

# **Malignant bone tumors**

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# Osteosarcoma

Malignant osteoid

# *Epidemiology*

- 3 new cases /1 milion/ year
- 2. decade
- Metaphysis of long bones
  - 1/2 in knee region
    - distal femur
    - proximal tibia
    - proximal humerus

# *Classification*

- Primary

- Central

- High-grade

- Conventional high-grade (80 – 90%)

- Osteoblastic

- Chondroblastic

- Fibroblastic

- Telangiectatic

- Low-grade

# Peripheral

High-grade

Low-Grade

Parosteal

Periostal

- Secondary

- in Paget's disease of bone
- post radiation

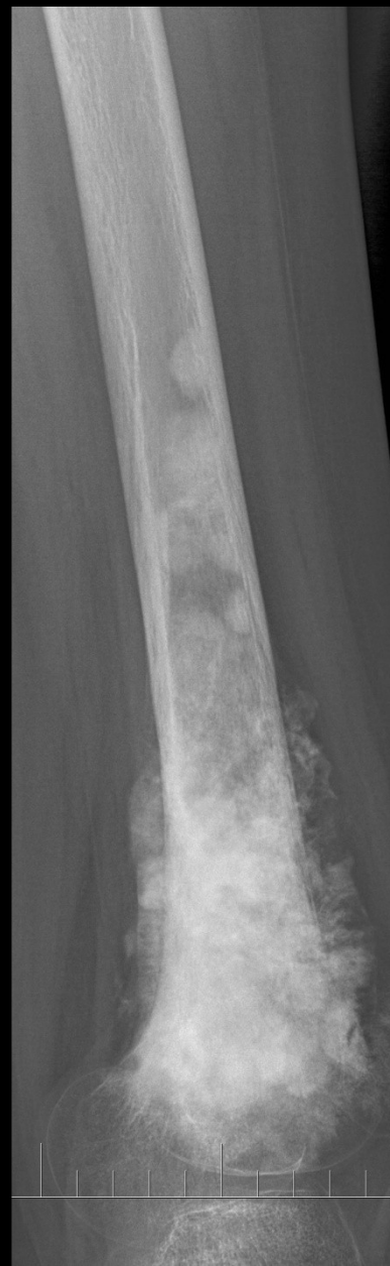
# *Symptoms*

- pain
  - during night, in rest
- swelling
- pathological fracture
- metastases in the time of diagnosis  
in 10-25 % of patients

# ***Diagnostics***

- X-ray
- CT / MRI
- Scintigraphy
- Chest X- ray or spiral CT
- Ultrasonography
- Biopsy – excisional, needle

# Conventional osteosarcoma



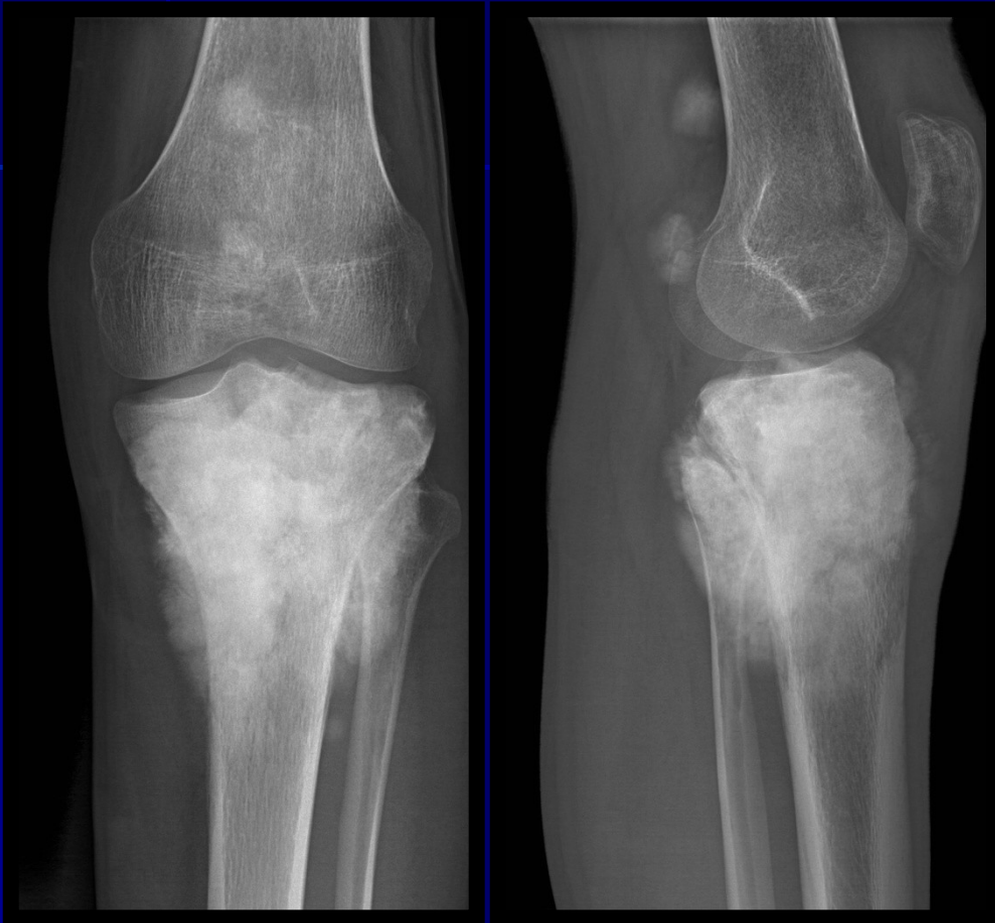
# Conventional osteosarcoma



# Parosteal osteosarcoma



# OSA



- Missed case
- Wrong prognosis

**Oncologic reflex**

# *Therapy*

- neoadjuvant chemotherapy
- surgery – radical resection / amputation
- adjuvant chemotherapy
- Metastasectomy in lungs
  
- Chemotherapy: (EURAMOS protocol)
  - metotrexat, doxorubicin, adriamycin, cisplatina, ifosfamid, etoposid.
- In low-grade OSA – only surgical treatment
  
- OSA is a radioresistant tumor

# *Prognostic factors*

- Metastases
- Size of the tumor
- Axial localisation
- Radicality of surgery
- Response to chemotherapy

# *Prognosis*

## *– 5 years survival*

- 70% - conventional high-grade OSA without MTS and with good response to chemotherapy (to 10 % of vital tumor cells)
- 90% - u low-grade OSA after radical surgery

