

Clinical Genetics

Renata Gaillyová



Clinical Genetics

- schoolroom, University Hospital Brno, Children's Hospital Brno, Černopolní 22
- Monday 8:00-12:30 Clinical Genetics
- Tuesday 8:00-10:30 DNA Diagnostics
10:00-12:30 Visit on the Department of Medical Genetics (Children's Hospital, Černopolní 9, Building G, 3th floor)

Bring a medical mantle and shoes in the laboratory,
please use the changing rooms for students

- Wednesday 8:00 - 10:00 Clinical Cytogenetics
- Writing test - terms are in IS

Clinical genetics

- Dept. of medical genetics
- Genetic prevention
- Genetic diseases
- Patients on the departement of clinical genetics
- Genetic counselling
- Chromosome abnormalities
- AD, AR, XR inheritance, disorders
- Multifactorial inheritance
- Teratogenes, Environmental hazards
- Prenatal diagnosis
- Reproductive genetics
- Hereditary cancer

Dept. of Medical genetics

- Genetic ambulance

genetic counselling

- Laboratory part

- Cytogenetic laboratories

Prenatal cytogenetics

Postnatal cytogenetics

Oncocytogenetics

Molecular - cytogenetics

- Lab. for DNA and RNA analysis
(clinical genetics and oncogenetics)

Characteristic of Medical Genetics

- Preventive Medicine
- Interdisciplinary cooperation
- Information from genetics (disease, possibilities of testing, prenatal analysis)
- Voluntary choice for patients
- Informed agreement

Primary genetic prevention

- **Before pregnancy**
- Folic acid (cca 0,8 mg/day, 3+3 months)
- Vaccination (rubella)
- Genetic counselling
- Contraception, family can opt for adoption or donor of gamets (oocytes, sperm)
- Pregnancy planning
- Rediction of environmental hazards (drugs, radiation, chemicals...)

Reproduction of the optimal age

- In women increases the risk of accidental congenital chromosomal aberrations in the offspring
- In men may increase the risk of de novo mutations in some monogenic diseases (Neurofibromatosis I, Achondroplasia..)

Prevention of spontaneous and induced mutations

- **Healthy Lifestyle**
- **The restriction of harmful substances - drugs, environmental hazards**

Vaccination, infection prevention

- Prevention of rubella embryopathie

Prevention of congenital toxoplasmosis

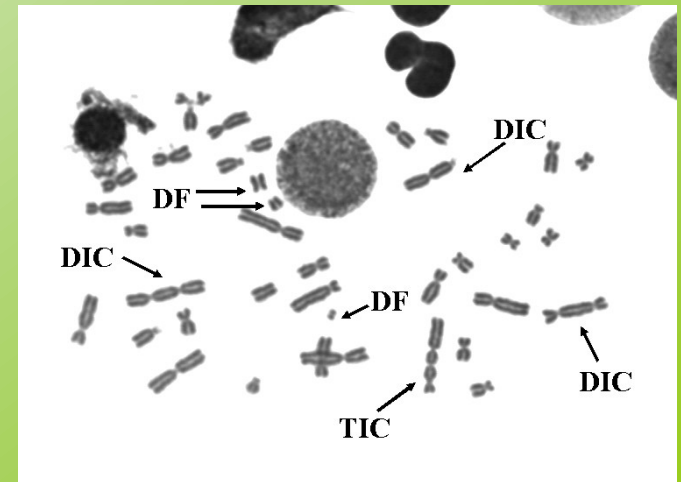
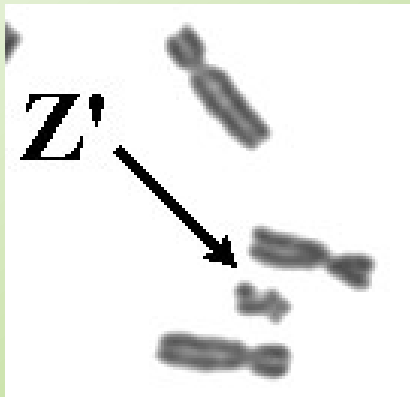
- Testing for infectious disease risk in mothers (CMV, varicella-zoster virus, ...)

Vitamin prevention of neural tube defects, anterior abdominal wall defects, clefts

- Folic acid at a dose of 0.8 mg daily (twice the dose in non-pregnant) for 3-6 months prior to conception and till the end of 12. week of pregnancy

Examination of acquired chromosomal aberrations

- Preventive examinations of persons exposed to environmental risks at work or persons with risk of long-term therapy (immunosuppressants, cytostatics,)
- The possibility of vitamin therapy to improve repair of DNA (3-6 months)



Contraception, sterilization

- **Contraception** - temporarily prevents conception in the limited impact of risk (treatment)
- **Sterilization** - the long-term inhibition of pregnancy in a high risk of disease in the offspring (Hereditary disease)

Adoption

- Alternative family care as an option at high genetic risk families

Donation

- of sperm, oocytes and embryos
- reduction in high genetic risk
- reproductive problems

Secondary genetic prevention

- Prenatal diagnosis
- Prenatal screening
- Prenatal tests
- Genetic counselling
- Termination of pregnancy (the law in Czech Republic- end of 24. week of gestation)
- Postnatal screening
- Newborn screening

Genetics diseases

- **Chromosome abnormalities**
 - about 0,6 - 0,7%

 - **Monogen diseases**
 - about 0,36%
- (study in 1 000 000 newborns)
- most then 90% of monogen diseases occur in childhood

 - **Multifactorial (polygenic or complex) disorders**
 - Occur in about 80% in the population

Patients on genetic departments

- Dead person
- Adults
- Pregnant women
- Fetuses
- Children

Patients on genetic departments

- Positive family history (chromosome abnormality, congenital malformations, mental retardation, diseases...)
- Pregnant women with encrease risk for the fetus
- Infertility - sterility, repeated fetal loss
- Donors (gamets)
- Patients with tumours

Children

- **Congenital malformations**

Children

- Suspicion of mongenic hereditary diseases or inherited metabolic disorders and their families

Children

- Suspicion on congenital chromosom aberations (children with congenital malformations, abnormal face, atypical visage, pre- or postnatal growth retardation, premature birth)

Children

- early or delayed puberty
- Malformations of the external or internal genitalia
- Low or high figure

Children

- Preventiv genetic examinatioun before adoption

Children or adults

- **Mental retardation**
- **Psychomotor retardation**
- **Developmental delay**

Children and adults

- Gender identity disorder

Children and adults

- people with long-term exposure to environmental pollutants
- (alcohol, cigarettes, drugs, radiation)

Children and adults

- patients with suspected hereditary cancer
- patients with cancer (sporadic occurrence)

Adults

- Donors of gametes
(preventive tests)

Adults

- Related partners
(increased risk for hereditary disease with
AR inheritance)

adults

- Infertility
- Repeated spontaneous abortions

Pregnant women

- With unfavorable family history

Pregnant women

- with adverse pregnancy history (chronic diseases with established therapies, acute disease in early pregnancy - temperature, drugs, X-rays, CT, vaccinations, toxoplasmosis, rubella, ...)

Pregnant women

- Prenatal biochemical screening
(Pathological results)

Pregnant women

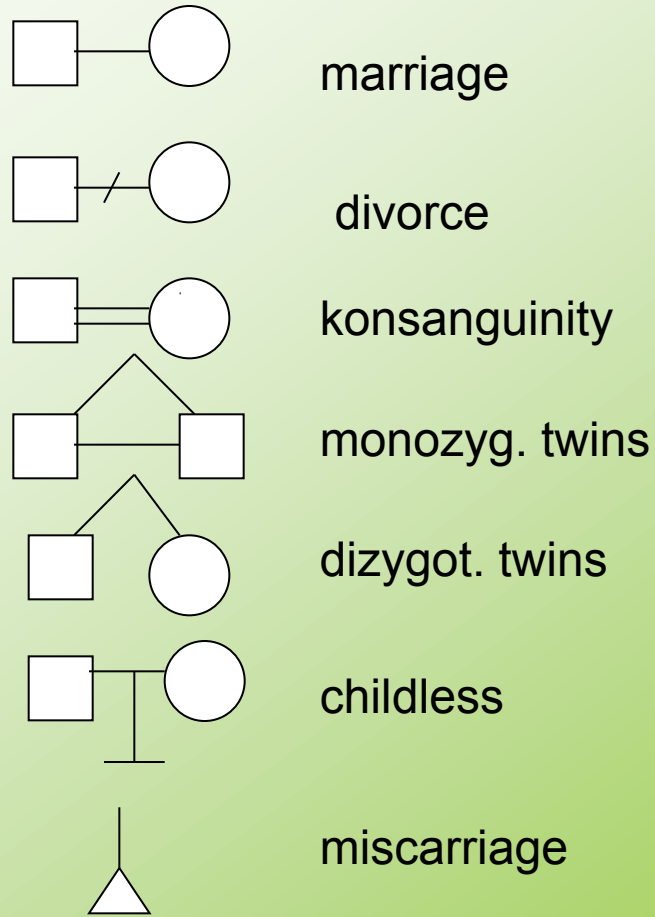
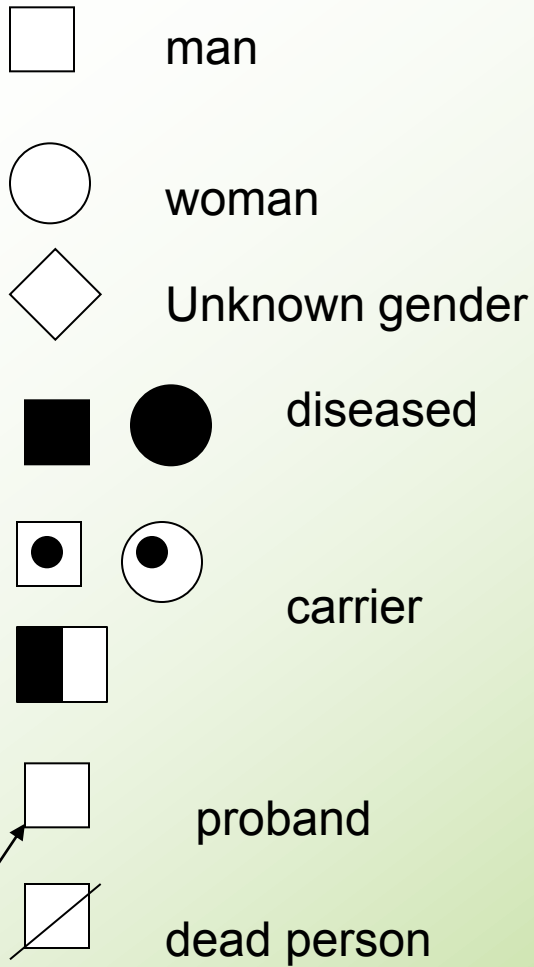
- Ultrasound prenatal screening
- pathological results
- Congenital malformations in the fetus
- Risk of chromosomal abnormality in the fetus

Genetic counselling

- Anamnesis
- Family history
- Pedigree analysis
- Examination of the patient
- Laboratory analysis
- Other examinations - neurology, psychology, hematology, CT, MRI ...

Three-generation pedigree

- Patient
- Siblings
- Children siblings
- Parents
- Parents siblings
- Children of parents siblings
- Parents parents



Clinical examination

Next steps

- Recommend the laboratory genetic testing
- Recommend other specialists if needed
- Require medical records
- Make photodocumentation

The result of genetic counselling

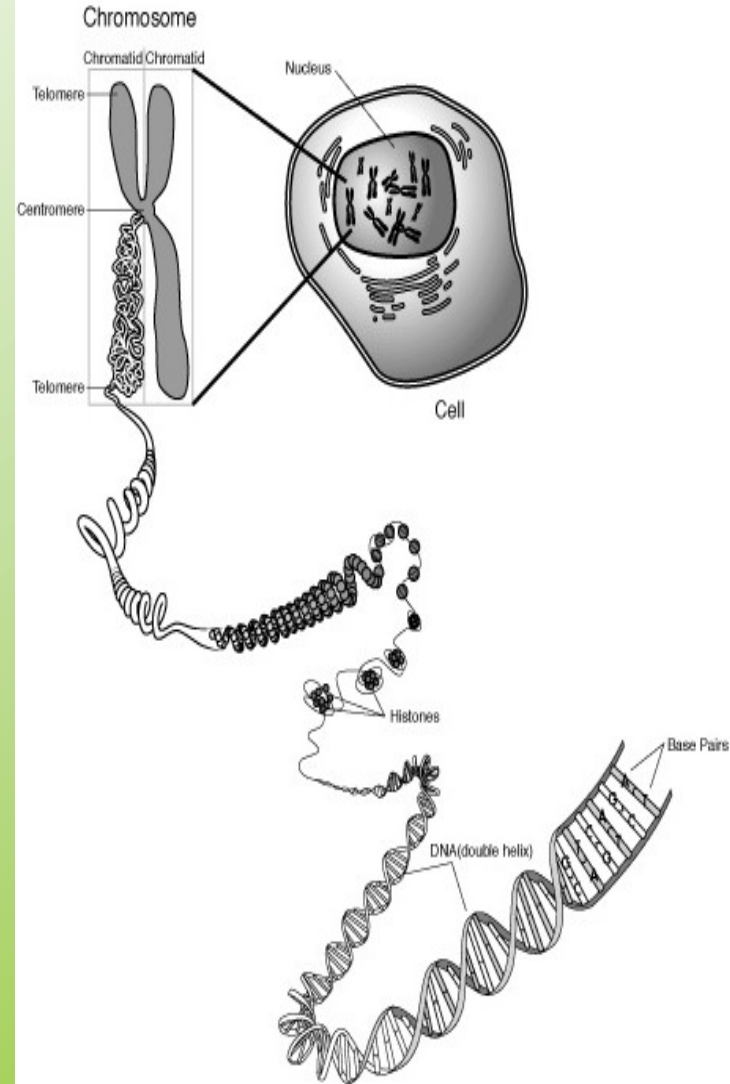
- Specify exact diagnosis (if possible)
- Determine genetic prognosis
- Is the disease hereditary?
- Type of inheritance
- Genetic risks for other family members
- Possibilities of treatment, prenatal analysis

Man



Cell

Chromosome

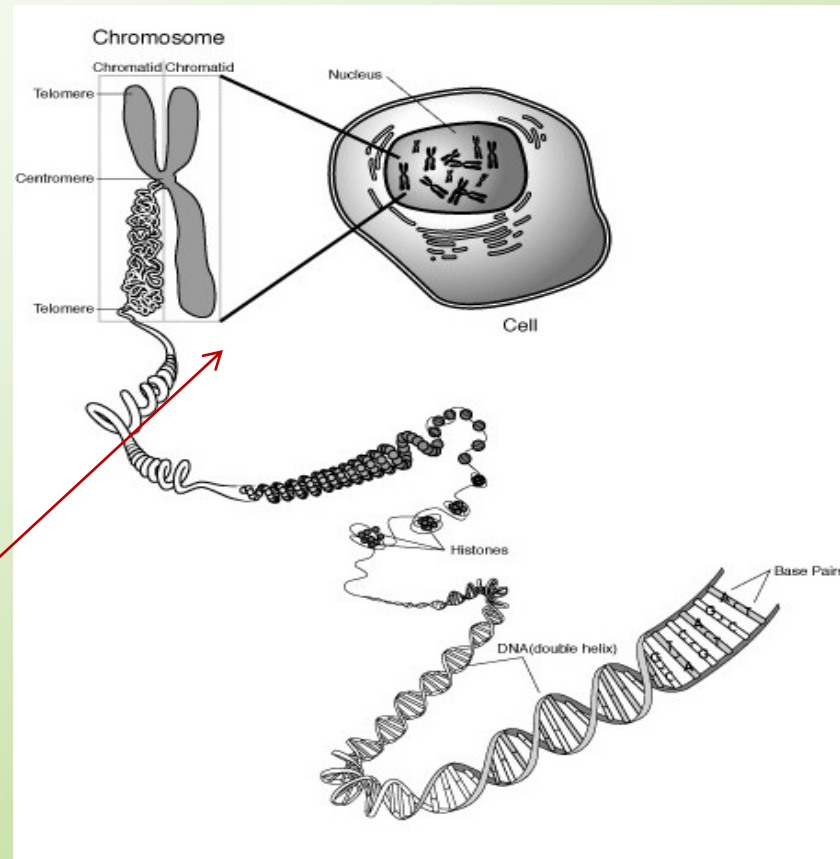


DNA

Man



Chromosome abnormalities



0,6-0,7% live born

Congenital chromosome abnormalities

- Autosomes
- Gonosomes

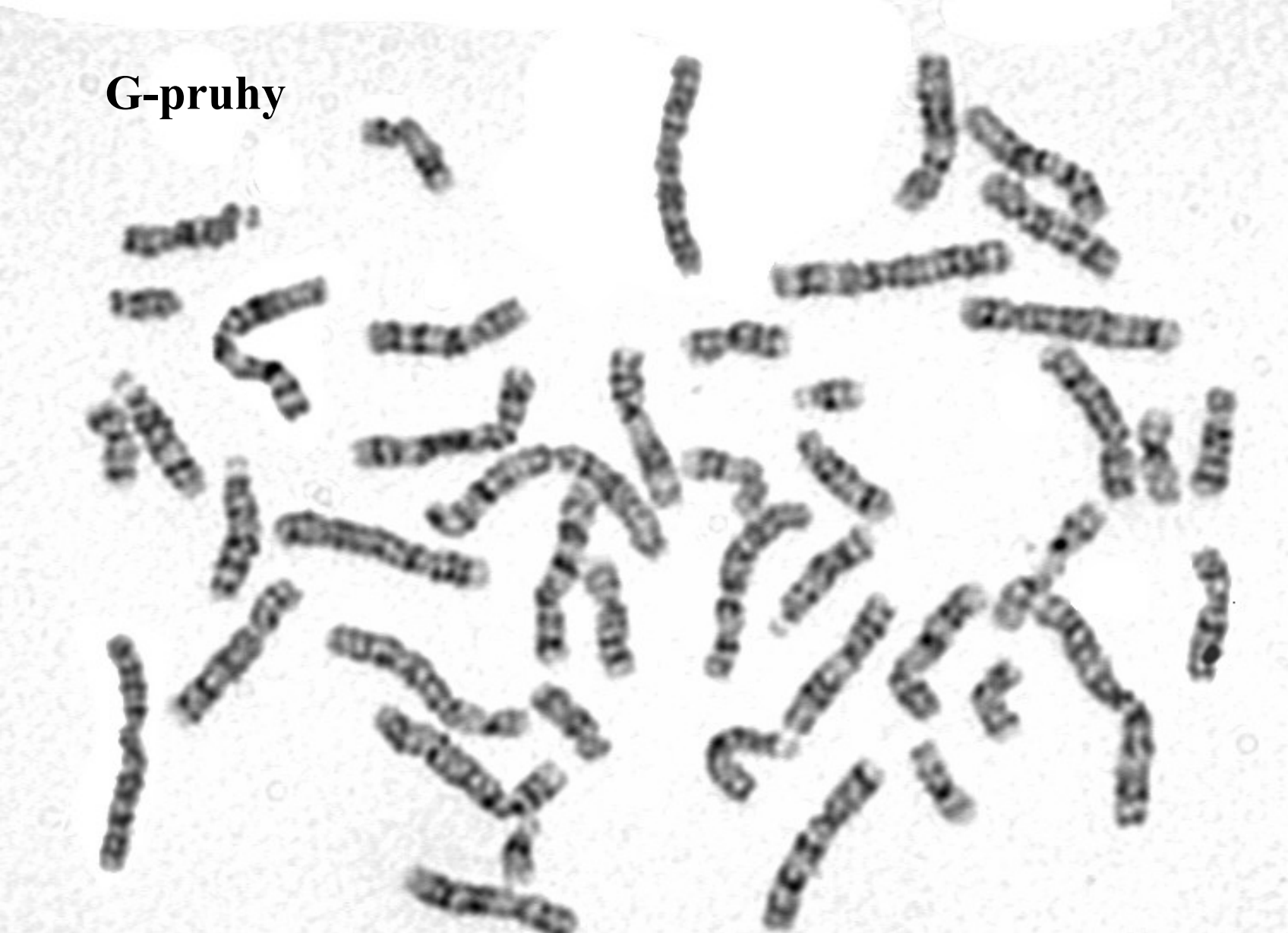
- Numerous
- Structural

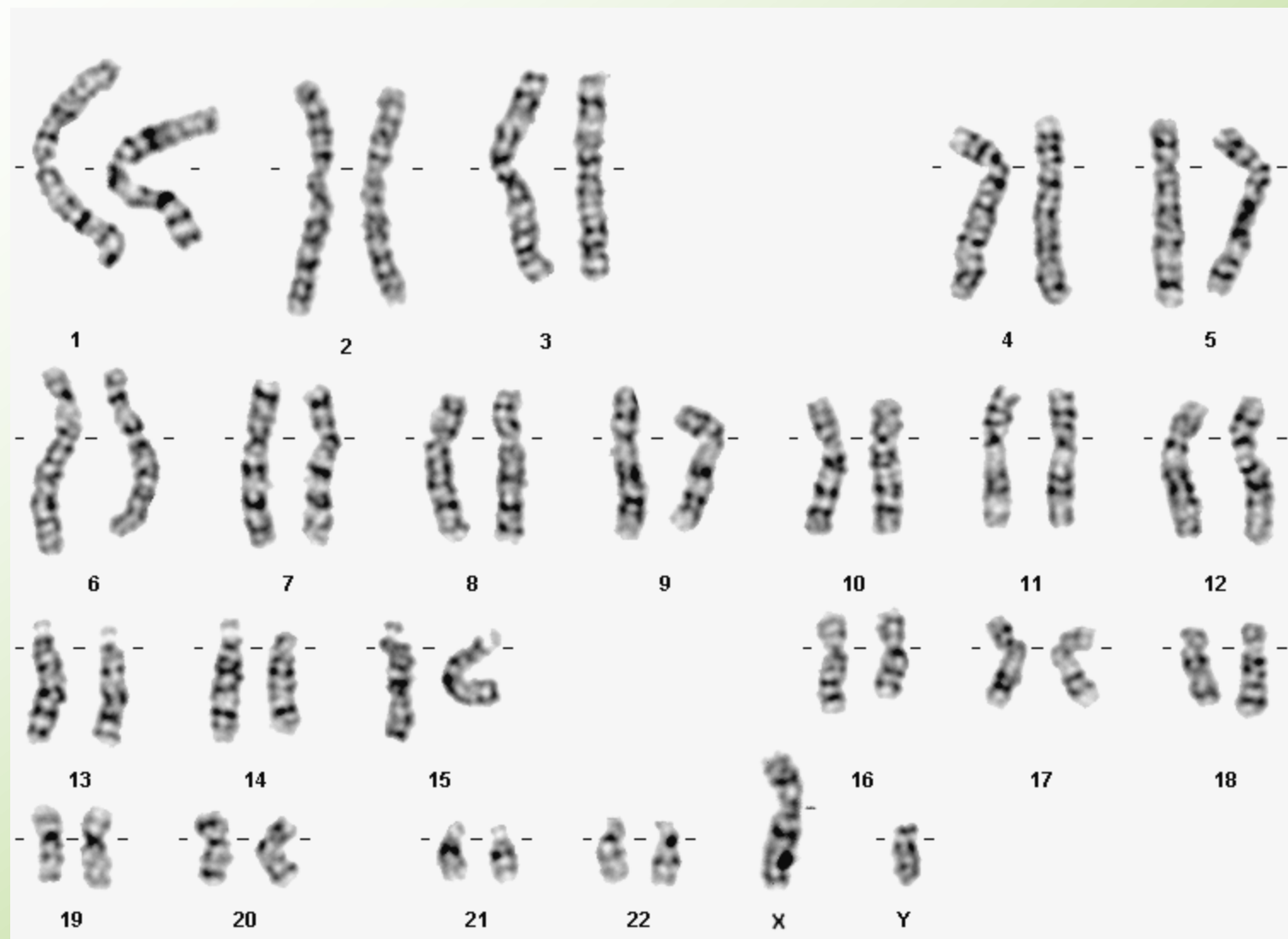
- Balanced
- Unbalanced

Populations frequency

Trisomy 21	1,5 per 1000 live births
Trisomy 18	0,12
Trisomy 13	0,07
Klinefelter syndrome	1,5
Turner syndrome	0,4
XYY syndrome	1,5
XXX syndrome	0,65

G-pruhy





Chromosome abnormalities in spont. abortions

All spont. abortions	50 %
Up to 12 weeks	60 %
12-20 weeks	20 %
stillbirths	5 %
trisomies	52 %
45,X	18 %
Translocations	2 - 4%

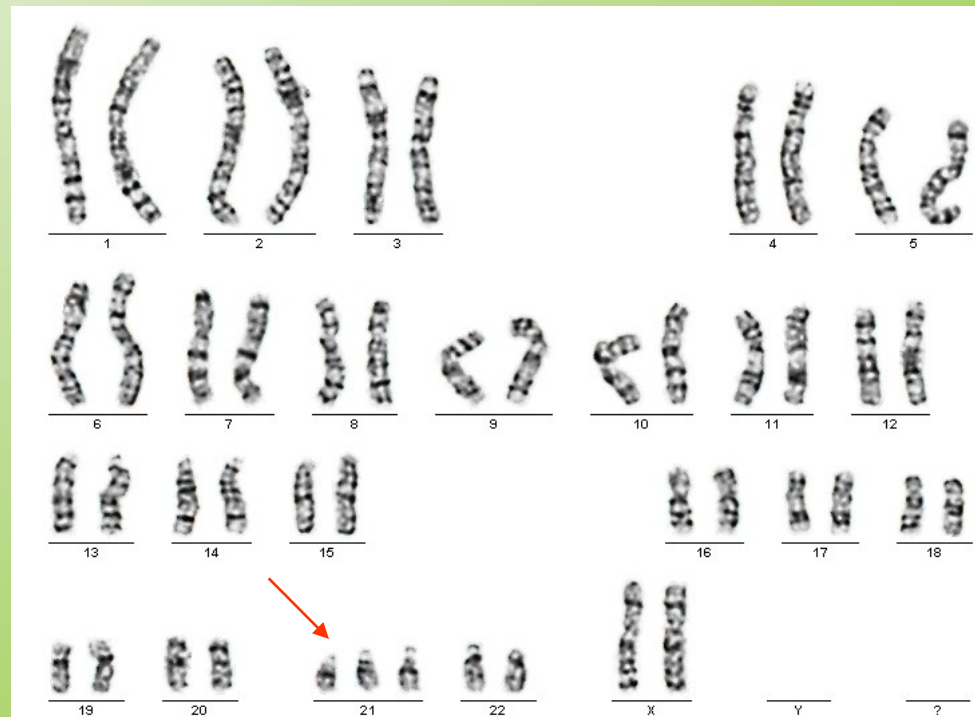
Maternal age and chromosome abnormalities in AMC (per 1000)

<u>years</u>	<u>+21</u>	<u>+18</u>	<u>+13</u>	<u>XXY</u>	<u>All</u>
35	3,9	0,5	0,2	0,5	8,7
37	6,4	1,0	0,4	0,8	12,2
40	13,3	2,8	1,1	1,8	23,0
43	27,4	7,6		4,1	45,0
45	44,2			7,0	62,0
47	70,4			11,9	96,0

Risk of Down syndrom (live births)

Maternal age (years)	Risk
15	1/1578
25	1/1351
35	1/384
40	1/112
45	1/28
50	1/6

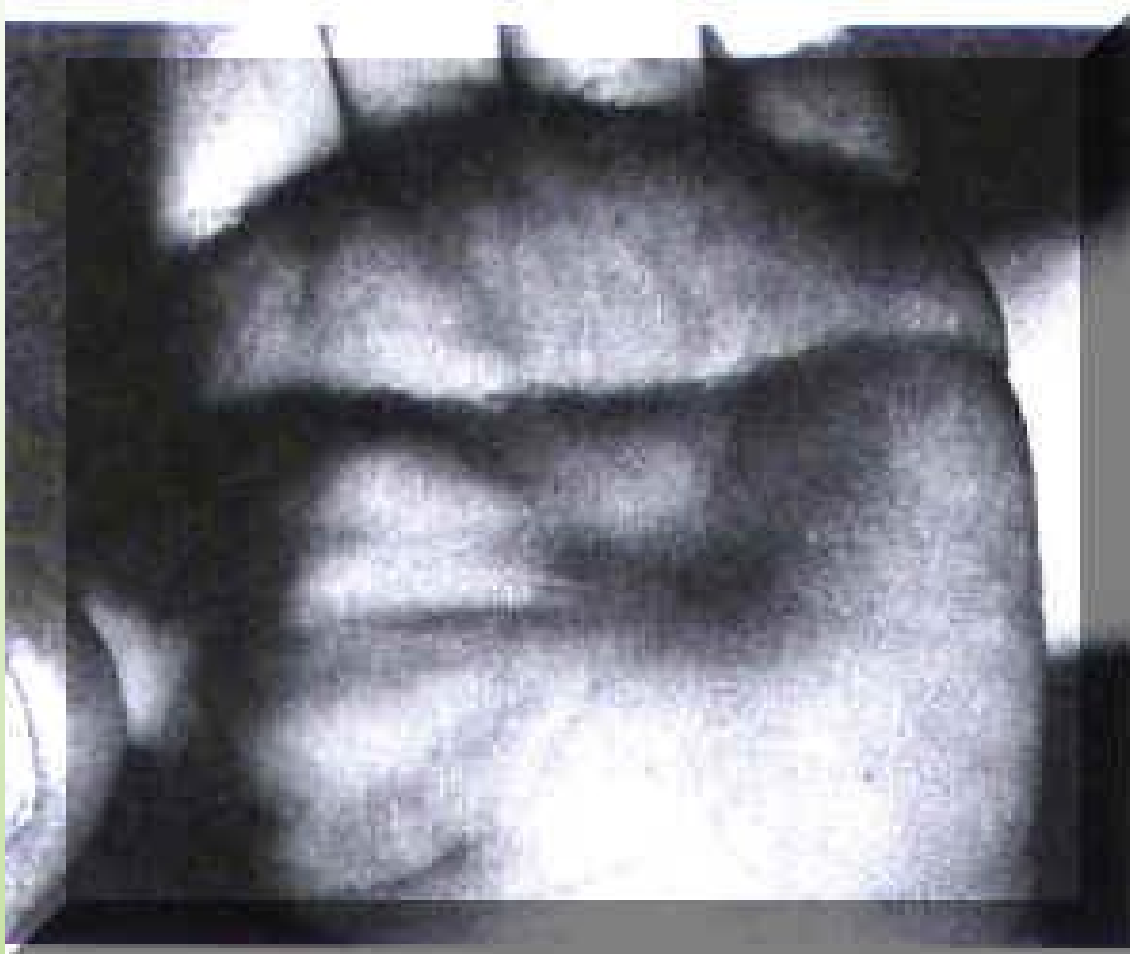
Down syndrome



Down syndrome



Typical grooves on the palms and soles

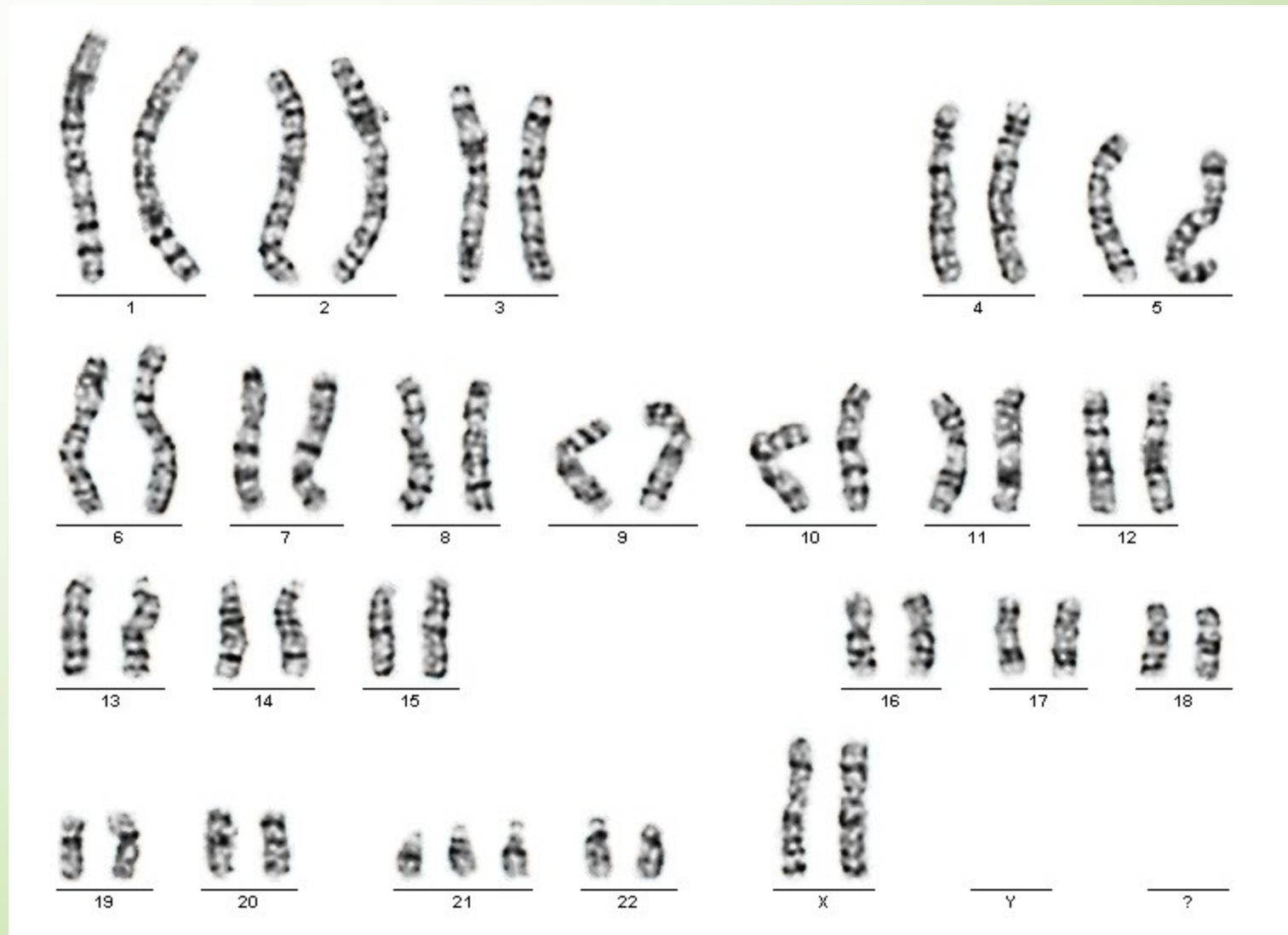


Down syndrome

- 47,XX,+21 or 47,XY,+21
- About 1/800-1000 newborns, 1/75 SA
- Hypotonia, joint laxicity, soft skin, flat face, prominent intercanthal folds, slanted palpebral fissurs, Brushfield's spots of the irides, small, down set ears, small nose, protruding tongue, simian crease in the hands (about 45%), short statue, mental retardation, congenital heart disease in about 50% of patients with DS, (atrioventricular canal)

