between 0.1 and 0.5 μm. Treatment of the SM fraction with NLS solubilized 97% of the protein and 64% of the phosphorus. Resolution on a discontinuous sucrose density gradient produced two major fractions: a light fraction floating on 1.0 M sucrose and a heavy fraction which banded between 1.4 and 2.2 M sucrose. Protein distributed equally between the heavy and light fraction and the phosphorus remained mainly in the light fraction (Table I). The other interfaces were relatively devoid of protein. The dense fraction contained relatively large amounts of bound formazan, judging from its deep purple color. It was typically
FIGURE 1  An electron micrograph of the SM fraction which was used as the source of PSDs. PSDs (arrows) are present as part of synaptic junctions. Bar is 1 μm. × 32,000.

FIGURE 2 A micrograph which illustrates the structures present in a pellet prepared from the 1.0 M sucrose fraction. Only membrane fragments are present. Bar is 1 μm. × 32,000.
TABLE 1
Distribution of Protein and Total Phosphorus on the Density Gradient after Solubilization of the SM, Myelin, or Mitochondrial Fractions with 3% NLS

| Fraction      | SM protein | SM phosphorus | Myelin, protein | Mitochondria, protein |
|---------------|------------|---------------|-----------------|-----------------------|
| 0.0–1.0 M     | 46.0 ± 3.0 | 97.0 ± 0.2    | 96.0 ± 1.1      | 19.7 ± 5.0            |
| 1.0–1.4 M     | 5.4 ± 1.3  | --            | <1.0            | 7.0 ± 0.4             |
| 1.4–2.2 M     | 45.5 ± 1.2 | 2.9 ± 0.2     | <1.0            | 3.7 ± 0.5             |
| 2.2 M pellet  | 3.3 ± 3.0  | --            | 2.7 ± 0.4       | 69.4 ± 5.1            |
| Total insoluble† | 2.3 ± 0.2 | 37.4 ± 1.6   | 6.1§             | 10.2 ± 0.7            |

* Data are means of two to four experiments except § which represents a single determination. Recovery of protein in gradient fractions for SM averaged 43% and for mitochondria, 80%. Recovery of phosphorus was 50%.
† Total insoluble refers to the percent of the protein in the starting material (SM, myelin, or mitochondria) which sediments at 44,000 g for 30 min (30 rotor) after treatment with 3% NLS.

obtained as aggregates that were quite difficult to resuspend and tended to adhere to glassware and centrifuge tubes.

Electron microscope examination of the fraction floating on 1.0 M sucrose showed that it contained exclusively membrane fragments (Fig. 2). By contrast, the 1.4–2.2 M fraction (Fig. 3) contained numerous electron-opaque barlike structures (P) which resembled PSDs seen in the SM fraction and in sections of brain tissue. These opaque barlike or platelike structures were approximately the same size as PSDs seen in the SM fraction or in tissue sections (200–600 Å wide × 0.1–0.5 μm long). Other structures were present which resembled PSDs in size and electron opacity but which were slightly more extended or amorphous in shape (P'). At high magnification (Fig. 3, inset) the barlike structures appeared to consist of a diffusely defined matrix which was similar to that seen in PSDs in the SM fraction. Few, if any, of those structures were attached to pre- or postsynaptic membranes, and only small amounts of plasma membrane were seen (M). On the basis of their morphological characteristics we tentatively identified the electron-opaque structures in the 1.4–2.2 M fraction as PSDs.

