Antibody analyses with BiacoreTM system convenient, sensitive, and versatile

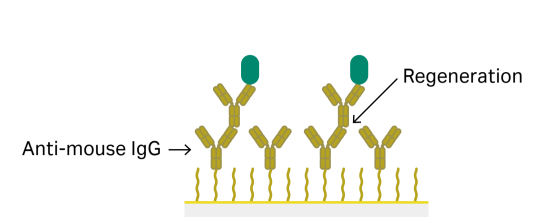
Eric Roush¹, Ewa Pol², Anna Moberg²

¹Cytiva, Marlborough, MA, USA, ²Cytiva, Uppsala, Sweden

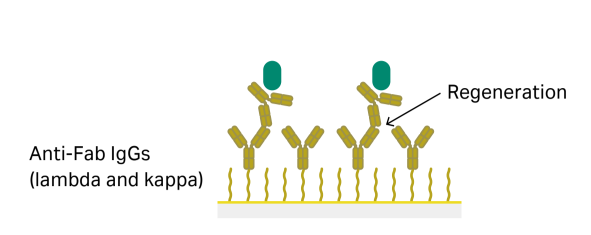
Convenience

Label-free interaction analysis on a Biacore™ system offers an array of possibilities for in-depth antibody characterization. A range of capture kits are available, ranging from antibody to different types of tag capture kits. Biacore sensor surfaces may also be used for convenient immobilization of a capture molecule of choice.

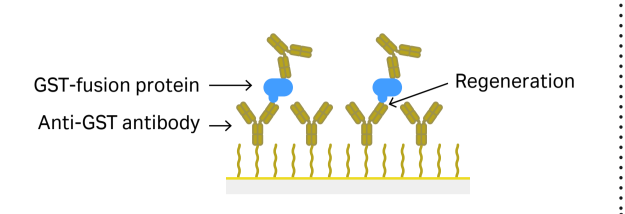
Mouse Antibody Capture Kit



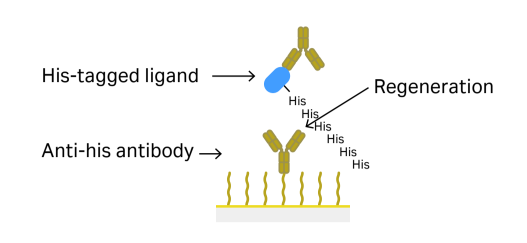
Human Fab Capture Kit



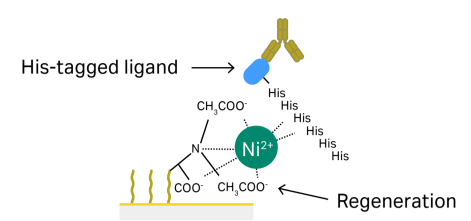
GST Capture Kit



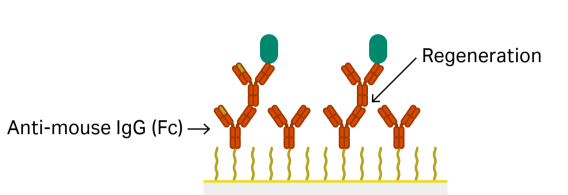
His Capture Kit



Sensor Chip NTA



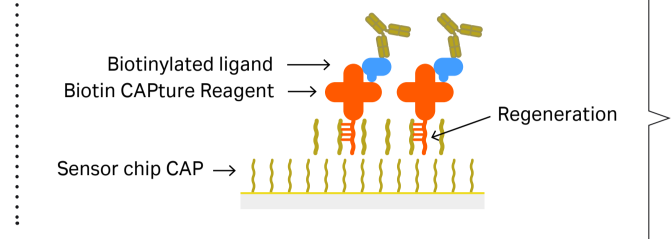
Human Antibody Capture Kit

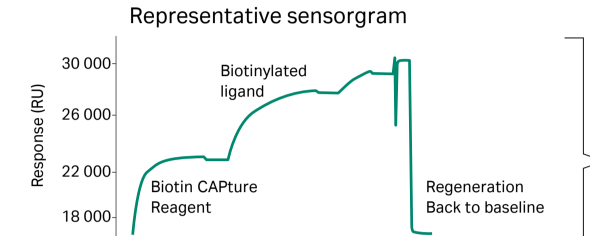


Benefits

- Orientated immobilization of ligand from complex solution
- · Captured ligand is easily changed
- Capture molecule specific to all human IgG subclasses
- High quality reagents and optimized protocols that save time and effort