Down syndrome (G-banding)





47,XX,+21

Happy nature

**Vision and hearing
disorders**

Hypothyroidism

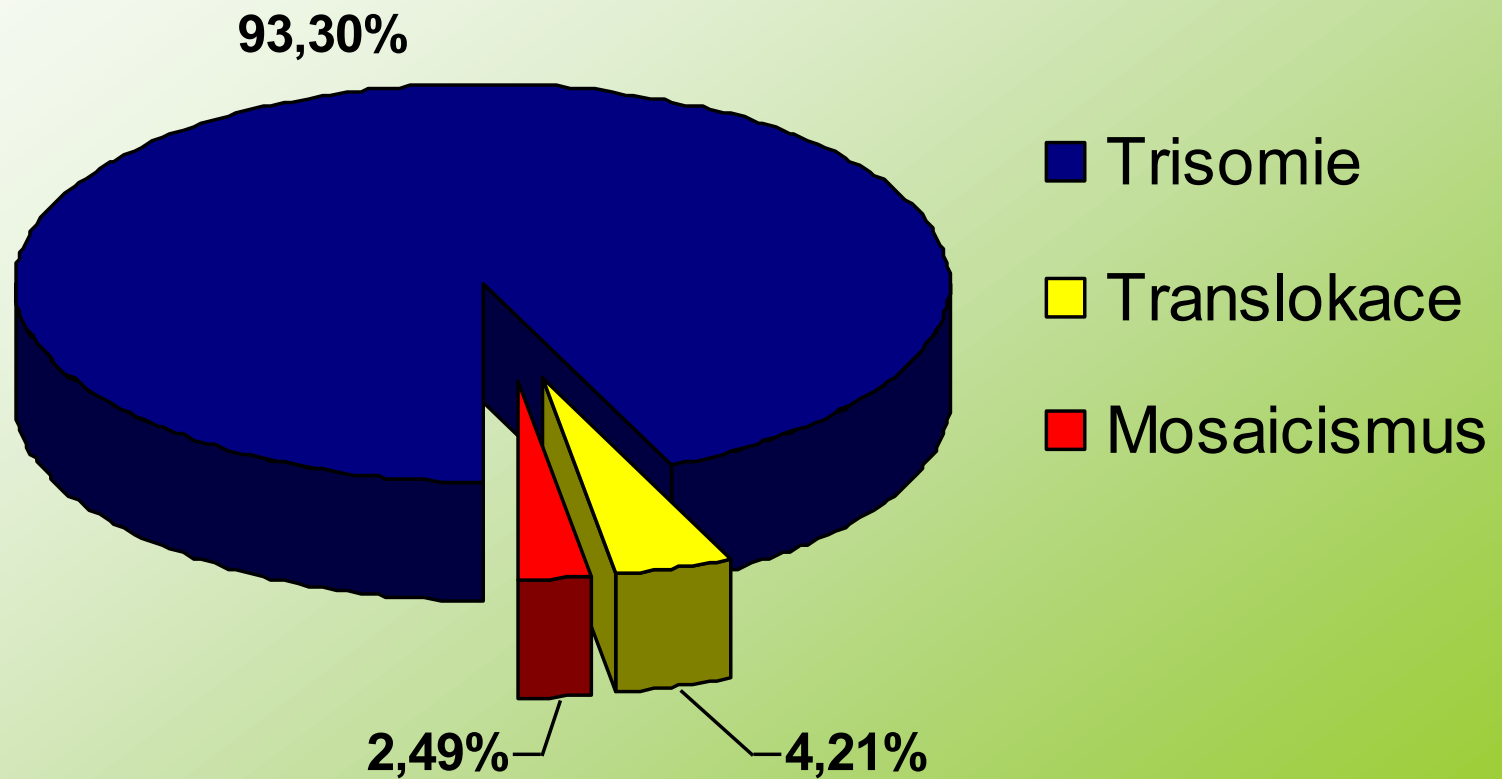
**Correlation between
positive stimulation and
height IQ**

Male sterility




**Alzheimer-like symptoms
in 40**

Cytogenetic findings in DS in Czech republic

1994 - 2001



Down syndrome- prenatal diagnosis

- I. trimester screening - combined screening
- 10.-14. week of gestation
- **Ultrasound**
- Nuchal translucency - NT ()
- (Absence of nose bone)
- **Blood**
- PAPP-A ()
- free-beta hCG ()
- Fals positive results less then 5%
- Reveals about 95% of fetuses with Down syndrome
- 1/100 - positiv - genetic counselling and karyotyping
- 1/100-1/1000 - US and genetic counselling
- 1/1000 - negativ - US

Down syndrome - prenatal diagnosis

- II. trimester screening - biochemical screening
- 16. -18. week of gestation
- AFP - alpha-fetoprotein (↓)
- total hCG - chorionic gonadotropin (↑)
- uE3 - unconjugated estriol (↓)

- Fals positive results about 5%

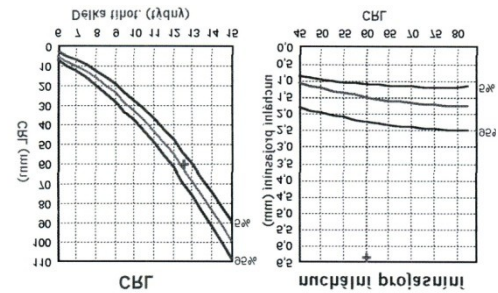
- Reveals about 70% of fetuses with Down syndrome

- 1/250 - positiv
- 1/250-1/350 - border
- 1/350 - negativ

Down syndrome - prenatal diagnosis

- Ultrasound
- 10.-14. week
- NT
- NB
- 20. week
- US- congenital heart disease and other malformations

NT+



Edwards syndrome

- 47,XX(XY),+18
- 1/5000-10 000 in newborns, 1/45 SA
- gynekotropie 4:1
- SA - 95%, death before 1 year mostly
- hypotrophy, atypical hands and feet, profil, prominent nose, small chin, congenital defects

Edwards syndrome

- 1:5000
- IUGR, hyopotrophie
- microcephalie
- dolichocephalie
- Cleft palate
- Down set ears
- micromandibula
- Hands, feets
- Other cong. malformations







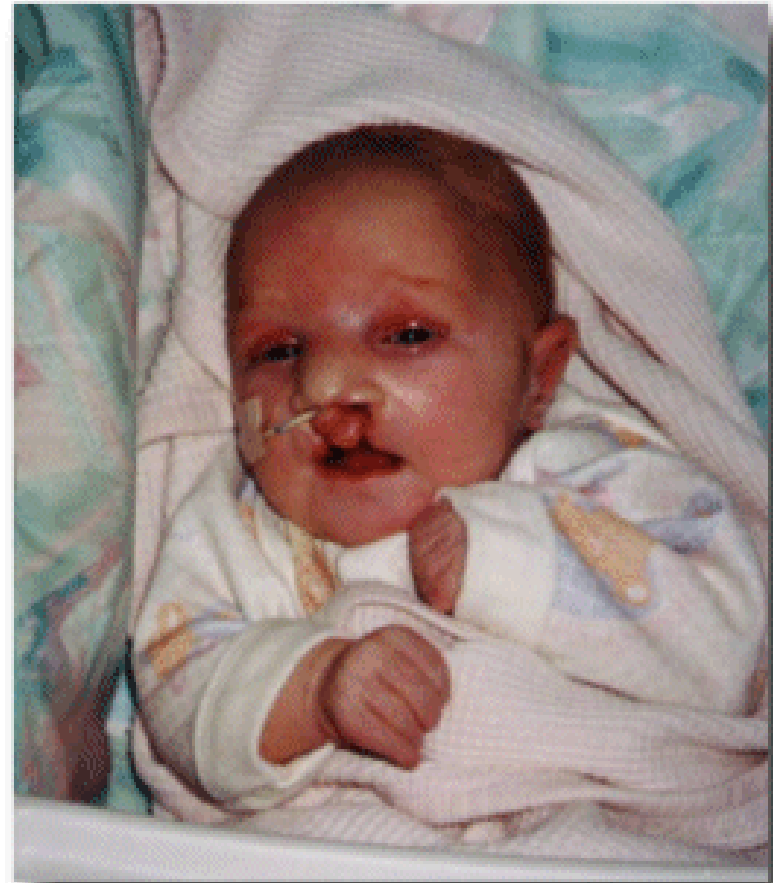
Patau syndrome

- 47,XX(XY),+13
- 1/5000-10 000 in newborns, 1/90 SA
- 95% SA
- death before 1 year mostly

- cleft lip and palate bilateral,
congenital defects (CNS, eyes,
postaxial hexadactily...)

Patau syndrome, + 13

- Microcephalie
- Trigenocephalie
- skin defects in the hairy part calva
- congenital defects of the brain (holoprosencephalie, arinencephalie)
- micro-anophthalmia
- Cleft lip, palate
- hexadactilie
- heart defects



Turner syndrome

- 45,X (in about 55%), mosaicism, structural abnormalities of X chromosome
- 1/2500 newborn girls, min. 95% SA
- prenat.- hydrops foetus, hygroma coli
- postnatal lymphedema on feet, pterygium coli, congenital heart defect coarctation of aorta, small stature, other congenital defects, hypogonadism, hypergonadotropins, sterility-infertility

Turner syndrom 45,X

- 1:2000
- hygroma colli
- hydrops
- Low weight in newborns
- Lymfoedema
- Pterygia
- Cubiti valgi
- Aortal stenosis
- Small statue
- Sterility



Klinefelter syndrome

- 47,XXY
- relatively frequent 1/600-1000 liveborn males
- tall stature
- hypogonadism, gynaecomastia
- sterility, infertility

Others gonosome abnormalities

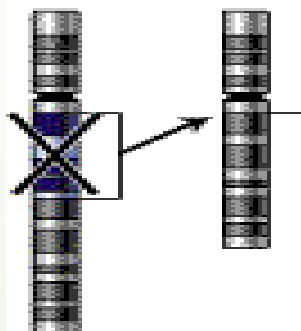
- 47,XXX
- 47,XY
- 48,XXXX
- 48,XXYY....

Structural chromosomal aberrations

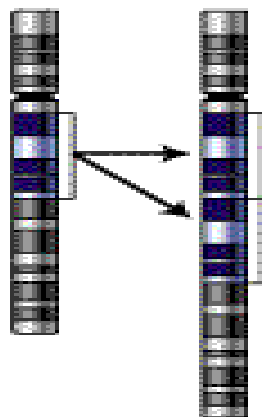
- deletion or a duplication of the genetic material of any chromosome, atypical structure - side by side to get the genetic material, which there normally is not - the effect of positional
- partial-partial deletions
- partial trisomy
- inversions, insertions, duplications

Types of mutation

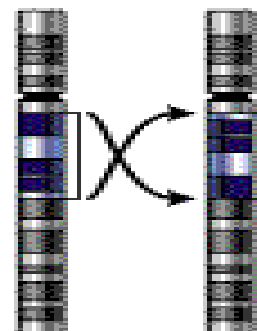
Deletion



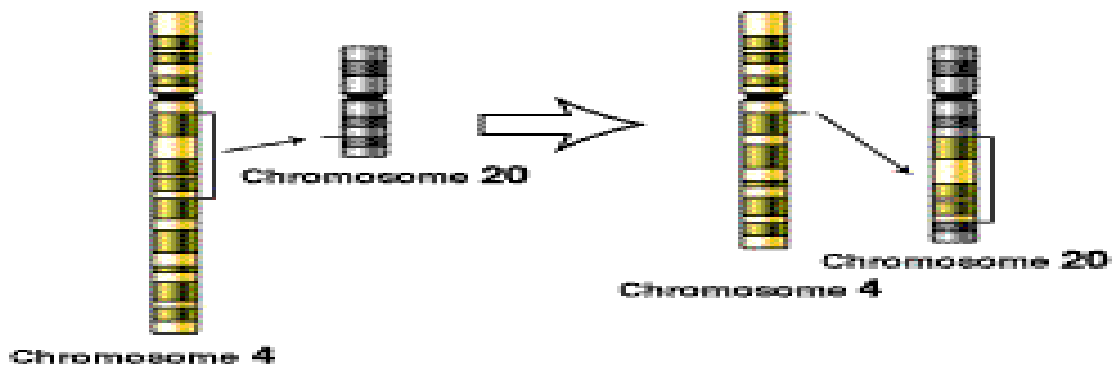
Duplication



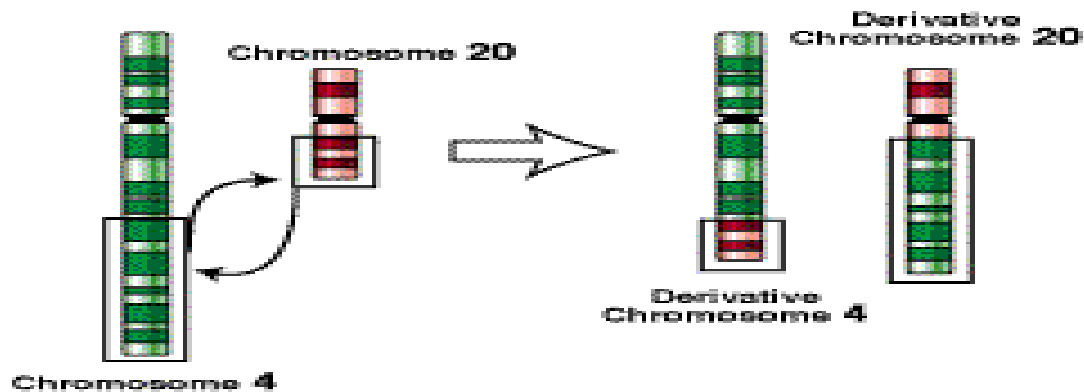
Inversion



Insertion



Translocation



Syndrom Wolf-Hirshorn

46,XX(XY),4p-

- severe mental retardation
- typical craniofacial dysmorphism - hypertelorism, pear nose, carp mouth,
- pre- and postnatal growth retardation,
- failure to thrive
- other associated developmental defects - heart, urogenital tract ...

Wolf-Hirschhorn syndrom (46,XX,4p-)

Incidence?

