Identification of excreted iron superoxide dismutase for the
diagnosis of Phytomonas

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An excreted iron superoxide dismutase (FeSODe) of pI 3.6 with a molecular weight of 28-30 kDa was detected in the in vitro culture of Phytomonas isolated from Euphorbia characias (SODeCHA) and from Lycopersicon esculentum (SODeTOM), in Grace’s medium without serum. These FeSODe excreted into the medium had immunogenic capacity: the positivity of the anti-SODeCHA serum persisted to a dilution of 1/30,000, and for the anti-SODeTOM to 1/10,000 by Western blot. In addition, cross reaction was detected between the anti-SODe serum of Phytomonas isolated from E. characias against SODeTOM, and the anti-SODe serum from L. esculentum with SODeCHA. This characteristic offers the possibility of its use to diagnose plant trypanosomatids. The validation of the test was confirmed by experimental inoculation of tomato fruits with Phytomonas isolated from L. esculentum. At 7, 10, 15, and 21 days post infection, it was possible to detect the presence of the parasites with the anti-SODe serum of Phytomonas isolated from L. esculentum at a dilution of 1/250. These serological results were confirmed by visualization of the parasites by optical microscopy. The data of this study confirm that the SOD is sufficient to identify a trypanosomatid isolated from plants as belonging to the genus Phytomonas.

Key words: superoxide dismutase - immunogenicity - molecular tool - Phytomonas spp.

Trypanosomatids belonging to the genus Phytomonas are fairly common in the latex, phloem, fruit sap, seed albums, and even in the nectar of many plant families. Also, their world distribution is widespread. They are etiological agents in devastating crop epiphytotics (coconut, oil palm, coffee, and manioc) but they also parasitize many plants without apparent pathogenicity (Dollet 1984).

Probably more than one genus of Trypanosomatidae is represented among plant parasites. Therefore, both the study of the circulation of parasites as well as the demonstration of infection in hosts require a precise identification of trypanosomatids. The distinctions between them is difficult on a morphological basis. Attempts to overcome this difficulty prompted the testing of various serological, biochemical, and molecular methods, but none to date have been very successful for genus distinction (Camargo 1999). In recent years, new polymerase chain reaction method for the amplification of the SL sequence has been successfully developed, and more recently new molecular markers have become available to help in the identification (Serrano et al. 1999, Dollet et al. 2001, Marín et al. 2004a).

Superoxide dismutases (SODs, EC 1.15.1.1) are metalloproteins that occur ubiquitously in nature and efficiently dismutate the superoxide anion into oxygen and hydrogen peroxide. SODs are assigned to different families, based on their metal cofactors – iron (FeSOD), manganese (MnSOD) and copper-zinc (Cu/ZnSOD) – which also differ in their location: cytosol, cell organelles, and even cell excretions. Thus, the Cu/ZnSODs appear primarily in the cytosol of eukaryotic cells; the MnSODs in the mitochondrial matrix; and FeSODs in the cytosol of plant cells as well as in some free-living protozoa and protozoan parasites (Fridovich 1995).

SOD activity has been detected in the main species belonging to the family Trypanosomatidae, in Trypanosoma cruzi (Ismail et al. 1997), in T. brucei brucei (Kabiri & Steverding 2001), in several species of the genus Leishmania (Ismail et al. 1994), in some lower trypanosomatids, and in plant trypanosomes (Quesada et al. 2001, Marín et al. 2004a,b). SODs from protozoan parasites are considered virulence factors that protect the parasites from the attack of the host cells by the action both of oxidant and anti-inflammatory agents (Paramchuck et al. 1997), and can even confer immunological capacities (Pérez-Fuentes et al. 2003). In general, all the parasitic protozoa studied to date have only FeSOD (Marín et al. 2004a,b).

Extracellular SODs (EcCu/ZnSODs) have been evidenced in invertebrates, in several species of nematodes, such as, Brugia pahangi (Tang et al. 1994), Onchocerca volvulus (James et al. 1994), and Caenorhabditis elegans (Fuji et al. 1998) and the crustacean Callinectes sapidus (Brouwer et al. 2003). Recently, Villagrán et al. (2005) have shown the presence of a iron SOD in T. cruzi. Also, in mammals, several researchers have studied extracellular SOD (EC-SOD), which is the major extracellular antioxidant enzyme and plays a critical role in the pathogenesis of a variety of pulmonary, neurological, and cardiovascular diseases (Suliman et al. 2001, Fukai et al. 2002).

