IMMUNOLOGICAL COMPARISON OF THE NADH:NITRATE REDUCTASE FROM DIFFERENT CUCUMBER TISSUES

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ABSTRACT

Soluble nitrate reductase from cucumber roots (Cucumis sativus L.) was isolated and purified with blue-Sepharose 4B. Specific antibodies against the NR protein were raised by immunization of a goat. Using polyclonal antibodies anti-NR properties of the nitrate reductase from various cucumber tissues were examined. Experiments showed difference in immunological properties of nitrate reductase (NR) from cotyledon roots and leaves.

KEY WORDS: nitrate reductase, antibody, dot-blot, immunological assays, Cucumis sativus.

INTRODUCTION

Most of the photosynthetic organisms require nitrate as their main inorganic source of nitrogen. Nitrate absorbed by the plant roots can be reduced in both, root and leaf tissues (Jolly and Tolbert 1978, Redinbourough and Campbell 1981). The first rate-controlling step of NO3 assimilation in higher plants is catalysed by nitrate reductase (NR, EC.1.6.6.1), which properties have been subject of a number of reviews (Beever and Hageman 1980, Campbell and Kinghorn 1990, Caboce and Rouze 1990). This multicenter enzyme reduces nitrate to nitrite using NADH as its natural electron donor.

Over the past 40 years, since the discovery of NR by Evans and Nason (1953), considerable attention has been focused on the method for isolation and purification of this enzyme. Since NR is a highly efficient catalyst, tightly regulated in plant tissues, it is not present in large quantities (Beever and Hageman 1983, Campbell and Sinarelli 1986). Moreover, NR is highly labile protein and consequently, difficult to purify. Very good results in purification of higher plant NADH-NR was achieved when affinity chromatography with activated Sepharose was used (Sherrard and Dalling 1979, Nakagawa et al. 1985, Nakamura and Ikawa 1993). This method enabled to obtain the pure enzyme through a specific elution with NADH. In protein characterization studies, the immunological techniques are important tools. Since nitrate reductase has been found as a good antigen, specific antibodies against the protein of NR from various higher plants were raised (Snapp et al. 1984, Hyde et al. 1989). Immunoassays were used to compare the nitrate reductase from different tissues (Nakagawa et al. 1986) and species (Notton et al. 1983, Cherel et al. 1986).

In the present work we purified the soluble nitrate reductase from cucumber roots and prepared antibodies against it to investigate the molecular properties of the enzyme from different tissues of cucumber seedlings.

MATERIAL AND METHODS

Plant material.

Cucumber seeds (Cucumis sativus L. var. Wisconsin) after germination were cultivated hydroponically for 5 or 10 days in N-free solution, as described previously (Buczek and Marciniak 1990). Nitrate reductase was induced by transferring the seedlings into one-third strength Hoagland solution.
Enzyme extraction.

Crude extract was prepared by grinding of roots in a chilled mortar with the extraction buffer containing 50 mM K-phosphate (pH 7.5), 1 mM EDTA, 1 mM DTT, 1 mM PMSE 10% glycerol and 0.5% PVP. The extract was filtered through four layers of cheesecloth and centrifuged at 20,000 g for 20 min. The supernatant was used as enzyme source.

Enzyme purification.

1. Precipitation with ammonium sulfate.

Solid ammonium sulfate was slowly added to the supernatant with continuous stirring. The fraction precipitated between 20-50% saturation was collected by centrifugation at 20,000 g for 10 min and dissolved in 10 mM K-phosphate (pH 7.5) containing 1 mM EDTA, 1 mM DTT and 1 mM PMSE.

2. Affinity chromatography.

a.) Cyanogen bromide method of Sepharose 4B activation and coupling of the ligand (according to Sherrard and Dalling 1979, with some modifications).

