Mechano growth factor (MGF) expression increased in secondary compared to primary foci in well neuroendocrine neoplasms

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Introduction
Insulin-like growth factor-I (IGF-I): important role in cell proliferation, differentiation, migration, and survival.
Alternative splicing → different IGF-I mRNA transcripts → proteins with different responses in different stimuli as IGF-I: muscle regeneration process + mitogenic and anti-apoptotic role in the pathophysiology of various types of cancer.
Prostate cancer: high grade cancers more autonomous and less sensible in IGF-1 compared to low grade malignancy + IGF-1: limited role in early development of the cancer but central role in cancer progression.

Objectives
The role of MGF in the pathophysiology of neuroendocrine neoplasms (NENs).

Materials and methods
We have used immunohistochemistry in 47 specimens of patients with NENs to show the expression status of MGF isoform.
Proliferation index ki-67 MIB1 (%) was evaluated.
We have studied 8 gastric, 17 pancreatic, 3 appendicenteral, 9 small intestine, 2 Colic and 1 retrococious, 1 gallbladder, 2 lung NENs, 1 well differentiated unknown primary (UPO), 3 poorly differentiated UPO, and 1 other.
We have studied the following subgroups:
1.specimens with negative staining (A), staining 1-10% (B), staining >50% (C)
2.specimens with Ki-67 ≤2% (I), 2-20% (II), >20% (III)

Results
No MGF staining was found in 24 (51%) specimens.
Cytoplasmatic staining was found in 23 specimens: 1% in 6 (26%), 10% in 9 (40%), 50% in 4 (17%), 80% in 1 (4%) and 100% in 3 (13%); focal staining in 15 (32% ≤65%); diffuse in 7 (15% ≤30%) and dot like in 1 (2% ≤5%) specimen.

IGF1Ec isoform F staining was more prevalent in specimens from metastatic foci compared to specimens from primary tumours (p=0.036).

Rate 0%: n=24 1-49%: n=24 >50%: n=8

MGF staining was more prevalent in specimens from metastatic foci (p=0.029).

Conclusions
Our preliminary data suggest that MGF expression may be involved in the pathophysiology of well differentiated NENs.
Further studies will shed light to the exact role of IGF1Ec isoform in the presentation or progress of NENs particularly in secondary sites.

References
Vitale et al 2008 Oncol Rep. 15:1249-56