

DISEASE:
Primary myelofibrosis

NAME:	Primary myelofibrosis
DESCRIPTION:	A rare myeloproliferative neoplasm characterized by stem-cell derived clonal over proliferation of mature myeloid lineages, such as erythrocytes, leukocytes, and megakaryocytes, with variable degrees of megakaryocyte atypia, associated with reticulin and/or collagen bone marrow fibrosis, osteosclerosis, ineffective erythropoiesis, angiogenesis, extramedullary hematopoiesis, and abnormal cytokine expression.
ORPHACODE:	824
SYNONYMS:	Agnogenic myeloid metaplasia Idiopathic myelofibrosis Myelofibrosis with myeloid metaplasia Osteomyelofibrosis
XREF(S):	Orphanet ICD-10 OMIM
ANALYTE(S):	JAK2 MPL TET2 CALR
CREATED:	13 May 2019 - 01:02

CHANGED:	22 Jun 2023 - 16:14
-----------------	---------------------

Source URL: <http://gentest-acc.healthdata.be/disease/2558>

RELATED CONTENT

Related Genetic Tests

- [Erythrocytoses, polycythémies, thrombocytoses et neutropénies congénitales \(gene panel\)](#)
- [Myeloid neoplasms with germline predisposition \(Hereditary MDS/Acute Leukemia\) \(gene panel\)](#)
- [Myeloid/lymphoid neoplasms with germline predisposition \(gene panel\)](#)
- [« Inherited bone marrow failures syndromes » with or without organ dysfunction](#)

Related Laboratories

- [Centre de Génétique Humaine - CHU Sart-Tilman](#)

Related Analytes

- [calreticulin](#)
- [Janus kinase 2](#)
- [MPL proto-oncogene, thrombopoietin receptor](#)
- [tet methylcytosine dioxygenase 2](#)

Related Gene Panels

- [Erythrocytoses, polycythémies, thrombocytoses congénitales \(gene panel\) - ULG](#)
 - [Hematologic Familiar Forms - ULG](#)
 - [Hereditary Myelodysplastic /Acute Leukemia Predisposition Syndromes \(gene panel\)](#)
-

Source URL: <http://gentest-acc.healthdata.be/disease/2558>