Proffered Papers

6560 POSTER

Prognostic factors affecting survival on pretreated patients with locally advanced or metastatic non-small cell lung cancer (NSCLC) — Subgroup analysis in a randomized Ph II study of pemetrexed  $500\,\text{mg/m}^2$  and  $1000\,\text{mg/m}^2$ 

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Background: The regimen with pemetrexed 500 mg/m<sup>2</sup> is used for patients with NSCLC in the 2nd or 3rd line settings in worldwide. In a Ph III study comparing pemetrexed with docetaxel in pretreated patients with NSCLC, good PS, early clinical stage, and more than 3 months from the last chemotherapy were identified as longer survival factors in pemetrexed arm. Results from latest randomized Phase II study with 2 dosages of pemetrexed 500 mg/m<sup>2</sup> (Pem 500) and 1000 mg/m<sup>2</sup> (Pem 1000) showed favorable response rates of 18.5% and 14.8%, respectively. We provide the survival outcome and examine the factors affecting longer survival. Materials and Methods: Pts with PS 0-2, measurable, stage III/IV NSCLC, 1 or 2 previous chemotherapy regimens, were randomized to either Pem 500 or Pem 1000 on day 1 of a 21-day schedule. The planned total sample size was 240 pts. The survival analysis was performed for both arms. Results: From October 2004 to October 2006, 244 pts were enrolled at 28 centers, 226 pts were randomized and treated, and 216 pts (Pem 500/Pem 1000: 108/108 pts) were evaluable for the survival analysis. Baseline patient characteristics (Pem 500/Pem 1000) were: Male 63%/64%; median age 62/62 years (range: 37-74/26-74); PS 1,2 61%/67%; Stage IV 81%/80%. One-year survival rates were 59.2%/53.7% and MST were 15.7M/12.6M, respectively. Among the variables examined for the Cox regression analysis (age, gender, histology, time from the last chemotherapy, dosage, with or without a prior platinum regimen, performance status, number of regimen, and clinical stage), factors showed a statistical significance were gender (male/female: HR 2.14), histology (non-adenocarcinoma/adenocarcinoma: HR 2.13), time from the last chemotherapy (≥3 months/<3 months: HR 0.56), performance status

**Conclusions:** Female, adenocarcinoma, a longer period from the last chemotherapy, good PS, and early clinical stage were identified as good prognostic factors. High dosage of 1000 mg/m² did not prolong survival longer than that of 500 mg/m², which supports the use of 500 mg/m² of pemetrexed in pretreated patients with NSCLC.

(PS 1, 2/0: HR 2.81), and clinical stage (IV/III: HR 1.81). There is no

difference in two dosages.

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Gated radiotherapy of lung cancer: interfractional changes in tumor volume and position during the treatment course

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**Background:** With the purpose of designing intelligent margins for curative radiotherapy, this study investigated the interfraction variations in tumor size and internal displacement with respiratory gating. The variations were also compared for different set up strategies.

**Methods and Materials:** During the treatment course, the patients underwent 3 respiratory gated CT-scans, equally spaced in time. The tumors were contoured on each CT-scan to evaluate the variation in volume and position. The primary lung tumors and the mediastinal tumors were contoured separately. The positional variations were measured as 3D mobility vectors and related to matching of the scans on the basis of bony landmarks and with skin tattoos.

Results: The median 3D mobility vector for the lung tumors was 0.52 cm for matching performed with bony landmarks and 0.81 cm for matching with skin tattoos. For the mediastinal tumors the corresponding vectors were 0.56 cm and 0.53 cm. The differences between the vectors were significant for the lung tumors. There was a significant reduction in tumor size from the first to the last CT-scan, both for lung (19%) and mediastinal tumors (34%). The interfractional overlap of lung tumors was 80–87% when matched using bony landmarks and 70–76% when matched using skin tattoos. The overlap of the mediastinal tumors were 60–65% and 41–47%, respectively.

**Conclusions:** The tumors varied considerably, regarding both tumor position and tumor volume. The variations in position were dependent on the set up strategy. Set up using skin tattoos was inferior to set up using bony landmarks.

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First-line treatment with vinorelbine (VNR) plus carboplatin (CBDCA) for patients with advanced non-small-cell lung cancer (NSCLC): MAP4/OP18 mRNA expression as marker predictive of response

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Background: Non-small-cell lung cancer (NSCLC) patients (pts) with locally advanced or metastatic disease at the time of diagnosis show marginal response to chemotherapy in terms of tumor shrinkage, time to progression and median survival. MAP4 and stathmin have been previously reported as potential markers of resistance to treatment based on microtubule-destabilizing agents.

Materials: 51 chemonaive pts with stage IIIB (with pleural effusion)-IV NSCLC and ECOG PS 0-1 were accrued at 7 sites between October 2003 and June 2006. Treatment consisted of CBDCA AUC5 IV day 1 plus VNR 25 mg/m² IV, days 1, 8 every 21 days. In this study, we have used quantitative PCR to analyze the expression of MAP4, stathmin, betatubulin III, BRCA1 and ERCC1 using mRNA isolated from peripheral blood samples.

Results: Median age 59 years (range 42-75); males 91%/females 9%; smokers: 72%; adenocarcinoma, 54%/squamous 33%; stage IIIB: 26%, IV: 74%. Median cycles: 3 (1-6). Hematological toxicities (%pts): grade 3/4 neutropenia, 18%/8%; grade 3/4 thrombocytopenia, 16%/6%; grade 3 anemia, 11%. Febrile neutropenia appeared in 2 pts (4%). Nonhematological toxicities (%pts): nausea/vomiting grade 3/4, 18%. Efficacy in evaluable population (n = 33): CR, 0%; PR, 39%; ORR, 66% (95% CI 47-73%); SD, 33%. With a median follow up of 7.2 months, median survival for the whole population was 7.75 (95% CI 6.98-8.51) months (mo), progression free survival 5.8 (95% CI 3.7-8.2) mo, 1-year survival 28.5%. In a preliminary set, 46 patients with stage IIIB and IV were analyzed. Lower levels of MAP4/OP18 mRNA expression are statistically associated with a response to vinorelbine-based treatment (p = 0.029). This significant relationship is maintained in a second analysis after 3rd cycle of treatment (p = 0.032). Higher levels of MAP4/OP18 were associated with a lower TTP (p = 0.05).

**Conclusions:** These preliminary results suggest that the ratio MAP4/OP18 may be a good predictor of response for NSCLC patients treated with vinorelbine-based chemotherapy.

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High-dose 130-nanometer albumin-bound paclitaxel in combination with carboplatin as first-line therapy in advanced non-small cell lung cancer

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Background: Carboplatin AUC of 6 mg/min·ml (C6) plus solvent-based (SB) paclitaxel (Taxol<sup>®</sup>) is a commonly used regimen in first line patients (pts) with advanced NSCLC resulting in response rates of 17–32%, median progression-free survivals (PFS) and survivals of 4–5 and 8–10 months, respectively. In these trials a median of 4 cycles were typically given and the rates of Grade (gr) 3 and 4 neutropenia and sensory neuropathy ranged between 35–63% and 3–15% respectively. We previously reported the results of solvent-free nab-paclitaxel (Abraxane<sup>®</sup>) 225–340 mg/m² and C6 both administered on day 1 every 3 weeks (q3w) in first line pts with NSCLC