# Chondrosarcoma

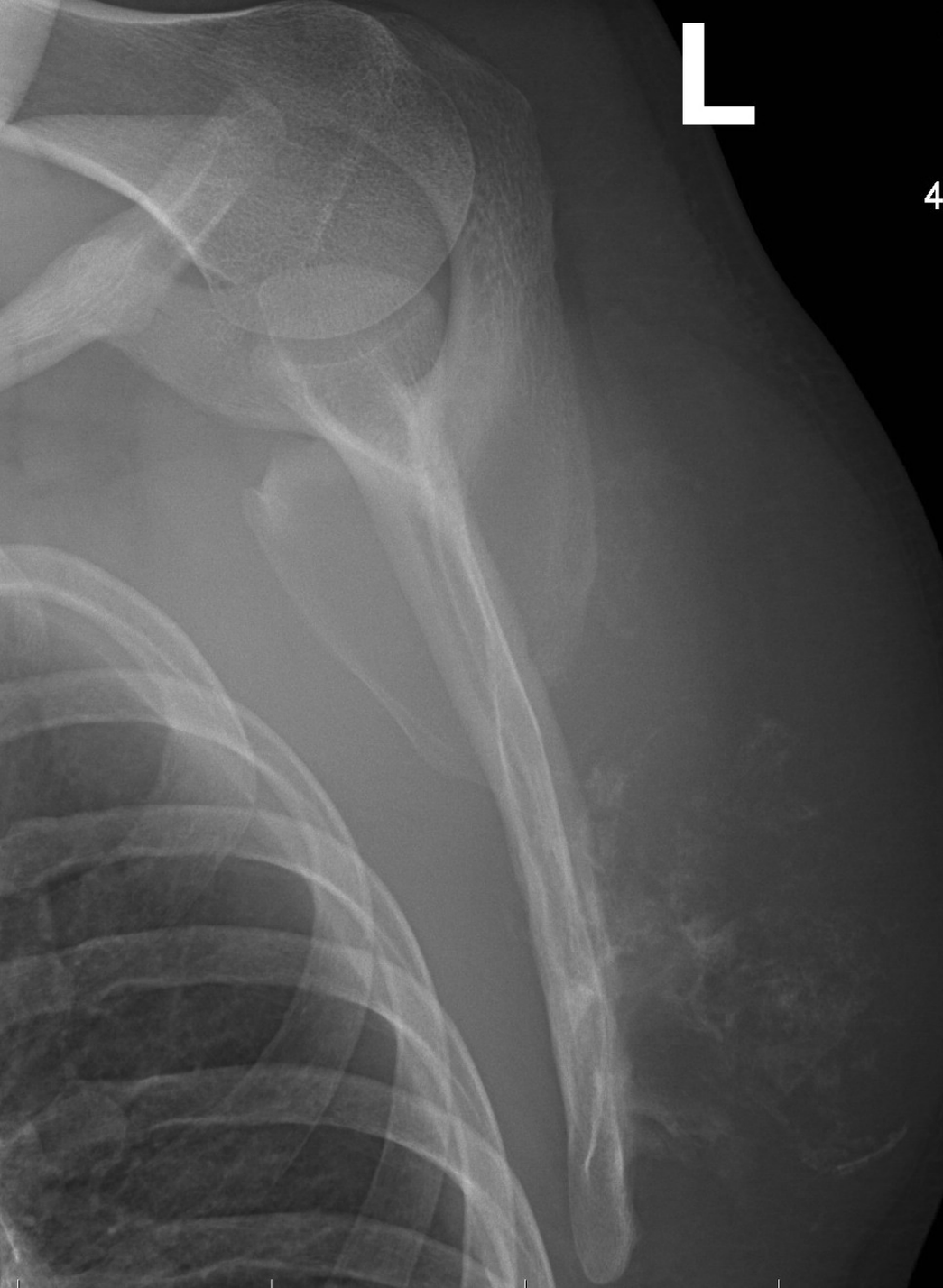
# *Epidemiology*

- 10% of primary malignant bone tumors
- Age:
  - primary: 40 – 60 years
  - secondary: 25 – 45 years
- Localisation-  
pelvis, proximal femur, proximal humerus

# ***Etiology***

## ■ **Secondary**

- **Multiple enchondromas** (M.Ollier, Maffucci sy)
- **Exostosis disease**  
cartilage over 2 cm
- **Chondroblastoma, chondromyxoid fibroma ...**



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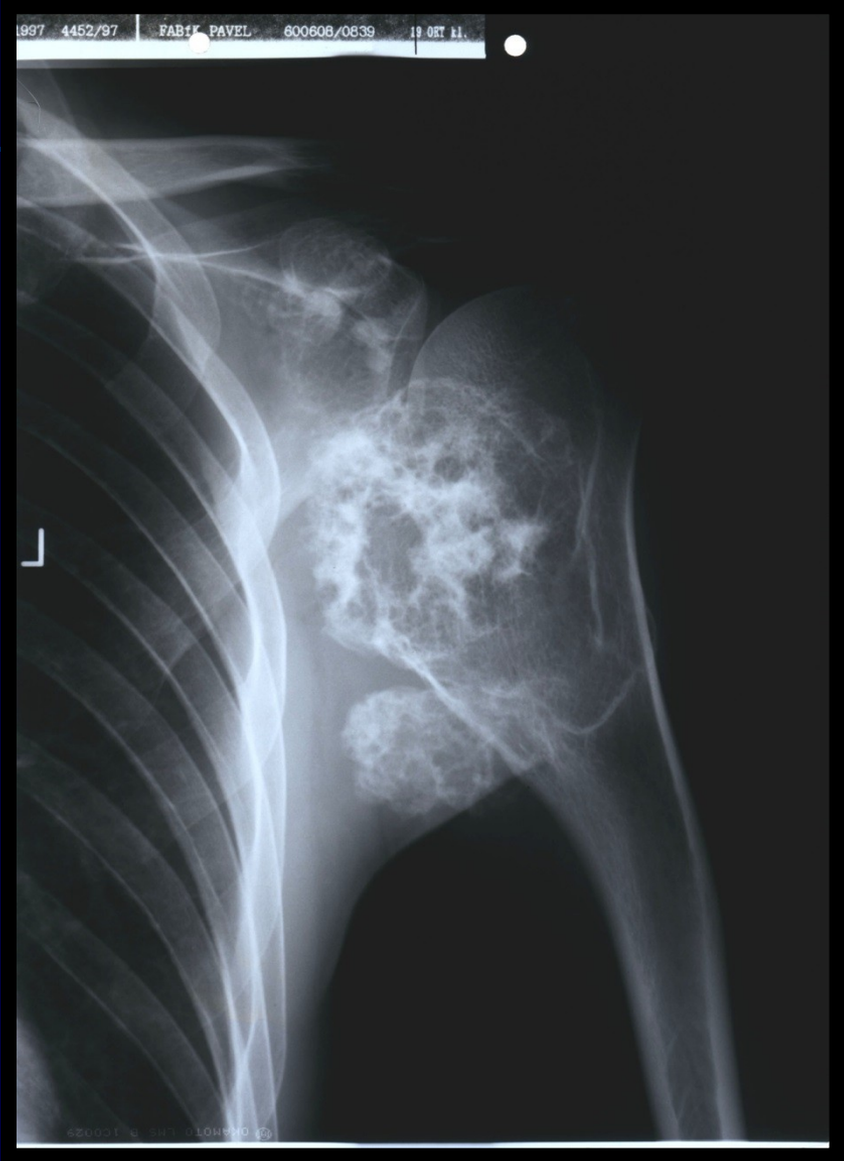
## Calcifications

# Chondrosarcoma





# Chondrosarcoma



# Chondrosarcoma



# Chondrosarcoma



# *Therapy*

- Radical resection – wide resection, amputation
- Metastasectomy in lungs
- Chemoresistant tumor
- Radioresistant tumor

# *Prognosis*

- **Prognostic factors:**
  - Radicality of surgery
  - Size
  - Histological grading
- **In intralesional surgery – 90% risk of local recurrence and lung metastases**
- **Prognosis:**
  - Conventional low-grade 90% 10 years
  - Conventional high-grade 20-40% 10 years
  - Dediferenciated sarcoma 15% 5 years

# Ewing sarcoma family

Group of high grade malignant round cells bone tumors with neuroectodermal differentiation and specific translocation.