**Cytochemical Characteristics of the PSD Fraction**

We sought to characterize these structures further by their cytochemical staining properties. PSDs stain selectively with E-PTA in tissue samples (Bloom and Aghajanian, 1966) or in fractions (Cotman et al., 1971; Cotman and Taylor, 1972; Jones and Breartley, 1972; Davis and Bloom, 1973) and with BIUL in tissue sections (Pfenniger, 1971). As shown in Fig. 4 the barlike structures (P) stained with E-PTA and resembled those seen in the specimens fixed with osmium tetroxide and stained with uranyl and lead ions (OsO₄-UL) (compare Figs. 3 and 4). In addition, the dense, irregular-shaped structures (P') stained with E-PTA. By contrast, the structures in the 1.0 M sucrose fraction did not stain. This indicates that the staining of structures in the PSD fraction was specific and further suggests that the specificity of E-PTA was not affected by NLS treatment. Structures in the PSD fraction also stained with BIUL. We found, however, that BIUL staining of subcellular fractions was somewhat less selective than has been reported for tissue sections. In the SM fraction, in addition to the synaptic junction and membrane coats, small circular structures (possibly condensed synaptosomes) stained with BIUL. Nonetheless, the stainability of the barlike structures in the PSD fraction with BIUL and E-PTA provided strong support for the identification of these structures as PSDs.

Evidence that all structures in the PSD fraction were PSDs would be gained if it could be shown that all structures, except membranes, stained with E-PTA. Qualitatively this appeared to be the case. The packing density of structures staining with E-PTA (Fig. 4) was similar to that of the structures seen in samples prepared by a general...
Figure 3 A survey micrograph of a typical field of the structures present in a pellet prepared from the 1.4–2.2 M sucrose fraction. Many electron-opaque structures which appear like PSDs are present and range in size from 200 to 600 Å wide and from 0.1 to 0.5 μm long. Other structures which appear similar in electron density and size, but which are less barlike in their overall structural appearance, are present (P'). A few fragments of membrane (M) can be found. A possible contaminant (C) is shown. The inset shows the predominant structure in the fraction at high magnification. The PSD consists of a relatively dense matrix. A plasma membrane is not evident along the length of the PSD. Sample was fixed with glutaraldehyde and OsO₄-UL. Bar is 1 μm. × 32,000. Inset, × 108,000.
FIGURE 4 A survey micrograph of a typical field of the 1.4–2.2 M sucrose fraction after staining with E-PTA. The barlike structures are stained (P) and are very similar in appearance and size to the barlike structures seen in a preparation fixed with glutaraldehyde and OsO₄-UL (compare insets of Fig. 4 to Fig. 3). Besides the barlike structures, other structures which are of a similar size but more irregular in their shape also stain with E-PTA (P'). Bar is 1 μm. × 32,000. Inset, × 108,000.
staining procedure using OsO₄-UL (Fig. 3). Furthermore, the relative number of barlike and irregular-shaped PSDs was similar in both preparations. In order to obtain direct evidence that all structures (except membranes) stained with E-PTA, serial sections from an E-PTA block were prepared. A ribbon consisting of a few sections was divided, and one set of sections examined directly. The other set of alternate sections was counterstained with potassium permanganate and lead ions (KMnO₄-L) in order to reveal any E-PTA-negative structures. KMnO₄-L is a general stain similar to OsO₄-UL, and thus reveals E-PTA-negative structures. For example, we found that if an E-PTA-stained SM fraction was counterstained with KMnO₄-L, membranes and other structures were revealed and the composition of the fraction appeared like that of an OsO₄-UL preparation. In the PSD fraction detailed comparison of identical areas on adjoining sections showed that the great majority of structures could be identified in both sections. The comparison of structures in the E-PTA- and E-PTA-KMnO₄-L-stained sections was like tracing these structures in two identically stained sections. We concluded that, except for membranes, nearly all structures in the 1.4–2.2 M sucrose fraction possessed the characteristic staining properties of PSDs. (Direct counterstaining of a pre-examined area of an E-PTA section with KMnO₄-L or other stains was not feasible since the damage from the beam of the electron microscope incurred during initial examination prevented subsequent staining of the area.)