The swollen Sepharose 4B (100 cm³) washed with 2,000 cm³ of distilled water by vacuum filtration was suspended in 200 cm³ of 2 M Na₂CO₃. The slurry was slowly stirred and 20 cm³ of CNBr dissolved in acetonitrile (g per cm³) was added. During activation pH of the slurry was near 11. After 3 minutes the activated gel was washed with 1,000 cm³ of 0.1 M carbonate buffer (pH 9.5) and immediately transferred to a ligand solution. Dextran blue 2,000 (2 g) dissolved in 200 cm³ of 0.4 M carbonate buffer (pH 10, a coupling buffer) was combined with CNBr-Sepharose 4B for 18 h in 4°C. Non-coated dye was removed with coupling buffer and the dye slurry was suspended in 0.1 M Tris-HCl buffer (pH 8.0) to block an excess of active groups. After 1 hour the gel was transferred onto coarse disk sintered glass funnel and rinsed out as follows with 500 cm³ of 0.1 M citric acid-NaOH and 0.5 M NaCl (pH 4.0); distilled water; 0.1 M citric acid-sodium citrate and 0.5 M NaCl (pH 8.0); distilled water. Washing procedure was repeated at least three times. Finally, the slurry was suspended in 0.05 M K-phosphate buffer (pH 7.5).

b.) Affinity chromatography on blue-dextran CNBr activated Sepharose 4B (chromatography was made by modification Redinbugh’s and Campbell’s method 1981).

The ammonium precipitated fraction was loaded immediately onto Sepharose 4B column (6 x 2 cm) equilibrated with 0.05 M K-phosphate (pH 7.5), 1 mM EDTA, 1 mM DTT and 20 μM FAD (buffer A). In some experiments the crude extract (125 cm³) was slowly stirred with 10 cm³ of blue Sepharose in buffer A. After 30 min, the blue Sepharose was collected by vacuum filtration and washed with buffer A without FAD, to the moment when A₂₈₀ was 0. Then Sepharose was suspended in about 30 cm³ of buffer A and packed onto 2 cm diameter column. Nitrate reductase was eluted from the gel with 70 cm³ of buffer A containing 100 μM of the activity of NR was determined. Fractions with the highest activity of nitrate reductase were combined, lyophilized and stored at -20°C.

3. Nitrate reductase assays.

Total NADH:NR (EC 1.6.6.1) activity was examined in crude extract and in fractions collected from Sepharose. The incubation medium (1 cm³) contained 25 mM K-phosphate buffer (pH 7.5), 10 mM KNO₃, 0.25 mM NADH and enzyme. After incubation (30 min at 27°C) zinc-acetate was added to stop the reaction and mixture was centrifuged. In supernatant nitrite was determined colorimetrically at 540 nm (Sanderson and Cocking 1964).

4. Protein determination.

Protein was determined according to the method of Bradford et al. (1972) using BSA as a standard.

5. Electrophoresis.

Poliacylamide gel electrophoresis was carried out according to Laemmli (1970). Concentration of the polyacrylamide native gel was 7%. The gel was stained with Coomassie Blue R-250 to protein detection. Visualization of NR activity in the gel was provided according to Heath-Pagliuso et al. (1984).

6. Preparation of antisera.

The pure, lyophilized protein of NR was used to raise a crude antiserum. Equal volumes of the enzyme (0.5 cm³ containing 100 μg of protein) and Freund’s adjuvant were emulsified and then injected (0.2 cm³ intradermally) into the goat every two weeks. The level of goat IgG anti-NR was controlled at regular intervals by standard method of ELISA. After 12 weeks the crude serum was collected by blood centrifugation.

7. Dot-blot assays.

Immunoblotting was carried out at room temperature. The samples (5 μl) were loaded onto a nitrocellulose sheet, air-dried and incubated 1 h with blocking solution containing 0.05 M K-phosphate buffer (pH 7.5) with 0.9% NaCl (PBS), 1% BSA, 1% casein, 2% PVP and 0.05% Tween 20. The nitrocellulose sheet was transferred to PBS containing goat anti-NR serum (1:10). After 1 h the blot was washed three times with PBS and incubated for 1 hr with horse-radish-peroxidase conjugated goat anti-goat IgG diluted 1:1000 in PBS. Anti-NR cross-reactive dots were identified by the dye with the mixture of 4 chloro-1-naphthol in 2 cm³ of methanol and 6 μl of 30% H₂O₂ in 8 cm³ of PBS.

