t(18;21)(p11;q11)
Adriana Zamecnikova
Kuwait Cancer Control Center, Department of Hematology, Laboratory of Cancer Genetics, Kuwait; annaadria@yahoo.com
Published in Atlas Database: June 2017
Online updated version: http://AtlasGeneticsOncology.org/Anomalies/t1821p11q11ID1791.html
DOI: 10.4267/2042/68936
This work is licensed under a Creative Commons Attribution-Noncommercial-No Derivative Works 2.0 France Licence. © 2018 Atlas of Genetics and Cytogenetics in Oncology and Haematology

Abstract
Chromosome translocation between the short arm of chromosome 18 and the long arm of chromosome 21 including the t(18;21)(p11;q11) is a rare event, reported only in sporadic cases.

Keywords
Chromosome 18; Chromosome 21; Acute erythroleukemia; AML-M6; Acute lymphoblastic leukemia; Follicular lymphoma.

Disease and pathology

Disease
Acute erythroleukemia (FAB type M6), acute lymphoblastic leukemia (ALL) and follicular lymphoma

Epidemiology
Only 3 cases to date: a 71-years old male diagnosed with acute erythroleukemia (Cigadusa et al., 2003), a female patient with follicular lymphoma (Lestou et al., 2003) and a 27-years old male with B-cell ALL (present case, personal observation).

Prognosis
Unknown (sporadic cases described). The ALL patient relapsed after 7 months of therapy and was alive in the last follow-up 2 years from the diagnosis.

Cytogenetics

Note
Breakpoints on 18p and 21q are difficult to ascertain in suboptimal preparations.

Additional anomalies
Associated with del(5)(q13q31) del(5)(q13q31), monosomy 7, hsr and complex karyotype in the AML case (Cigadusa et al., 2003), del(5)(q15q31), +7, t(14;18)(q32;q21) in the lymphoma case (Lestou et al., 2003) and with homozygous 9p deletion (70% of cells), detected by fluorescence in situ hybridization in the present case.

Variants
Genes involved are unknown.
t(18;21)(p11;q11)

Partial karyotypes with t(18;21)(p11;q11) (A). Fluorescence in situ hybridization with LSI TEL-AML1 probe (Vysis/Abott Molecular, US) probe showing relocation of AML1 (RUNX1) sequences from 21q22 to the short arm of chromosome 18 (B).

References


This article should be referenced as such: