Anti-flavin antibodies

Michael J. BARBER,*† Duane C. EICHLER,* Larry P. SOLOMONSON* and Brian A. ACKRELL†
*Department of Biochemistry, University of South Florida College of Medicine, Tampa, FL 33612, U.S.A., and †Department of Biochemistry and Biophysics, University of California, San Francisco, CA 94143, U.S.A.

Antibodies were elicited to FAD by using the hapten 6-6-(6-aminoethyl)-FAD conjugated to the immunogenic carrier protein bovine serum albumin. Cross-reactivity was determined by Ouchterlony double-diffusion analysis with 6-6-(6-aminoethyl)-FAD coupled to rabbit serum albumin. Anti-FAD IgG was partially purified by (NH₄)₂SO₄ precipitation followed by DEAE-cellulose/CM-cellulose and bovine serum albumin–agarose chromatography. The partially purified anti-FAD IgG fraction failed to inhibit the catalytic activities of the flavin-containing enzymes nitrate reductase, xanthine oxidase and succinate dehydrogenase, whereas enzyme activity could be inhibited by addition of antibodies elicited against the native proteins. However, the partially purified anti-FAD IgG fraction could be used as a highly sensitive and specific probe to detect proteins containing only covalently bound flavin, such as succinate dehydrogenase, p-cresol methylhydroxylase and monoamine oxidase, by immuno-blotting techniques. Detection limits were estimated to be of the order of femtomolar concentrations of FAD with increased sensitivity for the 8α-N(3)-histidyl linkage compared with 8α-O-tyrosyl substitution.

INTRODUCTION

Although antibodies are frequently used for the detection and characterization of macromolecules (Van Vunakis & Langone, 1980), techniques have been developed to enable antibodies to be elicited against low-M₆₆₆₆ haptons covalently linked to suitable immunogens such as BSA and haemocyanin (Kovac et al., 1985; Humphries & McConnell, 1976). The availability of antibodies elicited against a specific prosthetic group of an enzyme could facilitate their use both for the detection and localization of the enzyme, as well as other proteins bearing the same prosthetic group, in addition to possible use of the antibodies as site-specific inhibitors. (In complex redox enzymes containing a variety of different, independent, prosthetic groups, antibodies could prove to be inhibitors of extremely high specificity and affinity, potentially useful for delineating electron-transfer sequences.) In addition, antibodies directed against specific prosthetic groups could be utilized to provide topographical information concerning the structure of the enzyme, such as the subunit or polypeptide chain associated with the chromophore, as well as accessibility and information concerning prosthetic-group synthesis, modification and insertion.

The present paper describes the preparation and characterization of antibodies elicited against the FAD molecule, their use in identifying the location of covalently bound flavin chromophores within enzyme subunits and their application as a highly sensitive detection method for complex heterogeneous samples.

MATERIALS AND METHODS

Materials

BSA (fraction V, essentially fatty acid-free), RSA (fraction V), complete and incomplete Freund’s adjuvant, phenazine methosulphate and 2,6-dichlorophenol-indophenol were purchased from Sigma Chemical Co. HAWP nitrocellulose filters (2.4 cm; 0.45 μm pore size) were purchased from Millipore. DEAE-cellulose DE-52 and CM-cellulose CM-52 were obtained from Whatman. The immuno-blot assay kit containing goat anti-(rabbit IgG) antibody–horseradish peroxidase conjugate and nitrocellulose membrane (0.45 μm pore size) were purchased from Bio-Rad Laboratories. 6-AH-FAD was provided by, and is proprietary to, Miles Laboratories, and was synthesized by the procedure described by Morris & Buckler (1983).

Methods

Coupling of 6-AH-FAD to BSA. 6-AH-FAD was conjugated to BSA with the use of the coupling agent dimethyl adipimidate as described by Morris & Buckler (1983). The number of molecules of 6-AH-FAD conjugated per molecule of BSA was estimated in the following manner. The protein concentration of the conjugate sample was determined by the method of Lowry et al. (1951) and the concentration of 6-AH-FAD was determined from the visible-region spectrum of the conjugate sample by using an absorption coefficient of 11.3 mm⁻¹·cm⁻¹ at 450 nm (Whitby, 1953). The ratio of the molar concentrations of 6-AH-FAD and protein gave the number of conjugated flavin residues per albumin molecule.

