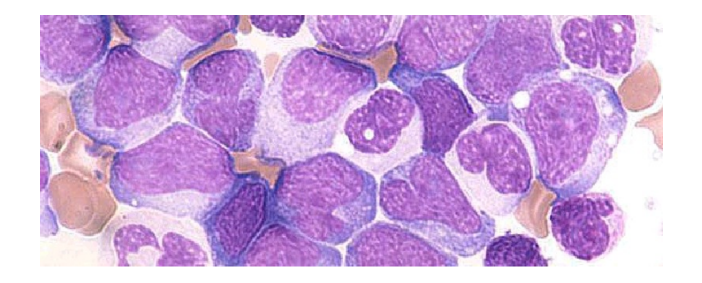
HEMATOLOGIC MALIGNANCIES











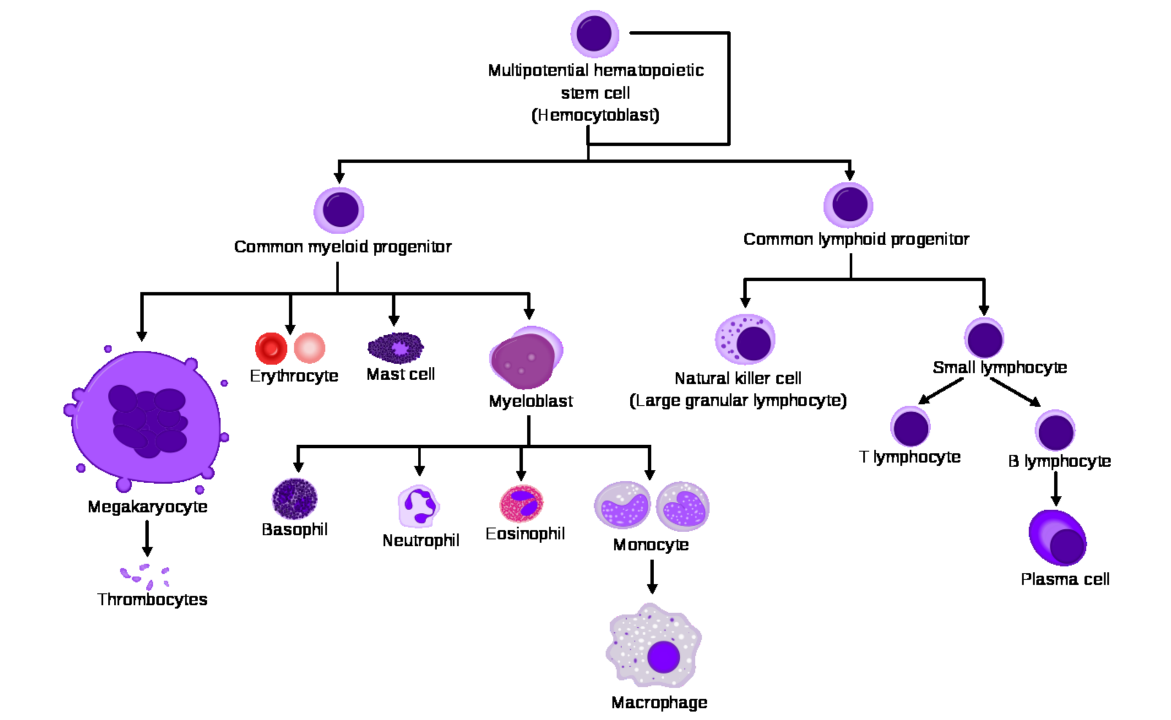


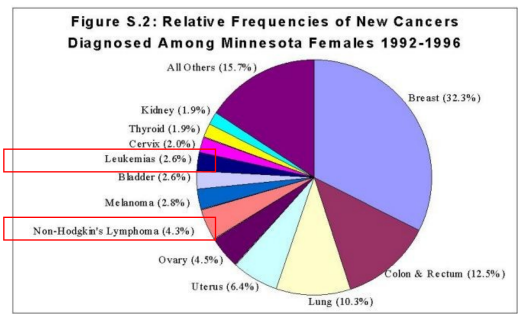
Hematologic malignancies

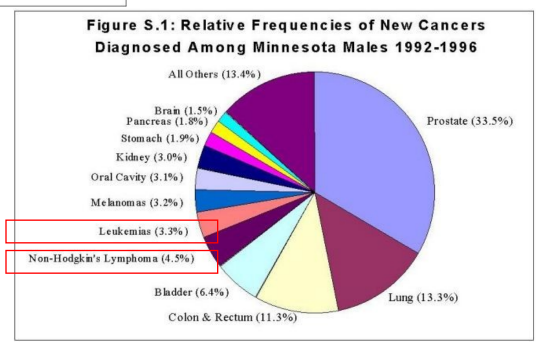
- **Origin hematopoietic cells**
- According to blood lineage
- lymphoid malignancies
- myeloid malignancies

Diseases

- Leukemias
- Lymphomas a lymphproliferative diseases
- Myeloproliferative diseases, myelodysplastic syndromes







Hematologic malignancies

CLONAL

disorders resulting from a mutation of DNA within a pluripotent marrow stem cell or very early progenitor cell.

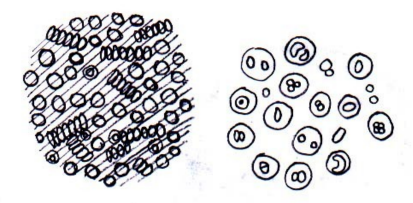
CLONAL POPULATION OF CELLS - cells with growth and/or proliferation advantage over against normal bone marrow cells.

Mutation of DNA can result in the expression of fusion genes that encode fusion proteins that are oncogenic or in the underexpression of genes that encode molecules critical to control of cell growth or progremmed cell death.

Symptoms

- Similar
- Often non-specific
- Bone marrow involvement (leukemia, myeloproliferative neoplasms)
- Lymphoid tissue involvement (lymphoma)

Leukemia







John Hughes Bennett: Two Cases of Disease and Enlargement of the Spleen, in which death took place from presence of purulent matter in the blood, 1845

Beilfunde,

Beifes Blut.

bunben ju fenn. In ben letten 8 Tagen waren endlich wie: In ben alteren Schriftfellern finden fich bier und ba ber febr gabtreiche, jum Theil blutige Durchfalle aufgetreten.

loren batte, baf es ber Mild, bem Chylus, Schleime (pi- taten; Leib voll, aufgetrieben, fluctuirend, bebeutende Bertuita) ober Citer verglichen murbe. (Haller, Elem. physiol. großerung und magige Schmerzhaftigfeit ber Dilg; baufiger, 1760. Tom. II. p. 14-16.) Die Mittheilung bes folgenben anhaltender Guften mit reichlichen geballten sputis, Raffel-Rrantbeitefalles wird biefe fdeinbar fabelhafte Ungabe be- geraufde auf ber Bruft; Appetit und Bunge gut; Bule 78 Schlage machenb; Sarn fparfam; große Erichopfung. (Inf. Rrantheite geichichte. (Musjug aus bem auf ber Colombo c. tinct. Cascarill. et Tinct. theb.). - In ben nachften Abtheilung geführten Journal.) Marie Straid e, Rochin, 50 Tagen beffert bas Befinden fich ber Durchfall nimmt ab, es Jahre alt, wurde am 1. Mary b. 3, in bie Charité aufae: ftellt fich enblich Stublverflopfung ein (Inf. Rhei c. Mell. nommen. Rach ihrer Ausfage batte fie bor einem Jahre bei Tarax.). Reue Diarrhoe (Emuls. comm. c. Aq. Amygd. amar.).

II. Weißes Blut (Leukamie).

Es giebt gewisse Wahrheiten, welche sich in der Wissenschaft nur sehr langsam und schrittweise Geltung verschaffen. So scheint es meinen Mittheilungen über weißes Blut (d. h. eine Vermehrung der farblosen Blutkörperchen in dem Maasse. dass die rothe Farbe des Blutes dadurch in eine röthlich-, gelblich- oder grünlichweiße verwandelt wird) und dem Zusammenhang desselben mit chronischen Milzanschwellungen zu ergehen. Bei der ersten Veröffentlichung des von mir beobachteten Falls (Froriep's N. Notiz. 1845. No. 780.) hob ich schon diesen Zusammenhang hervor und zeigte den Unterschied dieser Blutveränderung von der sogenannten pyämischen. Trotzdem übergeht Bischoff (Müller's Archiv 1846. Jahresber. p. 135.) in seinem Referat den ersteren ganz und bemerkt nur, dass eine chemische Untersuchung nicht angestellt sei und dass der Fall mit anderen, unter dieser Bezeichnung aufbewahrten Fällen nur die Aehnlichkeit des äußeren Ansehens



Rudolf Virchow: Weisses Blut. Frorieps Notizen, 36, s. 152 – 156, 1845

What leukemias are?

- Very different diseases
- Historical name: accumulation of white blood cells
- Not every accumulation of white blood cells is leukemia
 - Leukemoid reaction
 - Lymphoma leukemization
- Acute or Chronic
- Myeloid or Lymphoid

Common features of leukemia

- White blood cells accumulation
 - precursor cells (myelo-, lympho-) blasts
 - acute leukemia
 - myeloid lineage cells CML
 - mature lymphocytes (CLL)
- Leukocytosis (CML, CLL, AL)
- Normal WBC or leukopenia (AL)
- In almost all cases pathology in differential white blood count
- In all cases bone marrow involvement

LEUKEMIAS

Do you know differences between acute and chronic leukemias?

Briefly:

Acute leukemia - there is defect of proliferation, proliferation of <u>young</u> bone marrow cells (blasts) is increased!

Chronic leukemia - there is defect of apoptosis (programmed cell death), apoptosis of mature cells is decreased, <u>mature</u> cells are accumulated in the body!

CAVE: CL can switch to AL (CML in blast crisis, CLL in Richter s syndrome)

LEUKEMIA INCIDENCE

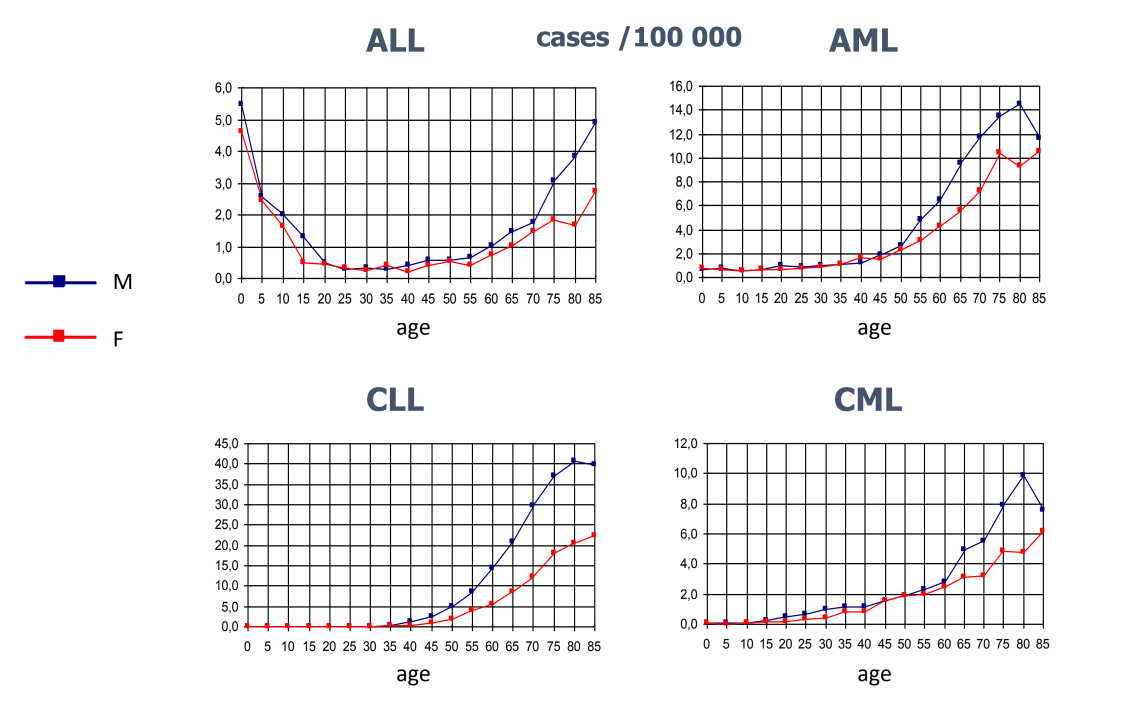
12,7/100 000 M 9,8/100 000 F

Slightly increasing incidence compared with 90's (except of CML)

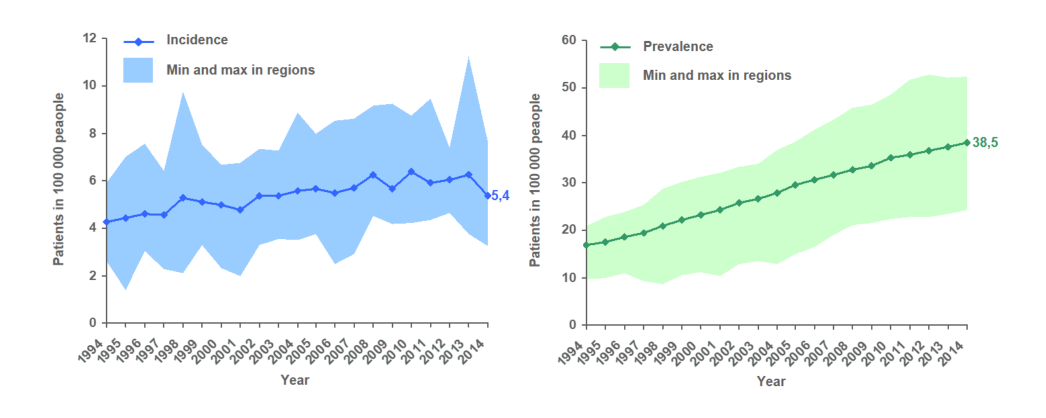
Europe:

40% CLL, 25% AML, 15% CML, 11% ALL, 2% HCL, 7% other

Myelodysplastic syndromes
1 - 2/100 000 (older 10-20/100 000)



LEUKEMIA INCIDENCE AND PREVALENCE - CLL as example



Clinical symptoms of malignant diseases of blood and bone marrow

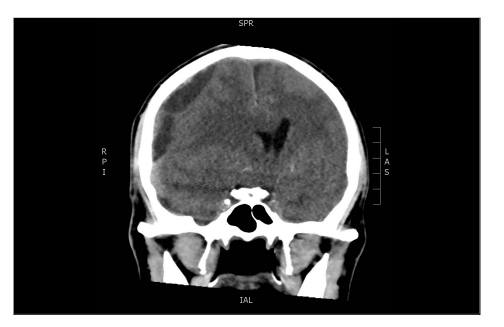
Symptoms affecting patients	Frequency
infection, fever bleeding thrombosis, DIC lymph nodes enlargement splenomegaly hepatomegaly mediastinal tumor CNS involvemet	36 % (all) 33 % (APL, AML) 10 % (APL, ET, PV) 57 % (ALL, CLL) 56 % (CML, CLL, PV, MF) 47 % (CML, AML) 14 % (ALL, CLL) 7 % (ALL, AML M5)
involvement of another organs	9 % (all)

CAVE: All symptoms of hematologic diseases are non-specific!

Bleeding in acute leukemia



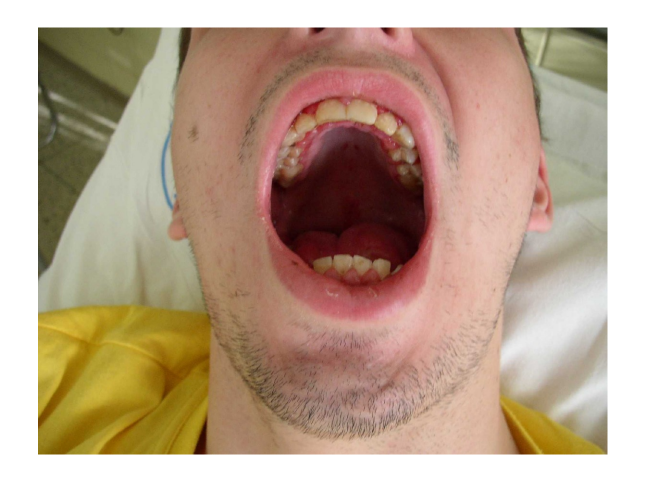






Bleeding in acute leukemia

AML – gum hyperplasia



AML – gum hyperplasia





PLL – skin involvement



ALL – skin involvement



Mastocytosis
- urticatia
pigmentosa

ON SOME

MORBID APPEARANCES

OF

THE ABSORBENT GLANDS

AND

SPLEEN.

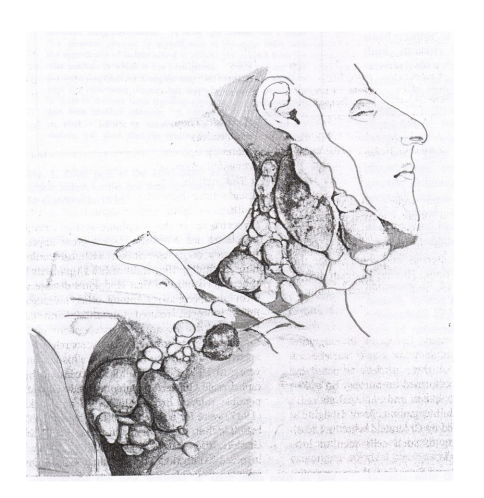
BY DR. HODGKIN.

PRESENTED

BY DR. R. LEE.

READ JANUARY 10TH AND 24TH, 1832.

