

## IMPROVED CLINICAL SPECIFICITY OF HEPARIN INDUCED THROMBOCYTOPENIA IMMUNOASSAY

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### OBJECTIVE OF THE STUDY

Immunoassays for Heparin induced Thrombocytopenia (HIT) offer a good sensitivity but poor clinical specificity as many of the antibodies including some IgGs, are asymptomatic.

### Improvement of HIT immunoassays is required for:

- Obtaining high Sensitivity/Specificity for « true » Heparin dependent Antibodies associated with clinical HIT.
- Avoiding interference of «false» antibodies («sticky samples»).
- Differentiate symptomatic from asymptomatic heparin dependent antibodies.
- The goal of this study was to evaluate which analytical factors could be associated with heparin dependent antibody pathogenicity, then to use these findings for developing more clinically relevant immunoassays for HIT.

### Background:

Affinity purified heparin dependent antibodies from 3 patients with HIT showed that only the high avidity IgG fraction, produces high positive SRA tests, whilst the lower affinity IgG fraction, although it bounds identically to H-PF4, was unable to activate platelets (SRA negative) (Amiral et al., BSH, 109,1-712000).

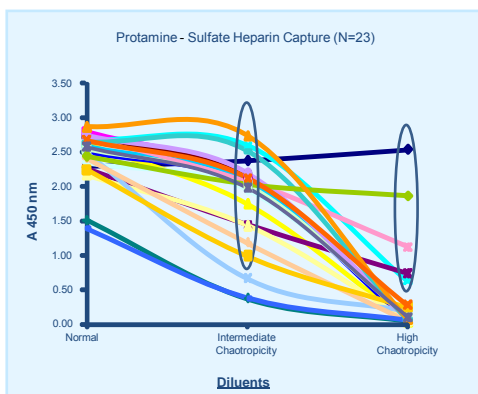
From this observation, we are developing an immunoassay approach by privileging binding of antibodies with the highest affinity to the heparin dependent antigen, mainly PF4.

### RESULTS:

#### Incidence of Sample diluent chaotropy on antibody binding to:

#### Heparin Protamine Sulfate

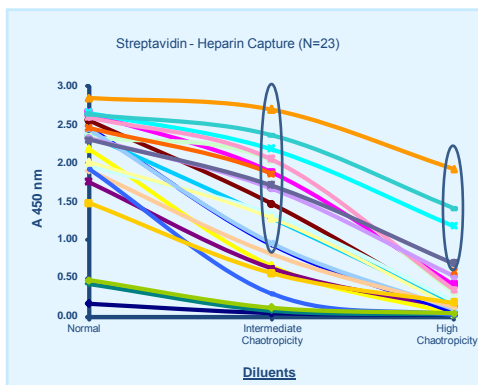
When excess of Heparin with Protamine sulfate is used for coating, all the samples have an OD>1.00 with diluent no.1 and only 5 remain positive with diluent no.3. These patients are within those presenting the most severe clinical symptoms and the highest HIT score. With buffer N°2, 19 out of 23 keep an OD >1.00. The OD doesn't decrease for 2 samples (these 2 samples were later demonstrated to have antibodies to Protamine Sulfate).



#### Biotinylated-Heparin-Streptavidin

With the Biotinylated Heparin Streptavidin system, only 20 samples have an OD>1.00 with the diluent no.1, but 2 are borderline (A450 of about 0.50) and 1 is negative. OD decreases for all samples with diluent no.3, but 4 specimen remain highly positive (OD>1.00).

Interestingly, with buffer N°2 of intermediate chaotropy, only patients with a definite HIT remain highly positive (A450>1.00), which represents 15 out of the 23 patients.