Specificity of the Isolation Procedure for PSDs

The effect of NLS on fractions devoid of synaptic junctions was examined to eliminate the possibility that structures similar in appearance to PSDs might be produced by such detergent treatment. A mitochondrial fraction and a crude myelin membrane fraction were treated with NLS and resolved on a sucrose density gradient under conditions identical to those used to prepare the PSD fraction. When a mitochondrial fraction was treated with 3% NLS, protein distributed throughout the gradient, but almost no protein was obtained at the 1.4–2.2 M interface (Table I). The structures in this fraction appeared as membrane sheets in the electron microscope. No PSD-like structures were seen. Treatment of a myelin membrane fraction with 3% NLS also produced very little protein in the fraction of a density equivalent to that of PSD (interface at 1.4–2.2 M sucrose) (Table I). Therefore, the procedure was selective: detergent treatment of other fractions that did not contain PSDs did not generate material with similar sedimentation properties or ultrastructural appearance. In addition, in view of the low levels of mitochondria or myelin in the initial SM fraction before NLS treatment (Cotman and Taylor, 1972), very little of the protein in the PSD fraction could have arisen from mitochondria or myelin.

In contrast with fractions devoid of synaptic junctions which did not produce a PSD fraction, a SJC fraction which was enriched in PSDs yielded a highly purified preparation of PSDs after treatment with NLS. SJC fractions were isolated by treatment of the SM fraction with Triton X-100 followed by density gradient centrifugation (Cotman and Taylor, 1972). By quantitative electron microscope analysis the fraction was composed of about 60–70% synaptic junctions including intact clefts and PSDs with small attached membrane segments and 30–40% unidentifiable membrane fragments.* No other structures were visible. Incubation of the SJC fraction with 3% NLS and resolution of the suspension on a sucrose density gradient produced a PSD fraction banding at 1.4–2.2 M sucrose which in the electron microscope was indistinguishable from the PSD fraction obtained directly from the SM fraction. The major difference was the increased yield of PSD per milligram of SJC protein compared to that per milligram of SM protein. Whereas 2 mg of SJC protein provided 0.5 mg of PSD, 50 mg of SM protein was required to obtain this amount of PSDs. The increased yield of PSD from the SJC fraction was in reasonable agreement with the estimated content of PSDs in the SJC fraction. Thus a PSD fraction could be directly isolated from an SJC fraction in a yield compatible with the enrichment of SJC fraction in PSDs.

Enzymatic Characterization of the PSD Fraction

We also examined fractions obtained by NLS treatment for enzymatic activities to detect possible

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*Churchill, L., G. Banker, and C. W. Cotman. 1974. Protein and carbohydrate composition of central nervous system synapses: Analysis of isolated synaptic junctional complexes and postsynaptic densities. Manuscript submitted for publication.
ble contamination not apparent in electron micro-
graphs, as well as the possible presence of marker
enzyme activities. Since NLS at 3% severely
inactivated most enzymes studied, a lower deter-
genent concentration was used (0.4%). Preliminary
experiments indicated that this concentration pro-
duced a similar PSD fraction, but with slightly
more membrane contamination. Cytochrome ox-
dase activity indicative of mitochondrial contami-
nation was low in PSD fractions (Table II). Acid
phosphatase, Na,K-ATPase, and 5'-nucleotidase
activities were low, indicating the relative absence
of membrane and lysosomal constituents. Since
cytochemical studies have suggested that cyclic
3',5'-nucleotide phosphodiesterase activity may
be associated with PSDs, we analyzed the phos-
phodiesterase activity present in these fractions.
Although the levels were variable and recoveries
low, the reproducible presence of phosphodiester-
ase in the PSD fraction was consistent with the
suggestion that it is a constituent of the PSD. Of
the enzymes examined, only phosphodiesterase
gave any indication of association with the PSD
fraction. Thus these data support the morphologi-
cal data on the identification of PSDs. In them-
selves, however, the enzyme data must be con-
sidered with caution in view of the variable and
often low recoveries and the use in these experi-

