Conclusion: Despite the numerous publications on focal therapy in prostate cancer, primary FRT is largely unexplored. Radiotherapy appears to be particularly suitable as a focal approach, since it has an established biological basis, known tumouricidal activity, possibility of dose differentiation, large availability of high-precision dose delivery techniques, limited or no invasiveness and familiarity to radiation oncologists and urologists. However, when applied as primary FRT, its use remains investigational since numerous questions remain unmet: consensus on the initial diagnostic tools, the optimization of technical parameters for therapy delivery, follow-up exams and scheduling, tumour control and toxicity profile, response evaluation and failure definition, salvage therapy and cost-benefit.

EP-1381

ADC of prostate tumour and normal tissue during radiotherapy after neoadjuvant hormone therapy

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Purpose or Objective: Changes in prostate and tumour ADC values during radiotherapy (RT) may aid prediction of response to treatment. Intermediate and high risk patients are likely to receive neoadjuvant hormone therapy (NA-HT), causing reduction in prostate and tumour volume and changes in ADC values. It is unclear how this affects further ADC changes during subsequent treatment. We assessed ADC values in prostate tumour and normal tissue during RT after NA-HT.

Material and Methods: Fifteen patients with T2b disease who were due to receive RT (60 Gy in 20 fractions) were recruited after 3 months of NA-HT. Patients underwent three 1.5 T MRI examinations: post NA-HT (one week prior to RT), at the end of the third week of treatment, and eight weeks after RT completion. The imaging protocol included T2 weighted and diffusion weighted imaging, acquired using the cardiac coil (EPI with TR/TE 8000/70 ms, b = 100, 400, 800 s/mm²). ADC maps were processed offline (ADCmap for Osirix). Normal central gland (CG), peripheral zone (PZ) and tumour were outlined on T2w images by a radiologist expert in prostate MRI, with pre-NA-HT imaging (T2w and DWI) available in 12 patients to aid identification. If disease was not clearly visible, clinical findings and biopsy results were used to aid delineation. CG, PZ and tumour regions were transferred to the ADC maps and median values extracted along with interquartile ranges. A Mann-Whitney U test was used to analyse differences between tumour and normal tissue regions at the three time points.

Results: 13 patients completed all scans, 2 patients missed 1 and 2 scans respectively. After NA-HT, there was a significant difference between median tumour and PZ (p=0.009) and tumour and CG (p=0.002) (median values plotted in figure 1). At the other time points, there was no difference between tumour and normal tissue ADCs.

Conclusion: The ADC values display a similar pattern to that seen in previous studies for patients receiving RT alone. The difference between tumour and normal tissue was smaller at baseline than has been seen in other work without NA-HT. This may be due to a reduction in normal tissue ADC during induction therapy, whilst tumour ADC values could have increased due to tumour shrinkage. Variation in imaging protocol for ADC measurement compared to previous studies may also play a role. The reduced magnitude of changes in tumour ADC seen during RT after NA-HT may make its use as a predictive tool for treatment response more challenging in this group of patients.

EP-1382

PET/CT and MRI guided salvage radiotherapy after prostatectomy for prostate cancer

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Purpose or Objective: At the time of a biochemical recurrence after prostatectomy it is important to distinguish patients who have a local recurrence from patients with distant metastasis. PET/CT and MRI are important imaging modalities that can be used in this scenario. The purpose of this study was to investigate the outcomes and toxicities of patients in a large single-institution cohort treated with salvage radiotherapy (sRT) and dose escalation up to 72 Gy. Boost planning was based on MRI or PET/CT.

Material and Methods: From 2008 to 2012 290 patients who received sRT were included into the analysis. Patients with a PSA > 1 ng/ml or a PSA doubling time > 3 months received a Choline PET/CT before the start of radiotherapy. Additionally, in most patients MRI of the pelvis was conducted. If there was a macroscopic tumor recurrence, defined as local recurrence in the prostate bed in MRI or PET tracer uptake, radiation therapy to the prostatic bed was...