IUGR

Hypotonia

Charakteristic
face

Heart defects

Hypotonie

Hypotrophie

Severe mental
retardation

Syndrom Cri du chat

46,XX(XY),5p-

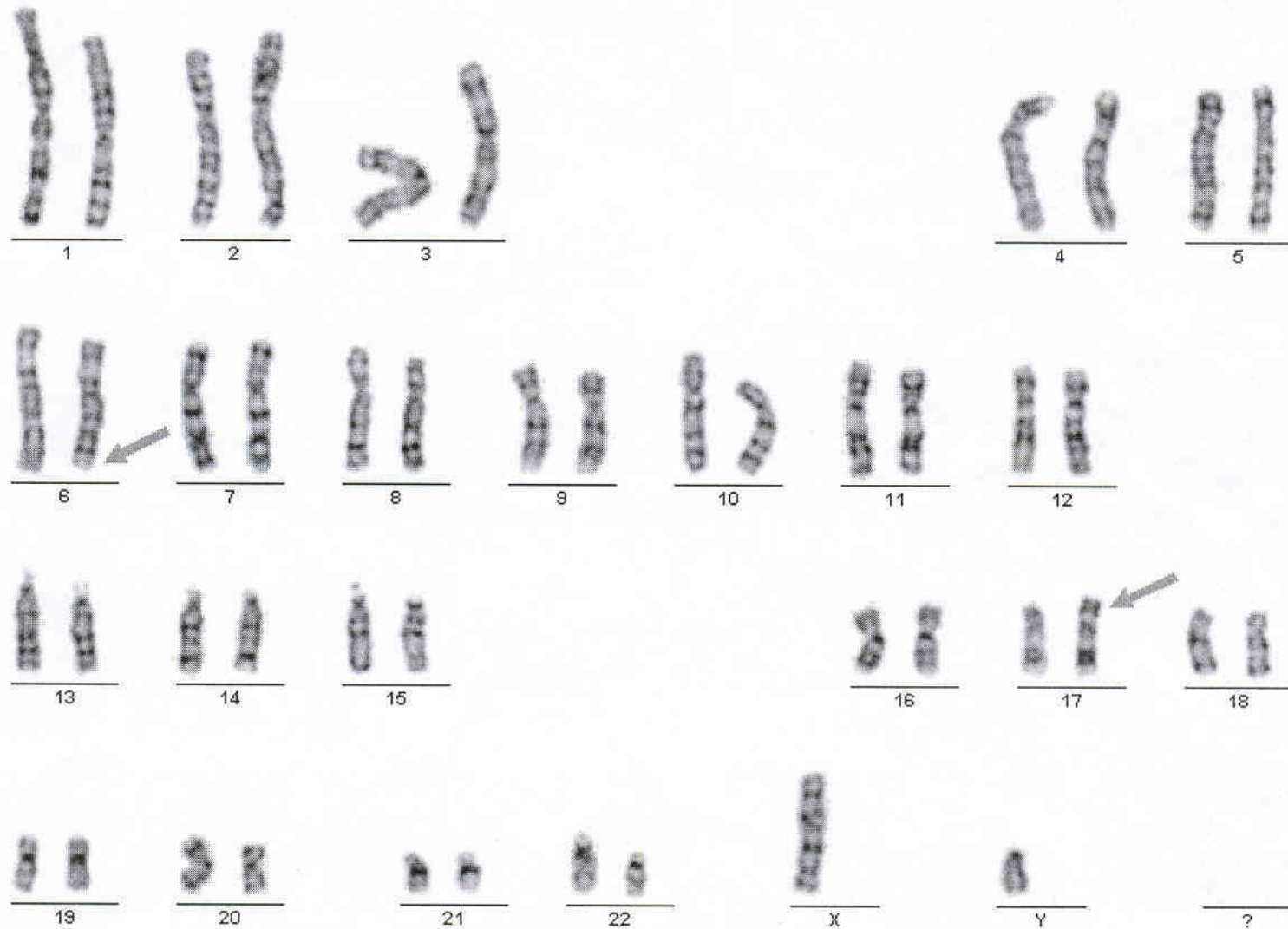
- anomalies of the larynx causes the characteristic cry of a similar feline meow (only in infancy)
- low birth weight and length
- mental retardation, short stature, failure to thrive, small moon shaped face, the position antimongoloid eye slits, mikrocephalie
- Other malformations and birth defects

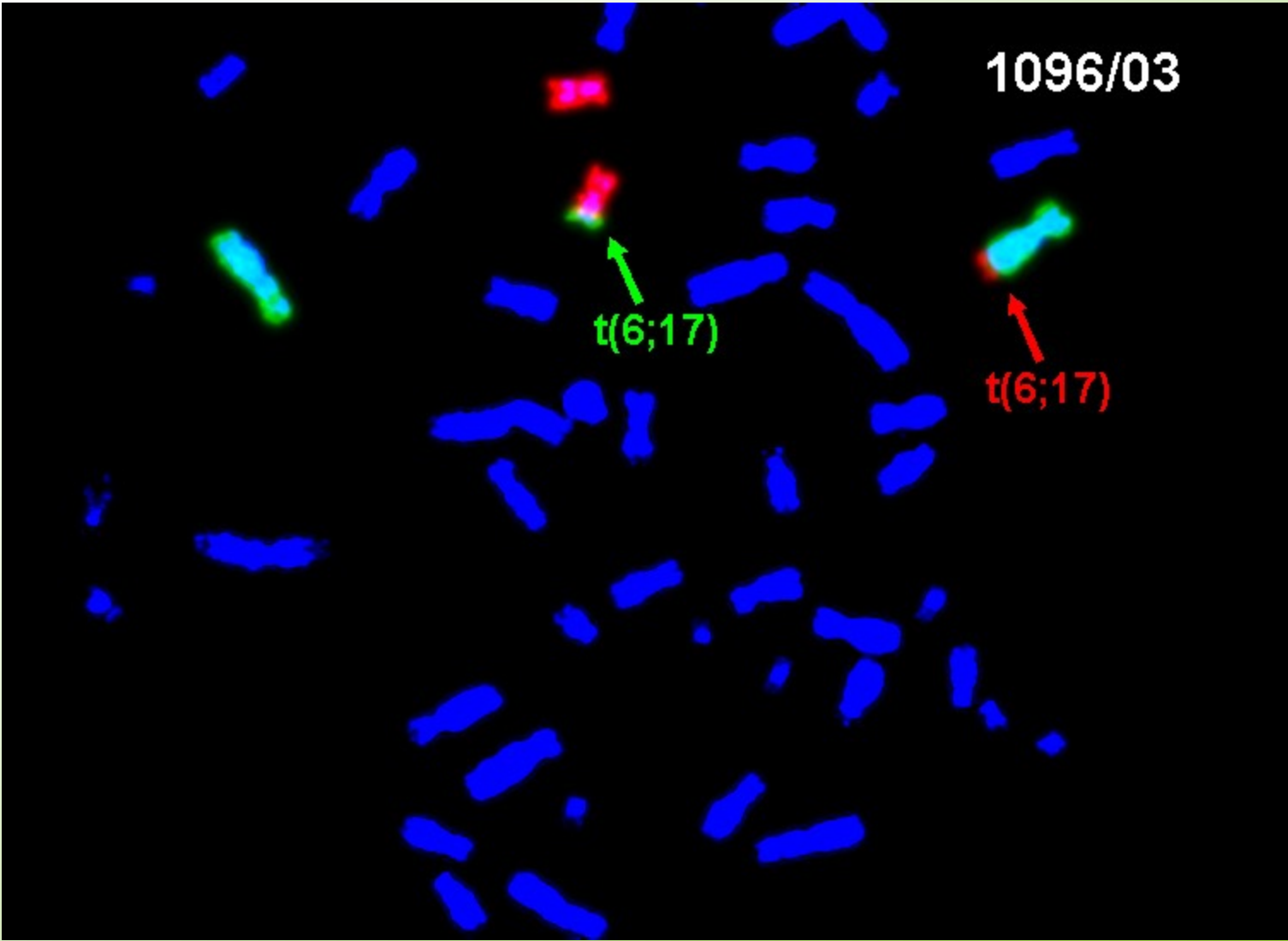
Cri du chat 46,XX(XY),5p-

- 1:50 000
- Typicaly cri in newborns
- laryngomalacie
- antimongoloid
- epicanthi
- hypotonie
- hypotrofie

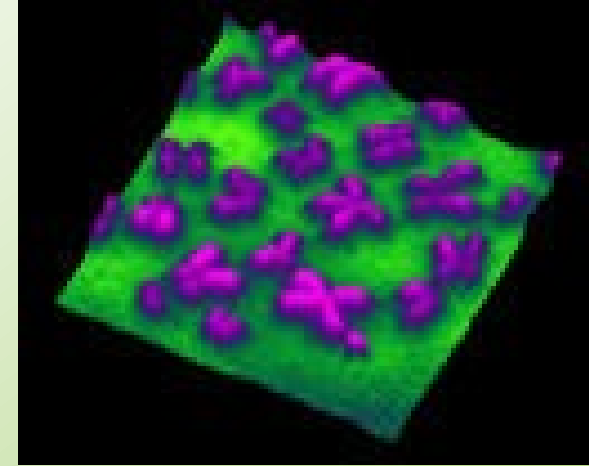
Other structural chromosomal aberrations

46,XY,t(6;17) – balanced translocation in a men with sterility

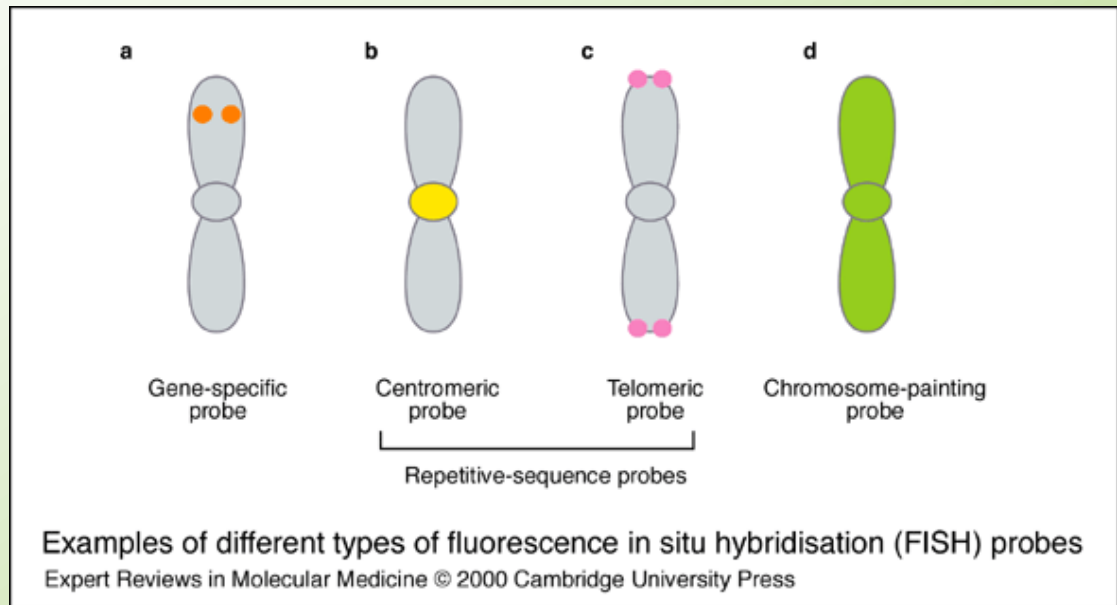




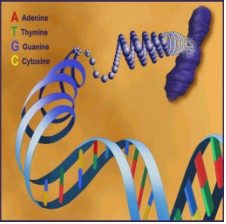
Mikrocytogenetic Molekular cytogenetic



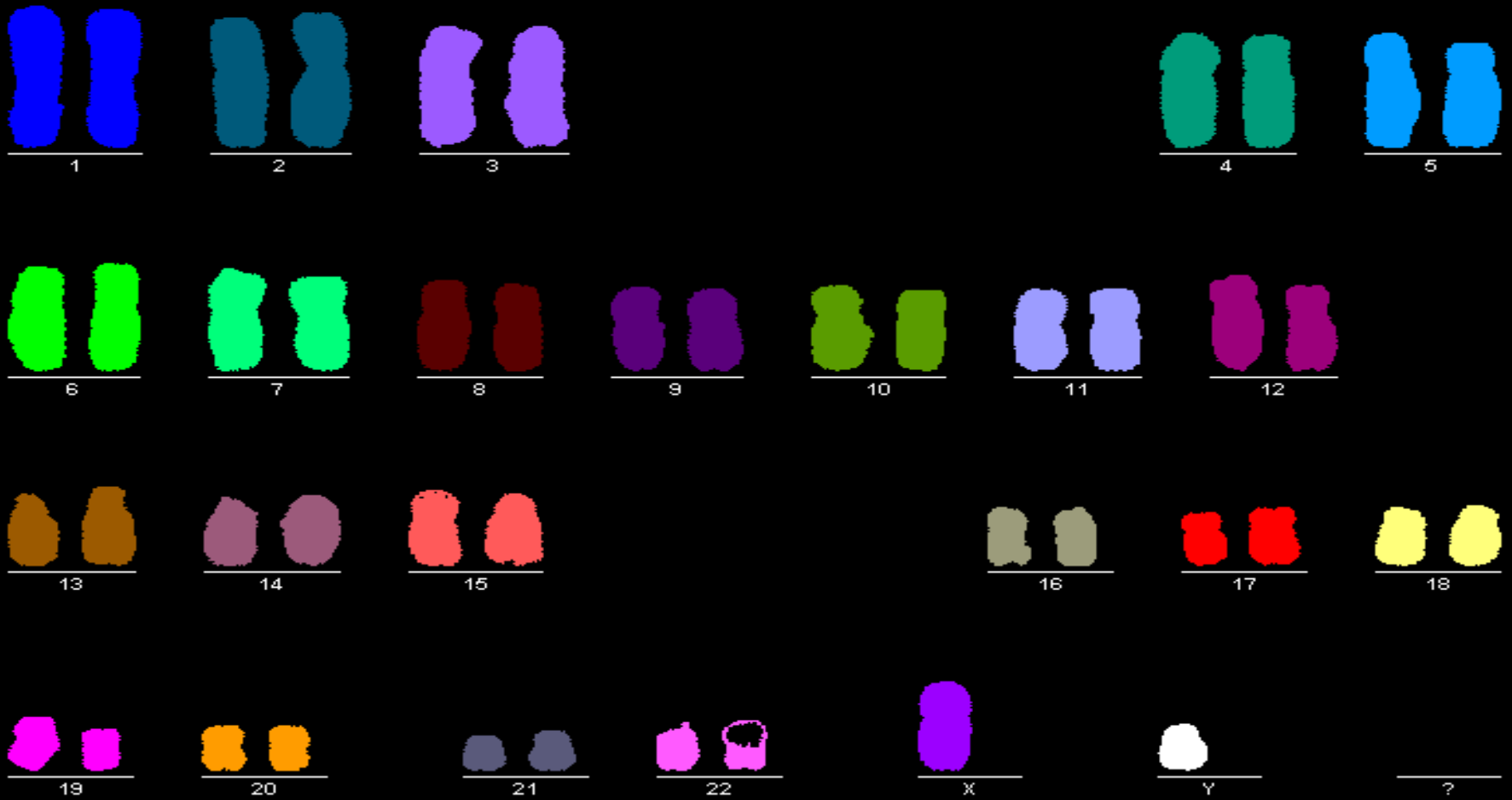
- FISH (fluorescenc in situ hybridisation), M-FISH, SKY (spektral karyotyping), CGH (komparativ genom hybridisation), MLPA
- mikroleletions or mikroduplikations, marker chromosomes, complex rearegements, oncology - oncocytogenetics, fast prenatal diagnostics ...)
- fast methods (possible for prenatal dg)
- metafase and intesfase examination