8. Antiserum titration.

Preamine control serum (10, 50 and 100 μl) or goat anti-NR serum (10, 40 and 80 μl) was added to 1 cm³ of crude enzymatic extract obtained from cucumber roots. Concentration of protein was 3 mg per cm³ or 2 mg per cm³ in serum and enzymatic extract, respectively. The samples were incubated for 1.5 h at 4°C, centrifuged at 13,000 x g 10 min and supernatant was used for determination of NR activity. All values selected for the figures and tables are representative for the results obtained in several (three to four) independent experiments.

RESULTS AND DISCUSSION

The pure, active nitrate reductase was isolated from cucumber roots using steps of purification. The first step was ammonium sulfate fractionation followed with an affinity chromatography with blue Sepharose (Table 1A and Table 1B). Finally, the nitrate reductase was purified 155 fold with only 4% recovery and specific activity of 0.9 units per mg protein (Table 1A).
Table 1B. For this reason in next experiments the crude extract from roots was mixed with blue Sepharose and then column was packed.

The elution profile of NR from the blue-Sepharose column is shown in Figure 1. The highest activity of the nitrate reductase was found in 4 fractions (fraction 10 to 14). Additionally, a large amount of protein was removed from column into the same fractions. It is worth of notice, that NR was eluted from the column always with a low concentration of NADH (7 μM). Because of low protein recovery the fractions exhibiting the nitrate reductase activity were pooled and concentrated by lyophilization.

After subjection of the purified enzyme preparation to polyacllamide disc gel electrophoresis under non-denaturing conditions, two protein bands appeared (Figure 2). Both bands were coincident with a specific staining for NR activity. It is suggested that during separation of the enzyme on blue Sepharose the most impurity have been lost.

The lyofilized samples of the pure enzyme were dissolved and used to immunization of the goat. The immunological answer was controlled by ELISA and results were summarized in Figure 3.

The amount of the polyclonal antibodies anti-NR in goat serum was highest after the second injection and after the fourth one was slightly reduced. However, the whole amount of antibodies in comparison with the basic level ("0") was satisfactory. Antiserum efficiency (probably the mixture of IgG and slight amount of IgM) was controlled by the dot-blot method on nitrocelullose paper (Figure 4) and by antiserum titration (Figure 5). After an immunochemical cross-reaction of anti-NR IgG with the enzyme and goat IgG conjugated with peroxidase, we observed the appearance of dye dots on nitrocelullose paper. In titration experiments the activity of nitrate reductase from cucumber roots decreased after 1.5 h pre-incubation with polyclonal antibodies (anti-NR). Preimmune control serum had no effect on the enzyme activity (Figure 5).

Results of dot-blot reaction and titration experiments confirmed the good affinity between the nitrate reductase and obtained antibodies. Polyclonal antibodies reacted not only with NR in crude extract but also with the pure enzyme.

Dot-blot technique was used to examine the immunological properties of NR from various cucumber tissues (Figure 6). We observed the positive reaction between goat NR-antibodies and crude extracts from roots, cotyledons and leaves of

Fig. 1. Separation of the soluble NR by chromatography on blue Sepharose 4B.
- NR activity (μmol NO₃⁻)
- protein (μg)
- NADH (μmol)
The crude enzymatic extract was loaded after centrifugation onto blue-Sepharose column. NR was removed with elution buffer containing 100 μM NADH (3 cm³ fractions). The level of NADH was controlled spectrophotometrically at 340 nm. Protein was determined according to Bradford (1976).

