

DISEASE:
Triple A syndrome

NAME:	Triple A syndrome
DESCRIPTION:	Triple A syndrome is a very rare multisystem disease characterized by adrenal insufficiency with isolated glucocorticoid deficiency, achalasia, alacrima, autonomic dysfunction and neurodegeneration.
ORPHACODE:	869
SYNONYMS:	2A syndrome 3A syndrome 4A syndrome AAA syndrome Achalasia-addisonianism-alacrima syndrome Adrenal insufficiency-achalasia-alacrima syndrome Allgrove syndrome Double A syndrome Quaternary A syndrome

XREF(S):	Orphanet ICD-10 OMIM OMIM MeSH MeSH
ANALYTE(S):	TRAPPC11 AAAS GMPPA
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