| Enzymatic Characterization of Membrane and PSD Fractions* Obtained after Treatment of the SM Fraction with 0.4% NLS and Resolved on the Discontinuous Density Gradient |
|---------------------------------------------------------------|
| Enzyme              | Fraction | Membrane | PSD | Percent total activity |
|---------------------|----------|-----------|-----|------------------------|
| Cytochrome oxidase   | Membrane | 93        | 7   | 97                     |
|                     | PSD      | 7         | 3   | 3                      |
|                     | % recovery | 16      | 185 | 156              |
| Acid phosphatase    | Membrane | 98        | 2   | 93                     |
|                     | PSD      | 2         | 7   | 7                      |
|                     | % recovery | 97      | 32  | 36                 |
| Na+ -K+—ATPase      | Membrane | 97        | 11  | 97                     |
|                     | PSD      | 3         | 4   | 1                      |
|                      | % recovery | 11      | 4   | 11                  |
| 5'-Nucleotidase     | Membrane | 79        | 20  | 94                     |
|                     | PSD      | 21        | 31  | 6                      |
|                      | % recovery | 20      | 31  | 64                  |
| Cyclic nucleotide phosphodiesterase | Membrane | 48        | 52  | 74                     |
|                     | PSD      | 52        | 26  | 78                     |
|                      | % recovery | 0.23 | 0.08 | 42                 |
|                      | % recovery | 0.23 | 0.08 | 3.9                     |
| Protein             | Membrane | 75        | 75  | 70                     |
|                     | PSD      | 25        | 30  | 30                     |
|                      | % recovery | 97      | —   | —

Each column represents data from an individual experiment.
* The membrane fraction refers in this case to the combined 1.0 M and 1.0–1.4 M fractions. The PSD fraction consists of
the 1.4–2.2 M fraction and the pellet.
† Percent total activity refers to the distribution of particulate enzyme recovered in the gradient fractions. The percent
recovery refers to the recovery of activity in the total particulate plus soluble fractions from the untreated SM fraction.
The activity of cyclic 3',5'-phosphodiesterase measured in the SM fraction was 4 µmol 5'-AMP formed/mg protein/h.
The activity levels of other enzymes were in the range previously reported (Cotman and Matthews, 1971).
§ The variability in the distribution of enzyme activity does not appear related to the different recoveries since the
lower percentages in the PSD fraction (26 and 22%) occur in the experiments with the lowest and highest activity.
Various attempts to improve the recovery were not successful.
ments of a detergent concentration which resulted in contamination.

Quantitative Electron Microscope Analysis

Quantitative electron microscope analysis of the PSD fraction in comparison with the SM fraction revealed an enrichment in PSDs with a decrease in membrane and other contaminants. As described previously, there were two types of structures which appeared like PSDs in the 1.4–2.2 M sucrose fraction. One type of structure was densely stained and barlike and was similar in length and width to PSDs in tissue sections (0.1–0.5 μm long x 200–600 Å wide). These were identified as PSDs. Another type also stained densely, but because it was thicker or branched slightly it was classified as PSD-like. These particles might have been damaged PSDs or PSDs sectioned so that their barlike structure was not recognizable. As shown in Table III, the PSDs and PSD-like structures represented 73–93% of the identifiable structures present in the PSD fraction compared to only 2% of the structures in the SM fraction. Inasmuch as estimates of composition based solely on the number of structures present are very likely misleading when marked size differences exist among different structures, we also measured the cross-sectional area occupied by each category. By this measure, PSDs and PSD-like structures made up 86–92% of the material in the PSD fraction compared to 4% in the SM fraction. Thus in the PSD fraction there was an enrichment of better than 20-fold in PSD structures. The membrane contamination of the PSD fraction was low (7–25% by number, 8–12% by area) as were other types of contaminants (0–2% by number or area).

Since the PSD-like structures stained with E-PTA, it seemed likely that they were damaged PSDs or PSDs sectioned so that their barlike structure was not evident. To test the latter possibility directly, an analysis of serial sections was undertaken. The results established that, of those structures that could be followed through three serial sections, 37% of the PSD-like structures appeared as a PSD in one out of three serial sections. Thus, variations in sectioning plane resulted in the presence of PSD-like structures. PSDs were found to be susceptible to damage. Care had to be taken in the handling of these fractions, otherwise more numerous extensively damaged PSD-like structures and fewer PSDs were observed. For example, prolonged contact

| Structure     | Number | Area |
|---------------|--------|------|
| PSD           | 30     | 34   |
| PSD-like      | 43     | 59   |
| Membrane      | 25     | 7    |
| Other contaminants | 2    | 0    |

Data are given in terms of the percent of the total number of structures in each category and by the percent of the area occupied by each structure. Each column represents a separate experiment.