Fig. 2. Electrophoresis of the pure nitrate reductase after blue-Sepharose separation.
The fractions with high activity of NR eluted from blue-Sepharose were analyzed by PAGE (Laemmli 1970). After staining with Coomassie Blue R-250 two bands of the proteins became visible (b). After developing NR activity according to Heath-Pagliuso et al. (1984) with 25% tetrazolium red, two bands of NR were visible (a).
Fig. 3. Demonstration of the level polyclonal antibodies anti-NR in goat serum by ELISA.

- ■ control serum (no immunized),
- □ 1st injection,
- ● 2nd injection,
- ○ 3rd injection,
- ▲ 4th injection.

Immunization of the goat carried out as described in "Material and Methods". It were five injection each containing 100μg of protein. In regular intervals blood samples were collected and the level of anti-NR estimated by the standard method of ELISA.

Fig. 4. Test of cross-reactivity of the goat anti-NR.

Nitrate reductase was obtained from cucumber roots and purified as described in "Material and Methods". 5 μl samples of lyophilized enzyme fraction (1), blue Sepharose fraction (2) and crude extract (3) were loaded onto nitrocellulose paper and labelled by goat anti-NR and then with anti-goat IgG conjugated with peroxidase.

Fig. 5. Titration of nitrate reductase by preimmune control and goat anti-NR sera.

Crude enzymatic extract from roots of the cucumber seedlings (1 cm³) was incubated 1.5 h in 4°C with preimmune serum (control — ○ —) or goat anti-NR (— ▲ —). After centrifugation (13 000 x g 10 min⁻¹) an enzyme activity was estimated as described in "Materials and Methods". Initial enzyme activity (100%) was 348 nmol NO₂⁻ x g⁻¹ FW.
Fig. 6. Dot-blot assays nitrate reductase in crude extracts from cucumber root, cotyledon and leaf.

5 µl undiluted samples of crude enzymatic extracts from roots (R), cotyledons (C) and leaves (L) (row 1) were loaded onto nitrocellulose paper. In row 2 each of the 5 µl samples contains 10 µg of protein. Dye dots were developed with 4 chloro-1-naphthol after staining with polyclonal anti-NR and goat anti-goat IgG conjugated with horseradish-peroxidase.

cucumber. These results indicate a similarity in antigenic sites of the nitrate reductase from different tissues. Also Sommers et al. (1983) tested NR from barley with Western-blot assay and showed that NR antibody from roots cross-reacted with NR from leaves. In our experiments different intensity of dye between dots in dot-blot assays was found. In row 2 (Fig. 5), when the amount of enzyme protein was the same in all dots, the dye reaction with cotyledon NR was slighter than with the other tissues.

After 1 h preincubation of polyclonal NR-antibodies at 4°C with crude extracts from all of the investigated tissues, differences in enzyme activity were observed (Table 2). With 1:100 dilution of antiserum, the activity of NR in roots and leaves was reduced by 30% and 45%, respectively. The decrease of NR activity in the presence of antibody was demonstrated also by Ferrario et al. (1983) for spinach nitrate reductase from both, root and leaf. However, these authors, basing on Ouchterlony test, suggested that the root and leaf NR of spinach was a different protein. On the other hand, Nakagawa et al. (1986) concluded that such result may be due to instability of the enzyme protein or properties of polyclonal antibodies. Our data have shown partial identity in immunological properties between the nitrate reductase protein in root and leaf tissues of cucumber seedlings. In contrast, the nitrate reductase of cotyledons demonstrated different properties from NR of roots and leaves. In the same dilution of antiserum (1:100) NR activity of cotyledons was stimulated whereas the enzymes from roots and leaves were inhibited. In the presence of NR antibody the nitrate reductase from cotyledon shows also a various intensity in dot-blot cross-reaction.

For this reason we suggest, that the nitrate reductase in storage tissues like cotyledons has slightly different molecular properties as compared with enzymes of other plant tissues.

TABLE 1. Purification of the nitrate reductase from cucumber roots.