In a previous paper, we reported in a digitonin-titration experiment that 20% of the SOD activity of Phy-
tomonas sp. was not solubilized even at high digitonin concentrations, indicating that part of the SOD could be associated with membranes or even excreted by the parasite (Marín et al. 2004b). Therefore, in the present study, we show the existence of a SOD excreted into the supernatant from the culture of Phytoomonas isolated from Euphorbia characias, and Phytoomonas isolated from Lycopersicon esculentum, cultured in Grace’s medium without serum. Also, we have studied the usefulness of excreted SOD in the identification and diagnosis of Phytoomonas infections using Western blot in the identification.

MATERIALS AND METHODS

Parasites and SOD excreted fraction extraction - Phytoomonas isolated from latex vessels of E. characias (Dollet 1984) and from L. esculentum (tomato) in Spain (Sánchez-Moreno et al. 1995), were grown in axenic Grace’s medium (Sigma) supplemented with 10% heat inactivated foetal bovine serum at 28ºC. Cells at the exponential growth medium (Sigma) supplemented with 10% heat inactivated (Sánchez-Moreno et al. 1995), were grown in axenic Grace’s distributed into aliquots of 5 × 10^9 parasites/ml in Grace’s medium without serum and grown for 24 h. The supernatant was collected by centrifugation at 1500 × g for 10 min, passed through a 0.45 µm pore size filter, and then precipitated with ammonium sulphate at 35-85%. This was centrifuged (9000 × g for 20 min at 4 ºC), redissolved in 2.5 ml of 50 mM potassium phosphate buffer (pH 7.8), and afterwards desalted by buffer exchange in a Sephadex G-25 column (Pharmacia, PD 10), previously balanced with the potassium buffer, bringing it to a final volume of 3.5-ml. Thus, we obtained the SODe fraction from Phytoomonas isolated from E. characias (SODeCHA) and SODe from Phytoomonas isolated from L. esculentum (SODeTOM). These fractions were finally concentrated by ultrafiltration in Centricon-50 tubes (Amicon) at 3000× g to a final concentration of 2 mg/ml. The protein content was determined in all fractions using the Bio-Rad test, based on the Bradford method (Bradford 1976), with BSA as a standard. For the extraction and purification of the P85 fraction, we followed the protocol described by Marín et al. (2004b).

Determination of molecular weights and isoelectric point of SODe - Apparent molecular weights of the semi-purified enzymes (SODeCHA and SODeTOM) were determined by the separation of 7 µg/lane on 20% homogeneous Native-PAGE gel in the Phast System (Amersham Pharmacia Biotech, Uppsala, Sweden). The isoelectric points were determined in the Phast System (7 µg/lane) in polyacrylamide Phast gel pl 3-9 as described Bécuwe et al. (1996). Proteins markers for molecular weights and pl were provided by Pharmacia (Uppsala, Sweden). The SOD activity on the gels was visualized by staining, following Beyer and Fridovich (1976), and for protein markers the lanes were stained with silver nitrate and coomassie blue, as described by Phast System handbook.

Polyclonal serum - To obtain the specific antibodies against the fractions SODeCHA and SODeTOM, we immunized two female 4-week-old Balb-C mice (concentration of proteins of 2 mg/ml). These fractions were separated by electrophoresis of IEF 3-9 in polyacrylamide gels as described elsewhere (Marín et al. 2004a). In this way, we obtained the anti-SODeCHA and anti-SODeTOM sera. In addition, the serum from the mouse that had not been immunised with the antigen fraction (control serum) was collected.

Experimental infection of tomatoes - Isolated tomato were infected by needle inoculation in two zones, with 5 × 10^5 culture forms of Phytoomonas isolated from L. esculentum in 50 µl. As controls, three tomatoes were inoculated with the same volume of axenic Grace’s medium supplemented with 10% heat-inactivated foetal bovine serum. At 7, 10, 15, and 21 days post-inoculation, three tomatoes, controls and infected, were processed to obtain the extracts for later studies. After the inoculation zones were collected, smears fixed with methanol and stained by Giemsa were used for light microscopy. Afterwards, the material was homogenized and centrifuged at 1500 × g for 10 min. The supernatant was filtered, precipitated with ammonium sulphate, passed through a Sephadex G-25 column, and concentrated as described above for obtaining the SODe fraction. Also, the zones opposite the inoculation from infected tomatoes were processed in the same way.