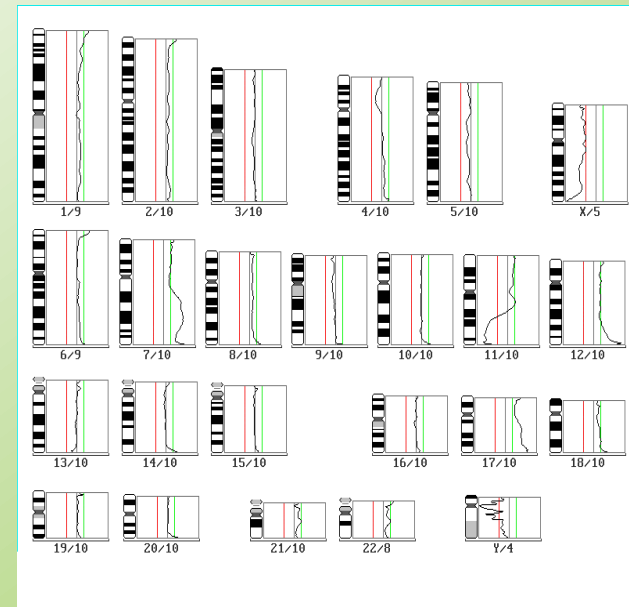
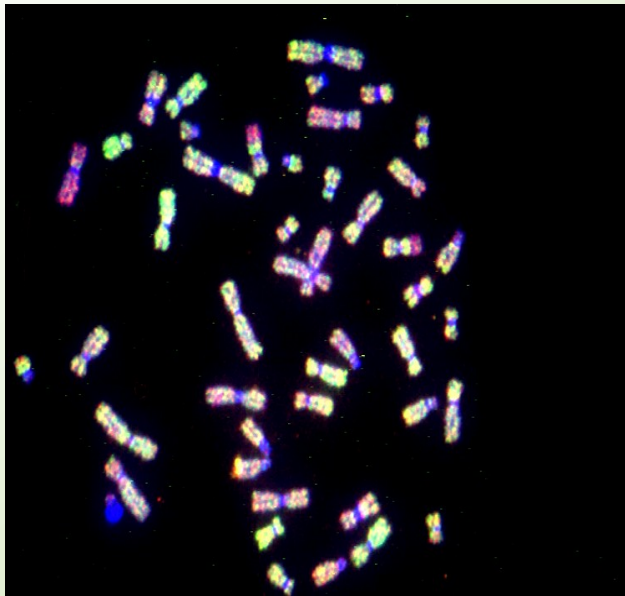
FISH



M-FISH (multicolor) Spektral karyotyping (SKY)

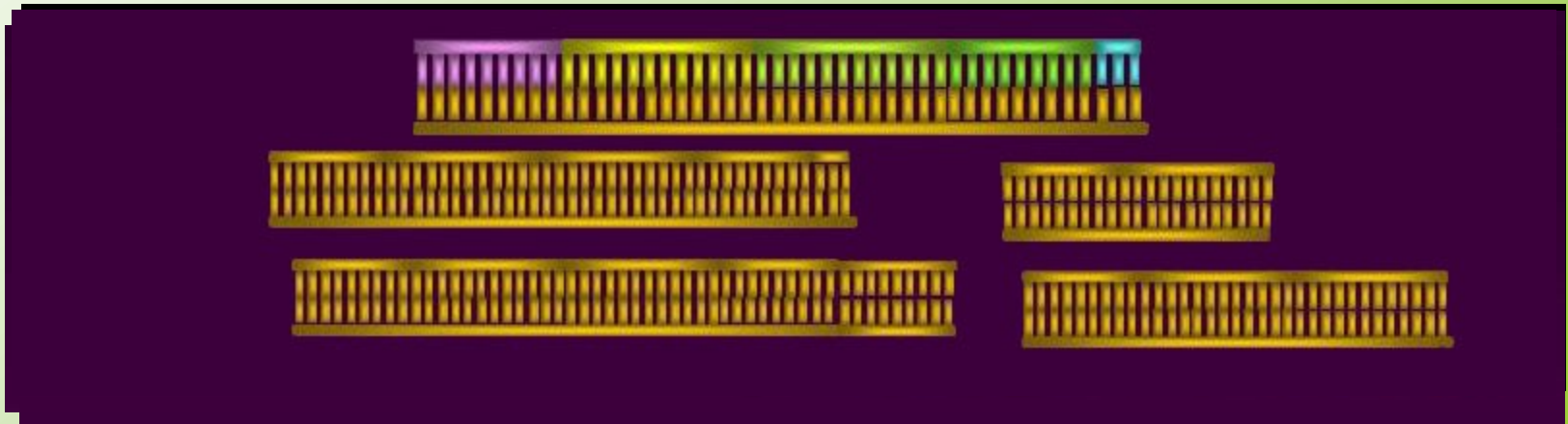


Comparativ genom hybridisation



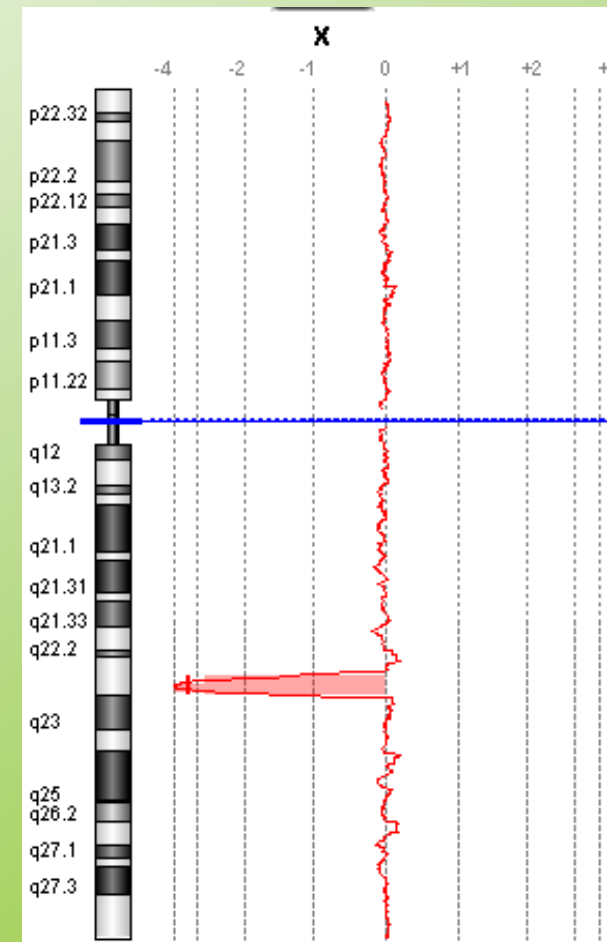
MLPA

Multiplex Ligation-Dependent Probe Amplification



array CGH

- DNA mikroarray
- Chip technology



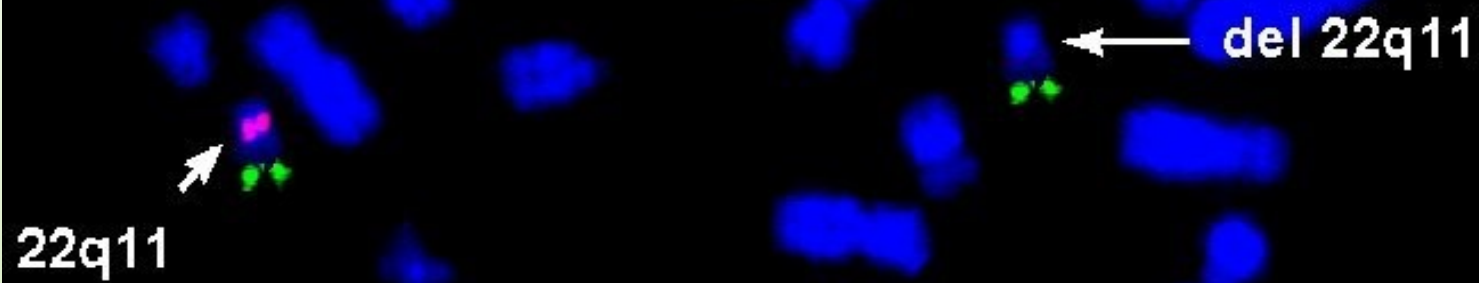
Microdeletions

- Di George syndrome
(del 22q11)
- Prader-Willi / Angelman syndrome
(del15q11-13)
- Williams Beuren syndrome
(del7q11.23)

Syndrom Di George

- Velo - Kardio- Facial syndrome
- CATCH 22
- Congenital heart disease - conotruncal, craniofacial dysmorfism, thymus aplasie, imunodeficient`cy, hypoparathyreoidismus

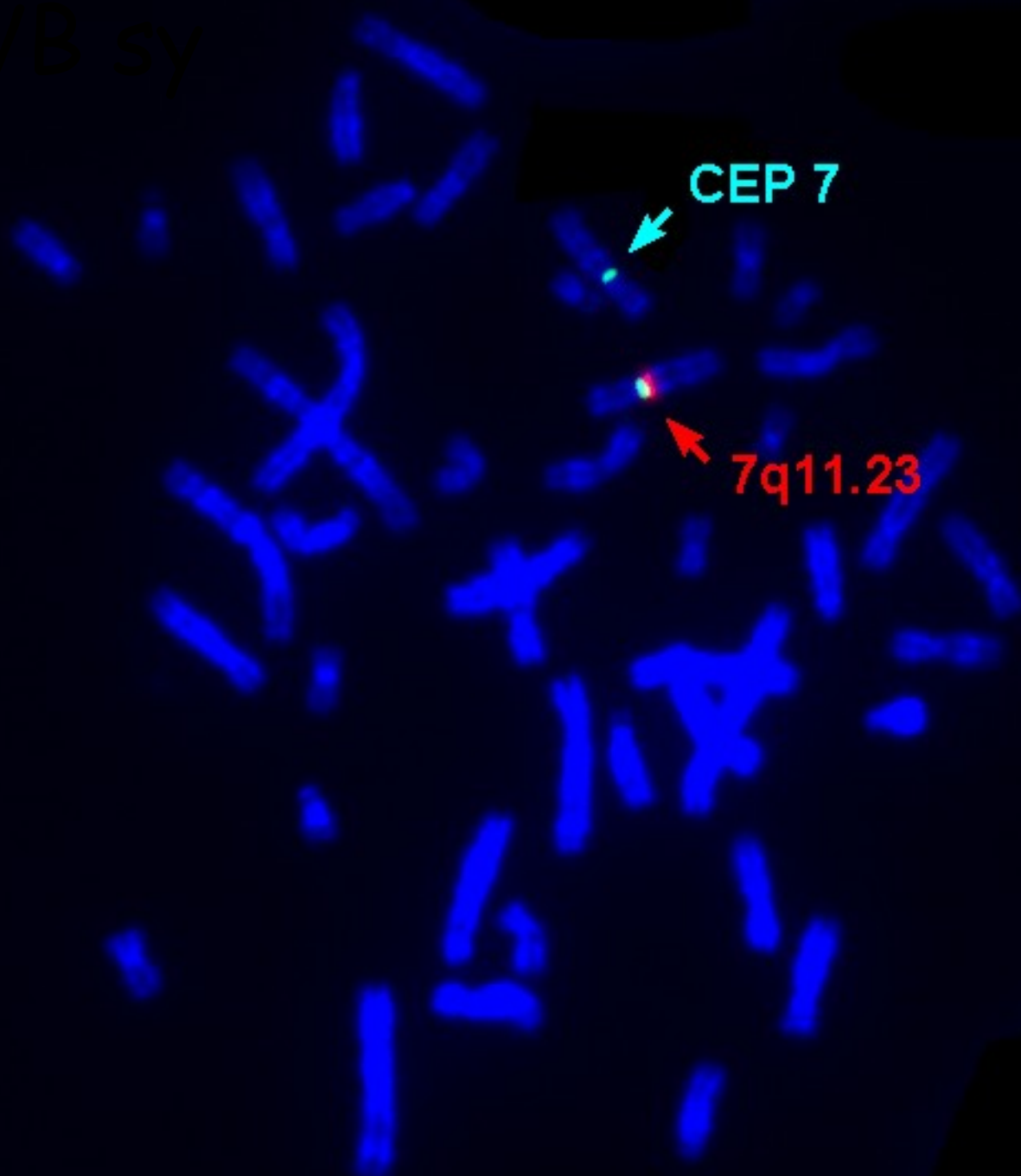
DiGeorge syndrom



Williams - Beuren syndrom

- del 7q11.23
- Facial dysmorfie - Elfin face, congenital heart disease, aortal or pulmonal stenosis, hypokalcemie, small statue, MR, hernie,...

Photo WB sy

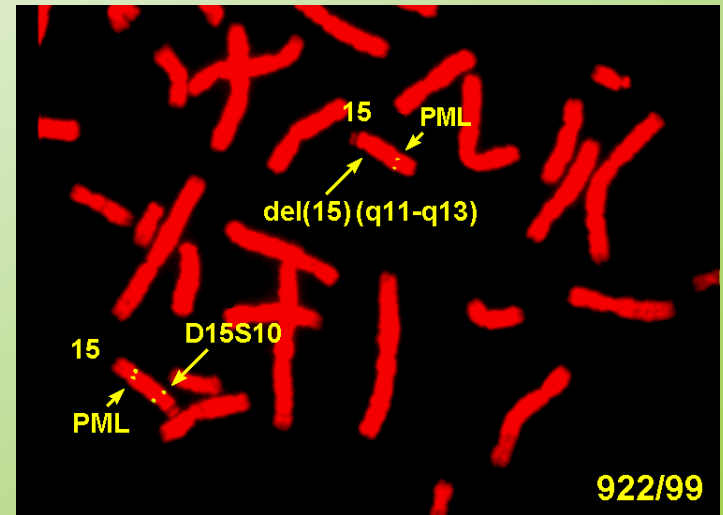


Prader-Willi syndrom

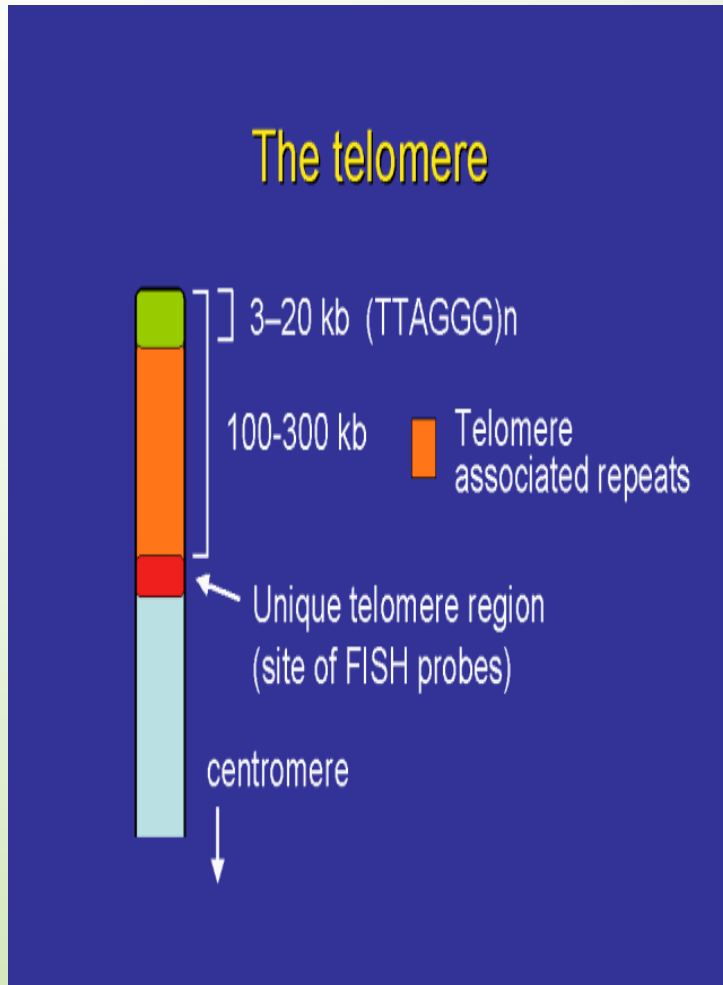
- Hypotonie, hypotrofie in small children
- PMR, small statue, obesity, hyperfagie, akromikrie, hypogonadismus
- mikrodeletion15q11-12 paternal

Angelman syndrom

- Severe mental retardation
- Epilepsie
- Laughter
- severely delayed speech development
- mikrodeletion 15q11-12 mat