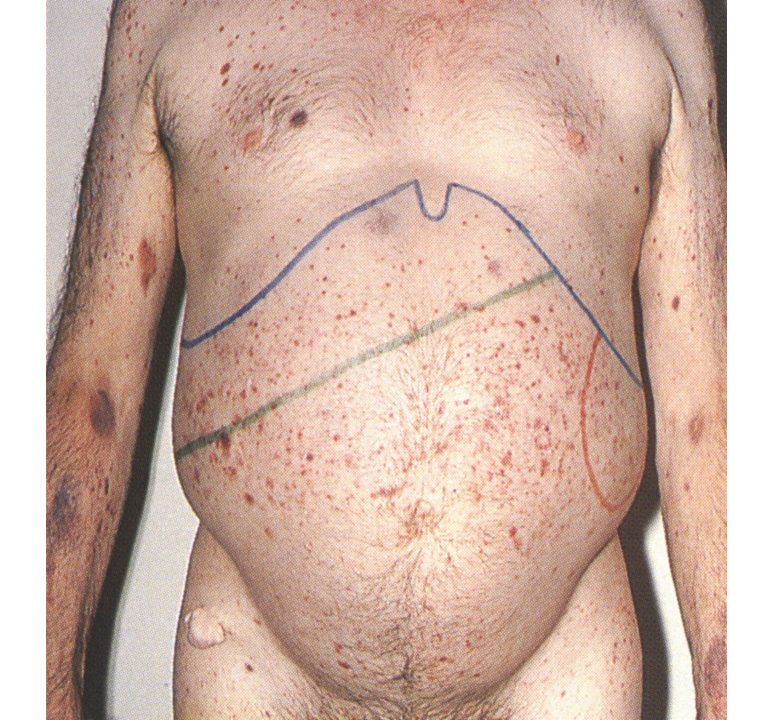
The morbid alterations of structure which I am about to describe are probably familiar to many







CLL - splenomegaly

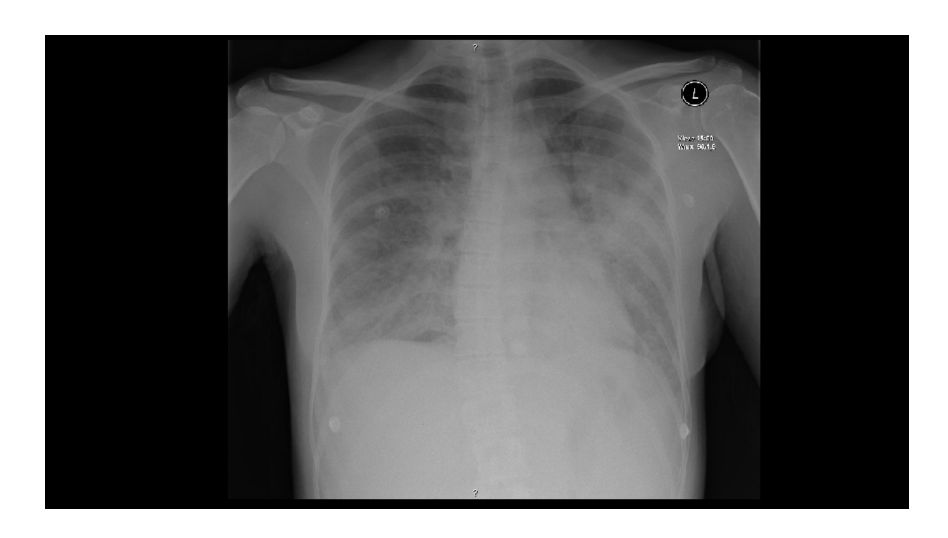


Myelofibrosis
- massive
splenomegaly



Lekocytes
- leukapheresis
bag

Lung infiltration in acute leukemia



Time from first symptoms to final diagnosis

TABLE II. Time from the First Symptoms (Analysis Only Performed Among Patients that Presented Symptoms) and from the First Medical Visit to a Definitive Diagnosis

	ALL	AML	APL	CLL	CML	HCL	Acute leukemias	Chronic leukemias	Total	
Time from the first symptoms to a definitive diagnosis ^a										
No. of analyzed pts.	90	305	59	125	68	22	454	215	669	
Days-median (range)	25 (3-194)	22 (0-226)	14 (3-90)	27 (3-274)	21 (1-256)	34.5 (4-370)	21 (0-226)	27 (1–370)	22 (0-370)	
Days—25–75% interval	14-43	12-36	8-22	14-52	11.5-48	14–77	12-35	13-60	12-42	
Time from the first medical visit to a definitive diagnosis										
No. of analyzed pts.	106	366 ^b	74	293	123	41	546	457	1003	
Days-median (range)	9 (0-108)	7 (0–171)	5.5 (0-71)	12 (0-343)	6 (0–119)	20 (0-355)	7 (0–171)	10 (0-355)	8 (0-355)	
Days—25–75% interval	3–16	3–16	2–12	4–22	2–16	8–36	3–15	3–23	3–19	

LEUKEMIAS – PREDISPOSING FACTORS

Increased risk of leukemia is in:

Genetic syndromes – M. Down, FA, ataxia telangiectasia, inherited germline mutations (*ETV6*, *RUNX1*, *DDX41*...)

Drugs (chemotherapy, alkylating agents)

Radiation (can cause all leukemias except CLL)

Socioeconomic factors

(increased incidence of childhood ALL in industrial countries, probably due to later contact of children with alergens or banal childhood infections)

Viruses (EBV, HTLV I, HIV)

Benzene, toluene, etc.

LEUKEMIAS – ETIOLOGY

Somatic molecular leasions involving:

Cell proliferation
Cell division
Cell maturation
Apoptosis
Cell self-renewal

LEUKEMIAS — ETIOLOGY

The most important somatic molecular changes in leukemia and myeloproliferative neoplasms:

BCR-ABL TP53 PML-RARa JAK2

LEUKEMIAS AND MYELOPROLIFERATIVE DISEASES Blood and bone marrow features

What can we found in periperal blood (WBC, RBC, platelets)?

- acute leukemia
- chronic leukemia
- myeloproliferative diseases

What can we found in bone marrow?

- acute leukemia
- chronic leukemia
- myeloproliferative diseases

Laboratory diagnostics

Peripheral blood count with differential WBC

Bone marrow

Flow cytometry (analysis of CD antigens)
(ALL, CLL, HCL)
Cytogenetic analysis (CML, AL, MDS, CLL)
Molecular genetic analysis (CML, APL, AL, CLL)
Cytology a cytochemistry
Histology (necessary in myeloproliferative diseases)

Do you know differences between trephine biopsy and sternal puncture? Sternal puncure - we can collect only marrow blood. SP fits for diagnostics of leukemias.

VÝSLEDEK VYŠETŘENÍ Z HEMATOLOGIE

Pacient:
Datum a čas odběru:

Datum a cas ouberu.	Hodn Wa	1	Todn	Maza/komant
Vyšetření ———————	Hodn. Výs	· · · · · · · · · · · · · · · · · · ·	Jedn.	Meze/koment.
Leukocyty	<.>		x10 9/1	(4 - 10)
Erytrocyty	<.>		x10 12/1	(3.8 - 5.4)
Hemoglobin	<l></l>	115.0		(120 - 160)
Hematokrit	<l></l>	0.320	1/1	(0.35 - 0.46)
Střední objem ERY	<l></l>	83.6	fL	(84 - 96)
Trombocyty	<.>	163.0	x10 9/1	(150 - 350)
Stř. množství HGB v	<.>	30.2	pg	(28 - 34)
Prům. koncentrace HG	<.>	362.0	g/1	(320 - 370)
Šíře distribuce ERY	<.>	13.5		(10 - 15.2)
Střední objem trombo	<l></l>	7.05	fl	(7.8 - 11)
Trombocytový hematok	<l></l>	1.15	ml/1	(1.21 - 3.5)
Šíře distribuce trom			00	(15.5 - 17.1)
Neutrofily %	<l></l>	19.80	00	(50 - 70)
Lymfocyty %	<.>	22.40		(20 - 40)
Monocyty %	<h></h>	57.00		(2 - 12)
Eosinofily %	<.>	0.28		(0 - 5)
Basofily %	<.>	0.60		(0 - 1)
Neutrofily (absolutn			x10 9/1	(2 - 7)
Lymfocyty (absolutní			x10 9/1	(0.8 - 4)
Monocyty (absolutní	<h></h>		x10 9/1	(0.08 - 1.2)
Eosinofily (absolutn	- Alamanda de la companya del companya de la companya del companya de la companya		x10 9/1	(0 - 0.5)
Basofily (absolutní	<.>		x10 9/1	(0.01 - 0.1)
Neutrofily mikroskop			%	(50 - 70)
Tyče mikroskopicky	<.>		00	(0 - 4)
Lymfocyty mikroskopi		23.0		(20 - 40)
Monocyty mikroskopic		2.0		(2 - 12)
Eosinofily mikroskop		0.0		(0 - 5)
Basofily mikroskopic		0.0		(0 - 1)
Metamyelocyty mikros			00	(0 - 0)
		0.0		(0 - 0)
Myelocyty mikroskopi		0.0		(0 - 0)
Promyelocyty mikrosk		54.0		(0 - 0)
BLASTY mikroskopicky		0.0		(0 - 0)
Prolymfocyty mikrosk				
Plazmatické buňky	<.>	0.0		(0 - 0)
Nedif.buňky	<.>	0.0		(0 - 0)
Nedif.blasty	<.>	0.0		(0 - 0)
Normoblasty mikrosko		5.0	/100 bb	
Hodnocení morfologie				1
Hodnocení morfologie				hypersegmentace neutrofilů,
Morfologie ERY	< >			· · · · · · · · · · · · · · · · · · ·
Morfologie PLT	< >			mírná anizo PLT,
KOMENTAR	< >			Změna oproti předešlému.
Neznamé vyšetření	< >			NRBC/100WBC :0.00





Laboratory diagnostics

Biochemical analysis of blood (elevated LD in myeloproliferative diseases)

<u>Coagulation – DIC, thrombophilia, bleeding</u> fibrinogen, aPTT, PT, AT III, DD, EGT

Other

(Chest X ray, abdominal sonography, ECG, heart sonography, serology — CMV...) - we have to exclude focal infections and to evaluate function of heart, kidneys, liver and lungs (chemotherapy is nephrotoxic, hepatotoxic or cardiotoxic)

CLASSIFICATION OF LEUKEMIA

FAB (1982)

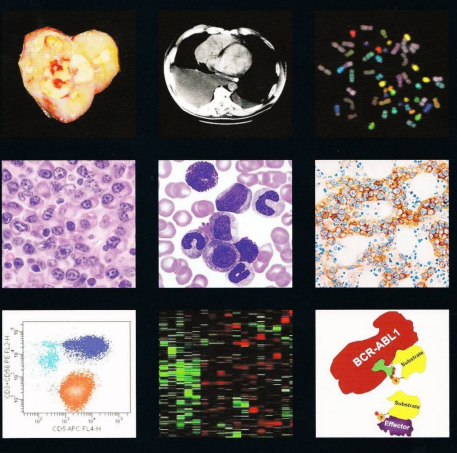
Classification according to morphology of malignant cells

WHO (1999-)

Classification according to morphology, cytogenetic features, flow cytometry, and molecular genetic features

WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues

Edited by Steven H. Swerdlow, Elias Campo, Nancy Lee Harris, Elaine S. Jaffe, Stefano A. Pileri, Harald Stein, Jürgen Thiele, James W. Vardiman





CHRONIC LYMPHOCYTIC LEUKEMIA (CLL)

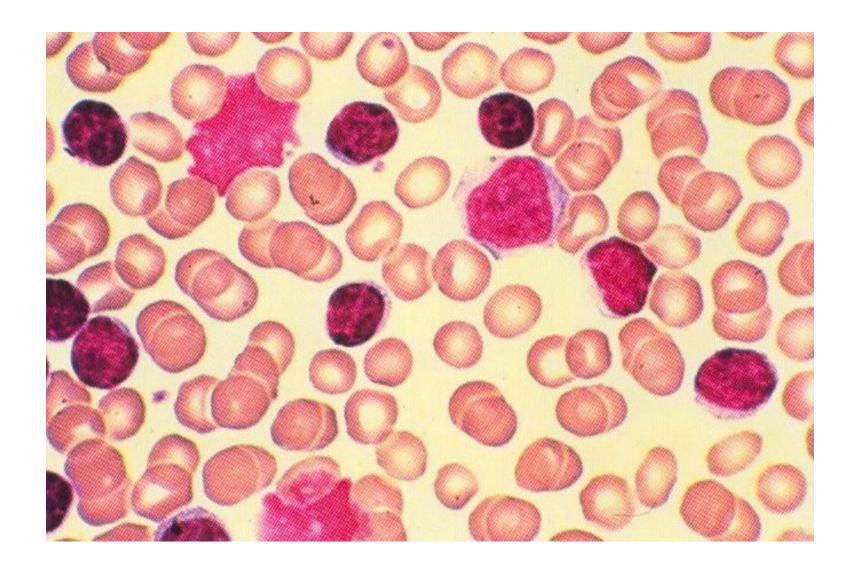
The most common leukemia of Caucasians. CLL is a disorder characterized by the accumulation of small mature-appearing lymphocytes in the blood, marrow, and lymphoid tissues.

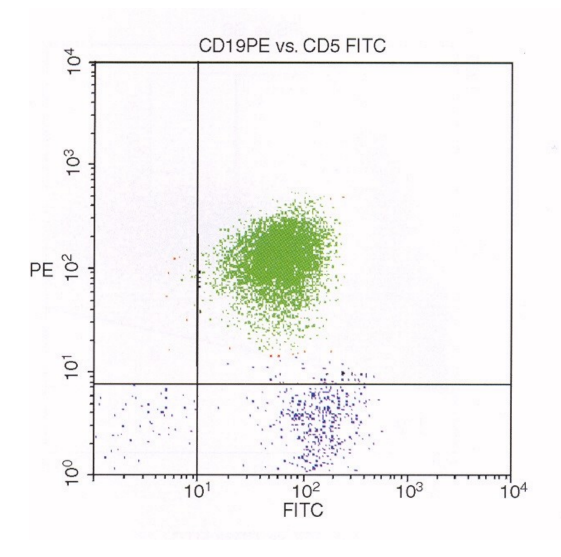
Laboratory and clinical features:

leukocytosis (absolute lymphocytosis), lymphadenopathy, splenomegaly, hepatomegaly, anemia, thrombocytopena, often autoimmune diseases (hemolysis).

Prognosis – different (better in CLL mutated genes for IgH or/and in CLL with del 13q14.

Median survival of CLL patients is 11+ years.



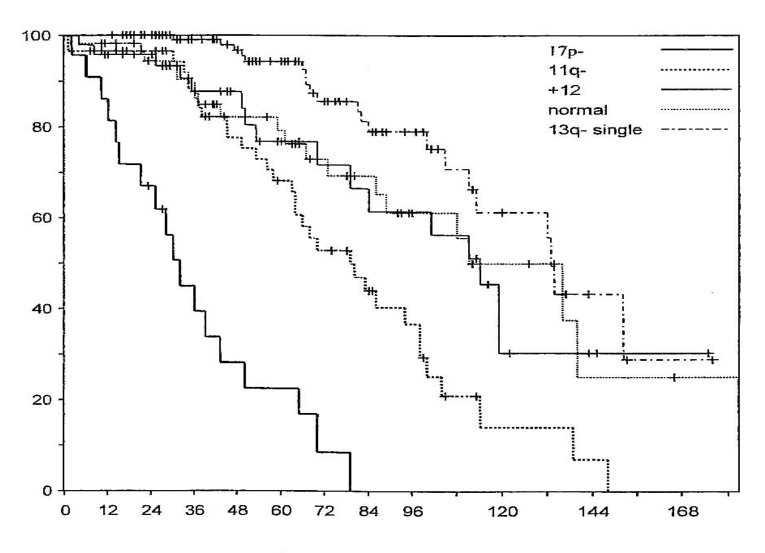


CLL staging

Treatment in stage Rai III or IV patients only (Binet C)

F		
Clinical stage (Rai)	Risk	Median survival
0 (lymphocytosis)	Low	>150 months
I (lymphocytosis + lymphadenopathy)	Intermediate	101
II (lymphocytosis + splenomegaly)	Intermediate	71
III (lymphocytosis + anemia Hb < 110 g/l)	High	19
IV (lymphocytosis + thrombocytopenia < 100x10 ⁹ /L)	High	19
Clinical stage (Binet)		
A (involvement <3 regions)	Low	Not reached
B (involvement ≥ 3 regions)	Intermediate	84
C (anemia and thrombocytopenia)	High	24

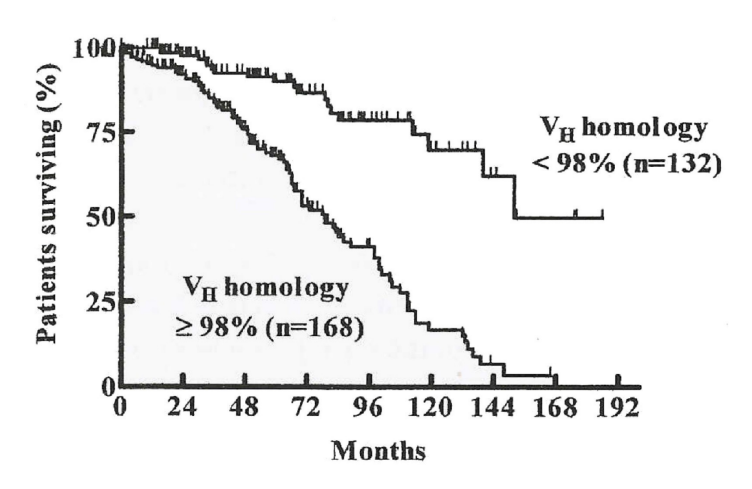
CLL prognosis based on cytogenetics



Overall survival (months)

CLL prognosis based on IgHV mutational status

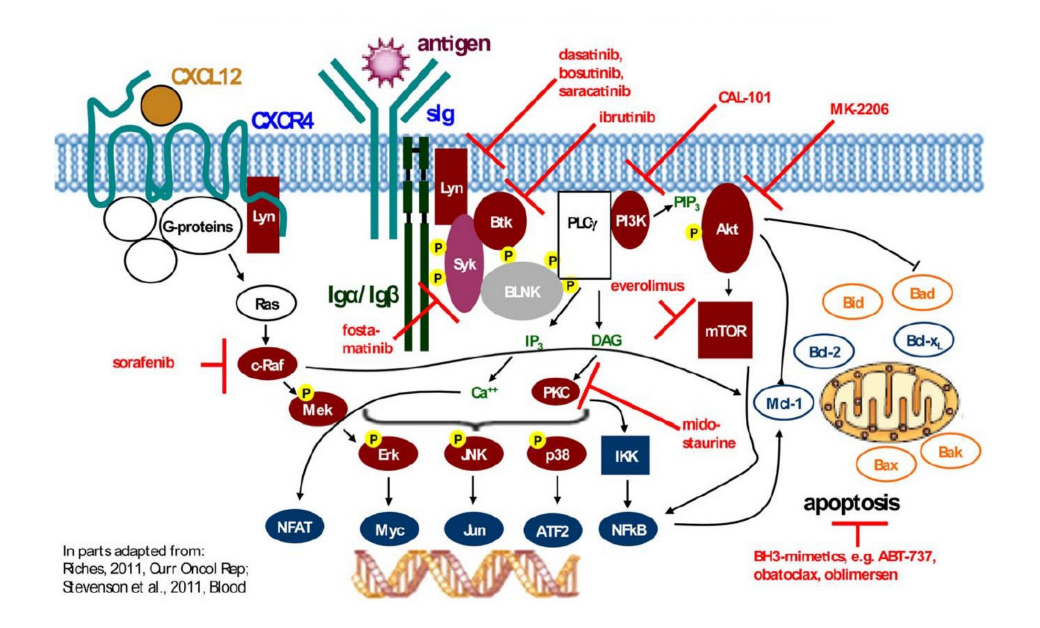
A



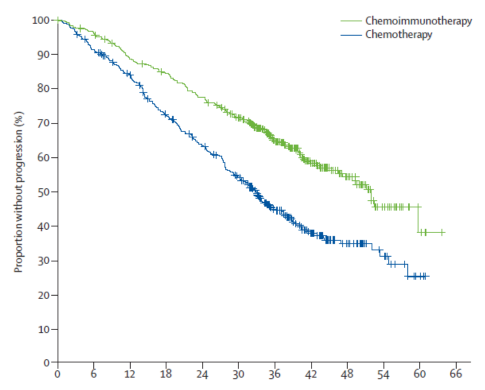
CLL THERAPY

Treatment for advanced stages only:

- fludarabine+cyclophosphamide+rituximab
 - bendamustine+rituximab
- chlorambucil + anti CD20 antibody (rituximab, obinutuzumab)
 - ibrutinib, idelalisib (BCR inhibitors)
 - venetoclax (Bcl2 inhibitor)
 - (allogeneic transplant)

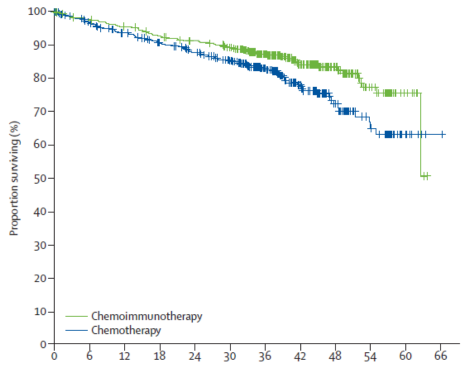


CLL – FCR regimen treatment outcome

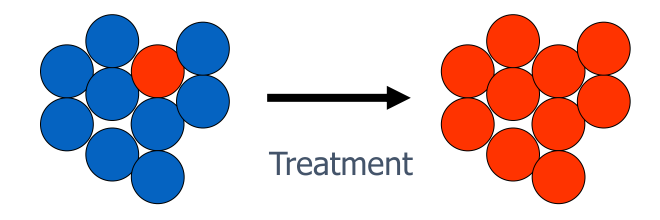


OS at 3 years 83% vs. 87% p = 0,012

PFS at 3 years 45% vs. 65% p < 0,0001

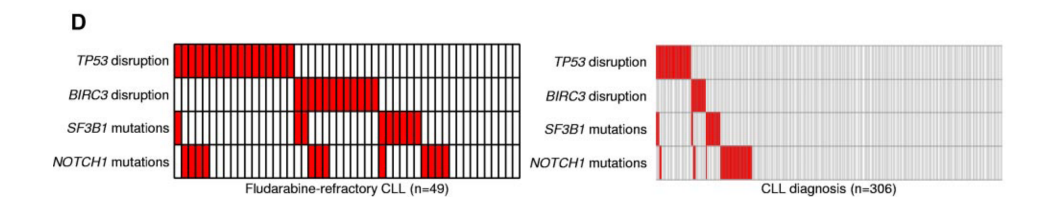


Clonal evolution in CLL – *TP53*



Before treatment

Unfavorable SFB3, NOTCH1, BIRC3 mutations



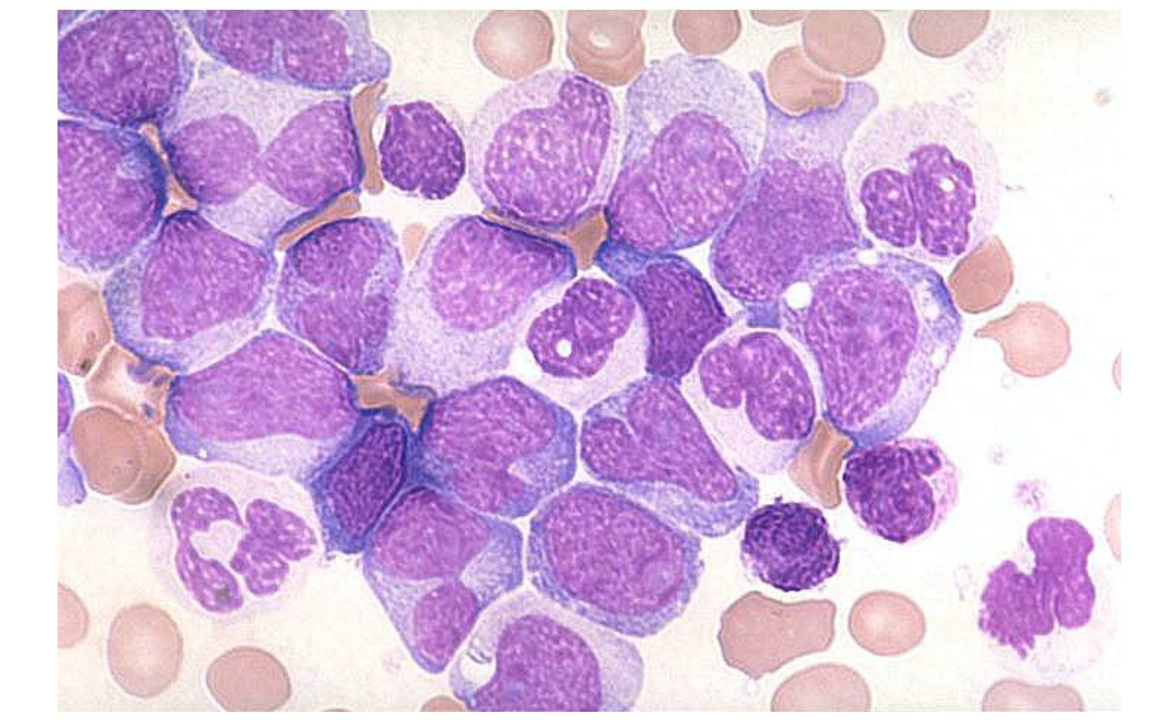
CHRONIC MYELOID LEUKEMIA (CML)

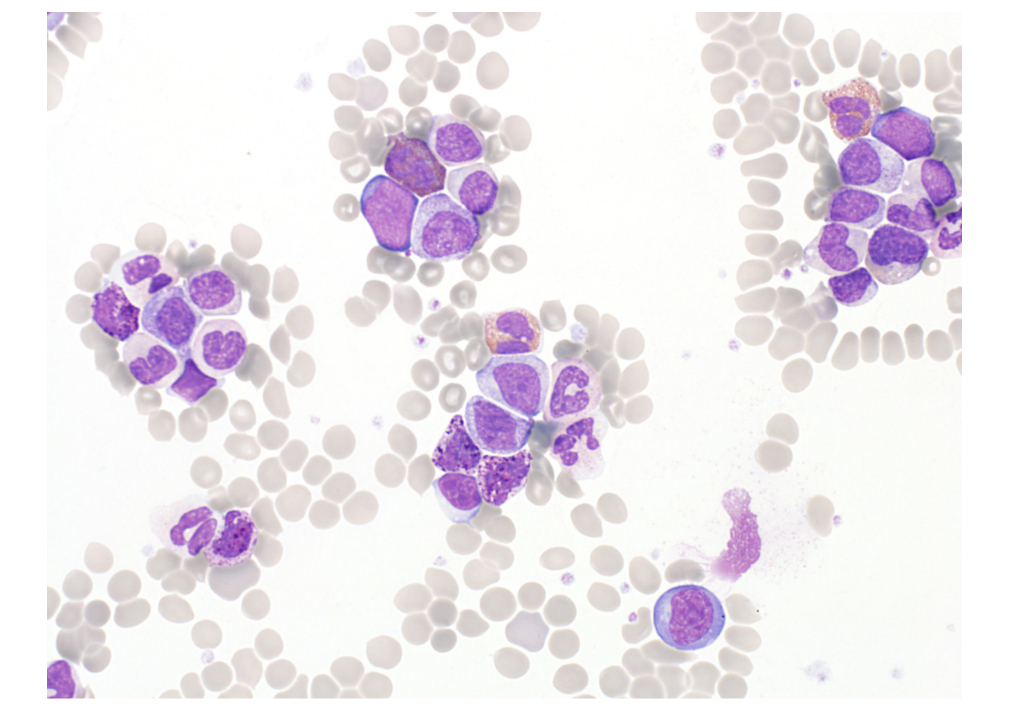
CML is a pluripotent stem cell disease that is characterized by extreme blood granulocytosis, basophilia, often thrombocytosis, anemia, and splenomegaly.

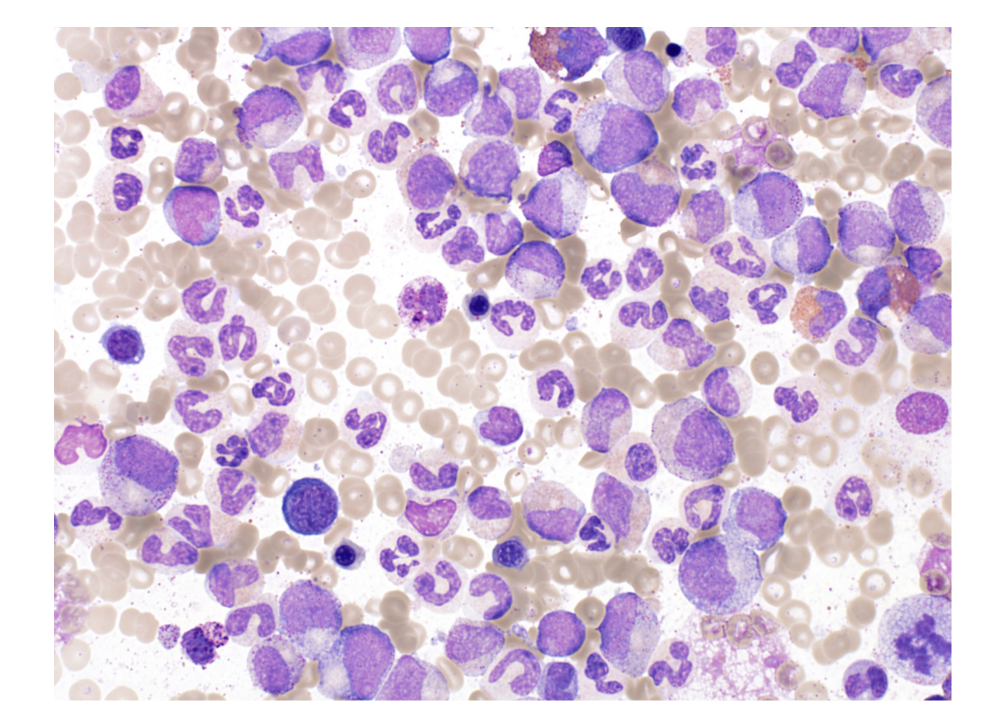
Stages of untreated CML:

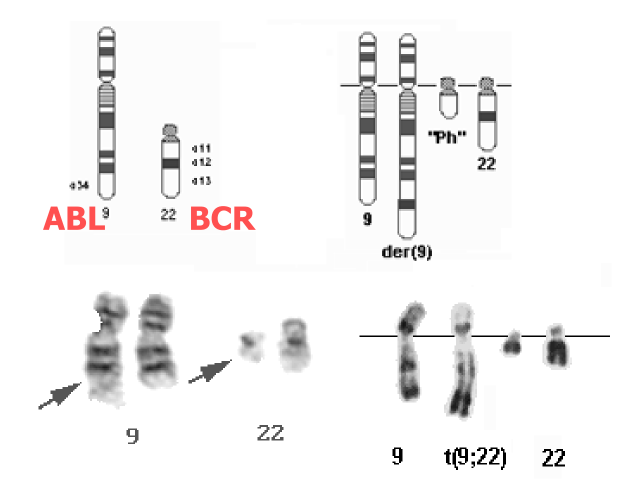
chronic phase, accelerated phase (rapid increase of WBC, worsening of thrombocytopenia, new cytogenetic features, resistence to treatment), blast crisis (resembles to acute leukemia)

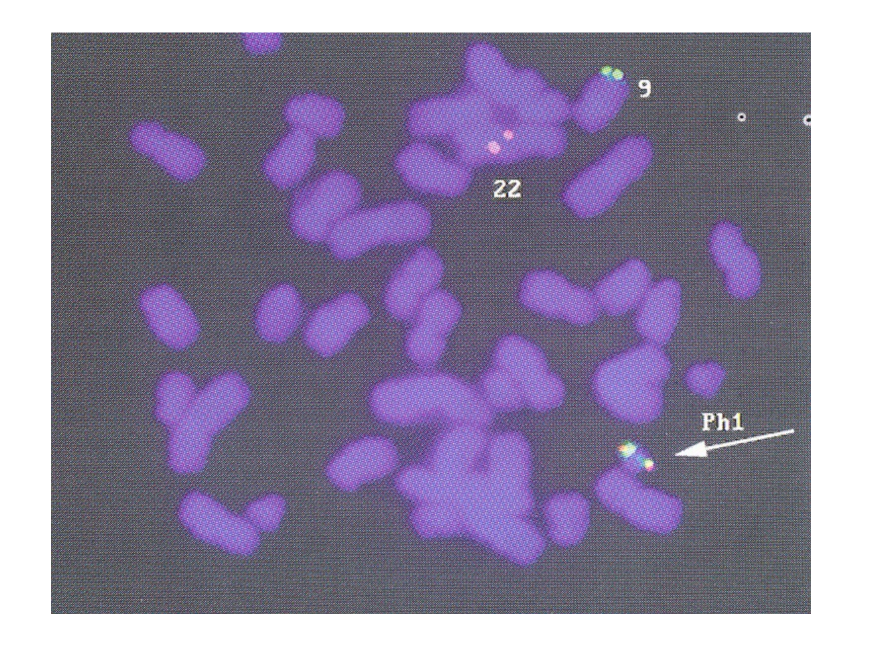
Etiologic role of chromosome discovered in Philadelphia - Ph chromosome











- Ph chromosome arises from t(9;22)
- chimeric gene BCR-ABL arises from Ph chromosome
- BCR-ABL gene produces BCR-ABL tyrosinkinase
- BCR-ABL tyrosinkinase induces defect of apoptosis

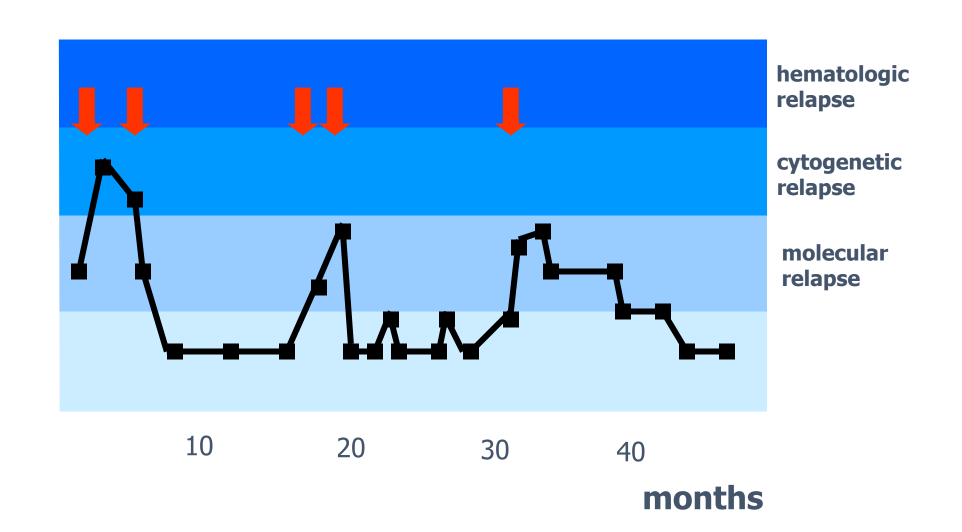
There is almost no BCR-ABL negative CML!

