

(Japanese tumor regression grade 2 or more). Post-CRT SUVmax in the responders was significantly lower than that of non-responders (median value 2.9 vs 6.2, respectively). Five-year overall survival was significantly better in patients with lower Post-CRT SUVmax (88% vs 47%, respectively, $p=0.036$). Five-year local re-recurrence free survival was significantly better in patients with higher SUVRR (80% vs 24%, respectively, $p=0.035$). **Conclusion:** Metabolic response assessed by PET/CT is useful for predicting tumor response and prognosis. The response might be utilized for post-operative adjuvant chemotherapy.

PP 9

Men and women display different proteomic diagnostic profiles in non small cell lung cancer

E. Izbicka, R. Streeper, A. Diaz, D. Campos, J. Michalek, C. Loudon, T. Long, S. Baek, R. Mussman. *CPC Ltd., San Antonio, USA*

Background: Plasma biomarker-based screening for lung cancer could provide substantial survival benefits in properly targeted high-risk populations.

Materials and Methods: Fifty-nine circulating proteins were analyzed using multiplexed immunoassays in plasma of patients diagnosed with non-small cell lung cancer (NSCLC; 245 men, 114 women), asthma (AST; 67 men, 112 women) and normal controls (NOR; 122 men, 165 women). Samples were split randomly into training ($N=402$) and test ($N=389$) data sets. A support vector machine (SVM) was used to identify discriminatory biomarkers in NSCLC and AST taking into account patients' gender. Mass spectrometry (MS) followed by data analysis using Mascot software was employed for biomarker discovery; validation of select biomarkers was achieved by immunodetection of target proteins in plasma. Pathway analysis was applied to characterize pathology- and gender-specific patterns of biomarker expression.

Results: We developed seven SVM models that classified subjects to NSCLC, AST or NOR for all 59 markers or subsets thereof, for both genders or single gender only, and for both pathologies and NOR or NSCLC and NOR only. When all biomarkers and genders were accounted for, SVM classified subjects to NSCLC, AST with an accuracy of 0.94 (SE: 0.012). Restricting to NSCLC versus NOR produced 4 markers [EGF, sCD40 ligand, IL-8 and MMP-8; sensitivity (SE) 0.93 (0.014), specificity (SP) 0.87 (0.02)]. Best subset of 5 variables for men (EGF, IL-8, sFAS, MMP-9 and PAI-1) and 3 variables for women (EGF, sCD40 ligand, IL-8) yielded SE and SP of 1 (0). MS identified 11 differentially expressed proteins including 3 putative gene products and yet unnamed proteins, a protein corresponding to chromosome X open reading frame 38, and several known proteins (syntaxin 11, cAMP-specific, rolipram-insensitive phosphodiesterase 7B, and interleukin-25), whose presence was independently confirmed by immunoblotting. Diagnostic biomarkers are products of genes residing on multiple chromosomes and are not limited to sex chromosomes.

Conclusion: The NSCLC-specific biomarkers and combinations thereof identified in this study warrant additional clinical validation to determine their role in screening targeted high-risk populations. The novel method for data mining is widely applicable to development of test kits for detecting biomarkers and combinations of biomarkers.

PP 88

C4.4A as a biomarker for poor prognosis in non-small cell lung cancer patients with adenocarcinomas

B. Jacobsen, T. Muley, M. Meister, H. Dienemann, I.J. Christensen, E. Santoni-Rugiu, O.D. Lærum, M. Ploug. *Finsen Laboratory, Copenhagen University Hospital, Copenhagen N, Denmark*

Background: Lung cancer is the most common cancer form in the world with a 5-year survival rate of only 15%. It is consequently relevant to search for and characterize new prognostic and predictive factors, providing a better basis for treatment decisions in this disease, ultimately leading to higher patient survival. The glycolipid-anchored membrane protein C4.4A, which is a structural homolog of the urokinase-type plasminogen activator receptor, is such a potential candidate. C4.4A is absent in the normal healthy lung, but it is induced in early precursor lesions of non-small cell lung cancer (NSCLC).

Materials and Methods: In the present study, we have undertaken an immunohistochemical, retrospective study on the expression of C4.4A in 229 cases of NSCLC. For each patient, one tissue section from the periphery and one from the center of the tumor were stained with our well-characterized polyclonal anti-C4.4A antibody. C4.4A levels were scored semi-quantitatively for intensity and frequency of positive tumor cells (range 0–16) and statistically correlated to survival.

Results: Expression of C4.4A was more pronounced in squamous cell carcinomas (SCC) compared to adenocarcinomas (AC), with median tumor center scores of 8.0 and 1.3, respectively. Consequently, statistical analysis of survival was performed separately for 88 AC and 104 SCC patients.