With 0.1 mM EDTA (overnight at 4°C) increased the prevalence of PSD-like structures. Thus, nonideal orientation of PSDs and structural damage gave rise to PSD-like structures. Previously it was shown that all PSD-like structures stained with E-PTA. These three facts provide strong support for the identification of all the PSD-like structures as PSDs.

In conclusion, we have prepared a highly purified PSD fraction and have documented its composition in several ways. The PSD fraction contained numerous barlike structures which closely resembled PSDs seen in tissue sections. Variations from the barlike structure appeared to arise from differences in alignment of particles with respect to the plane of section, as well as from partial damage due to the isolation procedure. Both damaged and ideally sectioned and preserved PSDs stained with E-PTA and BIUL. Contaminant particles such as mitochondria, myelin, and membrane fragments did not generate a PSD-type fraction when treated in a similar manner, so that the PSD was not a
detergent artifact generated from any subcellular fraction. In addition, as shown in the following paper (Banker et al., 1974), proteins characteristic of contaminating fractions were not seen in the PSD fraction. An SJC fraction greatly enriched in clearly identifiable PSDs can be used to isolate a PSD fraction in comparably greater yield per milligram of SJC than per milligram of SM protein. Cyclic nucleotide phosphodiesterase activity was prevalent in the PSD fraction whereas enzyme markers for possible contaminating structures sedimented primarily in other gradient fractions, leaving the PSD fraction devoid of such contamination. The enrichment of PSD from the SM fraction was about 20-fold and the resulting fraction was about 85% pure.

DISCUSSION

Our results establish that PSDs can be isolated from adult rat brain in high purity. The enrichment in PSD from SM is estimated at 20-fold and that from homogenate must be substantially greater. The isolated PSDs retain those properties known to characterize them in intact tissue. They stain with both E-PTA and BIUL and they contain phosphodiesterase activity, though apparently at reduced levels. There is little indication of contamination by a number of different criteria and we conclude that our procedure produces an extremely pure fraction of PSDs. At present, however, we do not know whether isolated PSDs retain all properties represented in vivo and we have not excluded all possible types of contamination.

Isolated PSDs are different from most subcellular fractions in their physical properties. PSDs are characteristically very adhesive. PSD fractions contain aggregates which are difficult to disrupt, and the PSDs readily adhere to glass or plastic surfaces in the absence of EDTA. It is possible that these properties are a reflection of the sequestering of Ca during isolation. This is compatible with earlier observations that Ca^{2+} aided in the isolation of well-preserved SJC (Cotman et al., 1971; Davis and Bloom, 1973). In our procedure PSDs carry considerable formazan, and this phenomenon is yet to be explained. At least a part of the formazan appears bound to PSDs since we have been unable to resolve formazan entirely from PSDs under a variety of centrifugation conditions. The formazan could arise from INT reduction by other structures and a preferential binding of released formazan by PSDs, or alternatively PSDs may themselves be capable of INT reduction. We should point out that formazan is not absolutely essential for the isolation procedure, since in one experiment we succeeded in isolating PSDs in the absence of INT, though the yield was lower than usual. The adhesive properties of the material and the presence of bound formazan are seen to a degree in SJC fractions (Cotman and Taylor, 1972), which are enriched in PSDs, but are generally less evident in the parent SM fraction.

PSDs in our fraction most often are not attached to plasma membranes, yet they closely resemble the PSDs found in intact tissue. In addition PSD structure is not greatly altered by a treatment that solubilizes most plasma membrane. Consequently the conception of the PSD as a "membrane thickening" is incorrect, as already noted (Akert et al., 1969). Rather, in agreement with van der Loos (1964), it is appropriate to consider the PSD as a separate organelle which is associated with the plasma membrane in situ, but which has its own distinctive existence.

