In vivo dynamic contrast enhanced MRI and histopathological assessment of tumor angiogenesis in luminal-like and basal-like breast cancer xenografts

E. M. Huuse1, S. A. Moestue1, T. F. Bathen1, A. Bofin1, G. M. Mælandsmo2, L. A. Akslen1, O. Engebraaten3,4, and I. S. Gribbestad1

1Department of Circulation and Medical Imaging, Norwegian University of Science and Technology (NTNU), Trondheim, Norway, 2Department of Laboratory Medicine, Children's and Women's Health, NTNU, Trondheim, Norway, 3Department of Oncology and Department of Tumor Biology, Oslo University Hospital, Oslo, Norway, 4The Gade Institute, Section for Pathology, University of Bergen, Bergen, Norway, 5Institute for Clinical Medicine, University of Oslo, Oslo, Norway

Introduction
Molecular sub-classification of breast cancer based on gene expression pattern represents clinically distinct patient groups (1). Two new breast cancer xenograft models, reflecting the luminal-like (MAS98.06, ER, low growth rate) and basal-like (MAS98.12, ER, high growth rate) subgroups, have recently been established by direct transplantation of primary tumor tissue (2). The purpose of this study was to investigate differences in tumor vasculature and angiogenesis, and the effect of tumor volume using dynamic contrast enhanced (DCE) MRI and histopathological measures in these two model systems.

Experimental
Tumors (n=27) from each model were orthotopically grafted in female BalbC nu/nu mice(3) to satisfying predefined volume criteria for medium-sized (volume = 450 mm^3) and large tumors (volume > 1400 mm^3), respectively (luminal-like: medium-sized tumors, n=6, 9 weeks of growth; large; n=7, 15-22 weeks of growth; basal-like: medium-sized; n=7, 8 weeks of growth; large: n=7, 11 weeks of growth, volume distributions are shown in Figure A) were examined using a BRUKER Biospec 7T animal scanner. Precontrast T1-values were measured using a series of T1-weighted spin-echo images, followed by a DCE-MRI sequence with 200 images, 4.8 sec temporal resolution and a voxel size of 0.32 x 0.32 x 0.6 mm'. During the 10th repetition, a dose of 0.3 mmol/kg gadodiamide (Omniscan TM, GE Healthcare, Oslo, Norway) was injected intravenously (4 sec). T1-weighted high resolution images were obtained after DCE-MRI (Figure A). Tofts model (4) was used to estimate Ktrans, ve, and vp for each voxel, and median values for each tumor were calculated. After MRI examination, a dose of 60 mg/kg body weight of pimonidazole (Hypoxyprobe-1, Massachusetts, USA) was injected intraperitoneally 30-45 minutes before tumors were harvested. Tumors were stained for both pimonidazole and dual stained for CD34 (Abcam, Cambridge, UK) and Ki67 (Dako, Glostrup, Denmark). Hypoxic fraction, microvessel density (MVD) and vascular proliferation index (number of dual stained vessels per area/MVD) were measured. Independent samples t-test was used for comparison of MRI extracted parameters and histopathological measures between the groups with different tumour volume and between the two models, and Spearman’s correlation between the MRI and histopathological parameters was calculated. The level of statistical significance was defined as p<0.05.

Results
The DCE-MRI results showed a significantly higher Ktrans in basal-like tumors compared to the luminal-like tumors. This was valid for both large and medium sized tumors (Figure B). ve were significantly higher in basal-like tumors than in luminal-like tumors. For basal-like tumors there was no difference in ve between medium-sized and large tumors, but a significantly lower vp in large compared to medium-sized luminal-like tumors (Figure D). VPI was significantly lower and hypoxia significant higher in luminal-like tumors compared to basal-like tumors. The ve and MVD were similar for all the tumors. The results showed a positive correlation between vp and MVD (p=0.51, P < .05) and a negative correlation between vp and hypoxia (p=-0.54, P < .05), and between Ktrans and hypoxia (p=-0.77, P < .05).

Discussion
Basal-like tumors are more aggressive than luminal-like tumors, which is in accordance with the higher Ktrans found for both medium-sized and large xenografts. The significant higher VPI in the basal-like tumors suggest a higher angiogenic activity in these tumors. The low vp combined with a high fraction of hypoxia in the large luminal-like tumors suggest a low angiogenic activity in these tumors that might contribute to the low growth rate of the luminal-like tumors. The significant correlation between in vivo measured vp and the ex vivo measurements of vasculature MVD combined with the correlation between the in vivo measured parameters Ktrans and vp, and the ex vivo measured hypoxia fractions, suggest that DCE-MRI provide in vivo assessment of tumor microenvironment in xenografts with different gene expression pattern.


Figure: A) The distribution of tumor volume in each animal group and high resolution post contrast images from a medium-sized tumor from each model, B-D) boxplots showing the distribution of Ktrans, ve and vp in each tumor group and parametric maps from two medium-sized tumors, and E) fraction of hypoxia in each groups and images of the histopathological staining. All images are of medium-sized tumors.