6-AH-FAD was conjugated to RSA by a similar procedure and purified by column chromatography on Sephadex G-25 equilibrated with 0.1 m-sodium phosphate buffer, pH 7.0, containing 0.1 m-NaCl, concentrated and characterized as for the BSA–6AH-FAD complex.

Immunization and antiserum preparation. Procedures used in the production of antisera to the albumin–6-AH-FAD conjugate were as described previously by Eichler & Glitz (1974). A 1 mg portion of conjugate
dissolved in 0.4 ml of 0.05 M-Tris/HCl buffer, pH 7.5, containing 0.1 M-NaCl and 0.1 mM-EDTA was mixed with 1.6 ml of complete Freund's adjuvant to form a uniform homogenate. Approx. 0.1 ml of this homogenate was injected into each toe-pad of a 3-month-old New Zealand White rabbit. After 2 weeks the rabbit was injected subcutaneously in the hindquarters with 0.5 mg of conjugate in 0.1 ml of buffer and mixed with 0.4 ml of incomplete Freund's adjuvant. At 4 weeks from the time of the initial injection, blood was collected from the ear vein, and the serum was tested by double diffusion on agar plates (Ouchterlony, 1968). Subsequently, at intervals of 7-8 days, the rabbit was bled from the ear vein, the blood was allowed to clot, and the serum was removed and clarified by centrifugation at 2000 g. The clarified serum was either processed for preparation of an IgG protein fraction or stored frozen at −70 °C. Similar procedures were used to elicit antibodies to purified nitrate reductase and xanthine oxidase.

**Purification of γ-globulin.** Preparation of IgG fraction from whole serum was carried out as described by Palacios et al. (1972). Antiserum (10 ml) was diluted with an equal volume of 0.05 M-Tris/HCl buffer, pH 7.5, containing 0.1 M-NaCl and 0.1 mM-EDTA. At 0 °C, 4.52 g of solid (NH₄)₂SO₄ was added slowly until 40% saturation was achieved. After 30 min the precipitate was collected by centrifugation at 20000 g for 15 min. The pellet was dissolved in 4 ml of 0.01 M-sodium phosphate buffer, pH 7.2, containing 0.015 M-NaCl and dialysed overnight against this same buffer. The dialysed sample was clarified by centrifugation, and applied to a DEAE-cellulose/CM-cellulose column (1.5 cm x 10 cm) that contained 5 cm of DEAE-cellulose overlayed with 5 cm of CM-cellulose, both equilibrated against 0.01 M-sodium phosphate buffer, pH 7.2, containing 0.015 M-NaCl. The flow-through fractions were pooled and concentrated by vacuum dialysis against 0.05 M-Tris/HCl buffer, pH 7.5, containing 0.1 M-NaCl and 0.1 mM-EDTA. The IgG fraction was further purified by passage through a BSA–agarose affinity column prepared by coupling BSA to CNBr-activated agarose in 0.05 M-sodium phosphate buffer, pH 7.2, to remove anti-BSA IgG.

**Immunodiffusion.** Double diffusion (Ouchterlony, 1968) was run in 1% agar made up in 0.05 M-Tris/HCl buffer, pH 7.5, containing 0.1 M-NaCl and 0.1 mM-EDTA. Standard antigen solutions were 1 mg/ml unless otherwise specified. Serum was placed in the centre well, and antigen in the outer wells. Development was allowed to take place overnight at room temperature. The immunodiffusion plates were then washed overnight in 0.9% NaCl, rinsed with distilled water, and dried in a hood after being covered with a moist strip of filter paper. Once the plates were completely dry, the filter-paper strip was removed and the plates were stained with 0.6% Amido Black suspended in destaining solution (methanol/water/acetic acid, 5:5:1, by vol.) followed by washing with distilled water, and subsequent destaining for 15 min.