Biotin CAPture Kit

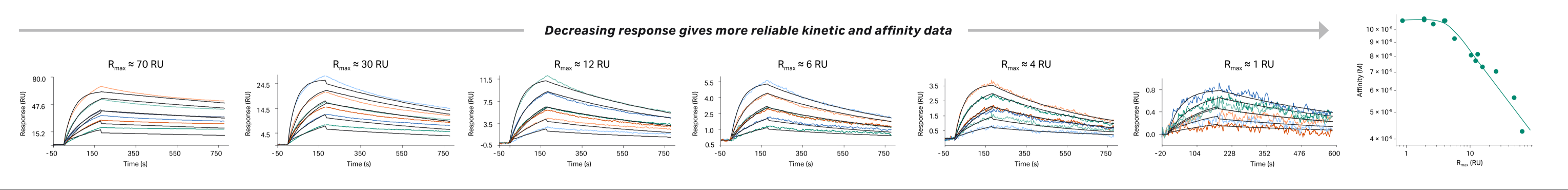




- Biotin CAPture Reagent is bound (captured) to the surface by oligo hybridization
- Biotinylated ligand is immobilized via biotin
- · Analyte interaction with ligand is studied
- The surface is regenerated and rebuilt in next cycle

Sensitivity

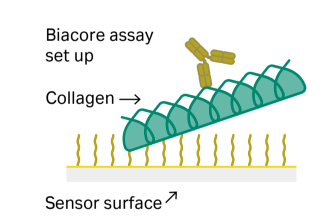
In many cases, a better mimic of the biological situation is to inject antibody and allow it to react with immobilized antigen. However, the bivalent nature of antibodies allow avidity effects to arise and kinetic analysis becomes challenging. Biacore system offers sensitivity that makes it possible to eliminate avidity effects by immobilizing very low amounts of antigen on the surface. Interaction analyses can be performed with confidence at extremely low response levels. As shown below, at the maximal observed response (R_{max}) below 6 RU, the avidity effects start to disappear and the interaction is described by 1:1 binding model.



