# FA1-MS12-T04

## Structural investigation of T cell Tolerance.

Stephanie Gras<sup>a</sup>, Zhenjun Chen<sup>b</sup>, John J Miles<sup>c</sup>, Yu Chih Liu<sup>a</sup>, Lars Kjer-Nielsen<sup>b</sup>, James McCluskey<sup>b</sup> & Scott R Burrows<sup>d</sup>, Jamie Rossjohn<sup>a</sup>. <sup>a</sup>Department of Biochemistry and Molecular Biology, Monash University, Clayton, Australia. <sup>b</sup>Department of Microbiology & Immunology, University of Melbourne, Australia. <sup>c</sup>Cellular Immunology Laboratory, Queensland Institute of Medical Research, Brisbane, Australia. <sup>d</sup>School of Medicine, University of Queensland, Brisbane, Australia. E-mail: Stephanie.gras@med.monash.edu.au

During selection of the T cell repertoire, the immune system navigates the subtle distinction between self-restriction and self-tolerance, yet how this is achieved is unclear. We studied how self-tolerance toward a trans-HLA (human leukocyte antigen) allotype shapes T cell receptor (TCR) recognition of an Epstein-Barr virus (EBV) determinant (FLRGRAYGL named as FLR).

The X-ray crystallography structure of the immunodominant LC13 TCR in complex with HLA-B8-FLR showed a very C terminally focuses docking of this TCR [1]. The LC13 TCR is self-reactive with HLA-B44, so the HLA-B8+/HLA-B44+ individuals use another T cell repertoire to recognize the HLA-B8-FLR complex. In order to understand how this new T cell repertoire lacks to self-react with HLA-B44, we were interested to study the atomic level of the interaction between one TCR of this new repertoire and the HLA-B8 in complex with the EBV antigen, FLR.

We have solved by X-ray crystallography the structure of the CF34 TCR, from HLA-B8+/HLA-B44+ individuals, in complex with the HLA-B8-FLR at 2.80Å resolution [2].

This structure shows that the CF34 TCR docked at the N terminus of HLA-B8-FLR, whereas the alloreactive LC13 TCR docked at the C terminus of HLA-B8-FLR.

The CF34 TCR docking onto the peptide-MHC coincided with a polymorphic region between HLA-B8 and HLA-B44 allowing CF34 to "see" the difference between the two MHCs. To further test this hypothesis we have performed an extensive mutagenesis study on the HLA-B8 molecule in order to map the energetic footprint of those two TCRs and we have also add to this study another TCR, named RL42 from HLA-B8/B44+, for which we solved the structure in complex with HLA-B8-FLR. All together those unpublished data shed some light on the T cell Tolerance mechanism.

[1] Kjer-Nielsen L et al., Immunity, 2003. [2] Gras S et al., Immunity, 2009.

# Keywords: immune system, TCRpMHC, Epstein Barrvirus

## FA1-MS12-T05

**Structure-based Discovery of Antivirals Targeting the Proteases of RNA Viruses.** <u>Rolf Hilgenfeld</u><sup>a</sup>, Lili Zhu<sup>a</sup>, Ksenia Pumpor<sup>a</sup>, Yuri Kusov<sup>a</sup>, Shyla George<sup>a</sup>, Thomas Peters<sup>b</sup>, Torsten Biet<sup>b</sup>, Sara Lacerda<sup>a</sup>, Jinzhi Tan<sup>a</sup>. *Institutes of <sup>a</sup>Biochemistry and <sup>b</sup>Chemistry, Center for Structural and Cell Biology in Medicine, University* 

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## of Lübeck, Ratzeburger Allee 160, 23528 Lübeck, Germany. E-mail: hilgenfeld@biochem.uni-luebeck.de

Current methods in structure-based discovery of antiviral lead compounds include de-novo design, virtual screening, fragment screening, and structure-guided medicinal chemistry, all of which require a detailed knowledge of the structure and function of the viral target proteins. We focus on the 3C and 3C-like proteases of enteroviruses and coronaviruses, resp., as targets for drug discovery [1-9]. The combined application of X-ray crystallography, STD-NMR spectroscopy, structureguided medicinal chemistry, and viral replicons in our laboratories proved essential to carry several antivirals through preclinical development. Some of the compounds that are in animal tests at this time exhibit very broad activities against a wide range of positive-strand ssRNA viruses and might be of use in case of future zoonotic transmissions of RNA viruses to humans. The structural studies also allow us to predict the emergence of resistance mutations that would undoubtedly arise during wide-spread application of the new antivirals and suggest ways to cope with them by modifying the inhibitors.

[1] Steuber H., Hilgenfeld R. Curr. Top. Med. Chem., 2010, 23, 323.
[2] Chen S. et al. Protein & Cell, 2010, 1, 59. [3] Verschueren K.H.G. et al., Chem. Biol., 2008, 15, 597. [4] Al-Gharabli S.I. et al., ChemBioChem, 2006, 7, 1048. [5] Tan J. et al., J. Mol. Biol. 2005, 354, 25. [6] Anand K. et al. Science, 2003, 300, 1763. [7] Yang H. et al., Proc. Natl. Acad. Sci. USA, 2003, 100, 13190. [8] Mesters JR et al., Curr. Opin. Struct. Biol., 2006, 16, 776. [9] Schmidt M.F. et al., Angew. Chem. Int. Ed. Engl., 2008, 47, 3275.

## Keywords: Antiviral Drug Design, Viral Proteases, Fragment Screening

## FA1-MS12-T06

Incorporation of the basal pilin FctB into the pilus of *Streptococcus pyogenes*. <u>Christian Linke<sup>a,c</sup></u>, Paul G. Young<sup>a,c</sup>, Richard D. Bunker<sup>a,c</sup>, Martin Middleditch<sup>a,c</sup>, Tom T. Caradoc-Davies<sup>d</sup>, Thomas Proft<sup>b,c</sup>, Edward N. Baker<sup>a,c</sup>. <sup>a</sup>School of Biological Sciences and <sup>b</sup>School of Medical Sciences, University of Auckland, New Zealand. <sup>c</sup>Maurice-Wilkins-Centre for Molecular Biodiscovery, Auckland, New Zealand. <sup>d</sup>Australian Synchrotron, Clayton, Victoria, Australia. E-mail: <u>clin180@aucklanduni.ac.nz</u>

Many pathogenic and commensal bacteria express pili, long hair-like protein polymers that enable adhesion to host tissues. The pili of Gram-positive bacteria are unique as they are assembled by specialised transpeptidases (sortases) through covalent linkages. In the common human pathogen Streptococcus pyogenes, the major pilin FctA forms the polymeric pilus fibre, while the minor pilin Cpa is the adhesin found at the pilus tip. The pilus is anchored to the streptococcal cell wall through the basal pilin FctB. Here, we present the crystal structure of FctB at 1.9 Å resolution. It reveals an immunoglobulin (Ig)-like N-terminal domain with an extended proline-rich tail. Surprisingly, the Ig-like domain is structurally homologous to the N-terminal domain of the major pilin FctA. This led to the identification of a conserved lysine residue as the putative site of covalent linkage to the pilus assembly, which we confirmed by mass spectrometry analysis of complete S. pyogenes pili. The C-terminal tail