- Ewing sarcoma
- PNET (periferal neuroectodermal tumor)
- Askin tumor of the chest wall
- Neuroblastoma in adults

# *Epidemiology*

- One new case /1 mil./ 1 year
- 5-25 years
- In metaphysis of long bones with extension into diaphysis and in flat bones (pelvis, scapula)

# *Symptoms*

- pain
- swelling
- Fever, redness,
- Leucocytosis, ESR elev.
- Biopsy- + identification of specific gene translocation t(11,22)q(24,12)

# Ewing sarcoma



# Ewing sarcoma



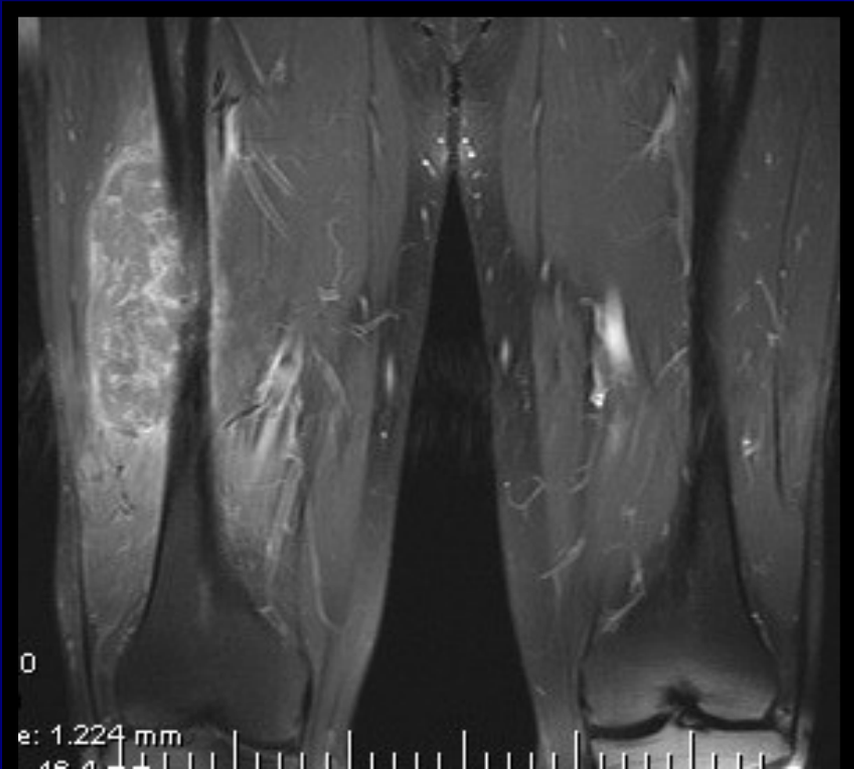
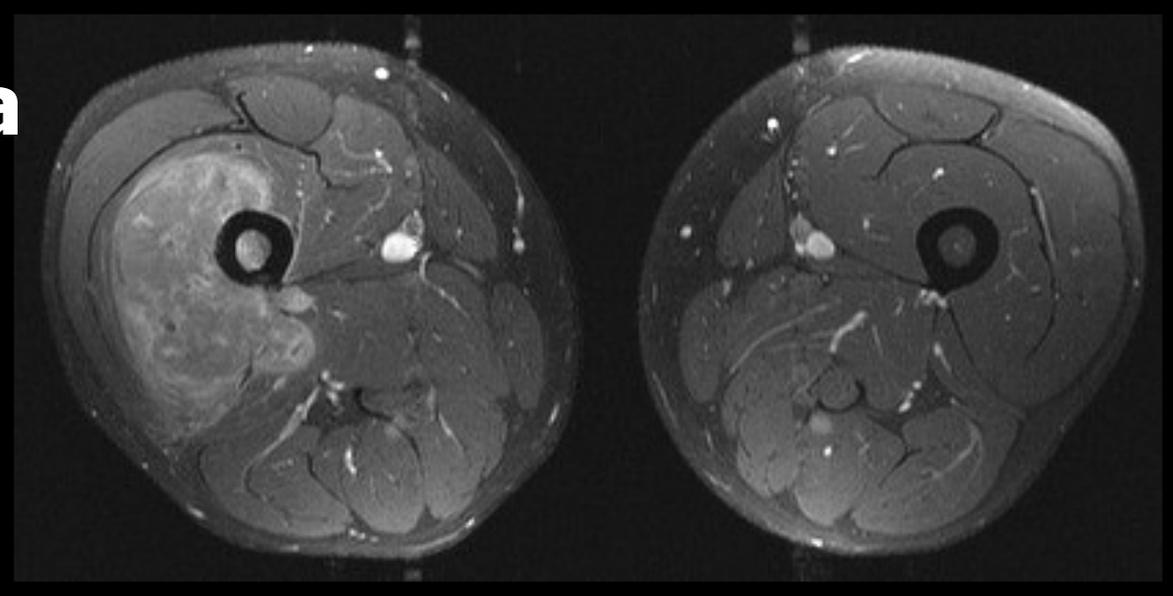
# Ewing sarcoma



# Ewing sarcoma



# Ewing sarcoma



# *Therapy*

- Chemo and radio sensitive tumor
- Neoadjuvant chemotherapy
- Local therapy:
  - Radiotherapy
  - Wide resection
  - Radiotherapy and wide resection
- Adjuvant chemotherapy
- In risk patients: transplantation of bone marrow
- Metastasectomy in lungs

# *Prognosis*

- Response to chemotherapy (systemic disease)
- **5-years survival in 60 % of patients**
- **Worse prognosis:**
  - metastases
  - **Size over 100cm<sup>3</sup>**
  - **Surgery not possible**
  - **Axial localisation**
  - **Local recurrence**
  - **Some genetic variants**

# Malignant fibrous histiocytoma in bone

- **In 5. decade**
- **In long bones – femur, tibia**
- **X- ray osteolytic lesion + cortical erosions, soft tissue mass**
  
- **Therapy: neoadjuvant chemotherapy + wide resection or amputation + adjuvant chemotherapy**
- **It is a radioresistant tumor**
- **Survival 35 % 5 years**



# Adamantinoma

- Very rare
- 90 % in tibia
- Therapy: radical resection
- Radioresistant tumor
- Prognosis – unclear



# Chordoma

- Axial localisation
- Osteolytic lesion
- Th- radical surgery or radiotherapy
- Prognosis- bad



# Malignant vascular tumors

- **Hemangioendotelioma**
- **Hemangiopericytoma**
- **Angiosarcoma**
  
- Osteolytic lesions
- Therapy: wide resection or amputation
- Chemotherapy in high grade
  
