

ANALYTE:
SLC6A19

NAME:	solute carrier family 6 member 19
SYMBOL:	SLC6A19
VERSION OF ORPHANET:	2023-06-22 14:14:43
SYNONYMS:	Hartnup disease
XREF(S):	<u>Orphanet</u> <u>Ensembl</u> <u>Genatlas</u> <u>HGNC</u> <u>OMIM</u> <u>Reactome</u> <u>SwissProt</u>
CREATED:	13 May 2019 - 01:01
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