Minimal residual disease during treatment

- Hematologic monitoring
- Cytogenetic monitoring
- Molecular genetic monitoring (RQ-RT-PCR, digital PCR, NGS)

Minimal residual disease during therapy

Molecular relapse is better manageable compared with cytogenetic or hematologic relapse



THERAPY

All patients treated!

- imatinib
- nilotinib, dasatinib, bosutinib
 - ponatinib
 - interferon
 - allogeneic transplantation

II. Zwei Fälle von Lencaemie.

Mitgetheilt

Dr. Lissauer in Bendorf.

Der in Nr. 31. dieser Wochenschrift von Dr. Valentiner mitgetheilte Fall von Leucaemie, bei welcher zur Coupirung des Fiebers Liq. arsenic. Fowler, angewandt wurde, brachte mir zwei Fälle derselben Krankheit in Erinnerung, die ich kurz nach einander im Landkrankenkause in Cassel zu beobachten Gelegenheit hatte, von welchen bei einem Liq. arsen. Fowler. eine Zeit lang versuchsweise von gutem Erfolge war. Ich theile beide Eälle hier kurz mit, theils als einen kleinen Beitrag zur Kenntoiss dieser im Ganzen immer noch selten diagnosticirten Krankheit, theils, um zur weiteren Anwendung obigen Mittels anzuregen.

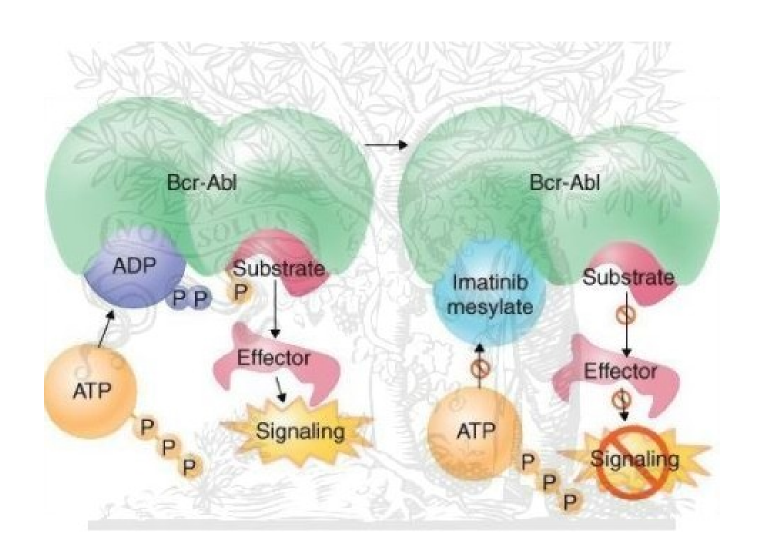
N. N., 32 Jahre alt, weiblichen Geschlechts, wurde im October v. J. aufgenommen. Sie gab an, früher stets gesund, mit 17 Jahren regelmässig menstruirt gewesen zu sein, und vor ungefähr einem Jahre ein uncheliches Kind geboren zu haben, das bald nach der Geburt gestorben sei. Von ihrem Liebhaber, der ihr die Ehe versprochen, hintergangen, habe sie sich sehr gegrämt und viel Sorgen gemacht. Zugleich will sie seit dieser

Asenic trioxide

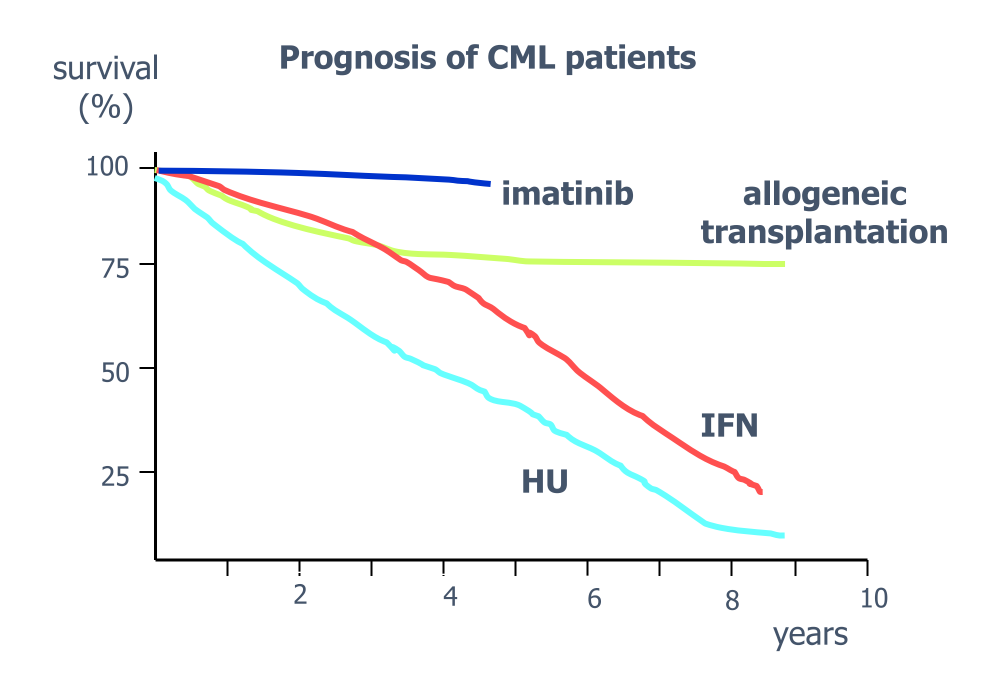
Lissauer: Zwei Fälle von Leucaemie. Berlin. Klin. Wochenschrift, 2, 1865, s. 403 - 404

Malgaigne I. c. p. 1004. Revue medic, chirurg., 1849, T. V.,
 p. 246.

Imatinib mode of action



CP-CML	Léčebná strategie
<u>1. linie:</u>	-Imatinib 400 mg
2. linie:	
IM-intolerance	-DASATINIB nebo NILOTINIB
IM-selhání	-DASATINIB nebo NILOTINIB -aloTKB (progrese do AP/BC, T315I)
IM-suboptimální odpověď	IM stejná dávka IM navýšení dávky DASATINIB nebo NILOTINIB



ACUTE MYELOID LEUKEMIA (AML)

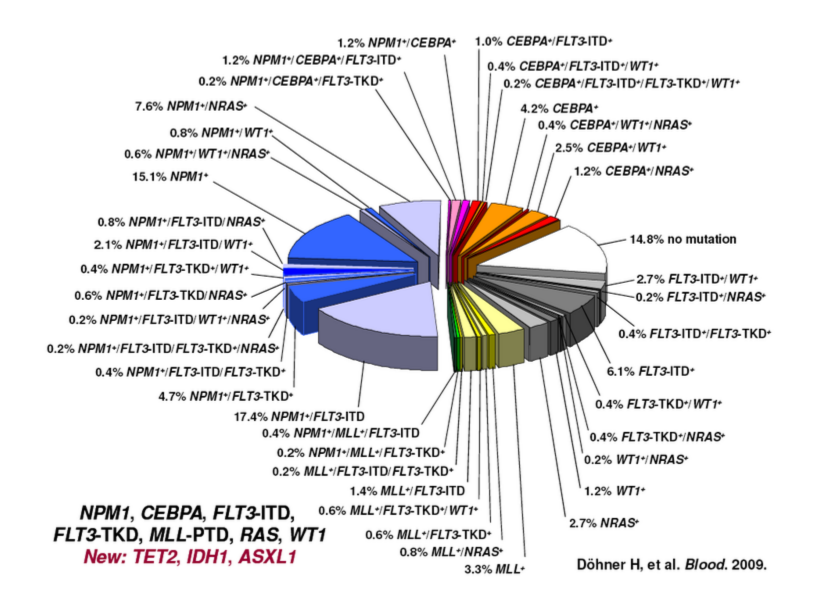
AML is clonal malignant disease that is characterized by the proliferation of abnormal (leukemic) blasts, principially in the marrow, and impaired production of normal blood cells.

Signs and symptoms of AML include pallor, fatigue, weakness, palpitations, bleeding, fever, and dyspnena.

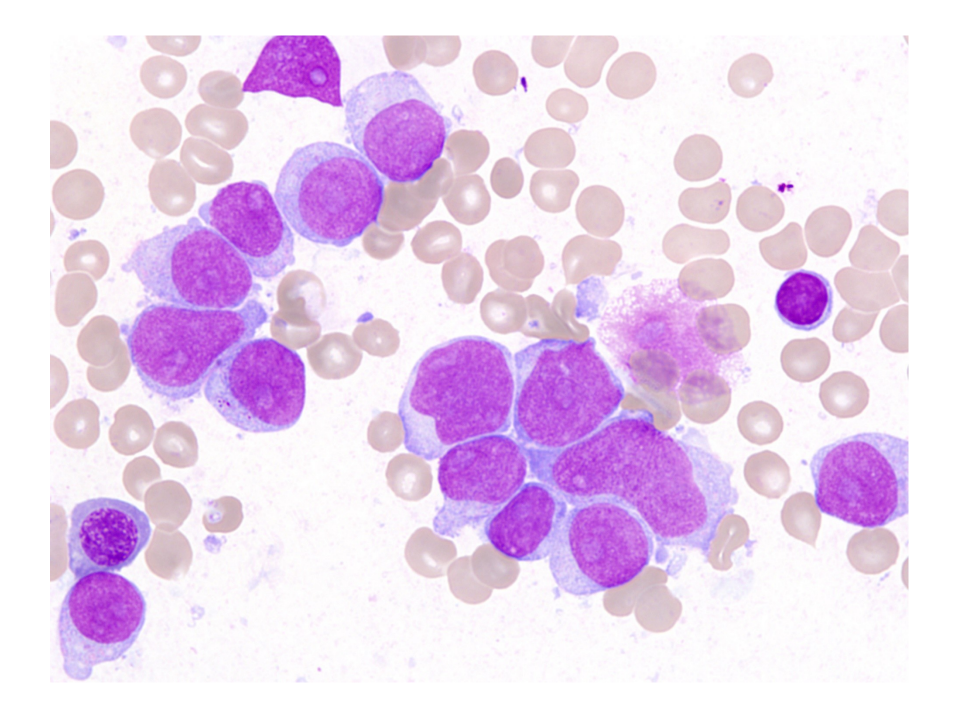
In bone marrow, there is more than 20% of blast cells. (less than 20% - myelodysplastic syndrome)

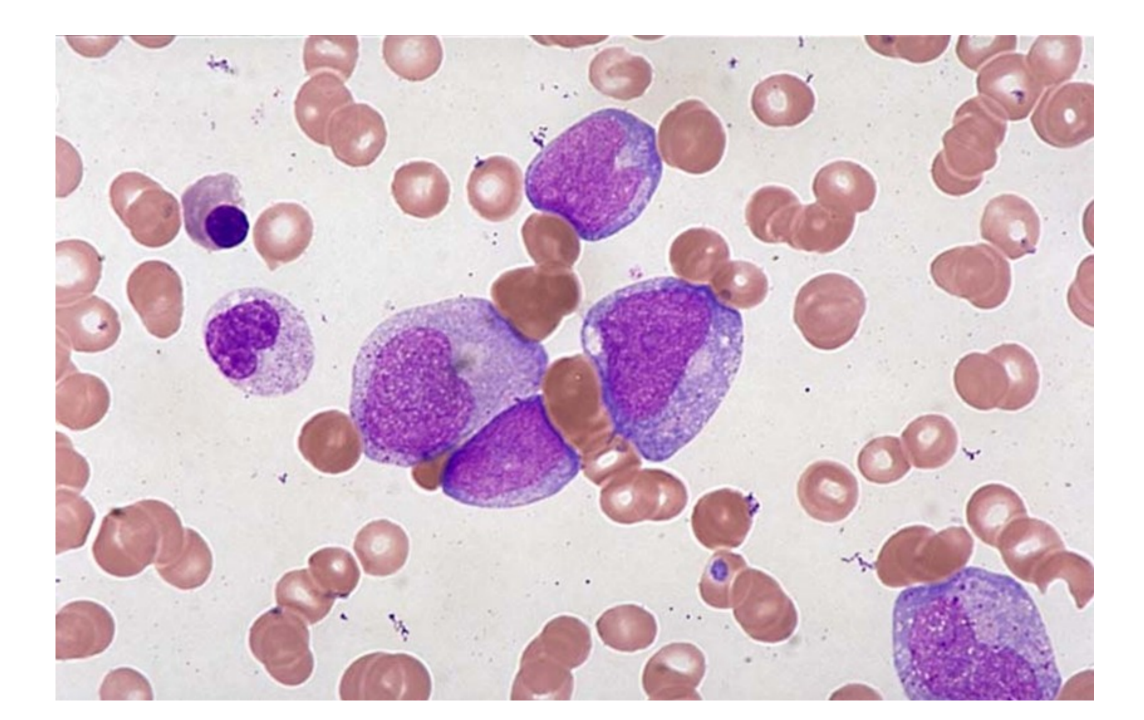
Median survival of untreated patients is 6 weeks.

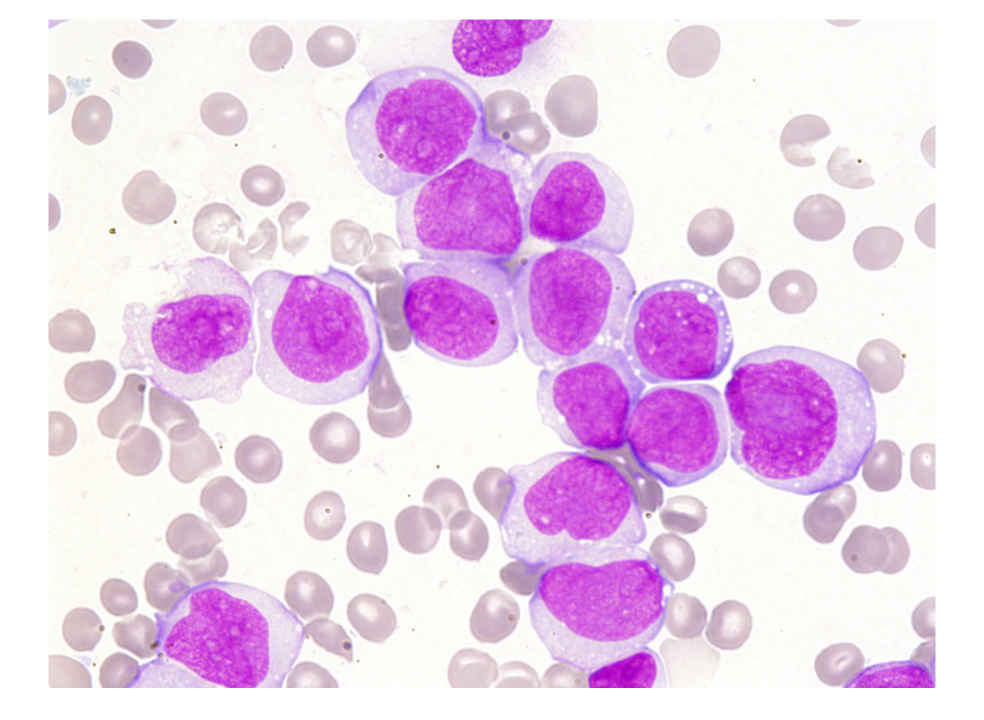
AML – heterogenous disease



Acute myeloid leukemia (AML) and related neoplasms
AML with recurrent genetic abnormalities
AML with t(8;21)(q22;q22.1); RUNX1-RUNX1T1
AML with inv(16)(p13.1q22) or t(16;16)(p13.1;q22); CBFB-MYH11
APL with PML-RARA
AML with t(9;11)(p21.3;q23.3);MLLT3-KMT2A
AML with t(6;9)(p23;q34.1);DEK-NUP214
AML with inv(3)(q21.3q26.2) or t(3;3)(q21.3;q26.2); GATA2, MECOM
AML (megakaryoblastic) with t(1;22)(p13.3;q13.3);RBM15-MKL1
Provisional entity: AML with BCR-ABL1
AML with mutated NPM1
AML with biallelic mutations of CEBPA
Provisional entity: AML with mutated RUNX1
AML with myelodysplasia-related changes
Therapy-related myeloid neoplasms
AML, NOS
AML with minimal differentiation
AML without maturation
AML with maturation
Acute myelomonocytic leukemia
Acute monoblastic/monocytic leukemia
Pure erythroid leukemia
Acute megakaryoblastic leukemia
Acute basophilic leukemia
Acute panmyelosis with myelofibrosis
Myeloid sarcoma
Myeloid proliferations related to Down syndrome
Transient abnormal myelopoiesis (TAM)
Myeloid leukemia associated with Down syndrome

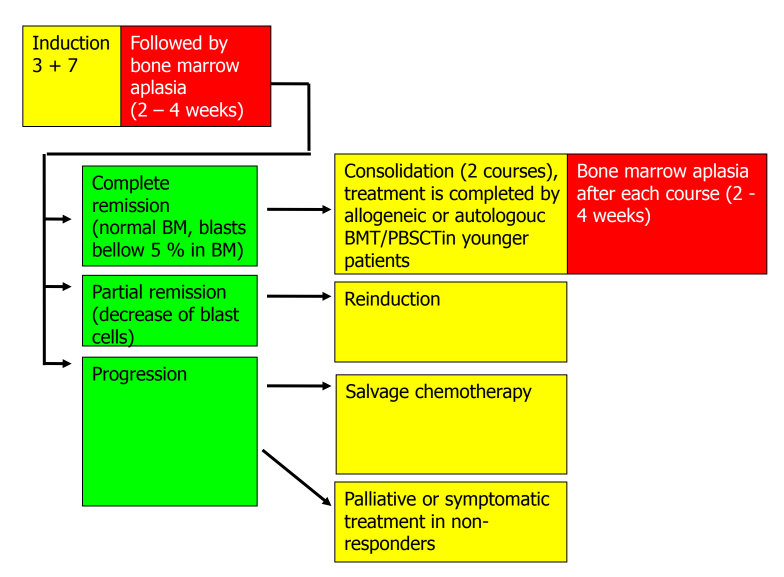






Treatment of AML

Treatment of choice of AML are courses of chemotherapy, the most potent drugs are cytosinarabinoside and anthracyclines.