**Enzyme preparation.** Succinate dehydrogenase was purified from bovine heart mitochondria as described by Baginsky & Hatefi (1969). NADH-linked nitrate reductase was purified to electrophoretic homogeneity from *Chlorella vulgaris* as previously described (Howard & Solomonson, 1982). Xanthine oxidase was isolated from bovine milk as described by Barber & Siegel (1982).

**Enzyme assays.** Succinate dehydrogenase activity was determined as described by Hatefi (1978) with succinate, 2,6-dichlorophenol-indophenol and phenazine methosulphate. Nitrate reductase activity was determined as the NADH-dependent reduction of NO₃⁻ as previously described (Solomonson et al., 1975). Xanthine oxidase activity was monitored at 295 nm by using the aerobic formation of uric acid from xanthine (AVIS et al., 1955). For the inhibition assays, purified enzyme was incubated with the desired volume of purified IgG fraction in 0.05 M-sodium phosphate buffer, pH 7.8, containing 0.1 M-NaCl and 0.1 mM-EDTA for 2 min at 25 °C and the assay was initiated by the addition of either substrate or acceptor. Control enzyme samples in the absence of the IgG fraction were subjected to identical treatments.

**SDS/polyacrylamide-gel electrophoresis.** Proteins were separated by SDS/polyacrylamide-slab-gel electrophoresis on linear gradients of 7.5–15% polyacrylamide with the buffer system described by Laemmlli (1970). The Mᵣ standards (BRL Laboratories) used were myosin (Mᵣ 205000), phosphorylase b (Mᵣ 97500), BSA (Mᵣ 68000), ovalbumin (Mᵣ 43000) and α-chymotrypsin (Mᵣ 25700). Proteins were stained with Coomassie Brilliant Blue.

**Immunoblotting.** Immuno-blotting of proteins transferred from SDS/polyacrylamide gels to nitrocellulose membranes (Towbin et al., 1979; Burnette, 1981) or after direct application of proteins to nitrocellulose membranes in a slot-blot apparatus (Schleicher & Schuell) was performed according to the procedures accompanying the Bio-Rad immuno-blot (goat anti-(rabbit IgG) antibody–horseradish peroxidase conjugate) assay kit. In the case of proteins separated by SDS/polyacrylamide-gel electrophoresis, proteins were electrophoretically transferred from the polyacrylamide slab gel to a nitrocellulose membrane with 25 mM-Tris/192 mM-glycine, pH 8.3, in 20% (v/v) methanol as the transfer buffer and a voltage of 60 V for 16 h at 4 °C. In the case of direct application of proteins to nitrocellulose, 200 µl of a protein solution (containing 0.1 ng to 10 µg of protein in 0.5 M-Tris/HCl buffer, pH 7.5, containing 0.1 M-NaCl and 0.1 mM-EDTA) was applied to each well of the slot-blot apparatus without vacuum. A vacuum was then applied and each ‘slot’ was washed with approx. 2 ml of Tris-buffered saline (0.02 M-Tris/HCl buffer, pH 7.5, containing 0.5 M-NaCl). Excess protein-binding sites on the nitrocellulose membranes were blocked by incubation for 1 h at room temperature with Tris-buffered saline containing 3% (w/v) gelatin. The blocked membranes were incubated with the first antibody or pre-immune IgG in Tris-buffered saline containing 1% gelatin for 2 h. The membranes were rinsed with water and then incubated with two changes of Tris-buffered saline containing 0.05% Tween-20 followed by one change of Tris-buffered saline for 20 min each. The membranes were then incubated with the second antibody, a 1:2000 dilution of horseradish-peroxidase-conjugated goat anti-(rabbit IgG) antibody (Bio-Rad Laboratories) in Tris-buffered saline containing 1% gelatin for 1 h. The membranes were rinsed after incubation of the second antibody and...
Anti-flavin antibodies