In addition to pathological stage, C4.4A score for the tumor center was a highly significant prognostic factor in the AC group both in univariate (p -value = 0.004; Hazard ratio (95% CI) = 1.44 (1.12–1.85)) and multivariate analysis (p -value = 0.0005; Hazard ratio (95% CI) = 1.65 (1.24–2.19)), demonstrating decreasing survival with increasing score. Only pathological stage was significant for the SCC group. These results consolidate earlier observations, now in a larger and independent patient cohort.

Conclusion: High expression of C4.4A is a significant, independent prognostic factor in AC of the lung and is also expressed in a fraction of atypical adenomatous hyperplasias, the putative precursor lesion of this histological subtype. Although the TNM classification still represents the gold standard for the management of NSCLC patients, C4.4A has a potential clinical value as a prognostic marker in pulmonary AC, which might be useful e.g. in decision-making regarding adjuvant radio- or chemotherapy in early stage patients.

PP 38

A fully automated molecular diagnostic system capable of point-of-care for personalized cancer treatment

S. Kitano, J. Myers, M. Amano, M. Ratain, W. Liu. *Life Science Research Laboratory, Technical Research Institute, Toppan Printing Co., Ltd., Kisarazu, Japan*

Background: KRAS, BRAF and PIK3CA mutations are strong predictors for efficacy of molecularly targeted agents such as cetuximab and panitumumab in metastatic colorectal cancer (mCRC). For mutation analysis, the current methods are costly, time-consuming, and not commonly available to clinicians. We have developed a novel, simple, sensitive and fully automated DNA mutation detection system (Toppan Genetic Analyzer, TGA) based on the Invader Plus technology for molecular diagnostics. This system includes the DNA extraction process from homogenized tissue sample. Here we report the results of comparison study between our detection system and direct sequencing (DS) in the detection of KRAS, BRAF and PIK3CA mutations. The effect of DNA purification/extraction process on mutation detection was also compared between the TGA system and the use of commercial kits.

Materials and Methods: Detection of KRAS, BRAF and PIK3CA mutations in mCRC samples were conducted by TGA and DS in a double-blind manner. DNA was extracted from a slice of either frozen tissue ($n=89$) or formalin-fixed and paraffin-embedded (FFPE) tissue ($n=70$) by using QIAamp DNA Micro Kit and EPICENTRE QuickExtract kit, respectively, and then used for TGA and DS experiments. For automated DNA extraction and mutation detection by TGA, a small slice (<1 mg) of frozen tissue ($n=5$) was homogenized in a glass homogenizer. The supernatant was then transferred to TGA for mutation detection.

Results: In the experiments with DNA extracted by commercial kit, all mutations ($n=41$ among frozen and 27 among FFPE samples) detected by DS were also successfully (100%) detected by the TGA. However, 8 frozen and 10 FFPE samples detected as wild-type in the DS analysis were shown as mutants in the TGA analysis. In the experiment testing for the automated DNA extraction and mutation analysis, TGA detected all mutations directly compared to the use of kit-extracted DNA samples. The fully automated reaction can be finished in 80 min.

Conclusion: We have developed a novel fully automated mutation detection system. Our data suggest that this system has the same accuracy as the DS but a higher sensitivity in mutation analysis. The system also has an excellent capacity of mutation detection in both frozen and FFPE samples. Meanwhile, TGA can rapidly detect mutations with simply crashed small amount of frozen tissue in a fully automated mode. These features highlight the great potential of our system for molecular diagnosis in personalized cancer treatment at the point of care.

PP 71

'Other' (non-activating, non-T790M) EGFR mutations and their clinical implications collected from various Tarceva trials in NSCLC

B. Klughammer, O. Spleiss, F. Hoffmann-La Roche Ltd., Basel, Switzerland

Background: EGFR activating mutations (exon 19 in-frame deletions and exon 21 point mutation L858R) in patients with NSCLC have been established as selection marker for 1st line treatment with TKIs like erlotinib and gefitinib (Azzoli et al 2009). The T790M mutation is characterized as resistance mutation (Pao et al. 2005). In many studies only EGFR activating and resistance mutations are assessed, e.g. OPTIMAL, EURLAC, IPASS. However, 'other' EGFR mutations could also contribute to clinical benefit from TKI treatment (Xu et al. 2009).

Materials and Methods: Exons 18–21 of the EGFR gene were amplified by polymerase chain reaction (PCR) using nested primers, and multiple independent products were directly sequenced on both strands. Data were collected from various Tarceva trials (BO18192 SATURN (Cappuzzo et al.