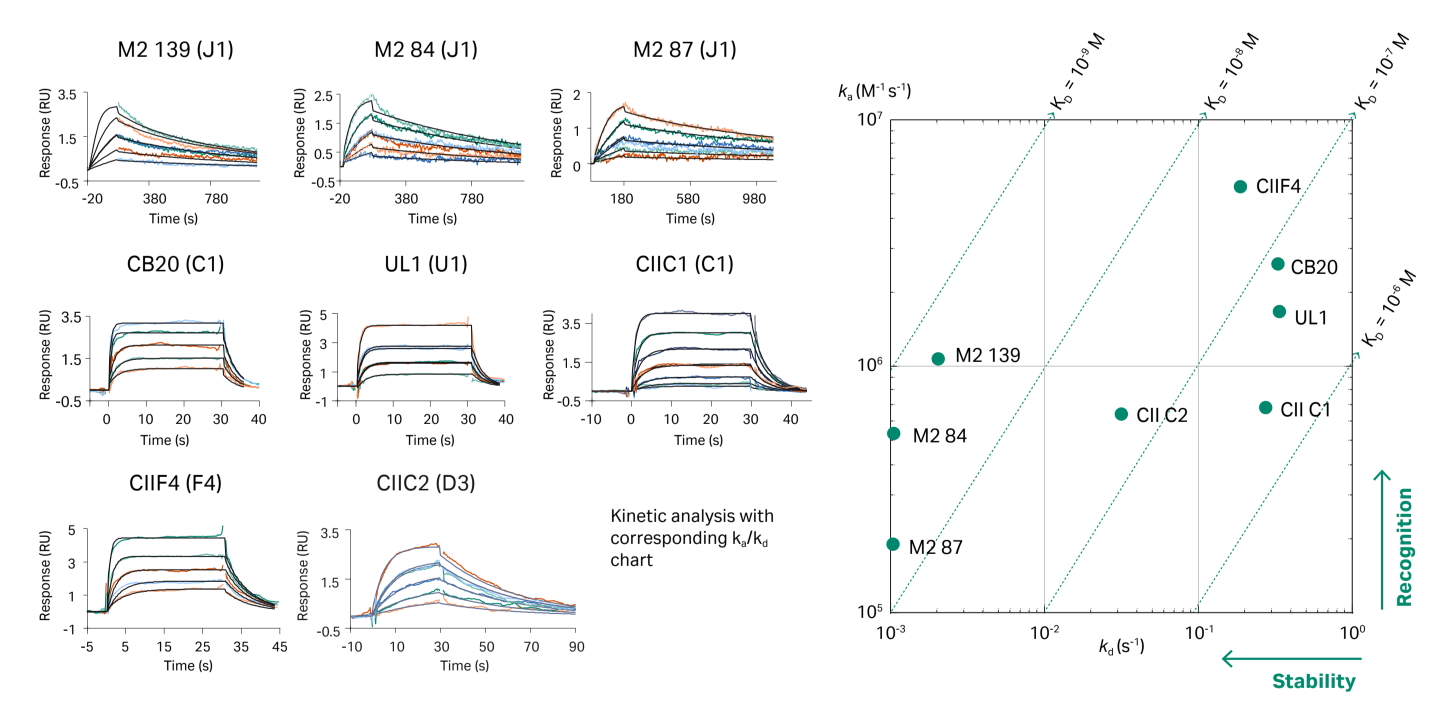
Versatility

Antibody characterization in the investigation of a disease mechanism

The autoimmune inflammatory disease rheumatoid arthritis (RA) is characterized by development of autoantibodies to collagen type II. The disease can be mimicked in a collagen induced arthritis animal model, in which the animal develops similar symptoms and autoimmune antibodies. To study the importance of specific collagen epitopes, we analyzed the interactions of antibodies, purified from mouse serum, with various epitopes on collagen: C1, U1, J1, D3 and F4.



In the Biacore assay, collagen was immobilized on the sensor surface because fibrillar collagen has a tendency to aggregate, and if used in solution (a reversed assay set up) the concentration would be underestimated, which would lead to an underestimation of the association rate constant and thereby also affinity.



M2 antibodies against J1 epitope are secondary strongly mutated antibodies. They are highly pathogenic, they induce strong clinical athritis, and they also display the highest affinity against J1 epitope.

CB20, UL1 and CIIC1 induces clinical and subclinical arthritis and are less pathogenic than M-antibodies. All three antibodies show considerably lower affinity against collagen than M antibodies.

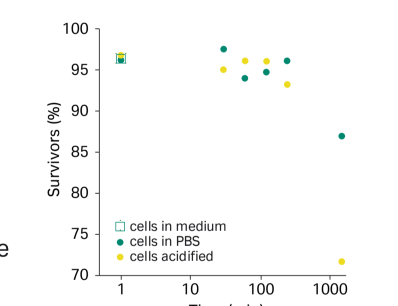
The pathogenicity of CIIF4 and CIIC2 antibodies is being investigated. According to these results, the higher affinity indicates the higher pathogenicity. Antibodies with longer residence times are likely to be stronger inducers of downstream destructive processes as for example immuno complex formation and complement activation.

Antibodies binding to surface-attached cells

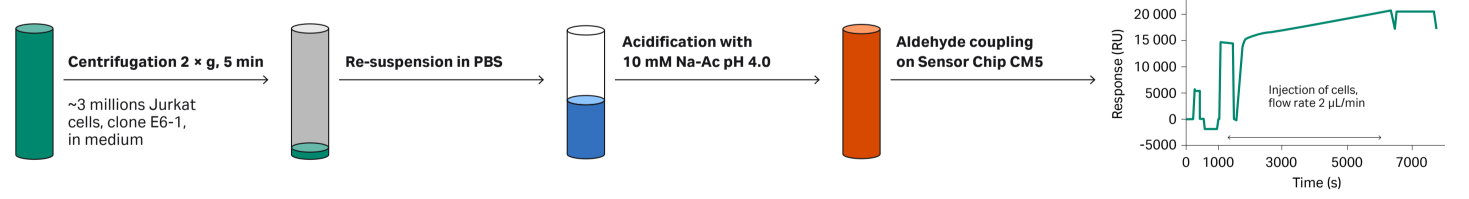
Studies of antibody interactions with cell surface proteins are hampered by the difficulties of creating stable and relevant conditions for proteins when isolated from the cell membrane. This could be circumvented by analyzing antibody binding to whole cell. Here we present inital results from the development of a cell assay.

Cell survival test Results from the examination of the cell survival ability when exposed to various steps of preparation procedure, binding of three antibodies, anti-CD25, anti-CD4 and anti-CD247, to the

glycoproteins on cell surface.



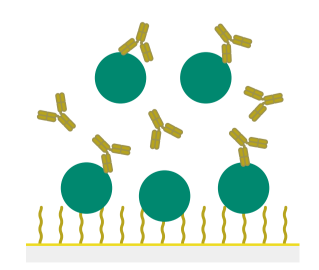




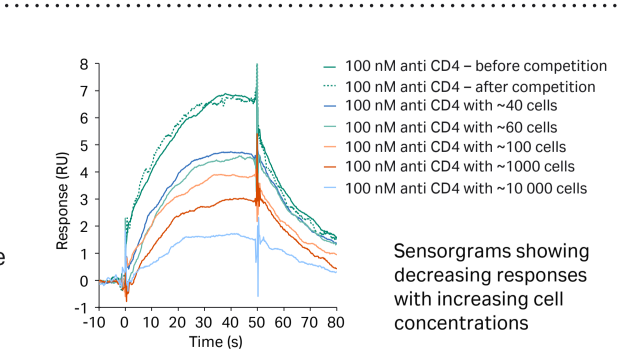
Three antibodies binding to glycoproteins on cell surface Anti-CD25 Ab Anti-CD247 Ab Anti-CD4 Ab **Negative** control 6-100 nM 3-100 nM 3-100 nM Anti-CD19 Ab 3-200 nM $k_a = 4.0 \times 10^5 \text{ M}^{-1} \text{ s}^{-1}$ $k_a = 7.8 \times 10^5 \text{ M}^{-1} \text{ s}^{-1}$ $k_a = 2.3 \times 10^5 \,\mathrm{M}^{-1} \,\mathrm{s}^{-1}$ $_{\rm D} = 6.5 \times 10^{-8} \, \text{M}$ $\zeta_D = 3.6 \times 10^{-8} \text{ M}$ 100 140 100 100 60

Confirming binding specificity by cell competition assay

will increase.



In the competition assay, the antibody is mixed with increasing number of cells and sequentially injected over the cell surface. If cells in solution are competing with cells on the surface to bind to the same antibody, the binding response will decrease and the concentration of cells in solution



Acknowledgements

Prof. Rikard Holmdahl and Dr. Christoph Kessel, Medical Inflammation Research, The Karolinska Institute in Stockholm and Dr. Andrew Sanderson, GlaxoSmithKline in Cambridge, are acknowledged for providing reagents and for valuable discussions on rheumatoid arthritis and cell assay, respectively.

Conclusion

With easy and efficient sample preparation, high sensitivity, and the comprehensive information in a variety of applications make SPR technology a key platform for antibody characterization.