Treatment of AML

Novel drugs for AML:

Midostaurin

Venetoclax

Gilteritinib

Acute promyelocytic leukemia (APL, AML M3)

APL is variant of AML (constitutes about 5-10% of AML in central Europe, about 25% of AML southern Europe, and 50 % of AML in eastern Asia).

There are prominent hemorrhagic complications (DIC, melena, hematuria, pulmonary bleeding, CNS bleeding)

Prognosis of APL was very poor 30 years ago (almost all patients died).

Nowadays, DFS is 80%.

Acute promyelocytic leukemia (APL, AML M3)

Promyelocytes are granular cells. In granula are coagulopathy-inducing factors (tissue factor...).

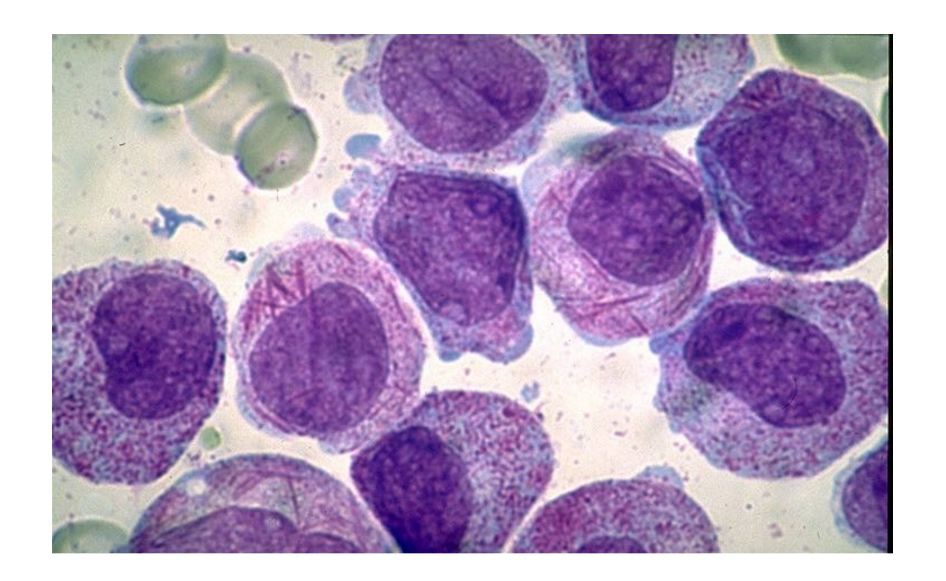
Majority of APL is characterized by t(15;17). A translocation between chromosome 17 and 15 results in chimeric fusion gene *PML-RARa*.

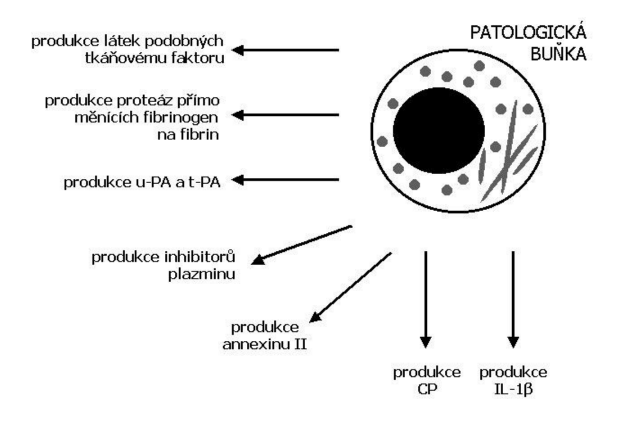
PML-RARa gene produces PML-RARa abnormal recepror for retinoids. (Retinoids are necessary for normal bone marrow cells differentiation). In cells with t(15;17) normal differentiation is stopped.

We can restore differentiation by means of ATRA + ATO or chemo.

Chemotherapy or arsenic trioxide + ATRA is treatment of choice for APL!



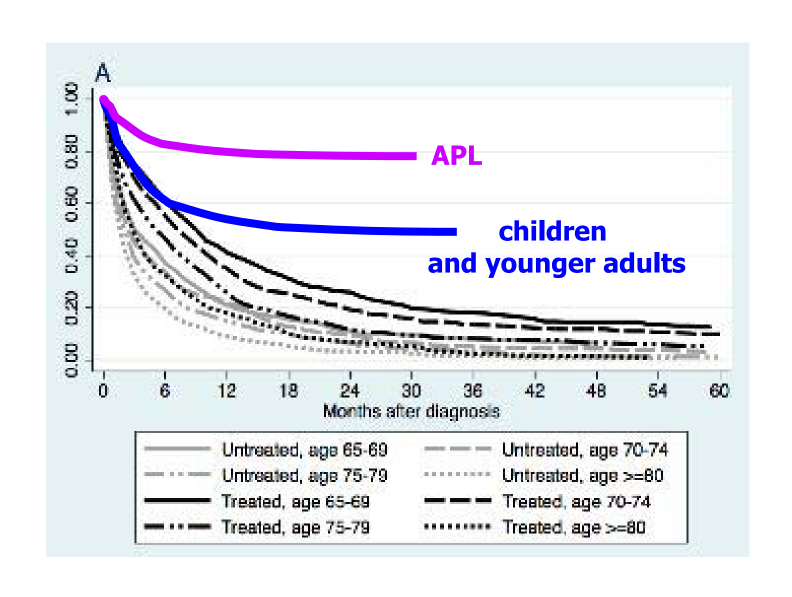




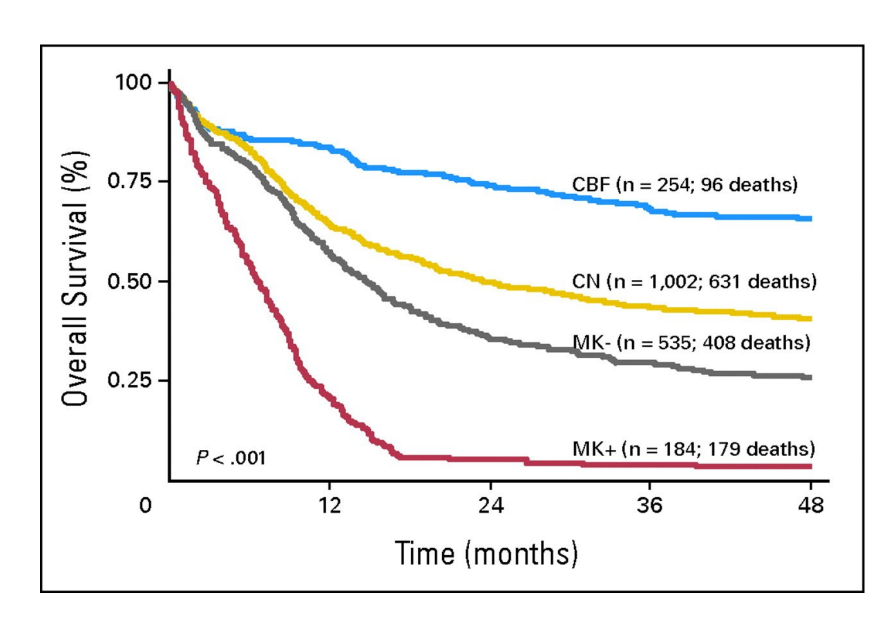
Bleeding diathesis in APL:

 $CP-cancer\ procoagulant,\ IL-1\beta-interleukin\ 1\beta,\ t-PA-tissue$ plasminogen activator, u-PA-urokinase.

Survival of AML patients



Survival of AML patients



good risk standard risk poor risk

t(8;21) inv(16)

normal karyotype

komplex. karyotype - 7

ACUTE LYMPHOBLASTIC LEUKEMIA (ALL)

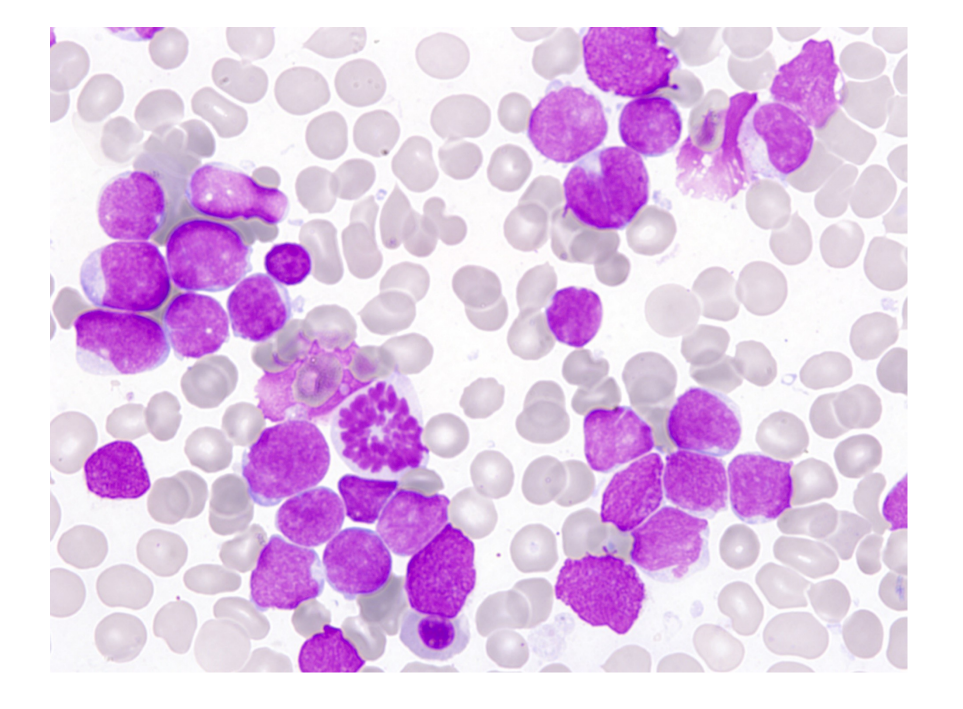
The most common leukemia in childhood.

In children - very good prognosis. In adults - poorer prognosis.

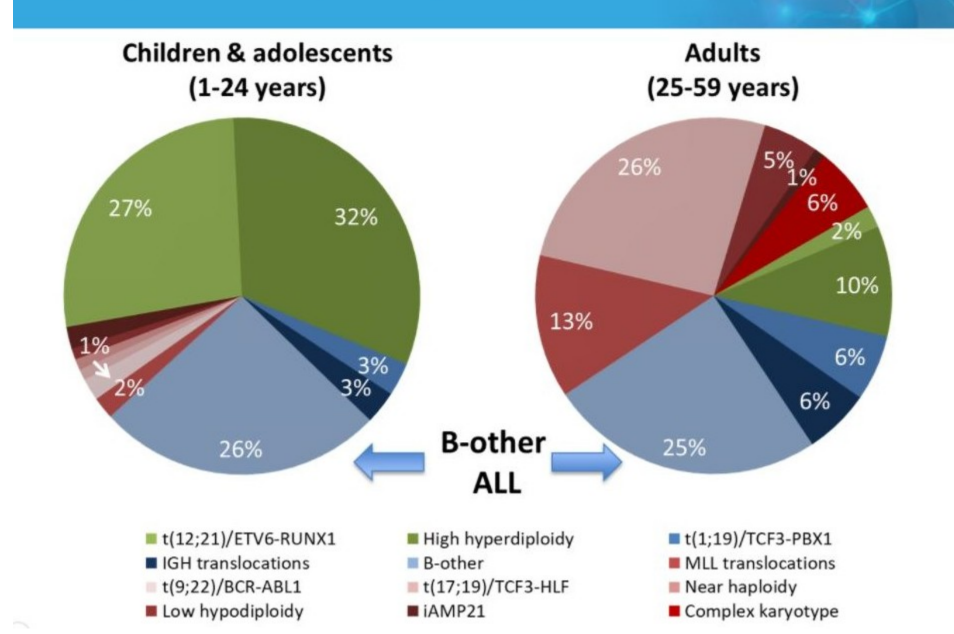
ALL is a neoplastic disease resulting from somatic mutation in a single lymphoid progenitor cell.

B precursor ALL vs. T precurosr ALL

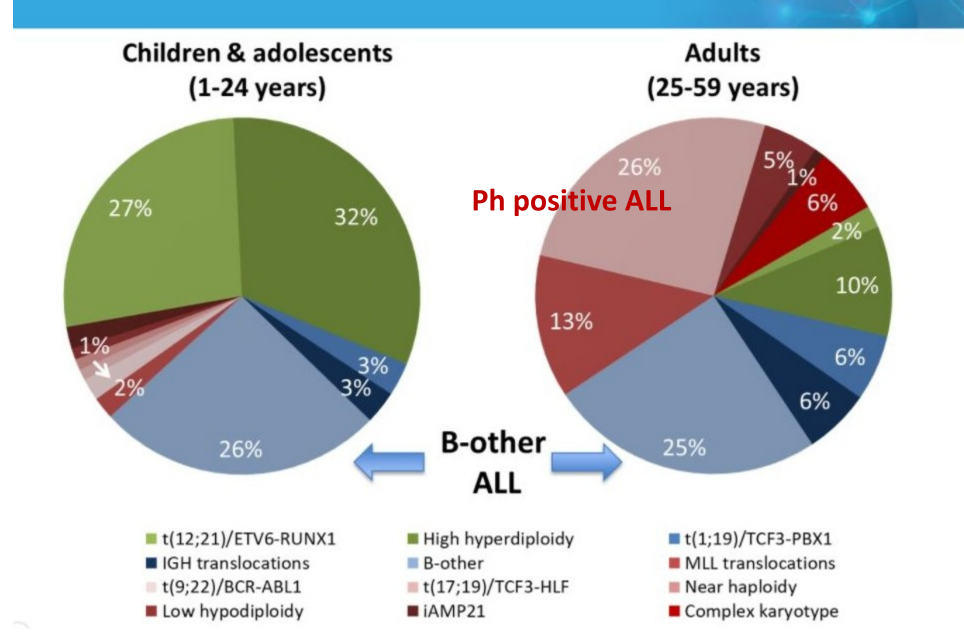
BM - more than 20% of lymphoblasts (usually 80 - 100%).



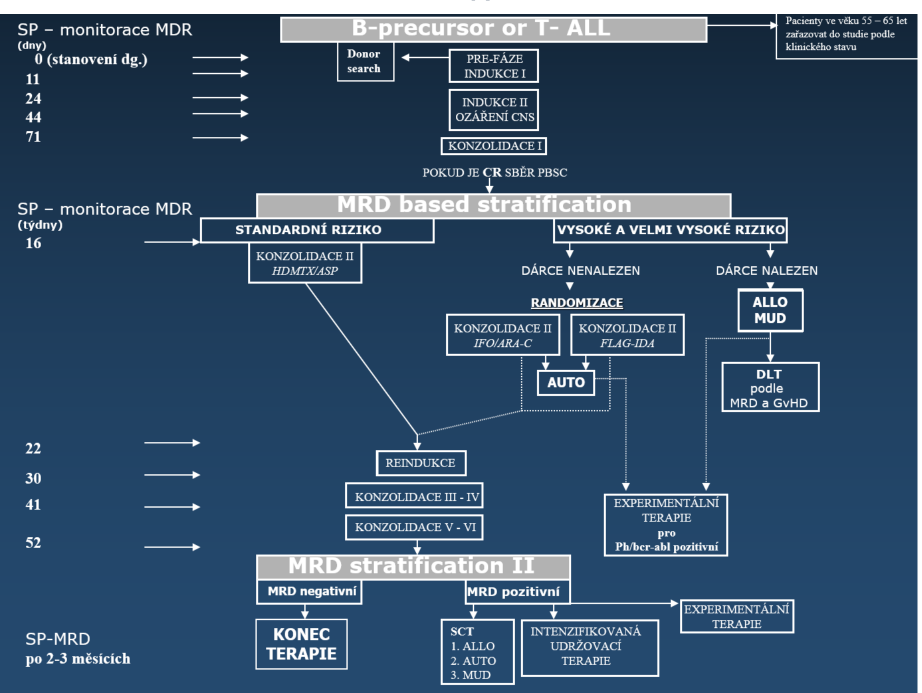
Age specific frequency of genetic subgroups in ALL



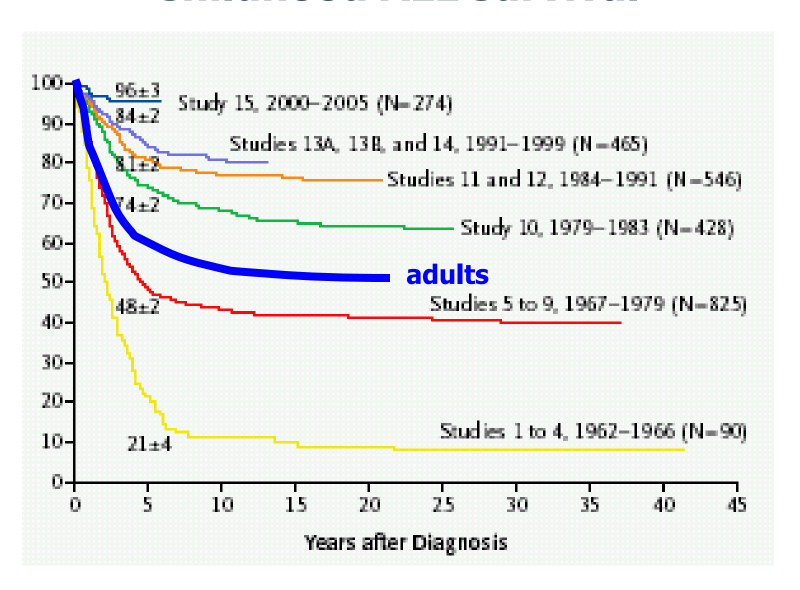
Age specific frequency of genetic subgroups in ALL



ALL – therapy overview



Childhood ALL survival



MYELODYSPLASTIC SYNDROMES

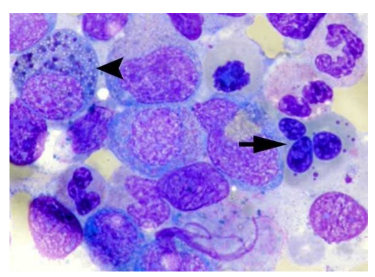
Heterogeneous group of malignant diseases with different prognosis – dysplasia of myeloid lineage.

