

comparison with pts treated with the same chemotherapy alone until PD suggests that it may be detrimental to stop chemotherapy after 6 cycles if disease did not progress.

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POSTER

Epstein-Barr virus quantification and aberrant host DNA methylation pattern as marker for nasopharyngeal carcinoma in non-invasive nasopharyngeal brushings

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Background: Nasopharyngeal carcinoma (NPC) is the most prevalent head and neck cancer in Indonesia. WHO type III, representing the majority of NPC, is 100% related to Epstein-Barr (EBV) infection. The viral DNA load is elevated in nasopharynx (NP) of most patient at diagnosis. A NP brush was used for in-situ sampling, allowing simpler and less invasive NPC diagnosis. We have shown EBV-DNA load as promising new diagnostic method and use it to screen NPC cases in high risk population. A growing evidence demonstrates that aberrant methylation in gene promoter is important in inactivating tumor suppressor gene (TSG) in NPC. This study aimed to quantify EBV-DNA load and determine methylation status of multiple TSGs in NPC, high risk individuals, and healthy EBV-carriers to evaluate whether methylation pattern may have additional value to identify early carcinogenic events.

Methods: NP brushing was taken from NPC, high risk patients presenting chronic problems in head and neck area, and normal EBV-carriers. Paraffin tissue of NPC patient was also included and subjected for DNA isolation in order to verify detection rate of methylation in brushing DNA. EBV-DNA load was measured using a quantitative real time PCR. DNA was modified using bisulfite treatment and amplified by methylation-specific PCR. Seven tumor suppressor genes were included (DAPK, CADM1, p16, RASSF1A, CHFR, RIZ1, and DLC1).

Results: All NPC patients showed elevated EBV-DNA and high frequency of methylated genes (DAPK 69.6%, CADM1 71.4%, p16 68.1%, RASSF1A 73.5%, CHFR 65.9%, RIZ1 41.7%, and DLC1 58.7%). Most of paraffin and brushing DNA revealed a concordance result of methylation status. The high risk individuals, who also demonstrated high EBV-DNA load, showed high frequency of methylated genes of DAPK (76.9%), CADM1 (61.5%), and DLC1 (61.5%), but low or undetected methylated genes of p16, RASSF1A, CHFR, and RIZ1. Healthy individuals showed low DNA load but similar methylation pattern as high risk population.

Conclusion: These results suggest that EBV infection and promoter hypermethylation might serve as useful markers to screen early NPC. At the time a prospective analysis in high risk group using non-invasive brushing samples to identify early stage NPC is in progress. In Indonesian normals, much abnormal methylation on certain TSGs probably reflects exposure to co-carcinogens in environment and food.

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POSTER

Pattern of locoregional failure after tomotherapy in head and neck cancer

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Background: Helical tomotherapy is a new radiation device delivering a highly conformal dose from a rotational gantry resulting in a more uniform target dose and better avoidance of organs at risk. Treatment failure patterns of head and neck cancer treated with helical tomotherapy and the adequacy of the target volume definitions and delivery techniques currently used were analysed.

Materials and Methods: Between June 2005 and March 2008, 76 consecutive patients with biopsy proven head and neck cancer were treated with helical tomotherapy (Hi-Art TomoTherapy[®], Madison, Wisconsin, USA) at the UZ Brussel. For patients with local or regional failure, the volume of failure (Vf) was determined on one or more diagnostic tools as computerized tomography (CT), magnetic resonance imaging or positron emission tomography obtained at the time of failure. The Vf is then contoured with Co-registration of the failure image (Vf) and the initial planning CT was performed. The failures were categorized as local or regional. The dose of radiation received by failure was calculated and analyzed using dose-volume histograms (DVHs) and accordingly it is classified as 1) In-field (InF): in which 95% or more of Vf was within the 95% isodose, 2) Marginal (MF), if 20% to 95% of Vf was within the 95% isodose, or 3) Outfield (OutF) if less than 20% of Vf was inside the 95% isodose. The mean, minimum and maximum doses received by each failure volume were displayed.

Results: Median follow up time was 14.8 months (3.5–38.8). Three-years overall survival, disease free survival and locoregional control were 69%, 47% and 59%, respectively. Twelve patients showed locoregional failure, 5 were local, 6 were regional and one showed both local and regional failure. With DVHs analysis, InF, MF and OutF were 9, 3 and 1, respectively. All MF had a history of surgery before radiotherapy.