Western-blot analysis - For the polyclonal serum titration, we used the SODe fraction (SODeCHA and SODeTOM) as the antigen fraction. These were run on IEF 3-9 gels (protein concentration of proteins of 2 mg/ml), and afterwards transferred to nitrocellulose, for 30 min, as prescribed in the Phast-System manual. The membrane was blocked for 2 h at room temperature using 0.4% gelatine and 0.2% Tween 20 in PBS, followed by three washes in 0.1% Tween 20 in PBS (PBS-T). Next, the membrane was incubated for 2 h at room temperature, either with negative serum at a 1/100 dilution or with anti-SODeCHA or anti-SODeTOM sera at dilutions of: 1/100, 1/500, 1/1000, 1/5000, 1/10,000, 1/20,000, and 1/30,000. After being washed as above, the membrane was further incubated for 2 h at room temperature with the second antibody, anti-mouse IgG (Fc specific) peroxidase conjugate (Sigma®) (dilution 1/1000). After washing, the substrate diaminobenzidine was added and the reaction stopped with several washes in distilled water (Marín et al. 2004a). For the cross reaction analyses, we followed the same protocol described above but with a dilution of the anti-SODeCHA and anti-SODeTOM sera of 1/5000.

Western-blot was also used to detect the SODeTOM in experimentally inoculated tomato fruits, as described above. For this, the extracts from the tomato of 7, 10, 15, and 21 days were used as the antigen fraction, as were the extracts of control tomatoes. The dilution of the anti-SODeTOM serum of 1/250 was used.

RESULTS

Isolation of an excreted FeSOD - When promastigote forms of Phytoomonas isolated from E. characias and L. esculentum (5 × 10^5 parasites) were cultured for 24 h in Grace’s medium without serum and the cell-free superna-
tant was collected, concentrated, separated by isoelectric focusing (pI 3-9) and 20% homogeneous native-PAGE and followed by SOD activity staining. We detected a single SOD band (Fig. 1A, lanes 2 and 4; Fig.1B, lanes 5 and 6), which we called SODeCHA and SODeTOM, respectively.

The isoelectric point of these bands of excreted SOD (SODeCHA and SODeTOM) was consistent with the isoelectric point (pI 3.6) of the SODII band detected in the partially purified fraction of *Phytomonas* promastigotes cultured under normal conditions (P85) (Fig. 1, lanes 1 and 3).

The two isoenzymes excreted had very similar molecular weights, of approximately 28-30 kDa (Fig. 1). In both cases, they were demonstrated to be FeSOD (data not shown).

Marker enzymes (pyruvate kinase and hexokinase) have confirmed that there was no lysis of the parasite during this culture period and that the presence of the SOD was due only to excretion by the parasite (data not shown).

**Immunogenicity of the SODe** - Polyclonal antibodies against the enzymes SODeCHA and SODeTOM were obtained from immunized Balb-C mice, and, by Western blot, we demonstrated their immunogenic capacity (Fig. 2). With the control serum (Fig. 2, lines 1 and 9), the reaction was negative at a dilution of 1/100, regardless of the antigen fraction. Meanwhile, in the case of the anti-SODE serum (Fig. 2; lanes 2-8 and lanes 10-16, respectively), the reactions proved positive up to a dilution of 1/30,000 for SODe-CHA and 1/10,000 for SODe-TOM.

We determined whether there was any cross reaction between the polyclonal sera antiSODeCHA and SODeTOM, by Western blot (Fig. 3). There was cross reactivity between the sera to a dilution of 1/5000.

**Detection of SODeTOM in infected tomatoes** - At 7, 10, 15, and 21 days after infection, the inoculation zones and opposite ones were processed as described to be analyzed by Western blot. The tomato extracts were used as the antigen fraction against the anti-SODETOM serum at the dilution of 1/250, and, the presence of SODeTOM was detected (Fig. 4). In addition, not only did the inoculation zone prove positive, but on day 21 post-infection, we demonstrated the presence of these parasites in zones opposite the inoculation in the infected tomatoes (Fig. 4, line 5). Thus, we showed that there is dissemination of the parasites in the tomato.