The PSDs isolated with our procedure are apparently derived from the asymmetric type of synapse (type I). They are generally similar in size to the PSDs from cortical asymmetric synapses (Colonnier, 1968), and no structures in these fractions resemble the thin sparse PSDs often present at symmetric (type II) synapses. On this basis, the purified PSDs most likely arise predominantly from excitatory axodendritic synapses. Thus our PSD fraction represents the distinctive organelle of this specific synaptic type in isolation.

PSDs have not previously been isolated as a pure fraction, and the addition of this fraction to the ensemble of synaptic structures which can be isolated by subcellular fractionation further refines the "dissection" of the synapse. It is now well known that synaptic boutons, often with an attached segment of postsynaptic membrane, can be isolated free from most other contaminating structures (Gray and Whittaker, 1962; de Robertis et al., 1962). These particles, called synaptosomes (Fig. 5 a), represent a complete synaptic unit in isolation. Synaptosomes can be disrupted and the plasma membrane freed from other structures and isolated as a synaptic plasma membrane fraction (Whittaker et al., 1964; Rodriguez de Lores Arnaiz et al., 1967; Cotman and Matthews, 1971; Morgan et al., 1971) (Fig. 5 b). The synaptic membrane is still quite complex, often consisting of plasma membranes from pre- and postsynaptic neurons as well as segments from membrane outlying the synaptic junction itself. Much of the

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membrane not part of the synaptic junction, as well as contaminating membrane, can be removed and SJC's prepared (de Robertis et al., 1967; Davis and Bloom, 1973; Cotman and Taylor, 1972). SJC's which are well preserved include a prominent PSD, short segments of pre- and postsynaptic membrane, and a synaptic cleft (Fig. 5 c). As described in this paper, PSD's can be isolated either directly from a SM fraction or from synaptic junctions (Fig. 5 d). These fractions, together with synaptic vesicle fractions, provide discrete components for analysis of the structure and function of central nervous system synapses.

Our PSD fraction affords a new approach to examine this part of a central nervous system synapse. In the past, the analysis of PSD has been based primarily on morphological and cytochemical techniques. The extensive enrichment and purity of our fraction provides an opportunity to analyze the properties of this structure directly. In the subsequent paper we characterize the proteins which are the major constituent of PSD's (Banker et al., 1974). The carbohydrate components of PSD's are reported elsewhere. We thank Mrs. Pat Lemestre for secretarial aid and Mrs. Carol Green for technical assistance.

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REFERENCES

Akert, K., H. Moor, K. Pfenninger, and C. Sandri. 1969. Contributions of new impregnation methods and freeze-etching to the problems of synaptic fine structure. Prog. Brain Res. 31:223.

Ames, B. N. 1966. Assay of inorganic phosphate, total phosphate and phosphatases. Methods Enzymol. 8:115.

Banker, G., L. Churchill, and C. W. Cotman. 1974. Proteins of the postsynaptic density. J. Cell Biol. 63:456.

Blackstad, T. W., and A. Kjaerheim. 1961. Special axo-dendritic synapses in the hippocampal cortex: Electron and light microscopic studies on the layer of mossy fibers. J. Comp. Neurol. 117:133.

Bloom, F. E. 1970. Correlating structure and function of synaptic ultrastructure. The Neurosciences: Second Study Program. F. O. Schmitt, editor. The Rockefeller University Press, New York. 729.

Bloom, F. E., and G. K. Aghajanian. 1966. Cytochemistry of synapses: selective staining for electron microscopy. Science (Wash. D. C.). 154:1575.

Breckenridge, B. M., and R. E. Johnston. 1969. Cyclic 3',5'-nucleotide phosphodiesterase in brain. J. Histochem. Cytochem. 17:505.

Caulfield, J. B. 1957. Effects of varying the vehicle for OsO4 in tissue fixation. J. Biophys. Biochem. Cytol. 3:827.