Conclusions: Target definition and coverage were adequate. The majority of locoregional failures were InF i.e. in the high dose regions. Future work on dose escalation to the highest risk regions is recommended. Special consideration for surgically manipulated patients must be taken in volume selections and coverage.

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POSTER

Clinical results and prognostic factors in radiotherapy for early glottic squamous cell carcinoma

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Background: The purpose of this study is to determine the prognostic factors for local control in T1a, T1b and T2N0M0 glottic squamous cell carcinomas.

Material and Methods: Data from 249 patients with T1–2N0M0 (T1a: 115, T1b: 48, T2: 86) Stage I–II glottic carcinomas, who were treated with definitive radiotherapy during 1976 to 2002 were analyzed retrospectively. Age, source, total dose, field size, overall treatment time, average fraction size, fractionation regimen, chemotherapy and etc. were set as variables in multivariate analysis.

Results: The 5-year local control rates (LCR) were 92%, 85% and 83% for patients with T1a, T1b and T2 glottic carcinomas, respectively. Only total radiation dose ($p=0.048$) was a significant prognostic factor for local control in multivariate analysis of T1b glottic carcinoma. Local control in the higher total dose group was better than that in the lower total dose group (5-year LCRs were 100% and 76% for the group of ≥ 66 Gy and the group of ≤ 66 Gy, respectively, $p=0.024$, logrank test). None of the treatment parameters were shown to be significant prognostic factors in multivariate analysis of T1a glottic carcinoma. In the analysis of T2 glottic carcinoma, OTT (overall treatment time of radiotherapy) ($P=0.0003$) and Total dose ($P=0.0036$) were the significant prognostic factors on local control in multivariate analysis. Higher total dose group (≥ 67 Gy vs. < 67 Gy) showed favorable prognosis (5-year LCR: 91% vs. 60%, respectively, $P=0.0013$; logrank test). And the shorter OTT group (≤ 54 days vs. > 54 days) showed favorable prognosis (5-year LCR: 87% vs. 71%, respectively, $P=0.023$).

Conclusions: Radiotherapy with a total dose of ≥ 66 Gy seemed to be required for local control in T1b glottic carcinoma. No significant benefit of total radiation dose > 64 Gy was shown in the analysis of T1a glottic carcinoma. Radiotherapy total dose of ≥ 67 Gy delivered with shorter period is required for T2 glottic cancer. The fractionation regimens of accelerated hyperfractionation is more effective than conventional fractionation in terms of shortening OTT and delivering high total dose with acceptable toxicity.

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POSTER

Prediction of clinical radiation induced toxicity through study of radiation induced apoptosis in peripheral blood lymphocytes (PBLs)

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Background: To analyze radiation induced apoptosis in PBLs of patients with head and neck (H&N) cancer, the application of a semilogarithmic model RID (Radiation Induced cell Death) = $\beta \ln(\text{Gy}) + \alpha$, to assess the association among the defined constants of this model and its utility as a prediction model for toxicity in patients treated with radiation therapy.

Material and Methods: A total of 79 patients with H&N cancer treated with radiation therapy, with or without surgery and chemotherapy were included. PBLs were obtained from peripheral blood samples using density gradient centrifugation (Ficoll Hipaque). Apoptosis was assessed by Annexin V and Propidium Iodide (IP) staining. Triple analysis at doses of 0, 1, 2, and 8 Gy were performed in all patients after 24 hours. Clinical toxicity was assessed by the RTG classification.

Results: RID was increased by the dose of radiation administered. α (initial value at x axis) and β (apoptosis increase due radiation dose–slope of the curve) constants, defined in the model, were statistically associated. β was associated with radiation induced toxicity, such as grade III or higher xerostomy bivariate ($p=0.035$) and multivariate analysis ($p=0.034$; EXP (B) 2.553, 95% CI (1.074–6.070)).

Conclusions: Radiation sensitivity of peripheral blood lymphocytes can be estimated using the Annexin IP staining to assess radiation induced apoptosis. The later adjusts to the α/β semilogarithmic model and allows to