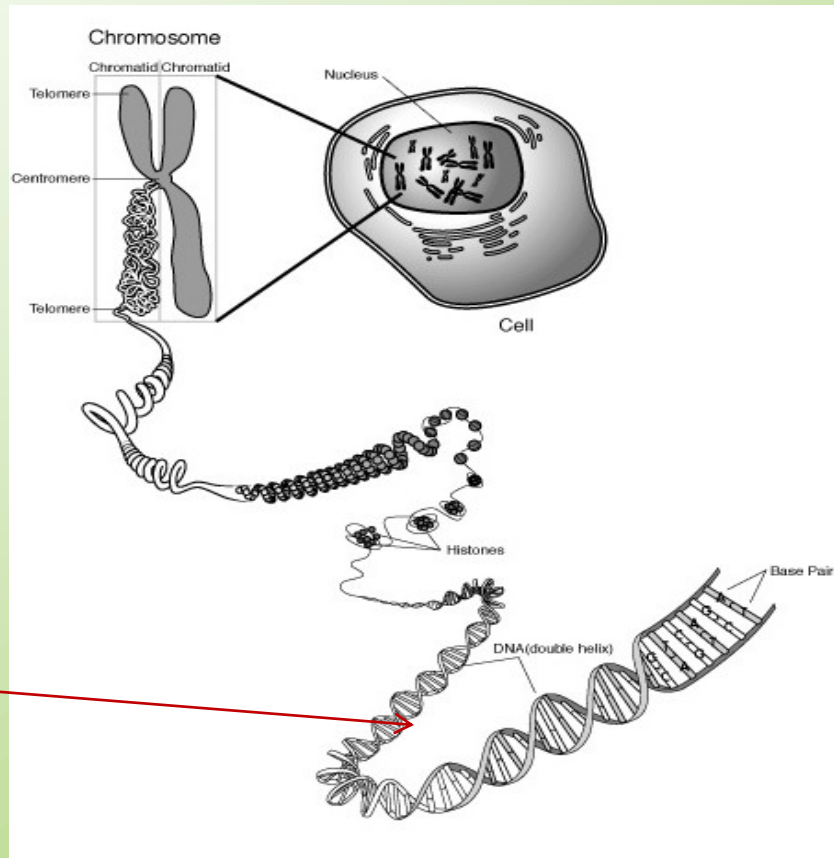
The telomere



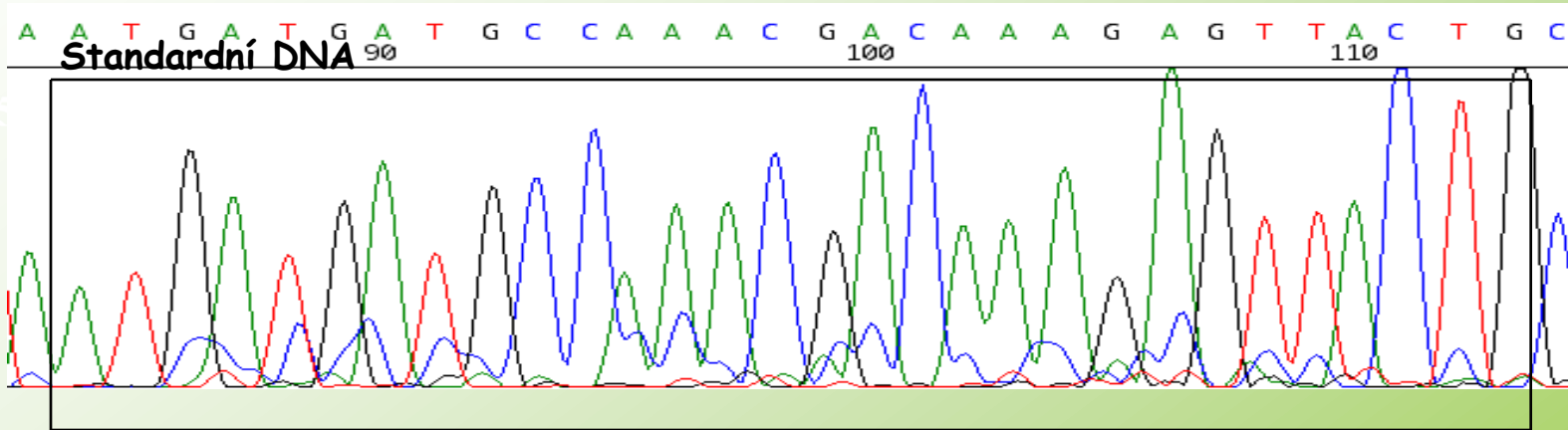
Rearrangement in about 6-8% children with mental retardation with or without congenital defect (FISH, HR-CGH, MLPA)

Mendelian inheritance

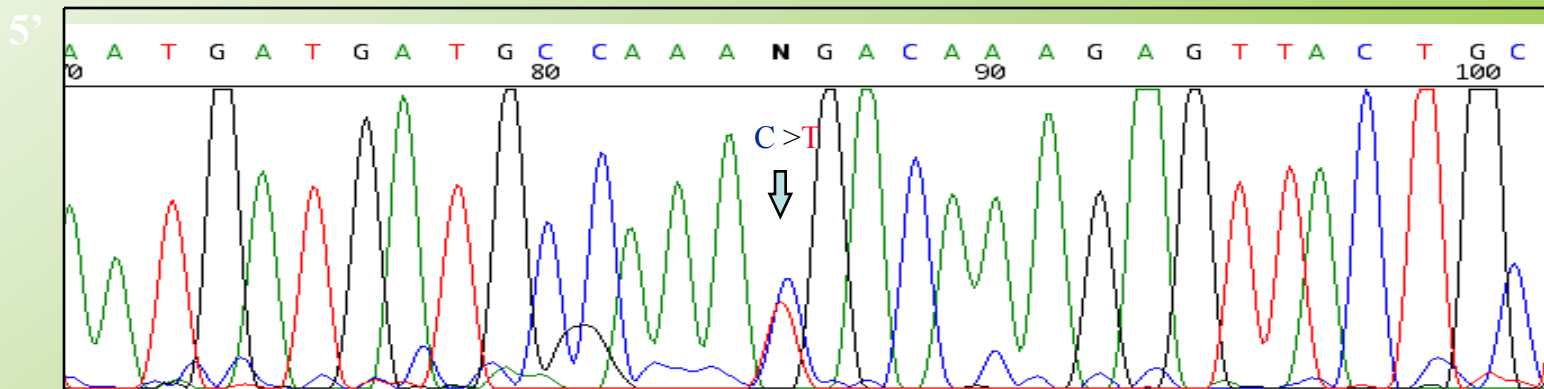
Monogenic diseases



DNA analysis



DNA NF1 pacienta, mt C5839T (Arg > STOP)

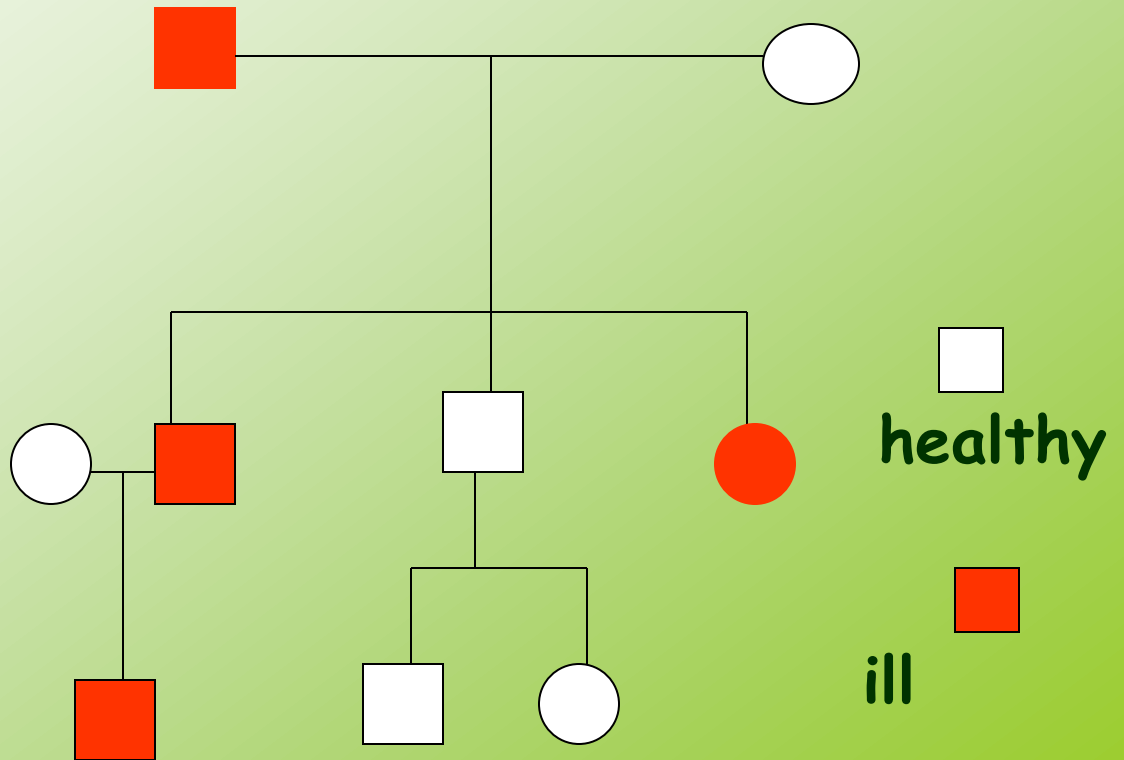


Autosomal Dominant

- The sexes are involved equally
- Heterozygotes are mostly affected clinically
- risk 50% for sibs and children
- new mutations
- penetrance, expresivity

Pedigree AD inheritance

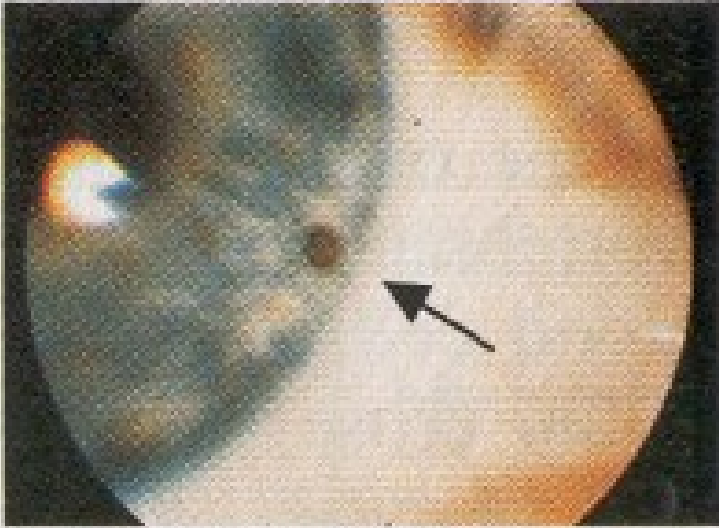
- the risk **50%**



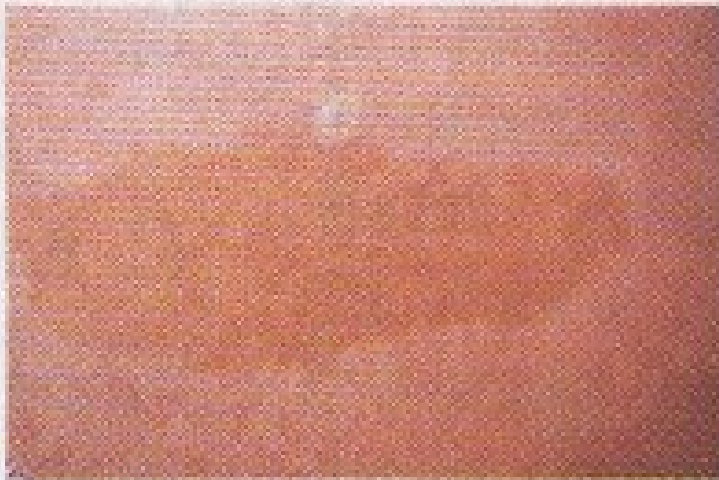
AD - diseases

- Neurofibromatosis 1 and 2
- Achondroplasia
- Huntington disease
- Marfan syndrome
- Myotonic dystrophy

Neurofibromatosis 1



1. Lisch nodule

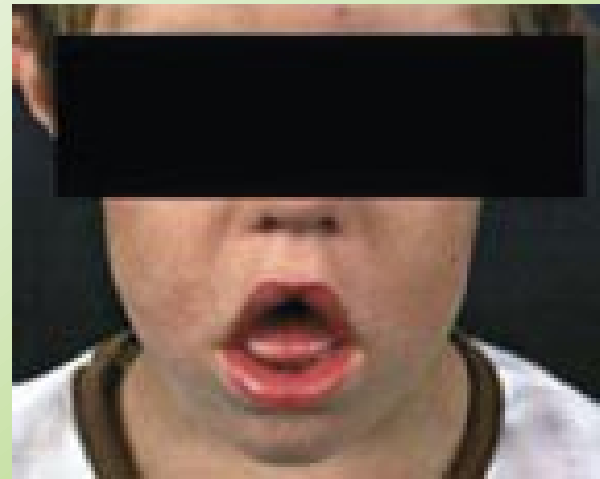


2. Café-au-lait spot

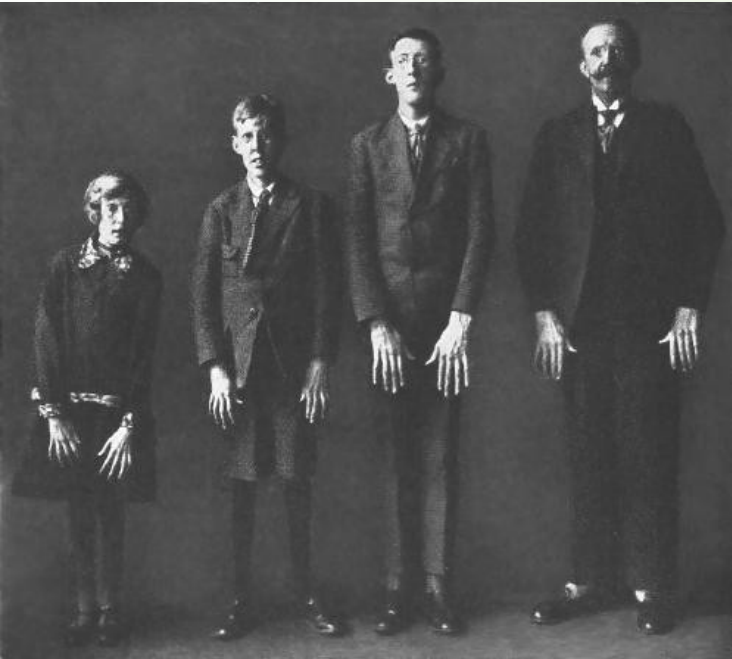


3. Neurofibromas

Myotonic dystrophy



Marfan syndrom

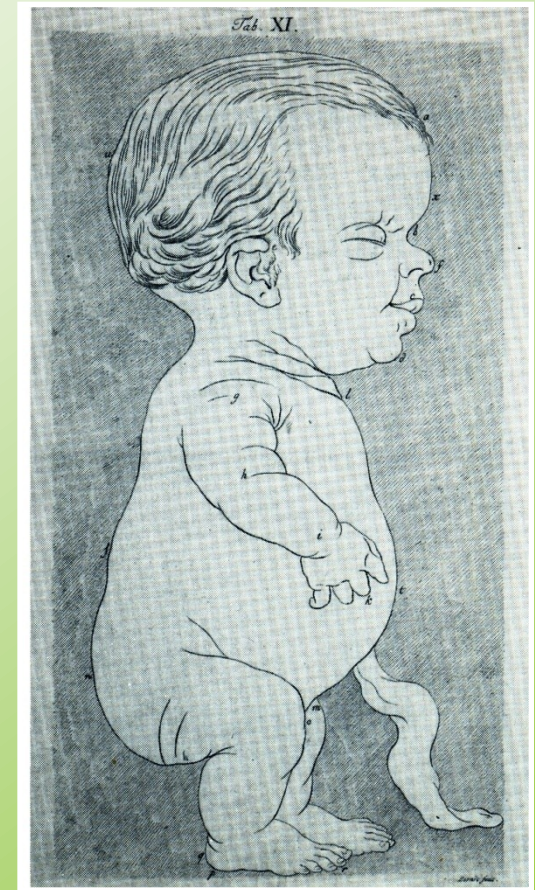


Marfan syndrom

arachnodactily



- Achondroplasia (ACH)
- 2 mutations in FGFR3 gene
- Paternal origin on new mutations