Fig. 1. Immunodiffusion of anti-FAD serum and IgG fraction

The centre wells contained anti-FAD serum collected at 6 weeks from the initial toe-pad injection (a and b) and BSA-agarose-affinity-purified anti-FAD IgG (c). The peripheral wells contained the indicated concentrations (mg/ml) of antigen: BSA, bovine serum albumin; BSA–FAD, 6-AH-FAD conjugated to BSA; RSA, rabbit serum albumin; RSA–FAD, 6-AH-FAD conjugated to RSA. In each case BSA or RSA is present in only one well; dilutions of BSA–FAD or RSA–FAD were placed in the peripheral wells.

subsequently incubated in horseradish peroxidase colour development medium until blue bands appeared. This development solution was prepared by mixing 30 mg of horseradish peroxidase colour development reagent (4-chloro-1-naphthol) (Bio–Rad Laboratories), dissolved in 10 ml of methanol, with 50 ml of Tris-buffered saline containing 40 μl of 30% (v/v) H₂O₂ immediately before use. Colour development was terminated by rinsing the membranes with distilled water. The stained membranes were incubated with 50% (v/v) glycerol for approx. 10 min, blotted dry and stored under clear plastic film.

RESULTS

Preparation and characterization of antigen

The coupling of a specific ligand to a suitable immunogenic carrier protein, such as BSA, has become an established method of eliciting antibodies to a desired ligand. Immunization of host animals with BSA–FAD conjugates with an average stoichiometry of one FAD molecule bound per BSA molecule and examination of the subsequent sera failed to reveal the production of IgG molecules directed against flavin as determined by Ouchterlony immunodiffusion. In contrast, use of the 6-AH-FAD derivative, which interposes a long hydrocarbon chain between the functional, reactive, amino group and the adenine moiety, facilitated the preparation of a BSA–flavin derivative that contained an average of four molecules of FAD per molecule of BSA. A similar stoichiometry was observed on spectroscopic analysis of the RSA–6-AH-FAD conjugate.

Characterization of the antibodies

Sera were monitored for antibody content by Ouchterlony double-diffusion analysis and exhibited cross-reactivity to 6-AH-FAD coupled to RSA as shown in Fig. 1. Sera collected from the sixth and seventh week after the primary injection were chosen for characterization. In the initial serum IgG preparation, a significant degree of cross-reactivity was detected against native BSA, indicating a relatively low titre of anti-FAD IgG and suggesting that the major antibody determinants were due to the immunogenic carrier protein. To decrease the amount of these non-specific antibodies, the IgG fraction was chromatographed on a BSA–agarose column to remove the high-affinity anti-BSA antibodies. This purified IgG fraction was used for all subsequent experiments.

The effects of anti-FAD IgG on the catalytic activities of a number of flavin-containing enzymes are shown in Fig. 2. Incubation of purified Chlorella vulgaris NADH-linked nitrate reductase, bovine milk xanthine oxidase and bovine mitochondrial succinate dehydrogenase with increasing concentrations of purified anti-FAD IgG resulted in no detectable decrease in the activity of any of the enzymes compared with control samples incubated...
either in the absence of or with the inclusion of pre-immune IgG. In contrast, to demonstrate that antibody-antigen interaction would be expected to result in a decrease in enzyme activity, both nitrate reductase and xanthine oxidase samples were incubated with purified anti-(nitrate reductase) IgG and anti-(xanthine oxidase) IgG respectively. In both cases there was a significant decrease in the catalytic activity.

**Detection of flavoproteins**

The purified anti-FAD IgG fraction was examined for its ability to cross-react with proteins containing either non-covalently (nitrate reductase) or covalently (succinate dehydrogenase) bound flavin. Purified bovine mitochondrial succinate dehydrogenase was applied to nitrocellulose over a 100-fold range of protein concentration. The nitrocellulose membrane was incubated with purified rabbit anti-FAD IgG, washed and subsequently incubated with horseradish-peroxidase-conjugated goat anti-(rabbit IgG) antibody. Detection was accomplished by using the peroxidase reaction. As shown in Fig. 3, the antibodies to 6-AH-FAD and the immuno-blot assay system showed this combination could be used to detect as little as 0.1 pmol of succinate dehydrogenase, corresponding to 0.1 pmol of FAD conjugated to protein. In contrast, the immuno-blot technique failed to detect any cross-reactivity between anti-FAD IgG and purified nitrate reductase (Fig. 4). However, limited experiments were performed to examine the specificity of the anti-flavin antibodies with respect to their ability to detect proteins containing FAD covalently bound through a linkage different from the 8z-N(3)-histidyl linkage present in succinate dehydrogenase (Walker et al., 1972). The results of immuno-blotting experiments comparing the detection limits of succinate dehydrogenase, p-cresol methylhydroxylase and monoamine oxidase are shown in Fig. 4. Different detection limits were observed among the three proteins, corresponding to