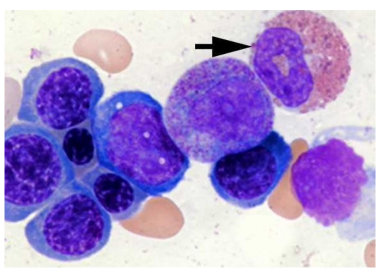
In BM - blasts bellow 20 % and dysplastic features (hypogranular cells, cells with atypical shape of nucleus, hypergranular cells, cells with abnormal plasma)

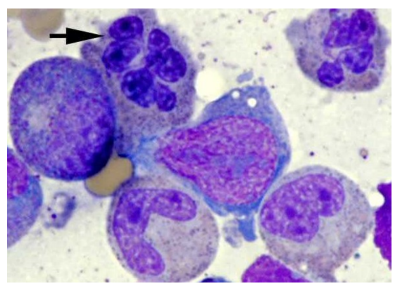
The only curative option is BMT/PBSCT in high risk patients.

Patients asyptomatic or without donor - only symptomatic treatment or watch and wait strategy.

MYELODYSPLASTIC SYNDROMES - dysplastic features







MDS – classification I

MDS type	Dysplasia	Cytopenia	Ring sideroblasts	Blasts in peripheral blood	Blasts in bone marrow	Cytogenetics
MDS with single lienage dysplasia (MDS-SLD)	1	1 or 2	<15%, < 5%	< 1%, no Auer rods	<5%, no Auer rods	Any except of del(5q)
MDS with mixed lineage dysplasia (MDS-MLD)	2 or 3	1 - 3	<15%, < 5%	< 1%, no Auer rods	<5%, no Auer rods	Any except of del(5q)
MDS with ring sideroblasts (MD	S-RS)					
MDS-SLD-RS	1	1 or 2	≥ 15%, ≥ 5%*	< 1%, no Auer rods	<5%, no Auer rods	Any except of del(5q)
MDS-MLD-RS	2 or 3	1 - 3	≥ 15%, ≥ 5%*	< 1%, no Auer rods	<5%, no Auer rods	Any except of del(5q)
MDS with isolated del(5q)	1-3	1-2	No or few	< 1%, no Auer rods	<5%, no Auer rods	del(5q) or 1 more except of -7 or del(7q)

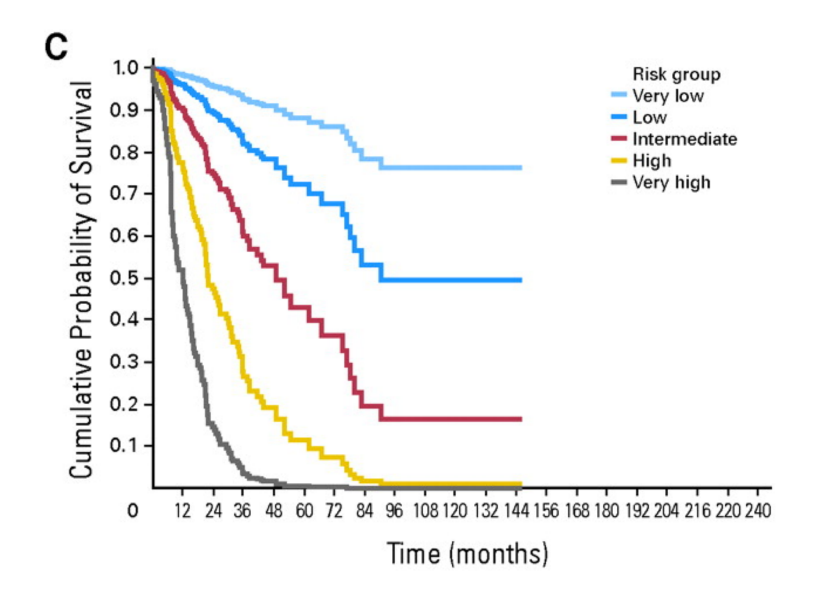
MDS – classification II

MDS type	Dysplasia	Cytopenia	Ring sideroblasts	Blasts in peripheral blood	Blasts in bone marrow	Cytogenetics
MDS with exces of blasts (MDS-EB)						
MDS-EB-1	0-3	1-3	No or few	2-4%, no Auer rods	5-9%, no Auer rods	Any
MDS-EB-2	0-3	1-3	No or few	5-19%, or Auer rods	10-19%, or Auer rods	Any
MDS unclassificable (MDS-U)						
With 1% of blasts in PB	1-3	1-3	No or few	1%, no Auer rods	< 5%, no Auer rods	Any
With 1 lineage dysplasia and pancytopenia	1	3	No or few	< 1%, no Auer rods	< 5%, no Auer rods	Any
With cytogenetic abnormality	0	1-3	< 15%	< 1%, no Auer rods	< 5%, no Auer rods	MDS typical feature
Refractory cytopenia in childhood	1-3	1-3	No	< 2%	< 5%	Any

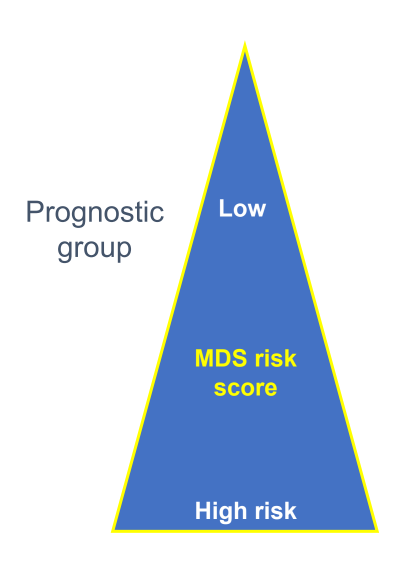
MDS - prognosis

	Score				
Prognostic marker	0	0.5	1.0	1.5	2.0
Bone marrow blasts (%)	<5	5–10		11– 20	21– 30
Karyotype	Good	Intermediate	Poor		
Cytopenia	0/1	2/3			

Score	IPSS subgroup	Median survival (years)		
0	Low	5.7		
0.5-1.0	Int-1	3.5		
1.5-2.0	Int-2	1.2		
> 2.5	High	0.4		

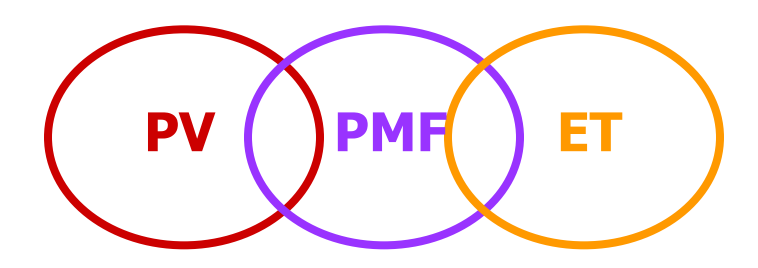


MDS THERAPY



- Supportive care, transfusions, prophylaxis of iron overload
- Erythropoietin
- Immnosupressive therapy
- Low-dose chemotherapy
- Epigenetic therapy (5-azacytidine)
- Allogeneic SCT, clinical trial

MYELOPROLIFERATIVE NEOPLASMS



MYELOPROLIFERATIVE NEOPLASMS

Proliferation of myeloid lineage

(granulocytic, erythroid, megakaryocytic)

MYELOPROLIFERATIVE NEOPLASMS

Myeloproliferative neoplasms (MPN)

Chronic myeloid leukemia (CML), BCR-ABL1⁺

Chronic neutrophilic leukemia (CNL)

Polycythemia vera (PV)

Primary myelofibrosis (PMF)

PMF, prefibrotic/early stage

PMF, overt fibrotic stage

Essential thrombocythemia (ET)

Chronic eosinophilic leukemia, not otherwise specified (NOS)

MPN, unclassifiable

Mastocytosis

Myeloid/lymphoid neoplasms with eosinophilia and rearrangement of PDGFRA, PDGFRB, or FGFR1, or with PCM1-JAK2

Myeloid/lymphoid neoplasms with *PDGFRA* rearrangement

Myeloid/lymphoid neoplasms with PDGFRB rearrangement

Myeloid/lymphoid neoplasms with FGFR1 rearrangement

Provisional entity: Myeloid/lymphoid neoplasms with PCM1-JAK2

POLYCYTHEMIA

Polycythemia is characterized by an increase of the total red cell volume.

Primary form (PV, clonal neoplastic disorder)
Secondary forms due to appropriate or inappropriate increases in levels of EPO (hemoglobins with high affinity to oxygen, high altitudes, pulomonary and heart diseases, tumours producing EPO)

PV is characterised by increases not only of the number of red cells but also of the granulocytes and platelets and splenomegaly.

POLYCYTHEMIA VERA

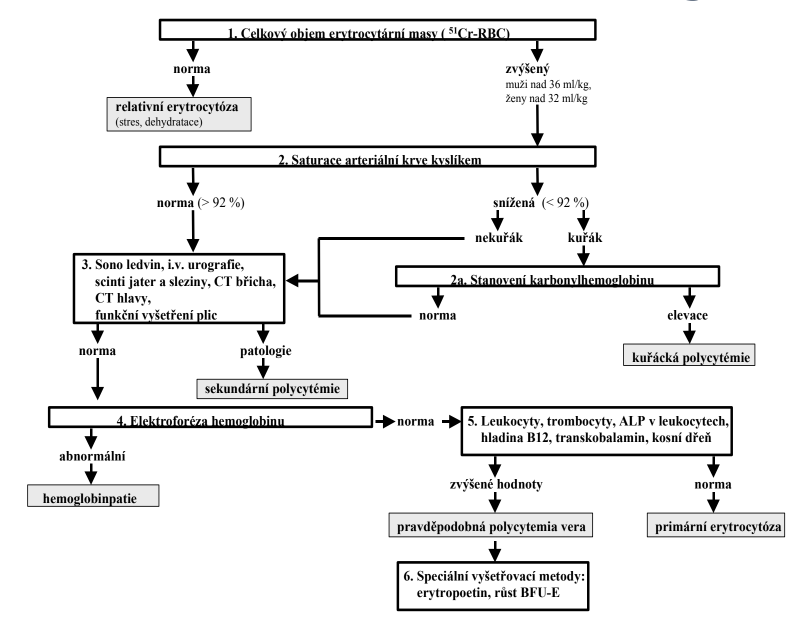
Diagnosis

Peripheral blood count
Histology of bone marrow
Total erythrocyte volum
Ertythropoietin level
JAK2 V617F mutation

We have to exclude all secondary polycythemias Secondary polycyhemias are more frequent than PV

Complications - bleeding, <u>thrombosis</u>, leukemia, bone marrow fibrosis

POLYCYTHEMIA VERA – differential diagnosis











POLYCYTHEMIA VERA

Therapy

Phlebotomy
Antiaggregant therapy of anticoagulation therapy

Interferon alpha Hydroxyurea Ruxolitinib (JAK2 inhibitor)

ESSENTIAL THROMBOCYTHEMIA

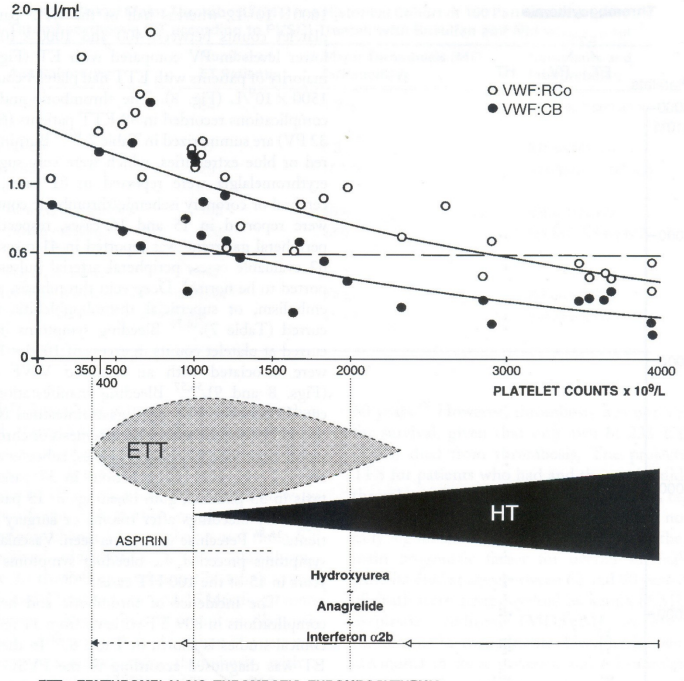
Clonal proliferation of megakaryocytes in bone marrow and incresed peripheral blood platelet count.

JAK2 V617F mutation, calreticulin mutation

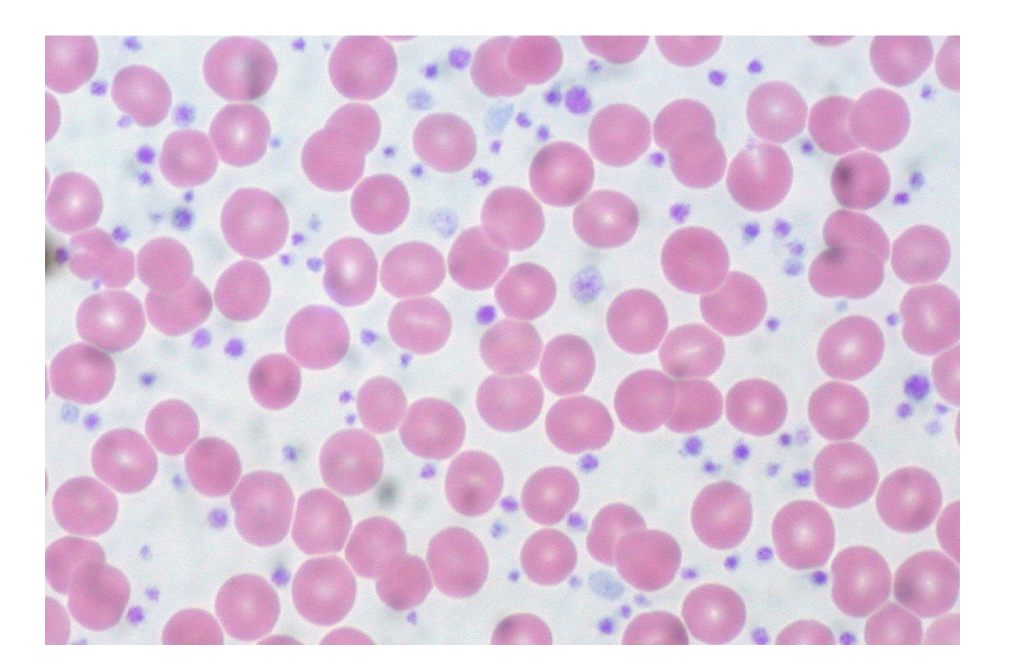
Differential diagnosis:

<u>Secondary thrombocytemias</u> (sideropenia, chronic infection, splenectomy, malignancies, bleeding, hemolysis). <u>Myeloproliferative disorders, MDS</u>

Complications - bleeding, <u>thrombosis</u>, leukemia, bone marrow fibrosis



ETT: ERYTHROMELALGIC THROMBOTIC THROMBOCYTHEMIA
HT: HEMORRHAGIC THROMBOCYTHEMIA



ESSENTIAL THROMBOCYTHEMIA

Therapy

Antiaggregant therapy of anticoagulation therapy

Interferon alpha Anagrelide Hydroxyurea

PRIMARY MYELOFIBROSIS

Clonal disorder chracterized by transformation of normal bone marrow to fibrotic and non-functional bone marrow. JAK2 V617F, CALR mutation, MPL muation

<u>Hyperplastic stage</u> - increased precurors of platelets in BM, increased WBC, RBC and PLT.

<u>Late stage</u> – fibrosis (extramedullary hematopoiesis leading to massive splenomegaly).

Prognosis – median shorter than in PV or ET.

PRIMARY MYELOFIBROSIS

Therapy

Interferon alpha Anagrelide Hydroxyurea

JAK2 inhibitors (ruxolitinib)

Supportive care

Allogeneic transplantation

Lymphoma

LYMPHOMA

Lymhoid tissue involvement (lymph nodes, other lymphoid tissue)

- Mature B cell neoplasms
- Mature T cell and natural killer (NK) cell neoplasms
- Precursor lymphoid neoplasms
- Hodgkin lymphoma
- Immunodeficiency-associated lymphoproliferative disorders

Local expansion symptoms

Systemic symptoms

Weight loss
Subfebrilia, fever
(>3 weeks)
Pruritus
Night sweat
Fatigue

Local expansion symptoms

Lymphadenopathy peripheral Lymphadenopathy mediastinal

(cough, feeling of pressure in the chest, upper vena cava syndrome)

Lymphadenopathy abdominal

(hydronefrosis, abdominal dyscomfort)

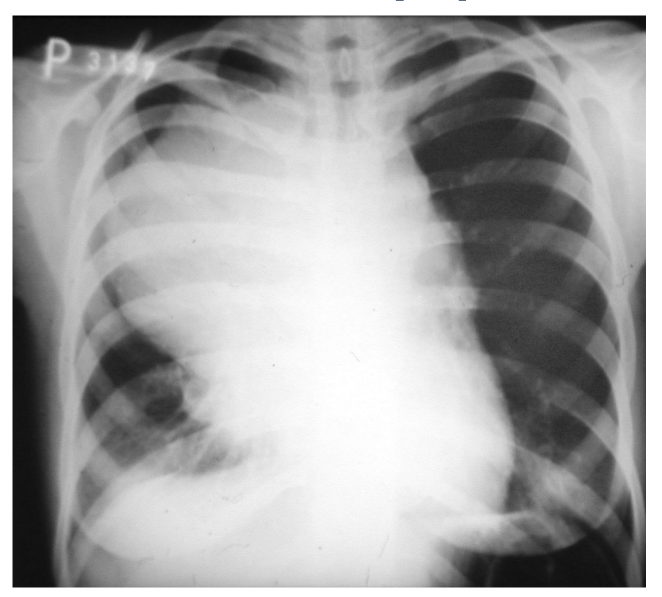
Splenomegaly

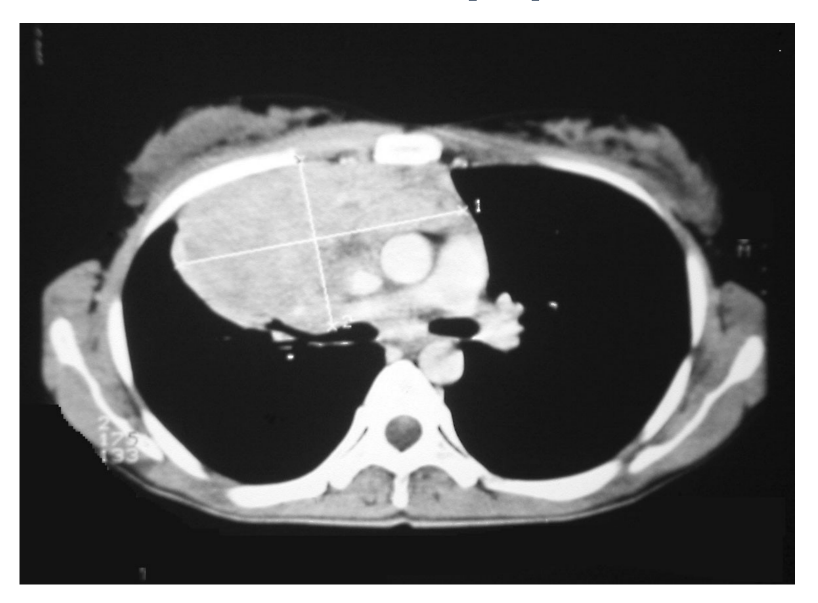
(abdominal dyscomfort, quick feeling of satiety)

Bone marrow involvment

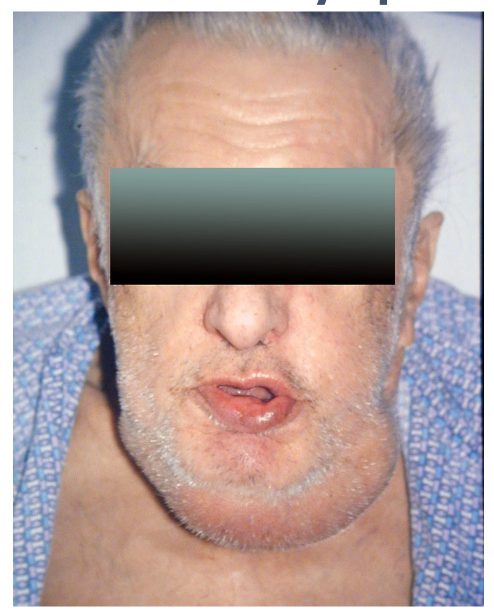
(cytopenia)

Osteolytic bone leasions

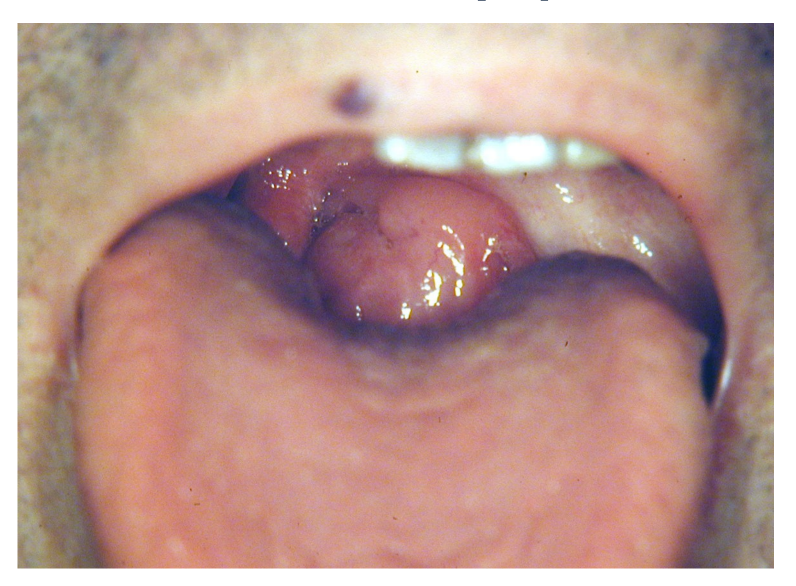




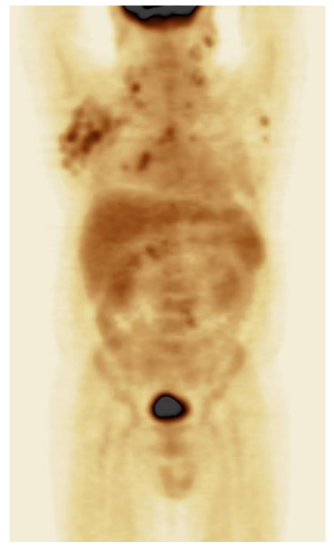














- Mature B cell neoplasms
- Mature T cell and natural killer (NK) cell neoplasms
- Lymph node involvment
- Extranodal lymphoma

Indolent NHL

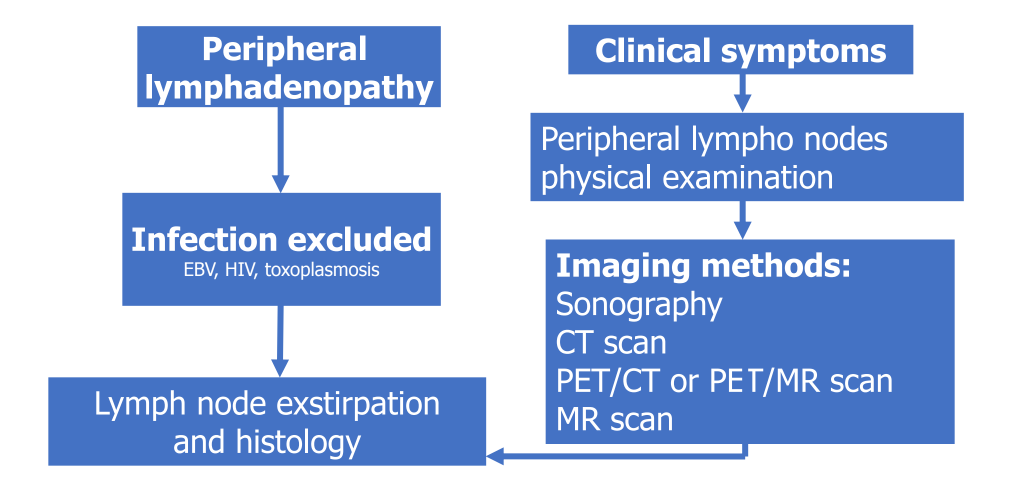
slow growth - remission possible, cure unlikely = start of treatment only with symptoms

Aggressive NHL

potentially curable, treatment start as soon as possible

Very aggressive NHL

Diagnostics



NON-HODGKIN LYMPHOMA Staging

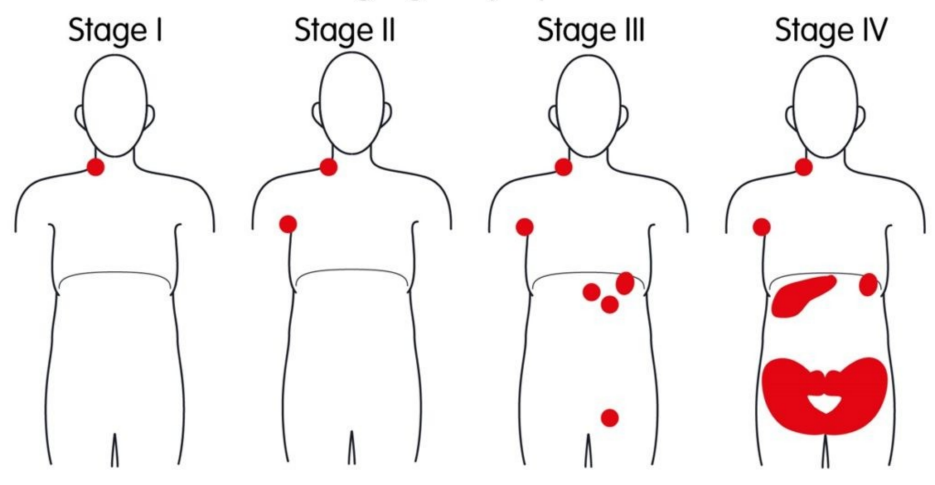
CT (neck, upper arms, chest, abdomen and pelvis)

- or MRI
- or now PET/CT, alternatively PET/MR

Trephine biopsy and bone marrow histology

Where appropriate, a specialized examination (gastroscopy, colonoscopy, lumbal puncture...)

Staging of lymphoma



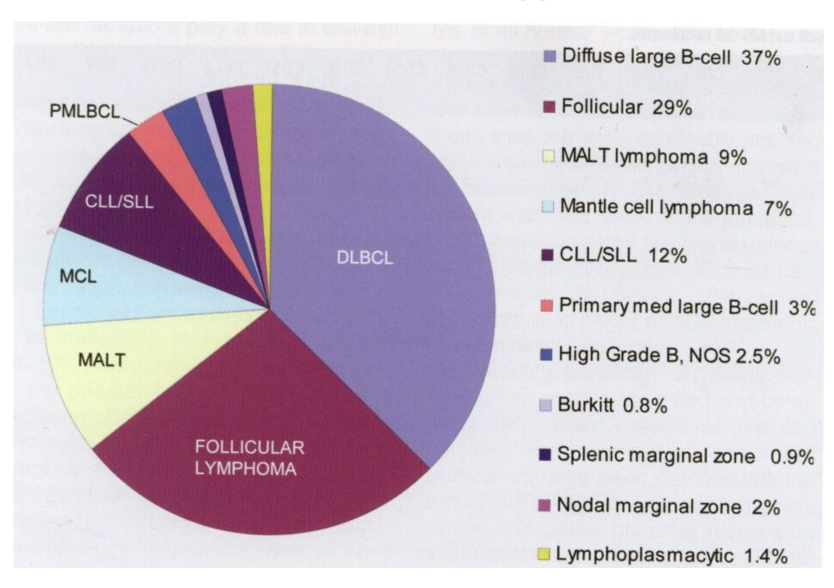
A: absence of B symptoms B: fever, night sweats, weight loss

Prognostication

Stage I and II = limited stage

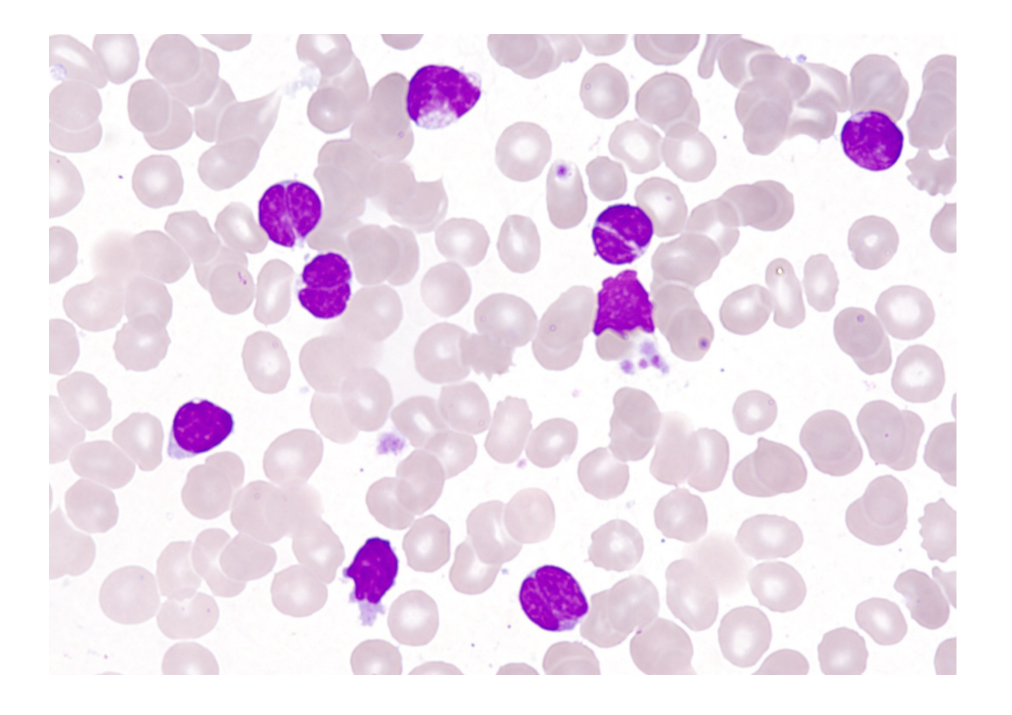
Stage III and IV = advanced stage (several prognostic indexes for adnaced stage – IPI, FLIPI...)

B-cell NHL subtypes

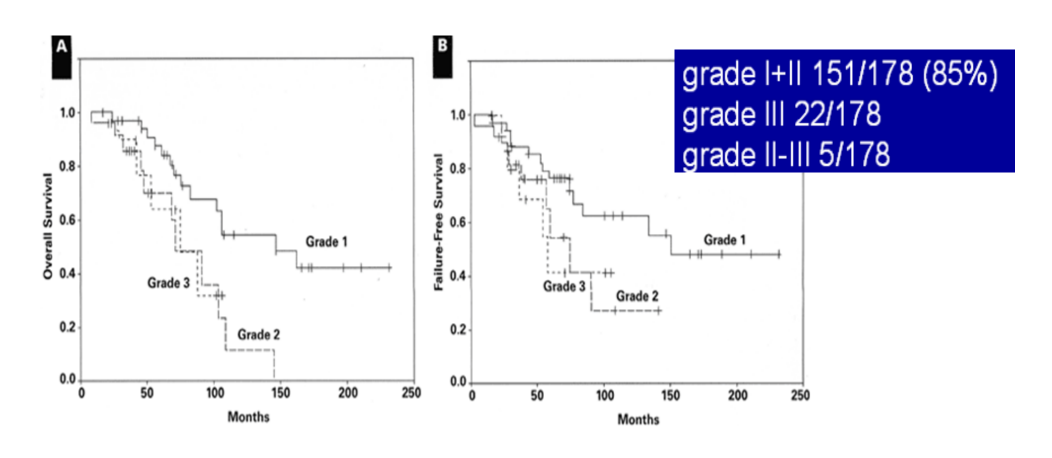


Indolent NHL - folicular lymphoma

- Survival without treatment several years in many patients
- Radiotherapy for limited stage (I.-II. st.) has curative potential
- Systemic treatment leading to remission, but no cure; repeatedly relapsed disease
- Systemic treatment in symptomatic patients only



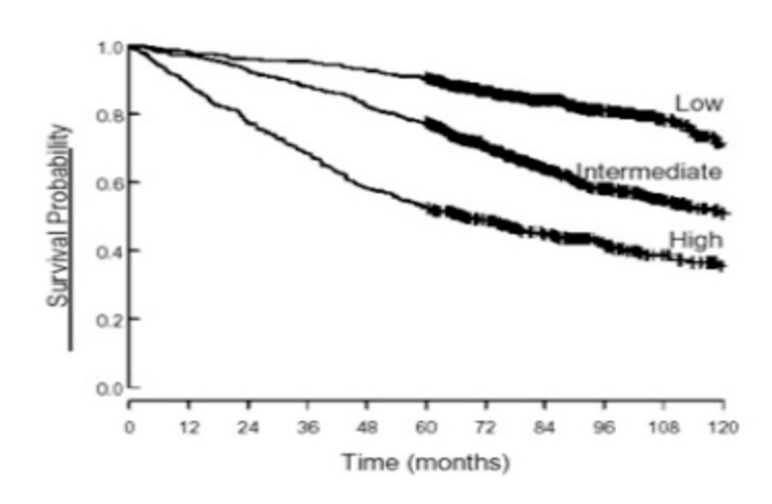
Folicular lymphoma prognosis according to histology



Folicular lymphoma prognostic index (FLIPI)

- Hemoglobin below 120 g/L
- Age over 60 years
- LDH above norm
- Stage II B or higher
- Involved lympho ode areas over 4

Low - FLIPI 0-1 Intermediate - FLIPI 2 High - FLIPI 3 and higher



Folicular lymphoma therapy

First-line therapy

- Limited FL (stadia I+II): IF RT 25-35Gy
- Advanced FL (stadia III+IV): anti-CD20 antibody + chemotherapy (R-CHOP regimen...)