- Radiotherapy in non oper. cases



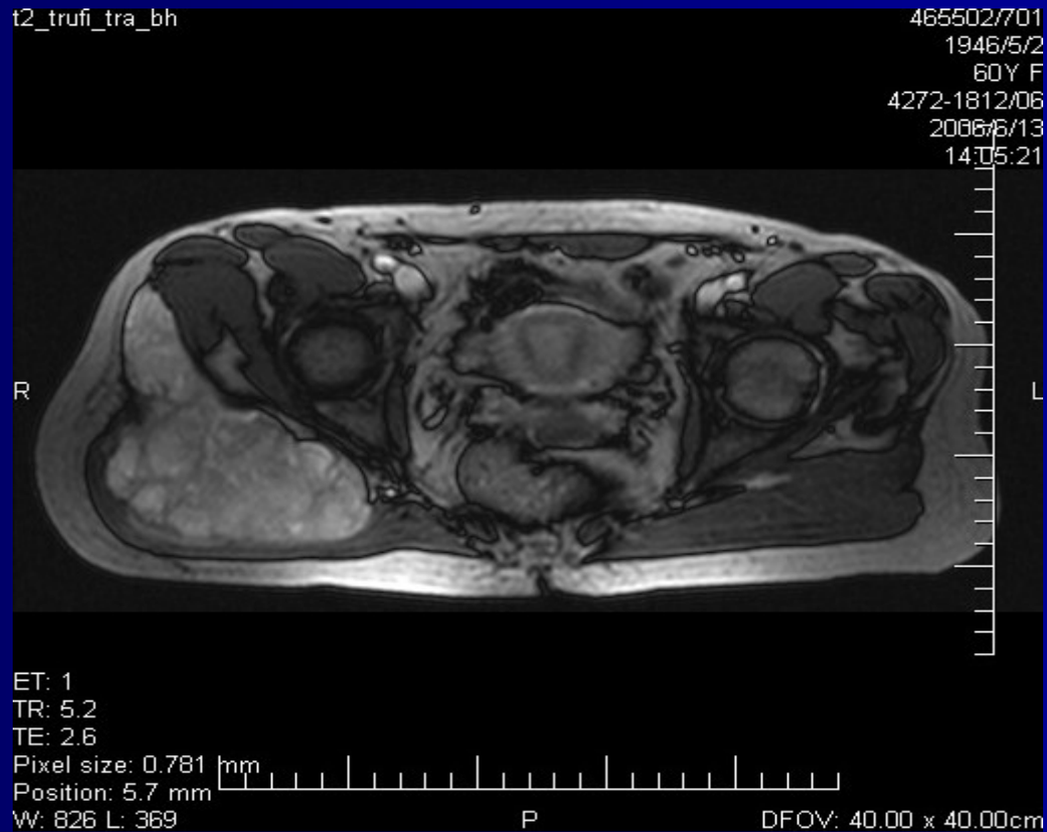
# Primary malignant soft tissue tumors - sarcomas

- MFH
- Synovialo-Sa
- Lipo-Sa
- Leiomyo-sa
- Fibro-sa
- Malignant schwannoma

# Primary soft tissue sarcomas

- Chondrosarcoma of soft tissue
- Lymphoma
- Malignant mesenchymal sarcoma

# Malignant fibrous histiocytoma MFH



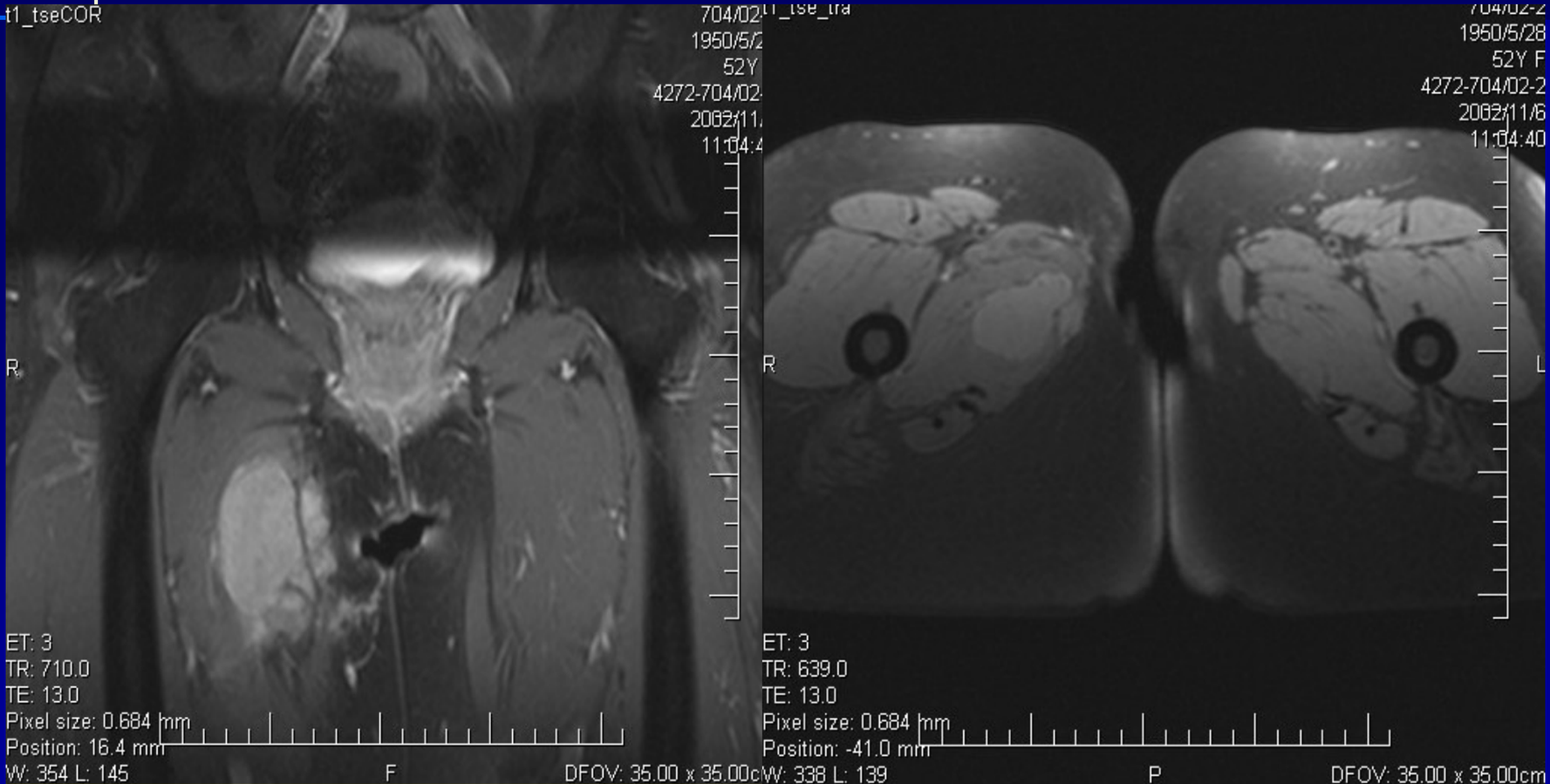
# MFH



# Therapy

- Neoadjuvant chemotherapy in G III
- Radical resection
- Adjuvant chemotherapy
- Radiotherapy in inoperable tumors