![Fig. 3. Immuno-detection of mitochondrial succinate dehydrogenase](image)

![Fig. 4. Immuno-detection of enzymes containing covalently bound FAD](image)
Anti-flavin antibodies

Fig. 5. SDS/polyacrylamide-gel electrophoresis of succinate dehydrogenase and immuno-detection of flavin-containing subunits

Purified succinate dehydrogenase (lanes 3–8) and nitrate reductase (lanes 2 and 9) were subjected to SDS/polyacrylamide-gel electrophoresis, transferred to nitrocellulose and either stained for total protein (lanes 2–5) or incubated with anti-FAD IgG (lanes 6–9) and examined for binding by anti-FAD IgG by using horseradish-peroxidase-conjugated goat anti-(rabbit IgG) antibody. Lanes 2 and 9 correspond to the application of 1 µg of total protein, lanes 3 and 8 to 5 µg of total protein, lanes 4 and 7 to 2.5 µg of total protein and lanes 5 and 6 to 1.25 µg of total protein. Lanes 1 and 10 correspond to pre-stained protein standards of the indicated Mr values.

Fig. 6. SDS/polyacrylamide-gel electrophoresis of heart mitochondria and the immuno-detection of flavin-containing proteins

Isolated bovine heart mitochondria were subjected to SDS/polyacrylamide-gel electrophoresis, transferred to nitrocellulose and either stained for total protein (lanes 1–4) or incubated with anti-FAD IgG (lanes 7–10) and examined for binding by anti-FAD IgG by using horseradish-peroxidase-conjugated goat anti-(rabbit IgG) antibody. Lanes 1 and 10 correspond to the application of 2 µg of purified succinate dehydrogenase, and lanes 2 and 9, lanes 3 and 8 and lanes 4 and 7 correspond to the application of 5 µg, 10 µg and 20 µg of total mitochondrial protein respectively. Lanes 5 and 6 correspond to pre-stained protein standards of the indicated Mr values.

differentially stained bands, even though the proteins were applied at equivalent flavin concentrations. Monoamine oxidase, which contains FAD bound via an 8α-S-cysteiny1 linkage (Kearney et al., 1971), was detected at concentrations directly comparable with those of succinate dehydrogenase (8α-N-histidyl-FAD), whereas p-cresol methylhydroxylase, which contains an 8α-O-tyrosyl linkage (McIntire et al., 1980), was detected only at higher concentrations. Note that the apparent decrease in binding of the anti-FAD IgG to monoamine
oxidase at high concentrations is due to inhibition of protein binding to the nitrocellulose by glycerol used to stabilize the protein during storage. This effect is lost on subsequent dilution of the samples.