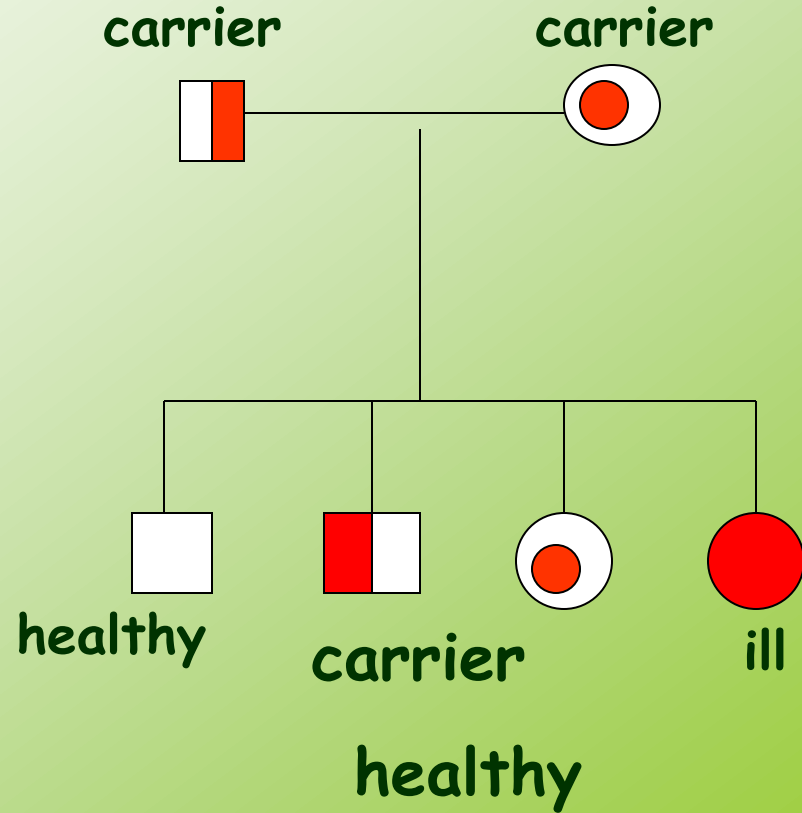


Autosomal Recessive

- Heterozygotes are generally unaffected clinically
- The sexes are involved equally
- An individual manifesting a recessive disorder usually has heterozygous parents
- Once a homozygote is identified, the recurrence risk for other child of some parents is 25%

Pedegree - AR inheritance

•The risk for next child 25%



AR - diseases

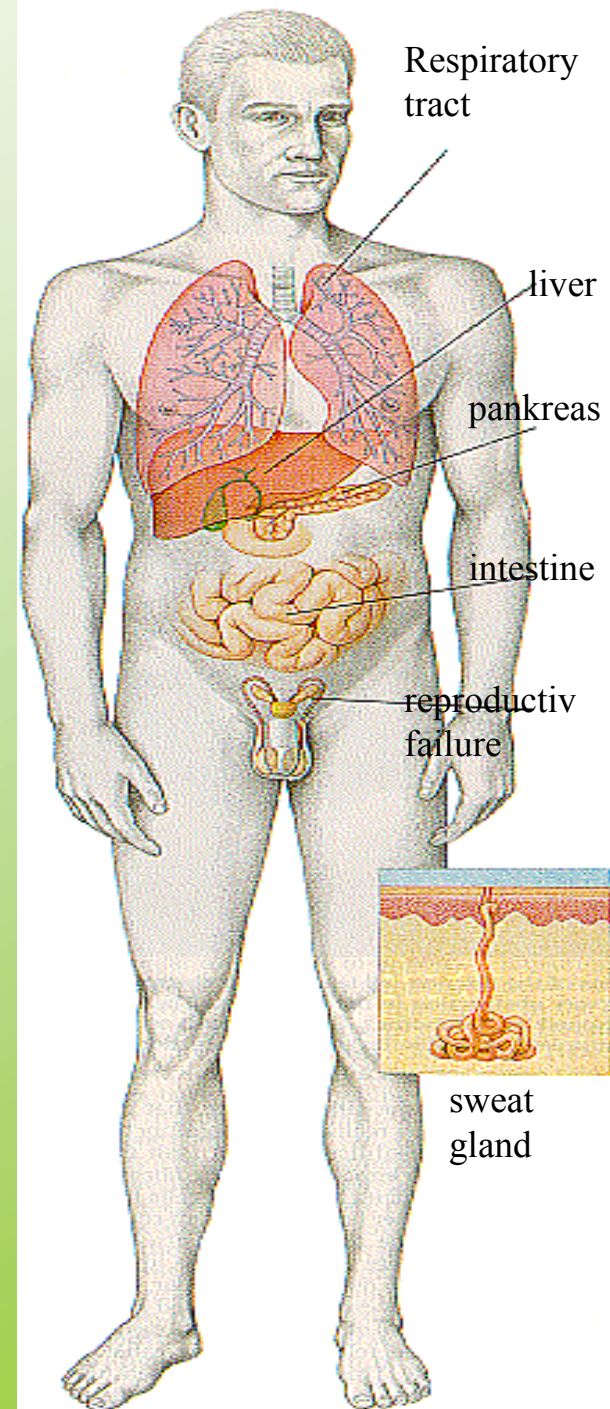
- Cystic fibrosis
(frequency of heterozygotes CR- 1/30)
- Phenylketounria (1/40)
- Congenital adrenal hyperplasia (1/40)
- Spinal muscular atrophy (1/60-80)

Cystic fibrosis

- Localized on chromosome 7q
- Frequency of Cystic Fibrosis in the Czech Republic: about 1/6000
- Frequency of heterozygots in the Czech Republic about 1/30
- About 1600 mutations in CFTR gene were identified

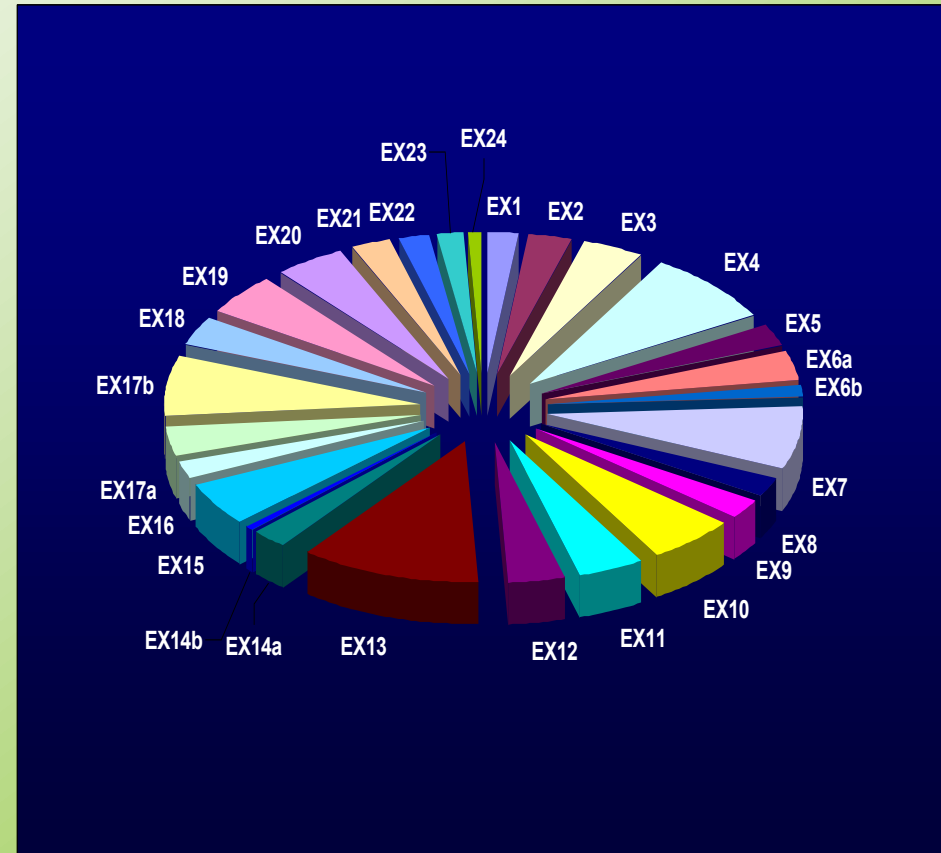
Cystic fibrosis

- disease affecting multiple organs



The reason for CFTR gene analysis

- Newborn screening in CR from 10/2009 (analysis of 50 mutations in CFTR gene)
- Suspicion on Cystic fibrosis in a patient
- Cystic fibrosis in the family
- Partners of heterozygotes for Cystic fibrosis
- Repeated fetal loss
- Sterility
- Relationship of the partners



CFTR gene - distributions of mutations

Most frequent CFTR mutations in Czech population

Mutation	Frequency in CR (%)
F508del	70,7
CFTRdele2,3(21kb)	6,4
G551D	3,7
N1303K	2,8
G542X	2,1
1898+1 GtoA	2,0
2143delT	1,1
R347P	0,74
W1282X	0,6

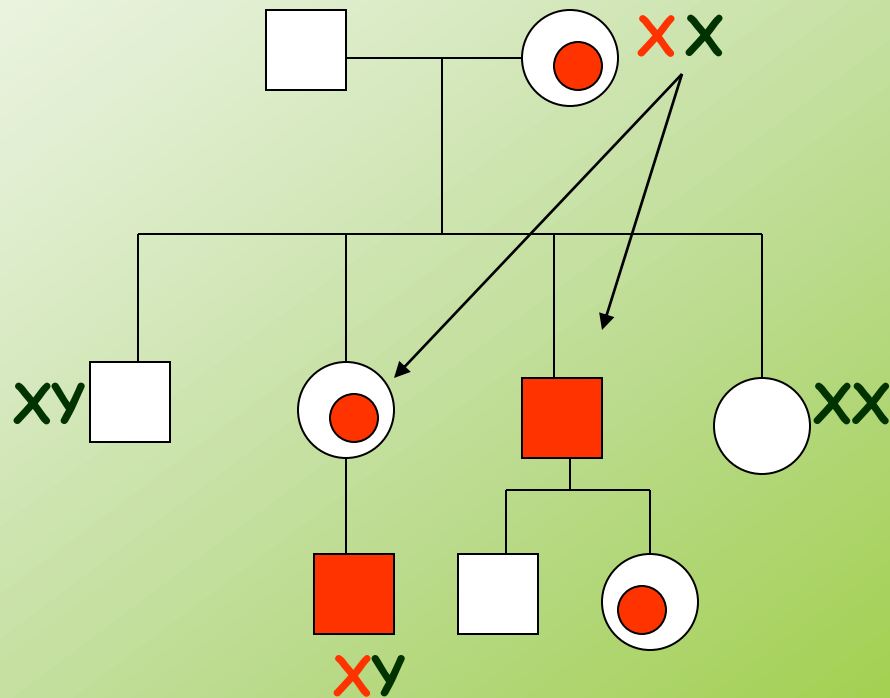
X-linked Recessive

- Females are not affected as severely as males or are not affected
- An affected male cannot transmit the trait to his sons, because the trait is on X-chromosome, and the father must necessarily transmit his Y-chromosome to a son
- All of the daughters of an affected male must be carriers, because the only X-chromosome that the father can give to a daughter contains the mutation

X-linked Recessive

- Risk for daughters of a carrier - mother
- 50% for carrier
- Risk for sons of carrier - mother
- 50% for disease

X- recessive inheritance



XR - diseases


- Hemophilia A and B
- Duchenne and Becker muscular dystrophy
- Fragile X chromosome - X-linked disease

Duchenn/Becker muscular dystrophy

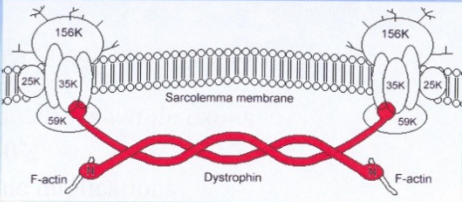
DMD Mutations

- Mutations of the gene fall in to three categories:
 - Deletions of one or more exons
65%
 - Small mutations *within* exons
30%
 - Intragenic duplications
5%
- So, exon screening will pick up 65% of DMD Mutations

Innovative screening solutions for human genetic analysis




•Dystrophin protein forms part of muscle structure (molecular glue)



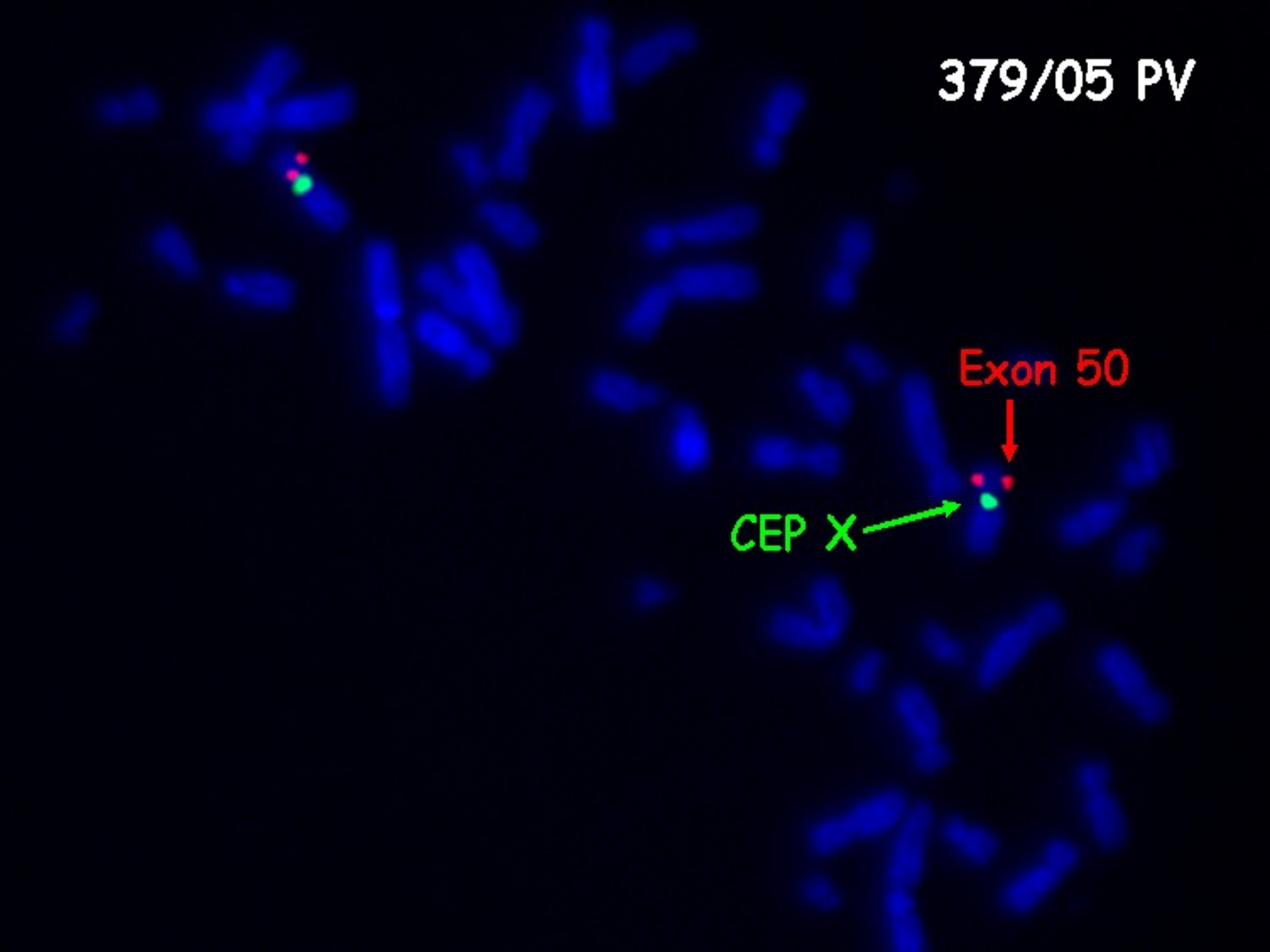
•Helps stabilize membrane during muscle contraction and relaxation

Innovative screening solutions for human genetic analysis

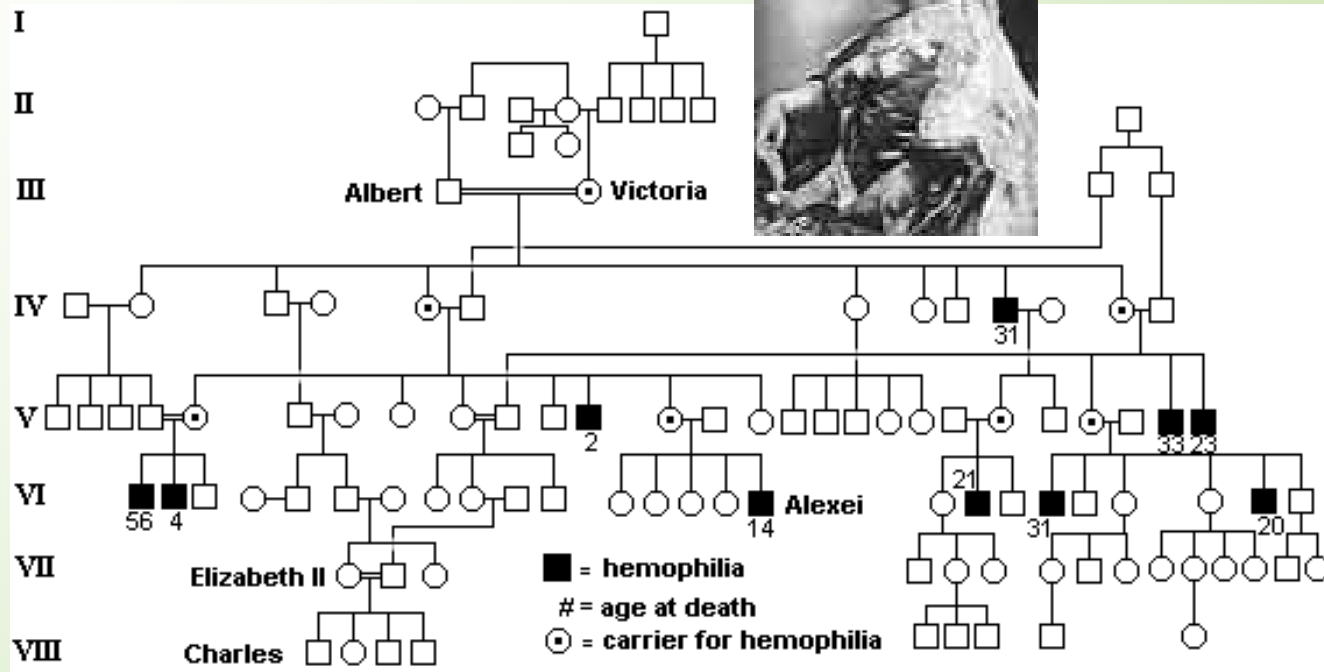


379/05 PV

Exon 50
CEP X

A fluorescence microscopy image of a chromosome spread. The chromosomes are stained blue. Two specific regions are highlighted with colored dots: a green dot labeled 'CEP X' and a red dot labeled 'Exon 50'. Arrows point from the text labels to their respective dots. The 'CEP X' dot is located on the centromere of a chromosome, while the 'Exon 50' dot is located on the arm of a chromosome. The background is black.