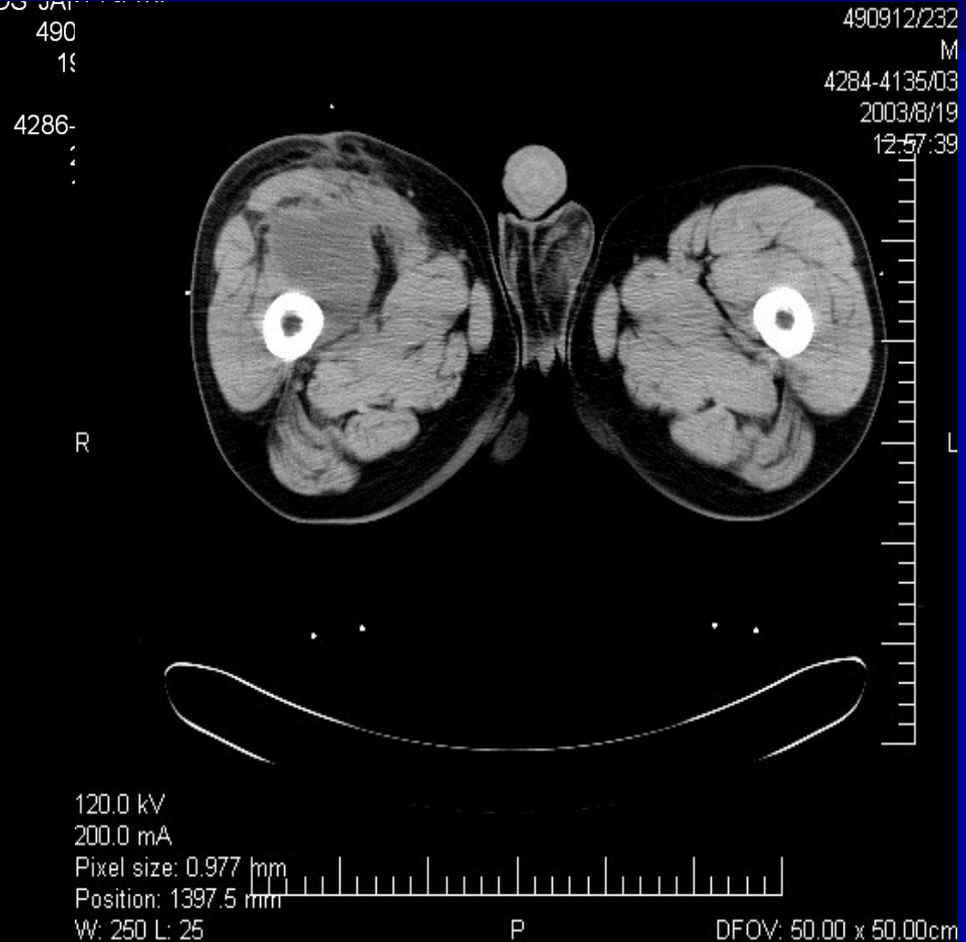
# Synovialo-Sa



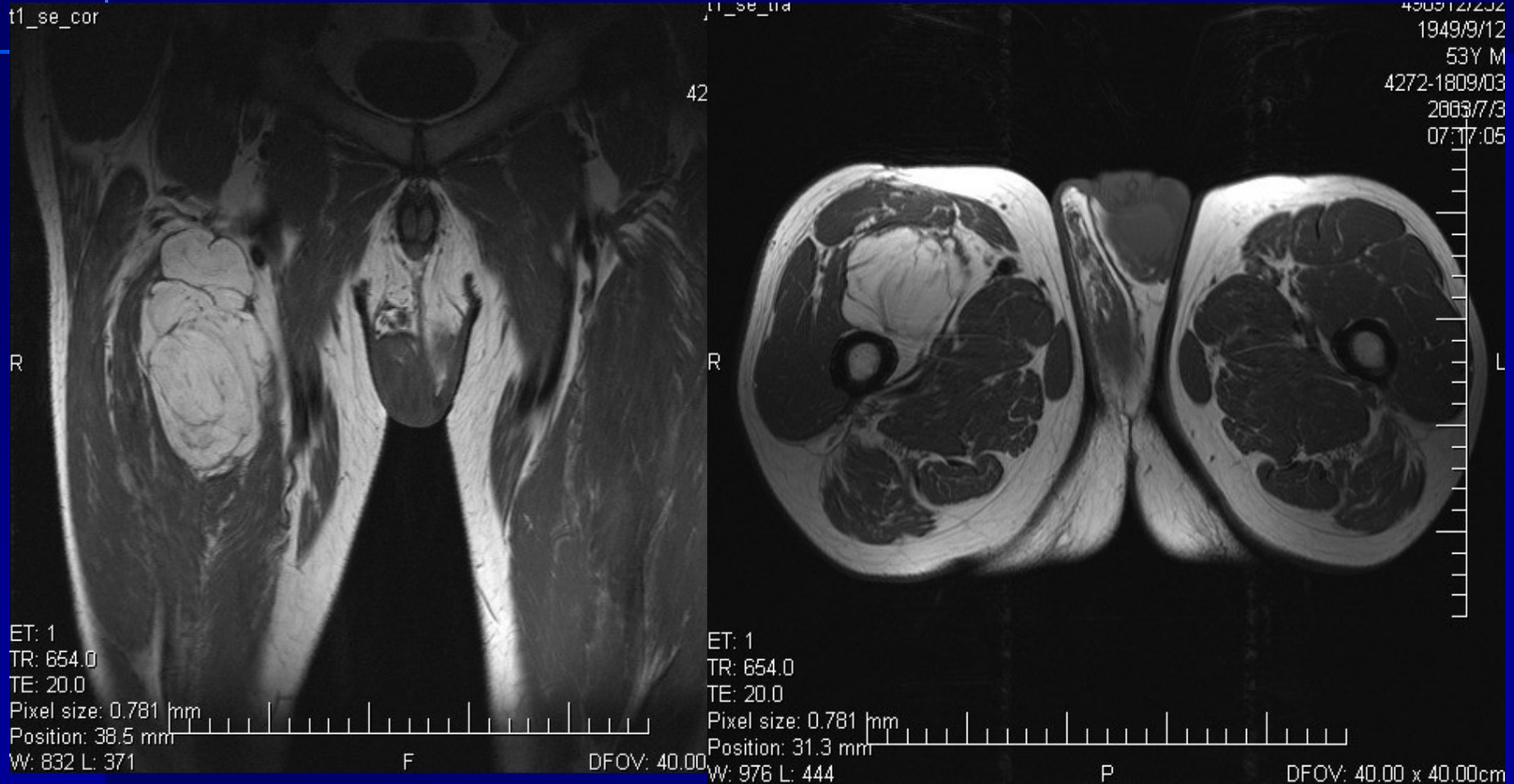
# Liposarcoma



64.5 kV  
Pixel size: 0.275 mm  
W: 826 L: 2094



# Liposarcoma



# Leiomyo-Sa

t2\_trufi\_tra\_bh

535909/118

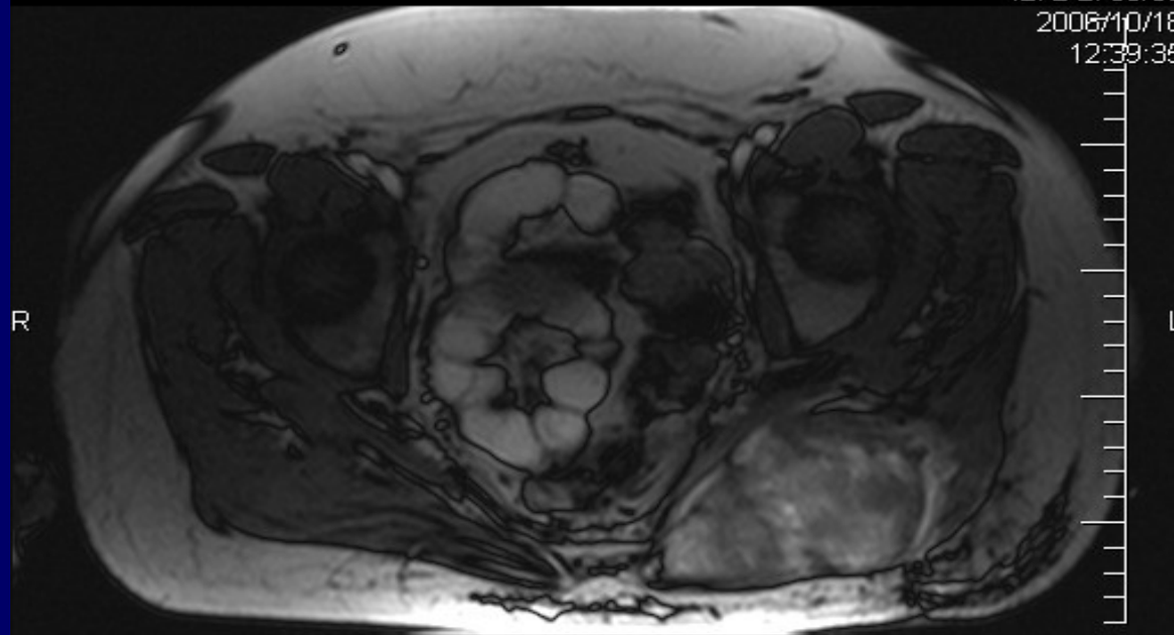
1953/9/9

53Y F

4272-2736/06

2006/10/18

12:39:35



ET: 1

TR: 5.2

TE: 2.6

Pixel size: 0.781 mm

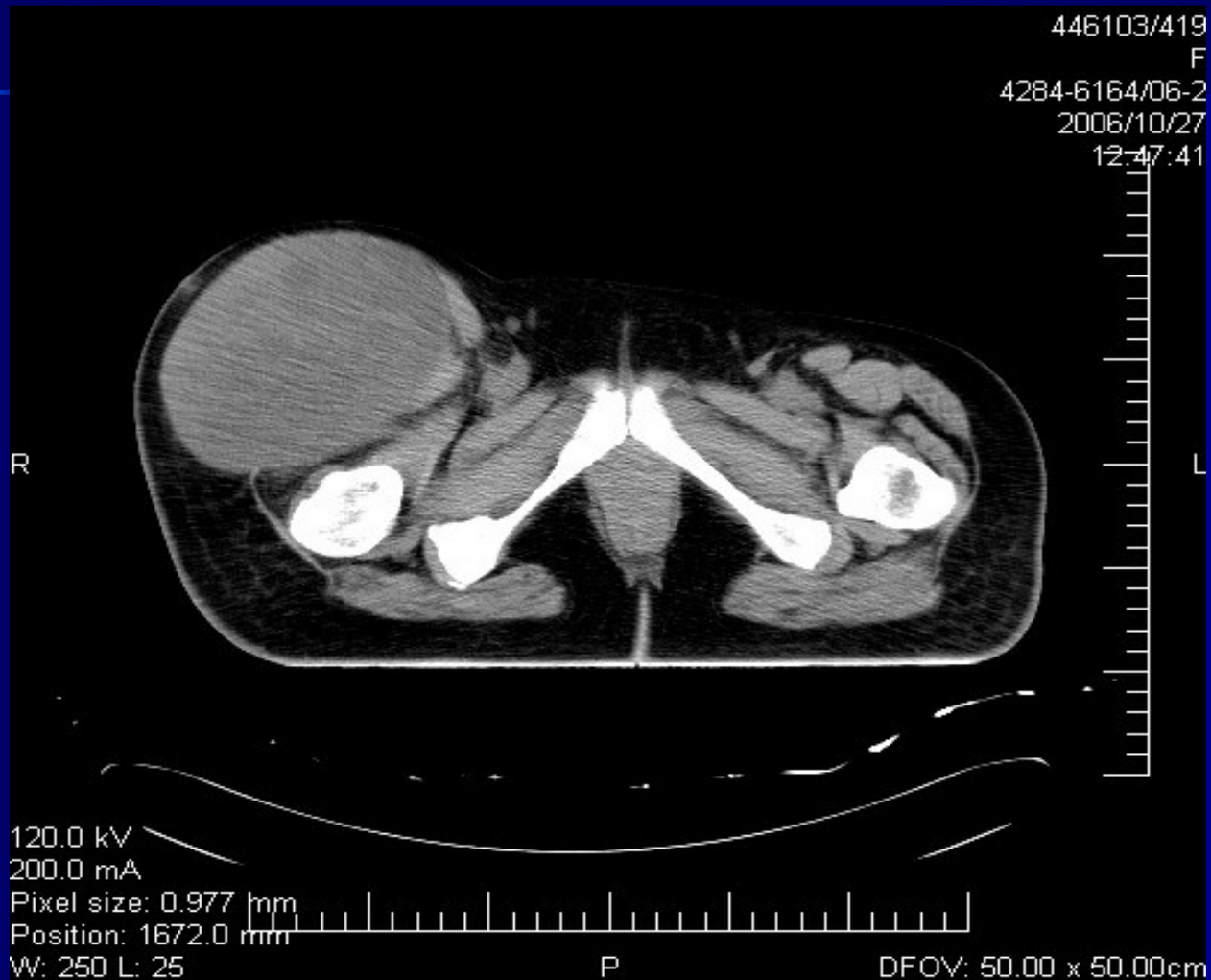