The antibodies to the flavin prosthetic group were also used to detect proteins containing covalently bound FAD after fractionation by SDS/polyacrylamide-gel electrophoresis. Succinate dehydrogenase was heat-denatured (3 min at 100 °C) in the presence of SDS (5%, w/v) and 2-mercaptoethanol (0.5 M) in 0.05 M-Tris/HCl buffer, pH 6.8, before being loaded on to the polyacrylamide gel. After electrophoresis and transfer to nitrocellulose membrane, the labelled protein was detected as described for the immuno-blots. As shown in Fig. 5, antibodies directed against 6-AH-FAD could be used to detect as little as 1.25 μg of succinate dehydrogenase applied to the gel, the antibodies detecting only the large subunit, Mr 70,000, known to contain the FAD prosthetic group (Davis & Hatefi, 1971), whereas the protein stain showed the presence of both large and small (Mr 30,000) units. The apparent difference in sensitivity between the immuno-blots and SDS/polyacrylamide-gel electrophoresis patterns may reflect the efficiency of protein transfer from SDS/polyacrylamide gels to the nitrocellulose membrane. The anti-FAD antibodies could also be used to detect protein bands containing covalently bound flavin in crude enzyme preparations. Heart mitochondria were isolated, subjected to SDS/polyacrylamide gel electrophoresis and transferred to nitrocellulose. The nitrocellulose was then either stained for total protein or incubated with anti-FAD antibody followed by goat anti-(rabbit IgG)–horseradish peroxidase conjugate and horseradish peroxidase staining (Fig. 6). Although multiple bands were revealed by protein staining, only a single band, corresponding to Mr 70,000, was detected with the antibody.

**DISCUSSION**

The preceding results demonstrate that it is possible to elicit antibodies to a protein prosthetic group provided that a suitable method exists for coupling the purified prosthetic group to an immunogenic carrier protein and that sufficiently high degrees of protein substitution can be obtained. In the current experiments we have shown these antibodies to be unreactive with flavin-containing enzymes in solution, presumably as a result of the fact that the flavin prosthetic group is buried within the protein and inaccessible to the antibody-binding site. However, the anti-FAD antibodies can be successfully used to detect denatured proteins containing covalently bound flavin after immobilization on nitrocellulose, and these antibodies exhibit some degree of specificity with respect to the nature of the flavin–protein linkage. These experiments have demonstrated that immuno-blotting techniques can be used to detect a covalently bound protein prosthetic group, such as FAD, at the picomole level. This provides a highly specific and extremely sensitive procedure for the detection of proteins, or protein subunits, containing a covalently bound prosthetic group at a level of sensitivity that far exceeds that which can be detected by visible-region or fluorescence spectroscopy.

The limited experiments performed with flavoproteins containing FAD bound via linkages differing from the 8z-N(3)-histidyl linkage found in succinate dehydrogenase suggested that the antibodies, though cross-reacting with the flavin chromophore present in p-cresol methylhydroxylase and monoamine oxidase, exhibited some degree of specificity with respect to the type of flavin–protein linkage. The FAD in monoamine oxidase, which is bound via an 8z-S-cysteinyl linkage (Kearney et al., 1971), was recognized with a sensitivity comparable with that for succinate dehydrogenase, but, in contrast, the FAD in p-cresol methylhydroxylase, which is present as an 8z-O-tyrosyl linkage (McIntire et al., 1980), was recognized with a significantly decreased sensitivity. These results may suggest that the level of cross-reactivity with the antibody could be influenced by the mode of co-ordination of the flavin moiety to the protein. These conclusions require the assumption that different proteins are sufficiently denatured upon binding to nitrocellulose and expose the FAD prosthetic group to equivalent extents. Any variation in the degree of unfolding could also result in apparent changes in antibody specificity.

We were also able to demonstrate that the antibodies to the 6-AH-FAD could be used to identify specifically proteins, or protein subunits, containing covalently bound flavin within complex protein mixtures after separation by SDS/polyacrylamide-gel electrophoresis. This approach should be particularly advantageous for identifying and analysing proteins containing covalently bound FAD or other flavin derivatives in complex protein systems.

We are indebted to Dr James Albarella and Dr. David Morris of the Ames Division, Miles Laboratories, for generously donating 6-AH-FAD and 6-AH-FAD–BSA conjugate and their interest in this research. We are further indebted to Dr. William McIntire for generously donating p-cresol methylhydroxylase and to Dr. Walter Weyer for the sample of monoamine oxidase. This work was supported by Grant 84-CRCR-1-1404 from the U.S. Department of Agriculture and Grants GM 32696 and ML 16251 from the National Institutes of Health.

**REFERENCES**

Anti-flavin antibodies


Received 28 May 1986/19 August 1986; accepted 13 October 1986