

DISEASE:
Peeling skin syndrome type A

NAME:	Peeling skin syndrome type A
DESCRIPTION:	A noninflammatory form of generalized PSS characterized by white scaling and superficial painless peeling of the skin.
ORPHACODE:	263548
SYNONYMS:	Generalized peeling skin syndrome type A Non-inflammatory generalized peeling skin syndrome type A. Non-inflammatory peeling skin syndrome type A PSS type A
XREF(S):	Orphanet OMIM OMIM ICD-10
ANALYTE(S):	FLG2 CHST8 SERPINB8
CREATED:	13 May 2019 - 01:02
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