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Everolimus and Transarterial Embolization (TAE) for Hepatic Metastasis in Neuroendocrine Tumors (NET)

Aman Chauhan¹, Tim Waits¹,2; Riham El Khouli¹,2; Steven Krohmer¹,2; Lowell Anthony¹,2

¹University of Kentucky; ²Markey Cancer Center

BACKGROUND: TAE of hepatic metastases is effective loco-regional therapy for neuroendocrine tumor (NET) patients. Transarterial chemotherapy (TACE) and transarterial radiotherapy (TARE) are other options in controlling hepatic metastases but deliver cytotoxic agents only at the time of the procedure. Systemic therapies such as everolimus or sunitinib are often held 2-4 weeks prior to and after the procedure. We hypothesize that concurrent oral treatment with everolimus and transarterial hepatic embolization is safe.

METHODS: Medical records of all NET patients (pts) treated at the Markey Cancer Center between July 2016 to July 2017 with concurrent transarterial embolization and everolimus were reviewed.

RESULTS: Total number of pts eligible for analysis was 15. Mean age of study cohort was 57.5 years (31-77). Seven pts were small bowel, 4 were pancreatic and remaining 4 were unknown NET primaries. Nine patients had grade 1 (Ki 67 < 2%) and 6 patients had grade 2 NETs (Ki-67 between 2-20%). Mean duration of treatment with everolimus prior to embolization was 3.8 months and mean dose was 10 mg orally every other day. Mean days of hospitalization were 1.46. Eleven (73%) pts were discharged the day after procedure. Ten (67%) pts reported nausea, 9 (60%) reported abdominal pain, 5 (30%) reported emesis. One patient had hematemesis immediately post procedure. Thirteen (87%) pts had no significant change in hemoglobin pre and post procedure (within one numerical point). No patient developed infectious complications.
CONCLUSION: Combining everolimus with TAE is safe and does not result in longer hospitalizations or toxicities greater than that expected from TAE alone.