These results were confirmed by direct examination of the parasite by light microscopy (Fig. 5). The flagellates
detected presented the typical promastigote form, with a twisted body and in most cases with a large flagellum, although the smallest promastigote forms without flagella were also visible.

**DISCUSSION**

In previous studies, we have purified and characterized two isoenzymes (SODI and SODII) from *Phytomonas* isolated from *E. characias* (Marín et al. 2004a,b), confirming that one of these (SODII), with a pI 3.6, presented immunogenic properties.

Currently, the direct way to detect the presence of flagellates belonging to the genus *Phytomonas* in a plant is by observation with the light microscope. This makes the identification and the diagnosis difficult and at times

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**Fig. 4:** Western blot of the polyclonal sera anti-SODeTOM at a dilution of 1/250, against the infected tomato extract. Lanes - 1: day 7; 2: day 10; 3: day 15; 4: day 21; 5: zone opposite the inoculation day 21; 6: tomato extract control; and 7: SODeTOM; marker proteins of pl, stained with Coomassie Blue.

**Fig. 5:** detection of *Phytomonas* isolated from *Lycopersicon esculentum* in tomato by light microscopy. A: day 7; B: day 10; C: day 15; and D: day 21. Bar = 1 µm.
even impossible, because promastigote stages of Leptomonas, Herpetomonas, Crithidia, and Blastocerithidia also occur in Phytomonas vectors (Wallace et al. 1992, Camargo & Wallace 1994, Podlipaev 2000), and have sometimes been detected in plant tissues. Thus, it becomes necessary to seek new methods to detect the parasite. Diverse authors have suggested that Western blot can be used as a complementary and alternative method for conventional serology tests (Ávila et al. 1993), and therefore may be useful in detecting and identifying parasitic protozoa using antigens from epimastigotes forms and SODe (Vissoci et al. 1998, Villagrán et al. 2005). Also, we have demonstrated that the SODII for its immunogenic characteristics is an excellent molecular marker of the genus Phytomonas, which could be applied to the different immunological techniques.

To our knowledge, this is the first evidence that members belonging to the genus Phytomonas excrete a FeSOD into the culture medium. These data confirm the findings of our earlier study (Marín et al. 2004a), showing that SODII would be formed in the cytosol and afterwards transported to the glycosomes, as occurs with other glycosomal enzymes (Parsons et al. 2001), and to the membranes, and part of this enzyme could even be secreted to the exterior by the parasite, as a defence mechanism of the parasite against toxic radicals generated by the host (Kabiri & Steverding 2001).

One of the aim in the present work was to study the immunogenic properties of this FeSODe, for which we obtained polyclonal antibodies against SODeCHA and SODETOM from immunized Balb-C mice. Western blot showed a positive reaction up to a dilution of 1/30,000. This result indicates that, FeSODE is a highly immunogenic protein, opening the possibility of its use for the diagnosis of parasitism by species belonging to the genus Phytomonas.

To confirm the usefulness of this excreted protein (SODE) as a marker in Western-blot diagnosis, we performed an inoculation experiment on tomato fruits. The presence of Phytomonas was detected at 7, 10, 15, and 21 days after infection in the inoculation zones with the anti- SODETOM serum at the dilution of 1/250, and also on day 21 in the zones opposite the inoculation. Hence, SODE is a specific and sensitive molecular tool and, used with Western blot, can be useful to the diagnosis of Phytomonas spp. The flagellates detected in the laboratory-infected tomatoes by light microscopy presented the typical promastigote form, although the smallest promastigote forms without flagella were also visible, these being very similar to others reported elsewhere (Jankevicius et al. 1989).

The data of this study together with those gathered recently in our laboratory (Marín et al. 2004a,b) confirm that the SOD (SODII or SODE) is sufficient to identify a trypanosomatid isolated from plants as belonging to the genus Phytomonas and to distinguish between a true Phytomonas and other trypanosomatids that can provoke transient infections in plants. Also, a certain amount of SOD is secreted by the parasites, which is highly immunogenic and by Western blot assay capable of detecting the presence of Phytomonas isolates from L. esculentum in crude preparations from tomato saps. Thus, we were able to detect this parasitism without the need for isolation, culture, or DNA extraction of flagellates. Of course, the validity of the SODEs by Phytomonas spp. as a diagnostic tool needs to be continuously evaluated as new isolates and molecular data became available.

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