Position: 2.5 mm

W: 967 L: 426

P

DFOV: 40.00 x 40.00cm

# Fibro-Sa



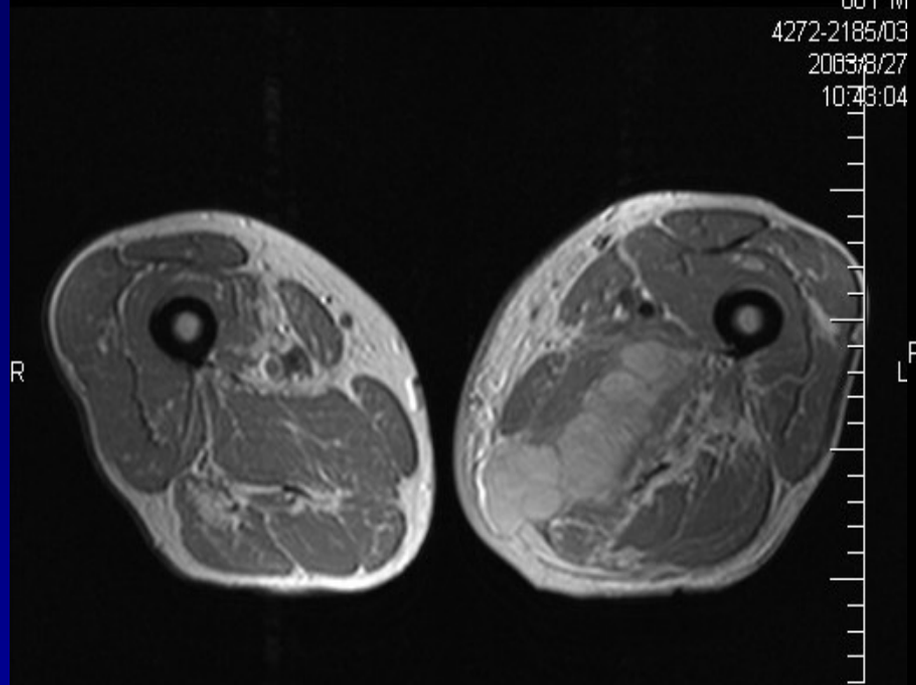
# Malignant schwannoma

AXIAL F->H  
pd+t2\_tse\_tra

REZNAK JAN  
230413/448  
1923/4/13  
80Y M  
4272-2185/03  
2003/8/27  
10:43:04

t2\_tse\_cor\_FS

230413/  
1923/4  
80Y  
4272-2185  
2003/8  
10:43



ET: 5  
TR: 4880.0  
TE: 14.0  
Pixel size: 0.781 mm  
Position: -8.6 mm  
W: 1087 L: 517

