Report from Plasma Proteome Project

REPORT ON THE HUPO HUMAN PLASMA PROTEOME PROJECT (HPPP) FOR 2009-2010

Co-Chairs: Gil Omenn (USA), Ruedi Aebersold (Switzerland), Mark Baker (Australia)

The Plasma Proteome Project met jointly with the Cardiovascular Initiative in Toronto, as an experiment in combining the interests of two initiatives. Jennifer van Eijk, MingMing Ning, and Peipei Ping organized the CV component of the program, and Gil Omenn, Ruedi Aebersold, and Mark Baker organized the Plasma component.

Terry Farrah of the Institute for Systems Biology in Seattle presented a major update on the Human Plasma PeptideAtlas. Eric Deutsch of ISB, Henning Hermjakob and Lennart Martens of EBI, and Gil Omenn of Michigan (absent due to cardiac surgery) highlighted the ProteomeXchange scheme for submission of well-annotated datasets to EBI-PRIDE, Tranche-ProteomeCommons, ISB-PeptideAtlas, and NCBI-Peptidome. The HPPP is in the lead in demonstrating this process for all HUPO initiatives and for the wider proteomics community. Bernd Wollscheid of ETH Zurich presented the background of N-glycosite proteome analysis and plans for an MRM Atlas. An ABI QTRAP 4000 and an Agilent CHIP QTOF platform were used to generate SRM assays for >5000 human N-glycosites, corresponding to specific peptides selected from experimental data, Unipep, and PeptideAtlas databases.

Michael Kuzyk of the University of Victoria - Genome BC Proteomics Centre in British Columbia, Canada, discussed an SRM/MRM study of 45 cardiovascular biomarker protein candidates analyzed in plasma from 60 patients with or without coronary artery disease; a set of 5 biomarkers differentiated the patients with and without CAD with up to 90% accuracy.

Discussion centered on these points: there is notable progress in enabling MS technologies and bioinformatics workflows, including studies of stroke, coronary heart disease, and myocardial dysfunction; the field is moving from discovery-driven workflows towards targeted experiments; and the SRM Atlas will be a good starting point for the community to download sets of transitions for proteins of interest in order to get a head start for their SRM measurements in clinical samples.

The plasma represents the final common pathway from organ proteomes as we seek to discover, confirm, validate, and utilize protein signatures and protein differential expression as biomarkers for diagnosis and prognosis. Extensive progress has been made during the past year in further updating the Human Plasma PeptideAtlas, with approximately 2000 protein IDs. A detailed set of thresholds has been devised at PeptideAtlas with increasing levels of stringency in terms decreasing likelihood of redundancy. Different thresholds are appropriate for comparisons with various external datasets. That work and the PeptideAtlas-2010 will be presented by Eric Deutsch in Sydney and is the subject of a manuscript expected to be submitted shortly.
As noted at the Toronto PPP/CVI Workshop due in part to the challenges of dynamic range and the dominance by highly abundant proteins in plasma, much attention has now been placed on targeted proteomics using selective or multiple reaction monitoring (SRM/MRM) of proteotypic tryptic peptides. An MRM Atlas is under development, led internationally by Ruedi Aebersold of ETH Zurich and Rob Moritz of ISB Seattle. Bernd Wollscheid of ETH will present major progress as part of the HPPP session on Wednesday 22 September of HUPO 2010 in Sydney. MRM adds the important dimension of absolute quantitation and avoids the dominance of abundant proteins.

As noted by the HUPO Human Proteome Project Working Group, the HPPP is a significant part of the foundation created by the HUPO Initiatives for the building of the Human Proteome Project.

There was one publication during 2009-2010 from the HPPP: