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Sinding properties of immunoglobulin-binding protein by double immunodiffusion (Ouchterlony) technique.

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- 1 The binding of SpLAG, SpLG and SpAG with animal sera, avian IgY, and purified mammalian IgG are investigated by double immunodiffusion.
- 2 Briefly, 1% agarose gels are prepared and wells cut into the gel using a template.
- 3 Initially, aliquots of 25 μ l each of the immunoglobulin-bacterial protein SpLAG, SpLG or SpAG at a concentration of one (1) μ g/ μ l are applied to the centre well.
- 4 The peripheral wells are filled with 25 μ l each of IgY (30 μ g/ μ l) or mammalian IgG or sera.
- 5 The gels are incubated at RT for 48-72 hours.

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6	After that the gels are examined for precipitin lines.
7	Human serum or human IgG are included as positive controls.
8	The positive results are taken as the presence of precipitin line/s and negative results, the absence of precipitin lines.
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