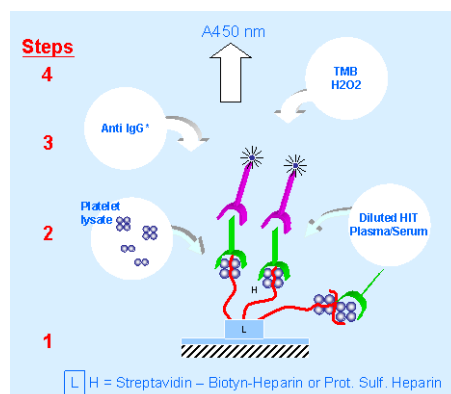
### ASSAY PRINCIPLE AND REAGENTS

The study was based on the dynamic assay that we recently developed (Zymustest HIA IgG,A,M or IgG). In this assay, the immune complexes between PF4 (provided by platelet lysates) and HIT antibodies (when present in the assayed specimen) are formed with immobilized heparin fixed onto the plate through Protamine Sulfate (heparin being in large excess), or through biotinylated heparin and Streptavidin. An assay variant was designed by increasing chaotropy of sample diluent, in order to favour the binding of antibodies with the highest affinity.

Only IgG isotypes were measured.

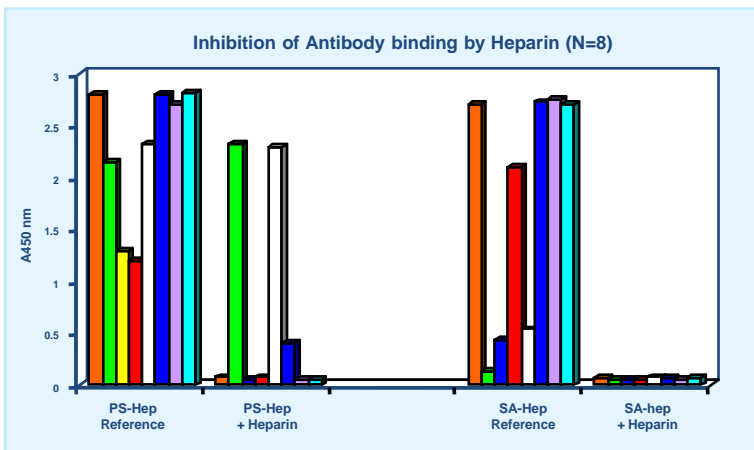
### Samples tested:

- 23 samples from patients with HIT were tested.
- All the patients were PAT and SRA positive, and platelets counts decreased by > 50%.



### Confirmation of specificity by inhibition study

The inhibition of antibody binding by heparin (2 IU/ml in the reactive well) were tested on 8 samples.



→ When the Streptavidin system is used for coating, antibody binding was completely inhibited in all the 8 samples, but inhibition occurred only for 6 samples for the Protamine sulfate coating.

→ The two samples non inhibited were further demonstrated to have antibodies to Protamine Sulfate.

### CONCLUSIONS

- HIT Antibodies «are» pathogenic when they bind to heparin dependent antigen (PF4 mainly) present in blood and onto blood cell surfaces. The most harmful are those exhibiting the highest avidity.
- Using biotinylated heparin immobilized onto the microELISA plate through Streptavidin, and in presence of platelet lysate, the most potentially pathogenic IgGs can be identified by using buffers with increasing chaotropy. This approach improves the clinical specificity of measured antibodies.
- New laboratory assays could be designed from this observation, but extensive clinical studies are required for documenting the correlation of analytical measurements with severity of HIT clinical context.
- Should we «disregard» asymptomatic antibodies?

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### DISCUSSIONS

This approach allows distinguishing between patients and selecting those having antibodies with high avidity and affinity, and potentially the most harmful.

- In these conditions, some heparin dependent antibodies became almost negative, whilst other remained highly positive. Presence of antibodies to Protamine Sulfate is evidenced in 2 of the patients.
- These preliminary results suggest that pathogenic effect of antibodies could be graduated by classifying them through their affinity for the heparin dependent antigen, testing preferentially their binding to biotinylated Heparin with Streptavidin.
- This could allow developing diagnostic approaches more specific for the immediate clinical risk of HIT in patients, or for a diagnosis correlating better with the clinical status.