Hemofilia A



**Multifaktorial -polygenic
inheritance**

**Diseases with complex
heritability**

Teratogens

Characteristic

- disease with multifactorial inheritance include not mendelian types of inheritance
- diseases exhibit familial aggregation, because the relatives of affected individuals more likely than unrelated people to carry diseases predisposing predisposition

Charakteristic

- in the pathogenesis of the disease play a basic role non-genetic factors
- disease is more common among close relatives and in distant relatives is becoming less frequent

Examples

- Congenital heart defects (VCC) 4-8/1000
- Cleft lip and palate (CL/P) 1/1000
- Neural tube defects (NTD, anencephalie, spina bifida,..) 0,2-1/1000
- Pylorostenosis
- Congenital hip dislocation
- Diabetes mellitus - most types
- Ischemic heart disease
- Essential epilepsy

Common congenital defects

Congenital heart diseases

- 0,5 - 1% in liveborn infants - population incidence
- etiology not known mostly
- about 3% combine with chromosomal syndromes (+21, +13, +18, 45,X, 18q-, 4p-, del 22q11 Di George sy)
- some mendelian syndromes associated with congenital heart disease (Holt-Oram, Williams, Noonan, Ivemark...)

Congenital heart diseases prenatal diagnosis

- For most serious congenital heart diseases
- Ultrasonography in 21. week of gestation - by specialists for prenatal kardiology

Congenital heart disease - genetic risks

condition	1 aff. sibling	1 aff. parent
Ventricular septal def.	3%	4%
Patent ductus art.	3%	4%
Atrial septal defect	2,5%	2,5%
Tetralogy of Fallot	2,5%	4%
Pulmonic stenosis	2%	3,5%
Koarctation of aorta	2%	2%

Congenital heart disease genetic risks

Risk in %

More than two affected firstdegree relatives	50
Sib of isolated case	2 - 3
Second-degree relatives	1 - 2
Offsprin- affected father	2 - 3
Offsprin - affected mother	5
Two affected sibs	10

Cleft lip and palate

- Incidence of CL in the population
1/500-1/1000
- Inheritance - multifactorial mostly
- Chromosomal trisomies (+13,+18)
- Syndromes associated with CL/CP/CLP
- (van der Woude sy, EEC sy, Pierre Robin sequence...)
- Prenatal diagnosis by ultrasonography
not sure

Cleft lip and palate- genetic risks

Relationship to index case	CLP	CP
Sibs (overall risk)	4%	1,8%
Sib (no other affected)	2.2%	
Sib(2 affected sibs)	10%	8%
Sib and parent affected	10%	
Children	4,3%	3%
Second-degree relatives	0,6%	

Patau syndrome, 47,XX,+13

EEC syndrome

Van der Woude syndrome

Sequence Pierre Robin

Neural tube defects

- Multifactorial inheritance (risk for 1. degree relatives about 2 - 4%)
- Maternal serum screening - elevated level of AFP
- Prenatal diagnosis by ultrasonography
- Raised AFP levels in amniotic fluid
- Primary prevention in pregnancies - folic acid
- Risk in the population - probably related to nutritional status

Teratogens

- teratogen is a substance whose effect on embryo or fetus may cause abnormal development

action may be direct or through the maternal organism

Human Teratogens

- Physical (radiation, heat (fever), mechanical impact)
- Chemical (chemicals, drugs)
- Biological (infection, fungus ...)
- Metabolic imbalance (disease mother)

The effect of teratogens depends on :

- dose
- length of the action
- contact time
- genetic equipment of the fetus and the mother

Critical period

- 14.-18. days after conception - the rule „all od nothing“
- 18.-90. day - organogenesis
- The most sensitive period for the emergence of developmental defects

Drugs

- Distribution of medicines practice into categories
 - A
 - B
 - C
 - D
 - X
- Food and Drug Administration, 1980

A

- in controlled studies have shown no evidence of risk to the fetus in the first trimester of fetal development or influence in the next period of pregnancy

product appears to be safe

B

- Animal reproduction studies demonstrate a risk to the fetus, but there's no controlled studies in women

Animal reproduction studies have shown adverse effects, but in controlled studies in women have not been confirmed

C

- Animal studies confirm the teratogenic embryotoxic or other adverse effects on the fetus,
- non-controlled studies in women
- lack of studies in animals and humans

product should be administered with caution and only in cases where the benefit for the woman of his administration exceeds the potential risk to the fetus

D

- risk to the human fetus is known
- medicine may be administered in a situation where its use for a woman needed (lifesaving)
- no other safer drug is available



- studies in animals and in humans clearly demonstrate a teratogenic effect
- drugs absolutely contraindicated in pregnancy

Drugs with teratogenic effect

- Thalidomid
- Hydantoin
- Valproic acid
- Anti coagulans - Warfarin
- Trimetadion
- Aminopterin
- Methotrexat
- Cyklophosphamid

Drugs with teratogenic effect

- Retinoids
- Lithium
- Thyxreostatic drugs
- Androgens
- Penicilamin
- Enelapril, Captopril
- Antituberkulotics - Streptomycin

Thalaidomid

- congenital heart defects
- limb reduction anomalies
- Other congenital defects
(gastrointestinal, urogenital tract
orofacial - ears anomalies, CNS
defects..)

Hydantoin

- Atypical face, growth retardation, mild mental retardation, behavioral problems, hypoplastic nails and fingers

Aminopterin a Methotrexat

- folic acid antagonist
facial dysmorfism, cleft lip and/or
palate, small mandible, ears
anomalies, hydrocephaly, growth
and mental retardation, miscarriage

Warfarin

- coumarin antikoagulans
- facial dysmorfism - nasal cartilage hypoplasia, CNS - defects

Retinoids

- Cleft lip and palate, mikrognatia, eyes anomalies, ears dysplasia
- Defects of CNS
- Thymus hypoplasia
- Limb defects

Infection

- Toxoplasmosis
- Rubella
- Cytomegalovirus
- Herpesvirus
- Others (parvovirus, antroprozoonosy, chlamydia..)

• TORCH

Toxoplasmosis

- chorioretinitis
- hydrocephaly or microcephaly
- intracranial calcification, mental retardation
- icterus, hepatosplenomegalia, carditis
- prematurity

- positiv IgM in the mother - treatment with Rovamycin
- Prenatal dg.: serology, DNA-PCR)

Rubella

- hearing and vision impairment (cataract, glaucoma, mikroftalmia, blidness)
- mental retardation
- Cong. heart defects
- icterus, hepatosplenomegalia

- prevention- vaccination

Cytomegalovirus

- Intrauterin growth retardation
- mikrocephaly, calcification in the brain, mental retardation,
- hepatosplenomegaly

- Repeated maternal infection is possible
- Prenatal dg.: serology, DNA-PCR

Varicella zoster

- Skin lesions and defects
- Brain damage, mental retardation
- Eye defects

- Prenatal dg. - serology, DNA-PCR

Metabolic dysbalance

- Fetal alcohol syndrom (FAS)
- Maternal Phenylketonuria
- Maternal Diabetes mellitus
- Maternal Hypothyreosis

Fetal alcohol syndrom

- Hypotrophy, growth retardation, mental retardation
- facial dysmorphism
- Congenital heart defects
- Limb defekts

- Abuse of 60g pure alcohol / day (longterm)
- Combine with malnutrition, folic acid deficit...

Maternal Phenylketonuria

- Low birth weight
 - hypertonia
 - mikrocefaly, mental retardation
 - Cong. heart defects
 - hyperaktivitiy
-
- newborn screening
 - (frequency 1/10 000 newborns
 - inheritance - AR)
 - initiation of treatment within three weeks to prevent mental retardation in the child

Reproductive Genetics

Preconceptional testing
Genetic counselling and analysis
in couples with reproductive disorders
Prenatal diagnosis
Preimplantation genetic diagnosis
Examination of potential donor gametes

Secondary prevention of genetic

- The procedures in pregnancy - prenatal diagnosis and early postnatal diagnosis

Prenatal diagnosis

- Non invasive methods- screening
- Invasive methods
- **CVS** - after the 10. week of gestation
- **AMC** - 15.-18. week of gestation
- **Cordocentesis** - after the 20. week of gestation

Prenatal diagnosis results

- CVS - karyotype - about 5 days
- AMC - karyotype - about 14-21 days
- DNA analysis (monogen diseases)
- About 5-15 days
- DNA from amniocytes after cultivation - exclusion contamination by maternal tissues

Prenatal analysis of most frequent aneuploidias QF PCR

- Examination of the most common numerical changes in chromosomes 13, 18, 21, X and Y
- The result for 24-48 hours

Prenatal screening (CR)

- Ultrasound (12. - 20. - 33. week)
- Ultrasound 20.week - cong. defect
- Ultrasound 20-22. week - cong. heart defect
- 10-14. week of gestation
- Free beta hCG, PAPP-A, US-NT, NB..
- 16.-18.week of gestation
- AFP, hCG, uE3

non-invasive prenatal testing NIPT

- NIPT (aneuploidie 21, 13,18,X/Y)
- Rh in the fetus
- SRY in the fetus - in X linked diseases in the family
- Some monogenic diseases in the fetus (achondroplasia)

Indications for prenatal examination / genetic counselling

- US screening - congenital defects
- Family history of known conditions for which diagnosis is possible (DNA analysis)
- Known chromosomal abnormality (de novo finding in previous child, structural change in parents)
- Positive prenatal screening for chromosomal abnormalities
- Advanced maternal age (38-40 years)
???

Preimplantation Genetic Diagnostics

- IVF - assisted reproduction
- **Preimplantation genetic screening**
- aneuploidias - array- CGH, chip technology
- FISH (13,18,21,X,Y, 15,16,22)
- **Preimplantation Genetic Diagnostics**
- Structural chromosomal aberrations
- (parents are carriers of balanced rearrangement)
- Monogenic diseases (known in family history)

PG Diagnostic

X

PG Screening

- PGD high genetic risk
- PGS (most common)
aneuploidies

Genetic counselling in infertility

Infertility

- Is the infertility one aspect of a genetic disorder that might be transmitted?
- Will correction of infertility give an increased risk of malformations in the offspring?
- Genetic testing before use of methods of assisted reproduction.

Infertility

- Pathological examination of the abortus where possible, this may identify major structural malformations.
- **Cytogenetic study of parents**, this is especially important where a structural abnormality is present.
- In general the finding of a chromosome abnormality in the abortus but not in parent is not likely to be relevant or affect the genetic risks.

Infertility

- A search for possible lethal mendelian causes (consanguinity- risk for AR diseases, X-linked dominant disorders lethal in male, myotonic dystrophy which gives heavy fetal loss in the offspring of mildly affected women)
- Inherited trombophilias in women with recurrent abortions (factor V Leiden, factor II - G20210A, hyperhomocystinaemia ? (MTHFR - C677T)

Factor V - Leiden

- frequency in the white European population of about **5 - 9%**
- AD inheritance
- increased risk of thromboembolism in homozygots for FVL 50-100x, in heterozygots 5-10x
- increased risk of fetal loss after the 10. week of gestation

Sterility in male

- AZF (azoospermia factor) deletions of the DAZ gene **Yq** (deleted in azoospermia)
- Infertile man - 4-5%
- Men with azoospermia - about 15%
- CFTR mutations and polymorphisms

Genetic risk in cancer

Genetic testing in oncologic patients

- Specification of the:
- Diagnosis
- Therapy
- Prognosis
- Monitoring of minimal residual disease

Genetic risks in cancer

- Tumours following mendelian inheritance (most AD, about 5%)
- Genetic syndromes predisposing to malignancy

Hereditary cancer syndromes

- AD inheritance
- Preventive, pre-symptomatic testing
- Prevention
- Associated problems

Hereditary cancer syndromes following AD inheritance

- Breast cancer - BRCA 1 and BRCA 2
- Familial Adenomatous Polyposis coli - FAP
- Von Hippel - Lindau syndrome - VHL
- Retinoblastoma
- Neurofibromatosis - NF1, NF2
- Li-Fraumeni syndrome
- Lynch syndrome - hereditary non polypous colon cancer - HNPCC

Genetic testing in Hereditary cancer syndromes

- Tests are voluntary
- Mostly in adults only
- In children only when prevention in childhood is present and when the risk of tumours is in childhood

Postnatal care and neonatal screening

- Early diagnosis

Dispensary

Specialized Care

Prenatal and perinatal management of pregnancies with malformation or genetic disease in the fetus

- Consultation with experts, who will continue to take care of the pregnant woman - ultrasound specialist, gynecologist, obstetrician, psychological support ..

Consultations with specialists, who will care after the birth of newborns with disabilities

The planned delivery of specialized care workplace - kardiocentrum, pediatric surgery, cardiology...

Newborn screening

Sampler card

0004305

Whatman 903[®] Lot 6272207/51 2009-05 SN

SN 0004305

**Kartičku vyplnit před odběrem
Nedotýkat se oblasti pro kapky krve
Při poškození kartičku nepoužít**

Požadavek (zaškrtnout): SKH CAH Jiný (vypsat): Odběr: První:
Opakovaný:

Jméno novorozence	
Jméno	Příjmení
Rodné číslo, pojišť'ovna <small>(dítě nebo matka)</small>	Porodní hmotnost g
Datum a čas narození <small>DD.MM.RRRR – HH.MM</small>	Datum a čas odběru <small>DD.MM.RRRR – HH.MM</small>
Kódové číslo odběru <small>Kód oddělení (AAA) • pořadí odběru (XXX) - AAAXXX</small>	Praktický dětský lékař Jméno, telefon
Jméno matky	
Jméno	Příjmení
Telefon matka (rodina) <small>Mobil i pevná linka</small>	Adresa matky (pobytu)
Odesílatel vzorku <small>Čitelné razítko, jménovka, podpis</small>	

CE IVD REF 10539735 Rev.0 LOT 6272207/51

Whatman GmbH
Hahnstraße 3,
37586 Dassel Germany

Screened diseases in CR from 10/2009

- Kongenital hypothyreosis
- Kongenital adrenal hyperplasia - CAH

(cumulative risk 1/2900)

Screened diseases in CR from 10/2009

- Inborn errors of metabolism
- Fenylketonuria (PKU, HPA)
- Leucinosis
- MCAD
- LCHAD
- VLCAD
- Def.karnitinpalmitoyltransferasis I a II
- Def.karnitinacylkarnitintranslocasis
- Glutaric aciduria
- Izovaleric acidurie

(cumulative risk 1/4000)

Screened diseases

- Cystic fibrosis

(1/4000)

- cumulative risk of all 13 screened diseases in CR - 1/1200