Chan-Palay, V., and S. L. Palay. 1970. Interelections of basket cell axons and climbing fibers in cerebellar cortex of the rat. Z. Anat. Entwicklungsgesch. 132:191.

Chan-Palay, V., and S. L. Palay. 1971. Tendril and glomerular collaterals of climbing fibers in the granular layer of the rat's cerebellar cortex. Z. Anat. Entwicklungsgesch. 133:247.

Colonnier, M. 1968. Synaptic patterns on different cell types in the different laminae of the cat visual cortex. An electron microscope study. Brain Res. 9:268.

Cotman, C. W., and D. Flansburg. 1970. An analytical micro-method for electron microscopic study of the composition and sedimentation properties of subcellular fractions. Brain Res. 22:152.

Cotman, C. W., W. Levy, G. Banker, and D. Taylor. 1971. An ultrastructural and chemical analysis of the effect of Triton X-100 on synaptic plasma membranes. Biochem. Biophys. Acta. 249:406.

Cotman, C. W., and D. A. Matthews. 1971. Synaptic...
plasma membranes from rat brain synaptosomes: Isolation and partial characterization. Biochim. Biophys. Acta. 249:380.

Coitman, C. W., and D. Taylor. 1972. Isolation and structural studies on synaptic complexes from rat brain. J. Cell Biol. 55:696.

Davis, G., and F. E. Bloom. 1970. Proteins of synaptic junctional complexes. J. Cell Biol. 47(2, Pt. 2):46 a. (Abstr.)

Davis, G. A., and F. E. Bloom. 1973. Isolation of synaptic junctional complexes from rat brain. Brain Res. 62:135.

De Robertis, E. D. P. 1964. Histophysiology of Synapses and Neurosecretion. Pergamon Press Inc., Elmsford, N. Y.

De Roberts, E., J. M. A. Zuccurra, and S. Fiszer. 1967. Ultrastructure and cholinergic binding capacity of junctional complexes isolated from rat brain. Brain Res. 5:45.

De Robertis, E., A. Pellegrino de Iraldi, G. Rodriguez de Lores Arnaiz, and L. Salganicoff. 1962. Cholinergic and non-cholinergic nerve endings in rat brain. J. Neurochem. 9:23.

Evans, W. H., and J. W. Gurd. 1972. Preparation and properties of nexuses and lipid-enriched vesicles from mouse liver plasma membranes. Biochem. J. 128:691.

Florendo, N., R. Barnett, and P. Greengard. 1971. Cyclic 3',5'-nucleotide phosphodiesterase: Cytochemical localization in cerebral cortex. Science (Wash. D. C.). 173:745.

Glairt, A. M. 1965. Techniques for Electron Microscopy. D. H. Kay, editor. F. A. Davis Company, Philadelphia, Pa. 2nd edition. 261.

Goodenough, D. A., and W. Stoeczenius. 1972. The isolation of mouse hepatocyte gap junctions. J. Cell Biol. 54:646.

Gottlieb, D. I., and W. M. Cowan. 1972. On the distribution of axon terminal containing spheroidal and flattened synaptic vesicles in the hippocampus and dentate gyrus of the rat and cat. Z. Zellforsch. Mikrosk. Anat. 129:413.

Gray, E. G. 1959. Axo-somatic and axo-dendritic synapses of the cerebral cortex. An electron microscope study. J. Anat. 93:420.

Gray, E. G. 1961. The granule cells, mossy synapses and Purkinje spine synapses of the cerebellum. Light and electron microscope observations. J. Anat. 92:345.

Gray, E. G. 1969. Electron microscopy of excitation and inhibitory synapses: A brief review. Prog. Brain Res. 31:141.

Gray, E. G., and V. P. Whittaker. 1962. The isolation of nerve endings from brain. An electron microscopic study of all fragments derived by homogenization and centrifugation. J. Anat. 96:79.

Guillery, R. W., and M. Colonier. 1970. Synaptic patterns in the dorsal lateral geniculate nucleus of the monkey. Z. Zellforsch. Mikrosk. Anat. 103:90.

