

DISEASE:

Leydig cell hypoplasia due to complete LH resistance

NAME:	Leydig cell hypoplasia due to complete LH resistance
ORPHACODE:	96265
SYNONYMS:	46,XY DSD due to complete LH receptor inactivation 46,XY DSD due to complete LH resistance 46,XY DSD due to complete luteinizing hormone receptor inactivation 46,XY DSD due to complete luteinizing hormone resistance 46,XY disorder of sex development due to complete LH receptor inactivation 46,XY disorder of sex development due to complete LH resistance 46,XY disorder of sex development due to complete luteinizing hormone receptor inactivation 46,XY disorder of sex development due to complete luteinizing hormone resistance Leydig cell hypoplasia due to complete LH receptor inactivation Leydig cell hypoplasia due to complete luteinizing hormone receptor inactivation Leydig cell hypoplasia due to complete luteinizing hormone resistance
XREF(S):	Orphanet ICD-10 OMIM
ANALYTE(S):	LHCGR
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Source URL: <http://gentest.healthdata.be/disease/3429>

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- Centrum Medische Genetica - UZ Brussel VUB

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- luteinizing hormone/choriogonadotropin receptor

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- Premature Ovarian Failure/Insufficiency (32 genes) - VUB

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