Therapy of relapse

- Chemoimmunotherapy with anti-CD20 antibody +/- maintenance with monoclonal antibody
- High-dose therapy and autologous bone marrow transplant
- Allogeneic bone marrow transplant
- Radioimmunotherapy
- Radiotherapy (limited forms)

Indolent NHL - MALT lymphoma

- MALT Mucosa Associated Lymphatic Tissuse lymphoma
- Ethiologic role of antigen stimulation, *H. pylori* infection
- Majority: MALT lymphomas of stomach
- Symptoms: non-healing stomach ulcers

MALT lymphoma therapy

Limited clinical stages (I or II)

Antibiotics, radiotherapy (surgery as alternative)

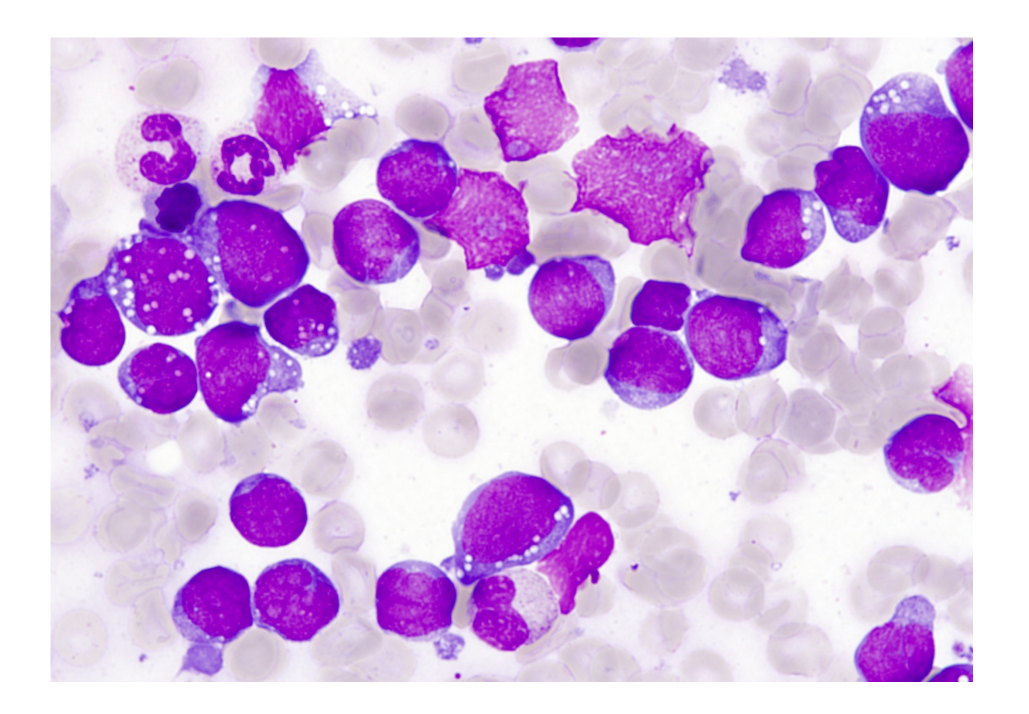
Generalized clinical stages III or IV

Chemoimmunotherapy (as in folicular lymphoma)

Aggresive NHL – principles of therapy

- Paliative
 - Mantle cell lymphoma

- Curative
 - DLBCL
 - Burkitt lymphoma



Aggresive NHL - DLBCL

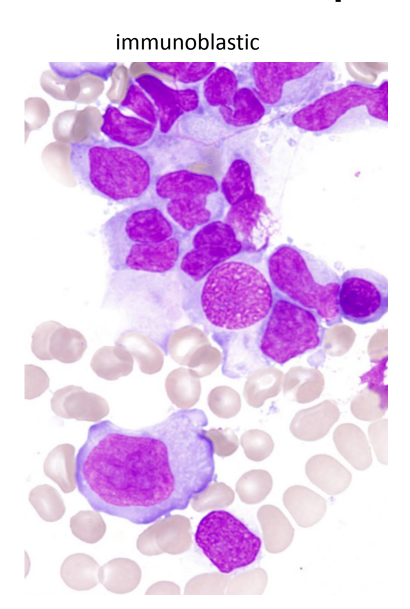
Diffuse large B-cell lymphoma
The most common lymphoma

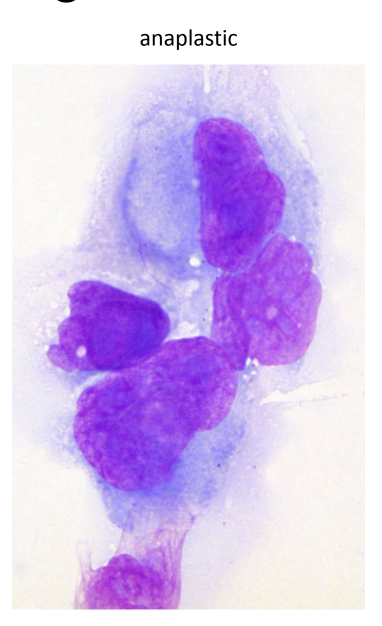
Symptoms

- Rapid local growth
- Large tumor mass
- Continuous generalization
- Frequent involvement of the central nervous system and bones

DLBCL – different morphological forms

centroblastic





DLBCL risk factors

- Age over 60 years
- Reduced physical fitness, ECOG higher than 1
- LDH level over upper limit of the norm
- Clinical stage higher than 2
- Extranodal involvement in more than 1 site

DLBCL therapy

First-line therapy

• anti-CD20 antibody + chemotherapy (R-CHOP regimen...)

Therapy of relapse

- Chemoimmunotherapy with anti-CD20 antibody
- High-dose therapy and autologous bone marrow transplant
- Allogeneic bone marrox transplant
- CAR T-cells

Very aggresive NHL

- Lymphoblastic lymphoma
 - acute lemphoblastic leukemia based protocols

- Burkitt lymphoma
 - agressive therapeutic regimens

HODGKIN DISEASE

 Lymphadenopathy with or without systemic symptoms fever, weigt loss, pruritus

Pathologic Hodgkin or RS cells

Two peaks of incidence: young adults and elederly

HODGKIN DISEASE

- Good risk goroup:
 Radiotherapy IF + 2 4 cycles of ABVD chemotherapy
- Intermediate risk group:
 BEACOP chemotherapy
- Poor prognosis:
 BEACOP
 Nivolumab
 Brentuximab vedotin (anti CD30)
 Autologous/allogeneic hematopoietic cell transplantation

HODGKIN DISEASE

Prognosis

- CR rate 95 %
- Progression free survival 90 % at 3 years

Multiple myeloma

MM

Proliferation of clonal malignant plasma cells in bone marrow

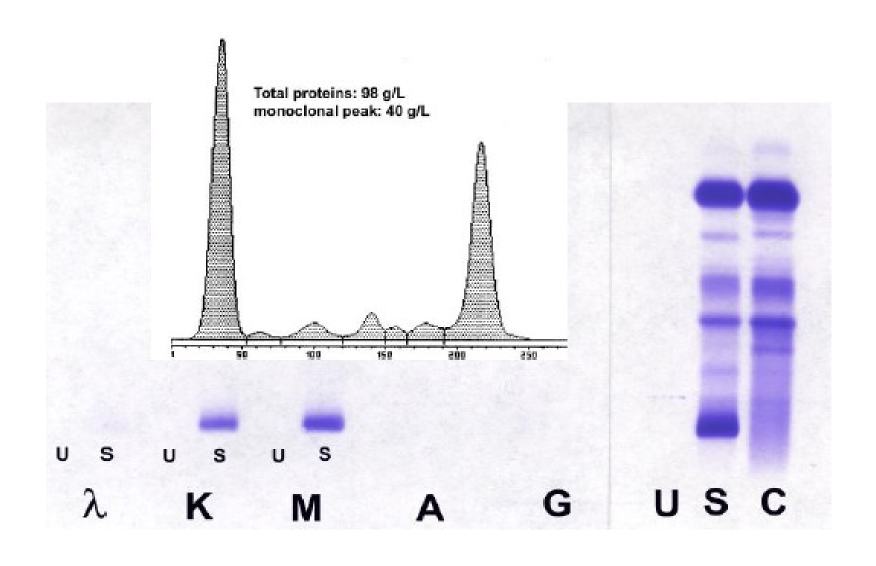
Complete monoclonal immunoglobulin molecule and/or kappa or lambda monoclonal free light chains produced by plasma cells

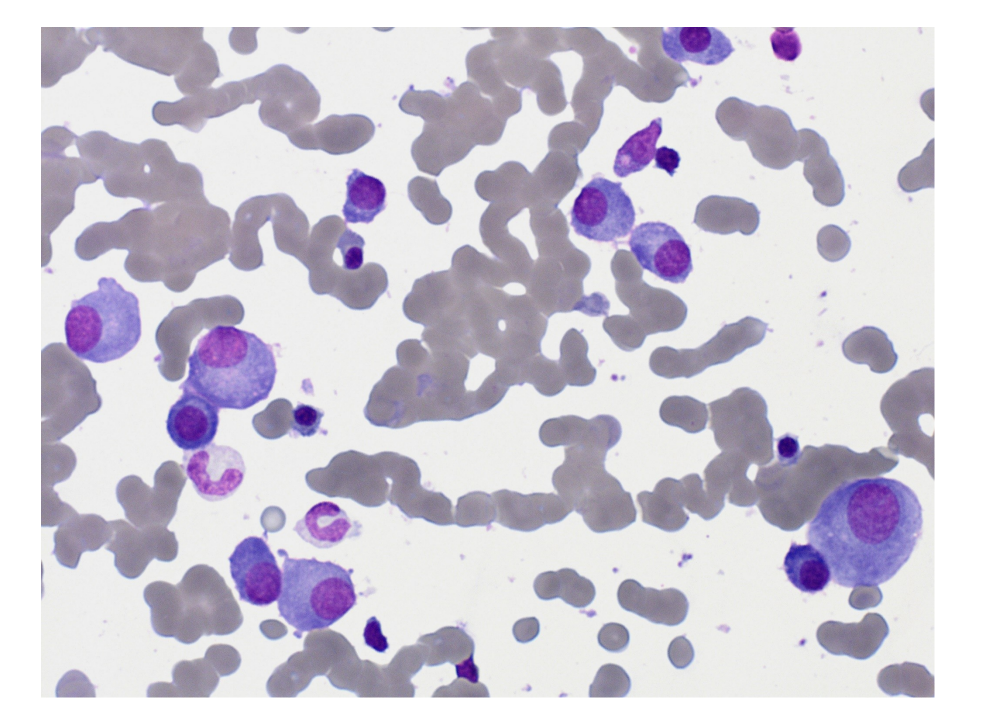
These changes lead to:

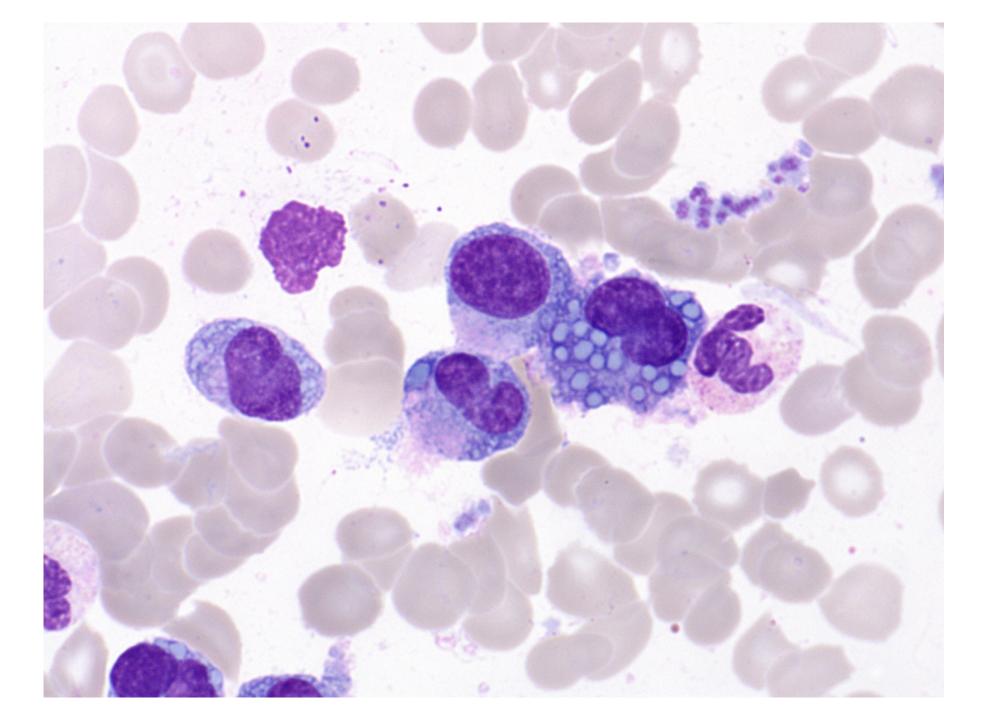
Osteolysis, osteoporosis, bone pain
Hypercalcemia
Hyperproteinemia
Renal failure
Coagulopathy
Neuropathy
Cytopenia

Incidence 4 / 100 000

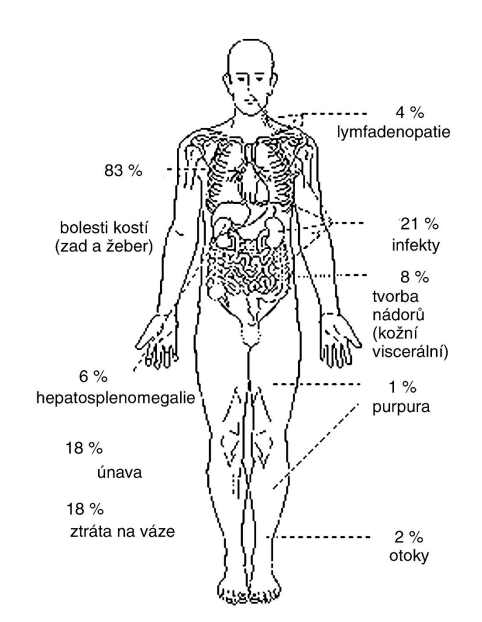
MM – immunofixation, electrophoresis, densitometry

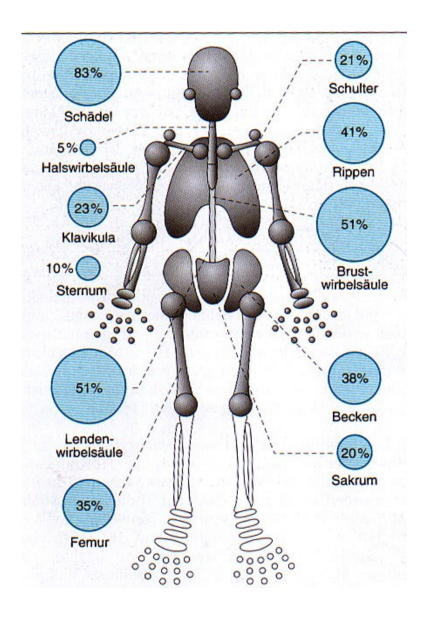




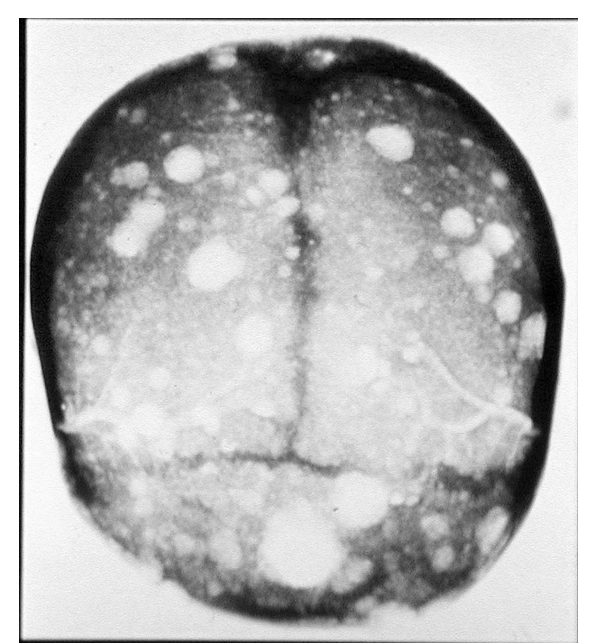
















Diagnostics:

Monoclonal immunoglobulin (or light chains) in peripheral blood and urine

Bone marrow histology/cytology

Imaging methods: X-ray, MR, PET/CT

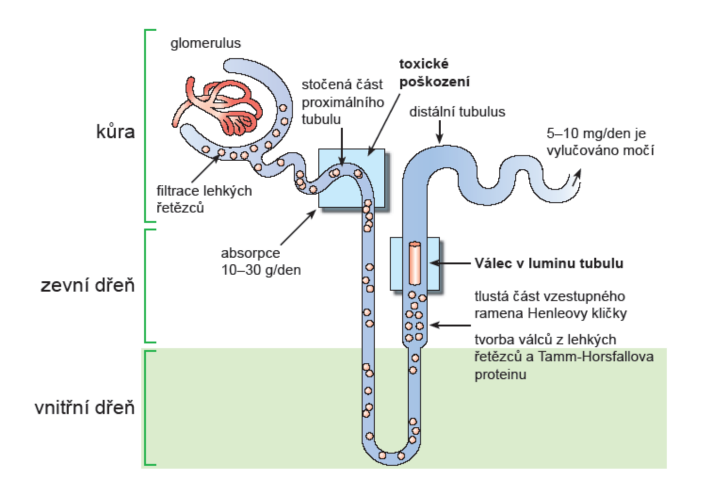
Serum immunoglobulins

Serum calcium level

Serum protein

Peripheral blood count

MM – kidney failure





MM - therapy

Indication for therapy:

Symptomatic patients: cytopenia, bone leasions, hypercalcemia, kidney failure...

Drugs:

Chemotherapy (vincristine, melfalan)

Corticosteroids (dexamethasone)

Proteasome inhibitors (bortezomib, ixazomib, carfilzomib)

IMIDs (lenalidomide, pomalidomide)

Anti-CD38 (daratumumab)

High-dose therapy + autologous hematopoietic stem cell transplantation

MM – supportive care

- Bisfosfonates
- Dialysis
- Plasmaferesis
- Radiotherapy
- Pain killers
- Prophylaxis of infection
- Tranfusions



positive mutation