<table>
<thead>
<tr>
<th></th>
<th>VOLUME (cm³)</th>
<th>TOTAL PROTEIN (mg)</th>
<th>TOTAL ACTIVITY (units⁺)</th>
<th>SPECIFIC ACTIVITY (units⁺ x mg⁻¹ protein)</th>
<th>RECOVERY (%)</th>
<th>PURIFICATION (fold)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>A.</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CRUDE EXTRACT</td>
<td>200</td>
<td>102.300</td>
<td>0.660</td>
<td>0.006</td>
<td>100</td>
<td>1</td>
</tr>
<tr>
<td>AMMONIUM SULFATE</td>
<td>10</td>
<td>59.440</td>
<td>0.261</td>
<td>0.044</td>
<td>67</td>
<td>68</td>
</tr>
<tr>
<td>BLUE SEPHAROSE</td>
<td>6</td>
<td>0.034</td>
<td>0.027</td>
<td>0.900</td>
<td>4</td>
<td>155</td>
</tr>
<tr>
<td><strong>B.</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CRUDE EXTRACT</td>
<td>200</td>
<td>124.000</td>
<td>0.724</td>
<td>0.006</td>
<td>100</td>
<td>1</td>
</tr>
<tr>
<td>BLUE SEPHAROSE</td>
<td>6</td>
<td>0.084</td>
<td>0.071</td>
<td>0.845</td>
<td>10</td>
<td>140</td>
</tr>
</tbody>
</table>

Units⁺ = µmole NO₂ x min⁻¹
25 g of cucumber roots were ground in a chilled mortar with extraction buffer (50 mM K-phosphate, 1 mM EDTA, 1 mM DTT, 1 mM PMSF, 10% glycerol, 0.5% PVPP, pH 7.5) and centrifuged at 20,000 g for 20 min. The supernatant named crude extract was used as the enzyme source. The ammonium sulfate precipitate (20%-50%) of the NR was loaded onto blue Sepharose column (6×2 cm³) as described in "Material and Methods".
TABLE 2. Preincubation of cucumber NADH-NR with goat polyclonal antibodies anti-NR.

<table>
<thead>
<tr>
<th>SERUM DILUTION</th>
<th>NR activity (µmol NO₃ x g FW x h⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Root</td>
</tr>
<tr>
<td>1 : 100</td>
<td>0.057</td>
</tr>
</tbody>
</table>

Crude enzymatic extracts from root, cotyledon and leaf were preincubated 1 h at 4°C without (control) or with 1:100 dilution of anti-NR, centrifuged (13 000 x g 10 min) and then activity of the enzyme was examined as described in "Materials and Methods". This experiment was repeated three times yielding similar results.

ACKNOWLEDGEMENTS

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LITERATURE CITED


IMMUNOLOGICZNE WŁAŚCIWOŚCI NADH:REDUKTAZY AZOTANOWEJ W RÓŻNYCH TKANKACH OGÓRKA

STRESZCZENIE

Praca miała na celu otrzymywanie przeciwciał skierowanych przeciwko reduktazie azotanowej (anti-NR) oraz sprawdzeniu ich interakcji z enzymem izolowanym z różnych tkank chogórka. Cytosolową formę reduktazy azotanowej uzyskano w wyniku ekstrakcji korzeni ogórka oczyszczonego przez filtrację na żel z blue-Sepharose 4B. Elektroforeza przeprowadzona w warunkach niedenaturujących białka potwierdziła czystość preparatu, który został następnie użyty do immunizacji kozy. Otrzymane przeciwciała anti-NR testowano metodą ELISA, miarzeczkowaniem immunologicznym oraz dot-blot, a następnie wykorzystano do identyfikacji reduktazy azotanowej w różnych tkankach ogórka. W ekstrakcjach z korzeni i w liści zaobserwowano wyraźne hamowanie aktywności reduktazy azotanowej w obecności przeciwciał anti-NR. Podobnego efektu nie znaleziono dla ekstraktu enzymatycznego otrzymanego z liścienni.

SŁOWA KLUCZOWE: Reduktaza azotanowa, przeciwciała, dot-blot, badania immunologiczne, Cucumis sativus.