

Rak neuroendokryny trzustki nieczynny hormonalnie

Kod Orpha: 506075 Kod OMIM:

Opis choroby *

Definicja

A rare neuroendocrine tumor of pancreas characterized by a well-differentiated epithelial pancreatic neuroendocrine neoplasm measuring at least 0.5 cm, without distinct hormonal syndrome. Tumors <0.5 cm are called microadenomas. Microadenomatosis is the multifocal occurrence of microadenomas. Histopathologic examination shows an organoid growth pattern and expression of synaptophysin and chromogranin A on immunohistochemistry. Tumors are often discovered incidentally, or patients may present with symptoms related to local or metastatic tumor spread. Microadenomas are considered benign, while larger tumors may behave in a malignant manner with extrapancreatic spread, metastasis, or recurrence.

Dane

Klasyfikacja

Choroba

Synonimy

Non-functioning PNET
Rak trzustki nieczynny neuroendokrynnie
Non-functioning pancreatic NET
Non-functioning pancreatic neuroendocrine tumor
Non-functioning well-differentiated NEN of pancreas
Non-functioning well-differentiated neuroendocrine neoplasm of pancreas
Non-functioning well-differentiated pancreatic NEN
Non-functioning well-differentiated pancreatic neuroendocrine neoplasm

Kod ORPHA

506075

Kod OMIM

-

Kod ICD10

C25.9

Kod ICD11

-

[*Źródło](#)

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.

Orphanet - internetowa baza danych dotyczących rzadkich chorób i sierochych leków. ©INSERM 1999 -
Dostępna na stronie www.orphanet.pl