DFOV: 40.00 x 40.00cm



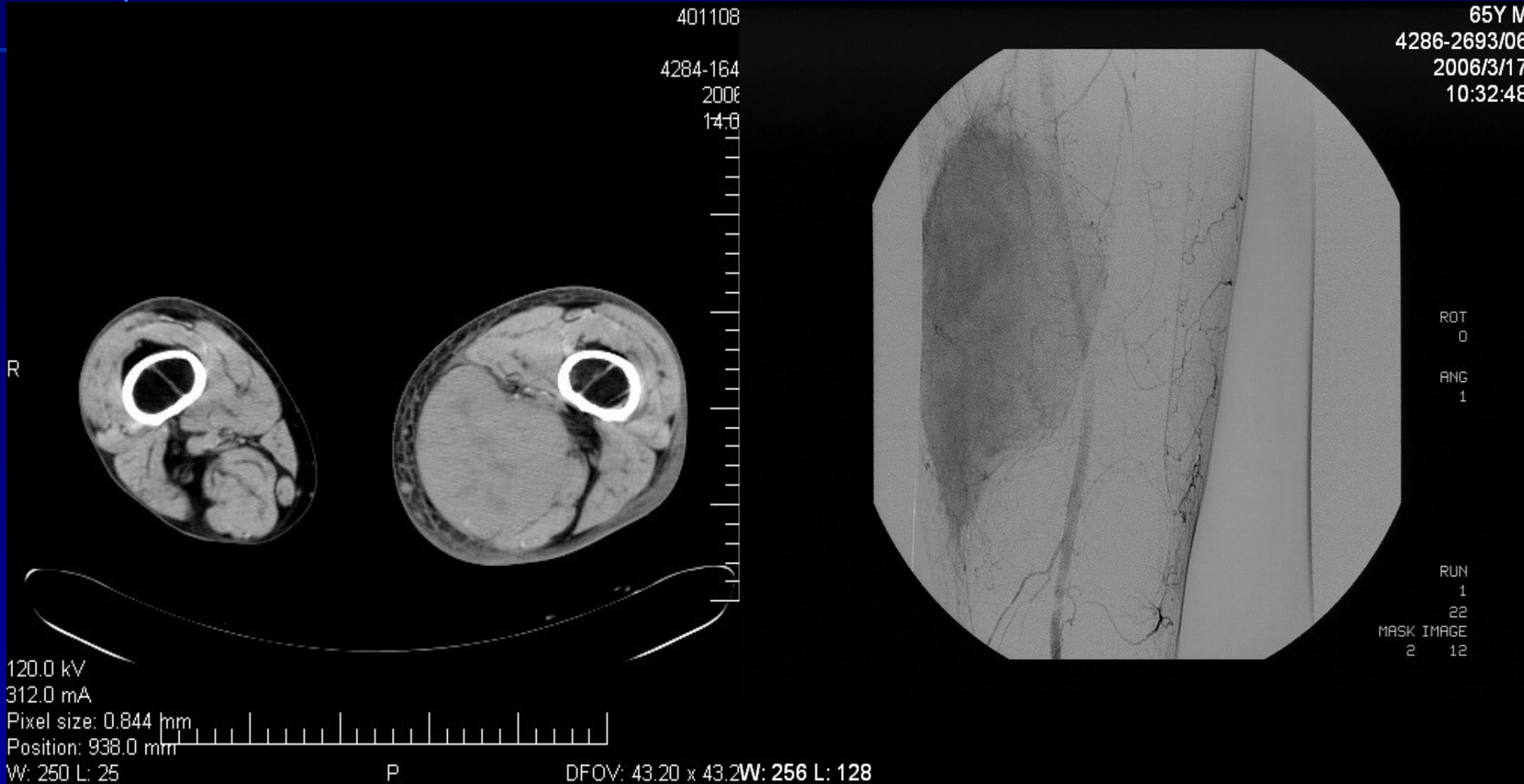
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TR: 3140.0  
TE: 108.0  
Pixel size: 0.879 mm  
Position: 10.2 mm  
W: 263 L: 94

DFOV: 45.00 x 45.00

# Extraskelletal chondrosarcoma



# Lymphoma



# Hemoblastosis in skeleton

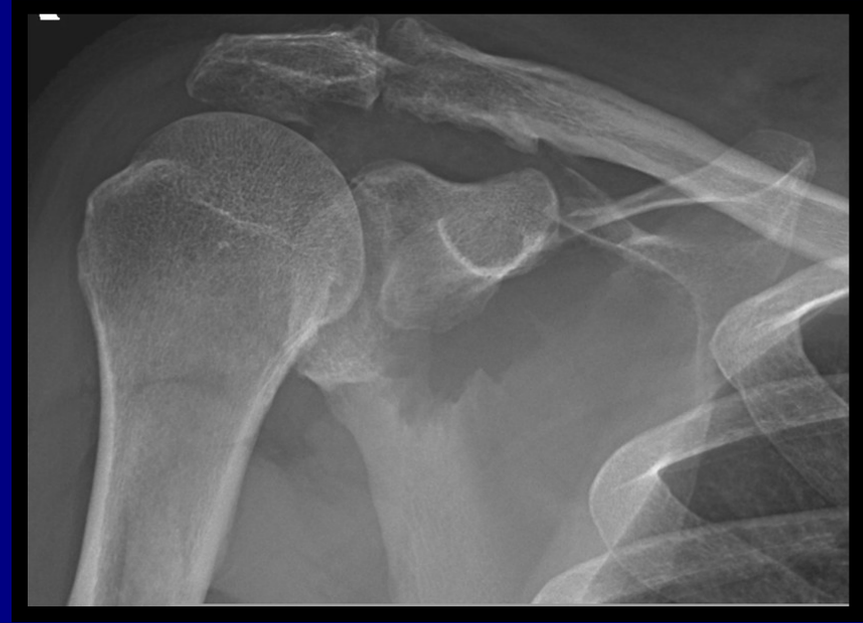
- **Primary bone tumors**
  - Multiple myeloma (plasmocytoma)
  - Solitary plasmocytom (myelom)
  - Primary bone lymfoma
  
- **Secondary lesions**
  - Hodgkin lymfoma
  - Non-Hodgkin lymfoma
  - Leukemia

Therapy- chemotherapy and radiotherapy  
in hematooncology

# Multiple myeloma

- Most frequent bone tumor
- 5 – 6. decade
- **Symptoms:**
  - pain
  - Pathological fracture
  - weaknes
  - letargy
  - infections
  - Renal failure
  - Headache





# Solitary plasmocytoma

- **Rare**
- **Osteolytic lesion**
- **Resection with replacement + chemotherapy**
- **Prognosis- better than in multiple myeloma**

# Primary bone lymphoma



# Skeletal metastases

# Carcinoma with MTS into the skeleton

- Breast
- Prostate
- Lung
- Kidney
- Thyreoidal gland

# Localisation

- Axial skeleton, pelvis, ribs, proximal femur and humerus

## X-ray

- Osteolysis, osteosclerosis, periosteal reaction



) mm  
9



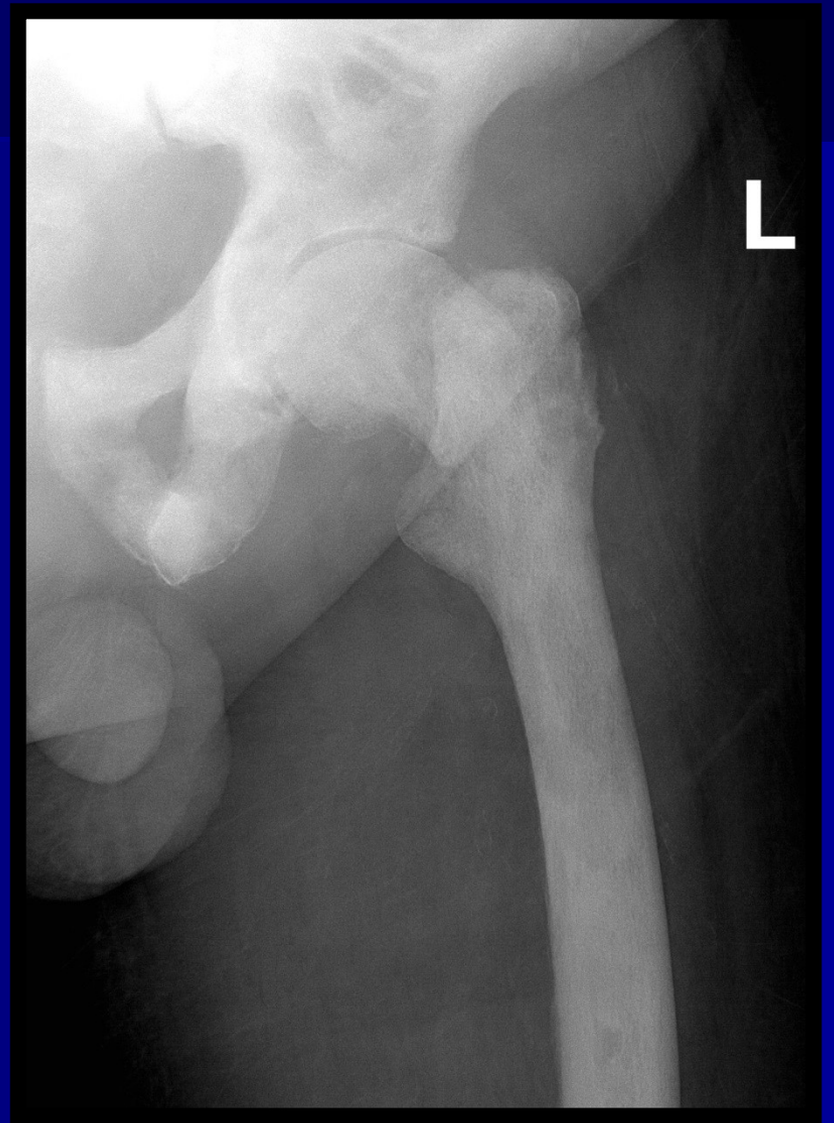
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3009 L: 4560

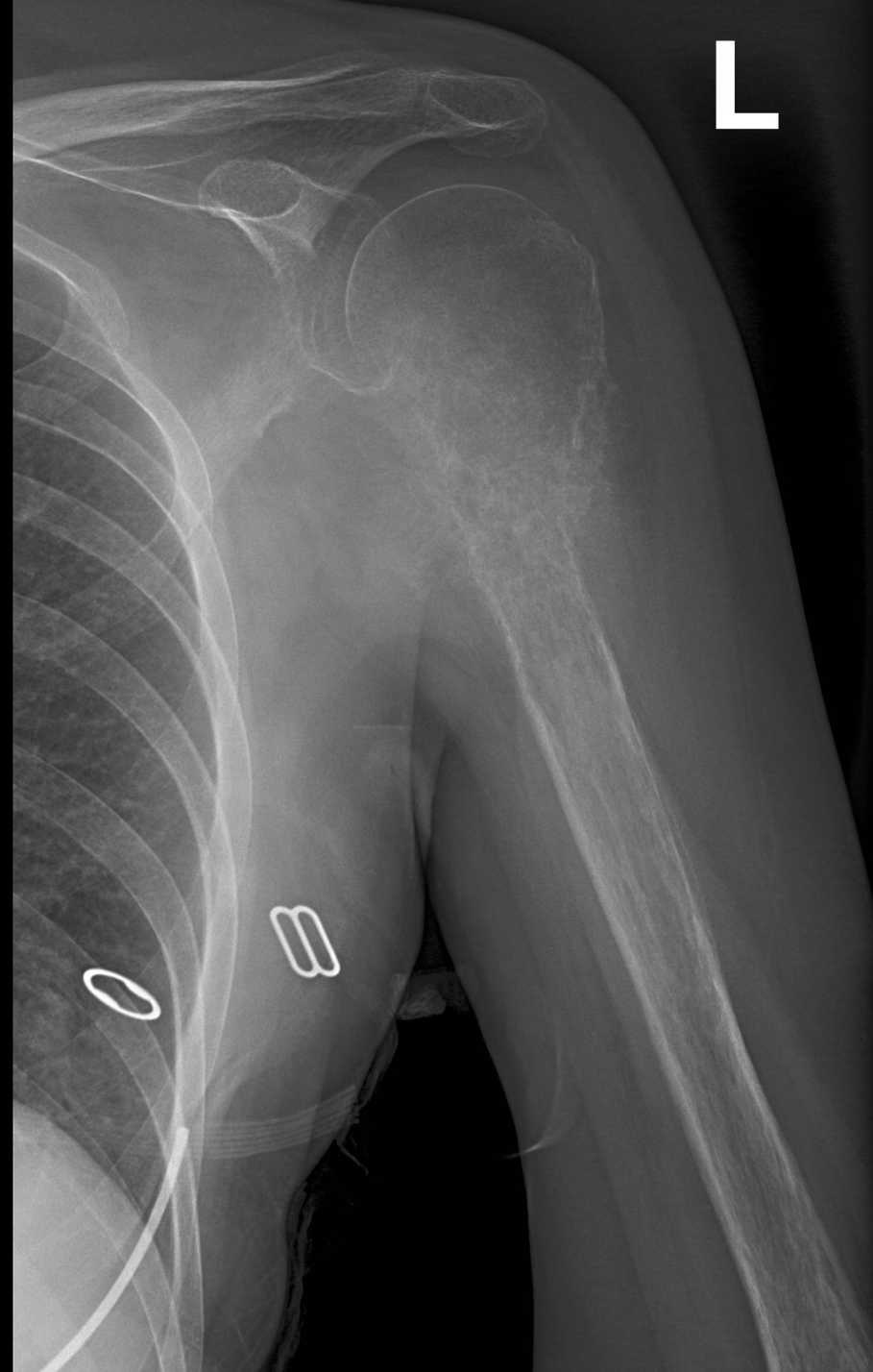
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kycle  
HIP



R



L



# Diagnostics

- History
- Radiological findings
- Scintigraphy
- Oncoscreening
- Biopsy

# Complications

- Pathological fracture
- Hypercalcemia
- Spinal cord lesion
- Anemia





# Mirel's score- risk of pathological fx

Points	1	2	3
Localisation	Upper extremity	Lower extremity	Trochanteric region
Pain	Mild	Moderate	Severe
Type	Osteoplastic	Mixed	Osteolytic
Size	<1/3 of diameter	1/3 – 2/3 diameter	>2/3 diameter
≤ 7 points	Risk 4%		Preventive OS not indicated
8 points	Risk 15%		OS ??
≥ 9 points	Risk 33% and more		OS is indicated



# Therapy

- Systemic therapy of carcinoma
  - Chemotherapy
  - Hormonal therapy
  - Immunotherapie, biological therapy ...
- Therapy of bone metastases
  - Bisphosphonates
  - Radioterapy
  - Surgery: radical, paliative
  - Conservative treatment
  - Others - RFA, embolisation ...
- Paliative management

# Surgery of bone metastases

Radical surgery – solitary MTS  
good prognosis

Simple surgery with mobilisation –  
multiple metastases  
• worse prognosis

# Types of surgery in MTS

	Removal of tumor	operation
No		Intramedullary nailing
Curretage		Cementoplasty + osteosynthesis
resection		Total replacement Intercalary spacer
Amputace		No
Spinal surgery		Instrumentation + fusion