Hamlyn, L. H. 1962. The fine structure of the mossy fibre endings in the hippocampus of the rabbit. J. Anat. 96:112.

Hendrickson, A. 1969. Electron microscopic radiography: Identification of origin of synaptic terminals in normal nervous tissue. Science (Wash. D. C.) 165:194.

Jones, D. G., and R. F. Breakley. 1972. Further studies on synaptic junctions. II. A comparison of synaptic ultrastructure in fractionated and intact cerebral cortex. Z. Zellforsch. Mikrosk. Anat. 125:432.

Kellenberger, E., A. Ryter, and J. Sechaud. 1958. Electron microscope study of DNA-containing plasma. II. Vegetative and mature phage DNA as compared with normal bacterial nucleoids in different physiological states. J. Biophys. Biochem. Cytol. 4:671.

Kornguth, S. E., J. W. Anderson, and G. Scott. 1969. Isolation of synaptic complexes in a caesium chloride gradient: electron microscopic and immunohistochemical studies. J. Neurochem. 16:1017.

Larramendi, L. M. H., and T. Victor. 1967. Synapses on the Purkinje cell spines in the mouse: an electron microscopic study. Brain Res. 5:15.

Lowry, O. H., N. J. Rosebrugh, A. L. Farr, and R. J. Randall. 1951. Protein measurement with the Folin phenol reagent. J. Biol. Chem. 193:265.

Morgan, I. G., L. A. Wolfe, P. Mandel, and G. Gomos. 1971. Isolation of plasma membranes from rat brain. Biochim. Biophys. Acta. 241:237.

Nafstad, P. A. J. 1967. An electron microscopic study on the termination of the perforant path fibers in the hippocampus and fascia dentata. Z. Zellforsch. Mikrosk. Anat. 75:532.

Palay, S. L. 1956. Synapses in the central nervous system. J. Biophys. Biochem. Cytol. 2(Suppl.):193.

Palay, S. L. 1958. The morphology of synapses in the central nervous system. Exp. Cell Res. 5(Suppl.):275.

Peters, A., S. L. Palay, and H. Webster. 1970. The Fine Structure of the Nervous System. Harper & Row, Publishers, New York. 138.

Penninger, K. H. 1971. The cytochemistry of synaptic densities. I. An analysis of the bismuth iodide impregnation method. J. Ultrastruct. Res. 34:103.

Price, J. L. 1968. The synaptic vesicles of the reciprocal synapse of the olfactory bulb. Brain Res. 11:797.

Raisman, G. 1969. Neuronal plasticity in the septal nuclei of the adult rat. Brain Res. 14:25.

Raisman, G., and P. M. Field. 1973. A quantitative investigation of the development of collateral reinnervation after partial deafferentation of the septal nuclei. Brain Res. 90:241.

Rall, W., G. M. Shepherd, T. S. Reese, and M. W. Brightman. 1966. Dendodendritic synaptic pathway for inhibition in the olfactory bulb. Exp. Neurol. 14:44.

Rodriguez, G. De Lores Arnaiz, M. Aberici, and E. de Robertis. 1967. Ultrastructural and enzymatic
studies of cholinergic and non-cholinergic synaptic membranes isolated from brain cortex. *J. Neurochem.* 14:215.

**Van der Loos, H.** 1964. Similarities and dissimilarities in submicroscopal morphology of interneuronal contact sites of presumably different functional character. *Prog. Brain Res.* 6:43.

**Venable, J. H., and R. Coggeshall.** 1965. A simplified lead stain for use in electron microscopy. *J. Cell Biol.* 25:407.

**Walberg, F.** 1968. Morphological correlates of postsynaptic inhibitory processes. In *Structure and Function of Inhibitory Neuronal Mechanisms*. C. van Euler, S. Skoglund, and U. Soderberg, editors. Pergamon Press Ltd., Oxford. 7–14.

**Whittaker V. P., I. A. Michaelson, and R. J. A. Kirkland.** 1964. The separation of synaptic vesicles from nerve-ending particles ("synaptosomes"). *Biochem. J.* 90:293.