

Diagnosis and Treatment of Patients with early and advanced Breast Cancer



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Osteooncology and Bone Health

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- **Versions 2002–2018:**
**Bischoff / Böhme / Brunnert / Dall / Diel / Fehm /
Fersis / Friedrich/ Friedrichs / Hanf / Huober /
Jackisch / Janni / Kolberg-Liedtke / Lux / Maas /
Nitz / Oberhoff / Schaller / Scharl / Schütz /
Seegenschmiedt / Solomayer / Souchon**
- **Version 2019:**
Diel / Kolberg-Liedtke

Bisphosphonates in Metastatic Breast Cancer

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- **Hypercalcemia**
- **Reduction of skeletal events (complications)**
- **Reduction of bone pain**
- **Increasing bone pain-free survival**
- **Treatment beyond osseous progression**
- **Use of bone resorption marker for therapy monitoring**
- **Bisphosphonates used alone for pain control**

	Oxford		
	LoE	GR	AGO
	1a	A	++
	1a	A	++
	1a	A	++
	1a	A	++
	5	D	++
	5	D	-
	5	D	-

Denosumab in Metastatic Breast Cancer

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- **Reduction of hypercalcemia**
- **Reduction of skeletal complications**
- **Reduction of bone pain**
- **Increasing bone pain-free survival**
- **Treatment beyond progression**
 - Progression while on bisphosphonates
- **Use of bone resorption markers for therapy monitorin**
- **Denosumab alone for pain control**

Oxford		
LoE	GR	AGO
1a	A	++
1a	A	++
1a	A	++
1b	A	++
5	D	+
4	C	+/-
5	D	-
5	D	-

Longer-Interval vs Standard Dosing of Zoledronic Acid

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- **¹ CALGB 70604 trial: n=1822 patients with metastatic breast cancer, metastatic prostate cancer, or multiple myeloma, 795 completed the study**

**SRE after 2 yrs: 29.5 % zoledronic acid every 4 weeks
 28.6 % zoledronic acid every 12 weeks**

- **² Optimze-2-trial: n=460 with metastatic breast cancer**

**SRE after 1 year³: 22.0% zoledronic acid every 4 weeks
 23.2% zoledronic acid every 12 weeks**

- ¹ Himmelstein et al. Effect of Longer-Interval vs Standard Dosing of Zoledronic Acid on Skeletal Events in Patients With Bone Metastases: A Randomized Clinical Trial. JAMA 317(1):48-58. 2017**
- ² Horobagyi GN et al. Continued Treatment Effect of Zoledronic Acid Dosing Every 12 vs 4 Weeks in Women With Breast Cancer Metastatic to Bone: The OPTIMIZE-2 Randomized Clinical Trial. JAMA Oncol 3(7):906-912, 2017**
- ³ Patients eligible for this trial had prior exposure to zoledronate or pamidronate for approx. 1 year or more**

Bone Modifying Agents for the Therapy of Bone Metastases

Oxford		
LoE	GR	AGO
1a	A	++
1a	A	++
1a	A	++
1a	A	++
1a	A	++
1a	A	+
1a	A	++
1a	A	++
4	C	-
5	D	--

- **Clodronate PO 1600 mg daily**
- **Clodronate IV 1500 mg q3w / q4w**
- **Pamidronate IV 90 mg q3w / q4w**
- **Ibandronate IV 6 mg q3w / q4w**
- **Ibandronate PO 50 mg daily**
- **Zoledronate IV 4 mg**
 - q4w
 - q12w
- **Denosumab 120 mg s.c. q4w**
- **Denosumab 120 mg s.c. q12w**
- **Other dosing or schedules, e.g. derived from adjuvant studies or therapy of osteoporosis**

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Skeletal Metastases

Treatment with Radionuclids

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- **Tumor progression after standard treatment of multiple / disseminated metastases and intolerable bone pain (Prerequisite: hot spots in the bone scintigraphy)**
 - ¹⁸⁶Rhenium-hydroxyethyliden-diphosphonat
 - ¹⁵³Samarium
 - ⁸⁹Strontium
 - ²²³Radium
 - ¹⁷⁷Lu-EDTMP

	Oxford		
	LoE	GR	AGO
	1b	B	+
	2b	B	+
	1b	B	+
	1b	B	+
	1b	B	+

Caveat: the potential benefits should be weighed against the risk of myelosuppression with pancytopenia

Metastatic Bone Disease of the Spine

Indications for surgery

Oxford LoE: 2b

GR: C

AGO: ++

- **Spinal cord compression**
 - With progressive neurological symptoms
 - With pathological fractures
- **Instability of the spine**
- **Lesions in pre-irradiated parts of the spine**

Bone Metastases Acute Spinal Cord Compression / Paraplegia

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	Oxford		
	LoE	GR	AGO
<ul style="list-style-type: none"> Decompression surgery, reduction of tumor volume, stabilisation surgery (< 24 h) and irradiation of the spine (RT) 	2b	C	++
<ul style="list-style-type: none"> Irradiation of the spine (< 24 h) +/- steroids 	3b	C	++
<ul style="list-style-type: none"> Immediate start of treatment 	1c	D	++

Clinical trials have included patients with different tumor entities!

Surgery for Bone Metastases

Technical Aspects

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Spine and limbs

Oxford LoE: 3b

GR: C

AGO: +

- Marrow splints
- Plate osteosynthesis
- Compound osteosynthesis (replacement by PMMA and osteosynthesis)
- Vertebral replacement by titanspacer
- Tumor-Endoprothesis
- Vertebroplasty / Kyphoplasty +/- thermoablation of the tumor
- Kypho-IORT (in studies only)*
- Resection of involved bone in oligometastatic disease (sternum, ribs, vertebrectomy and replacement with spondylodesis)

* Study participation recommended

Metastatic Bone Disease: Radiotherapy (RT)

	Oxford		
	LoE	GR	AGO
With fracture risk	1a	B	++
With functional impairment	1a	B	++
With bone pain	1a	B	++
Single dose RT = fractionated RT	2a	B	++
With neuropathic bone pain	1b	B	++
Asymptomatic isolated bone metastasis	5	D	+/-
Reduction of radiation induced pain flare by dexamethasone	1b	B	+
Radiotherapy in combination with hyperthermia	2b	B	+/-

Bone metastases

- With fracture risk
- With functional impairment
- With bone pain
 - Single dose RT = fractionated RT
- With neuropathic bone pain
- Asymptomatic isolated bone metastasis
- Reduction of radiation induced pain flare by dexamethasone
- Radiotherapy in combination with hyperthermia

Limited studies included breast cancer patients!

Metastatic Bone Disease

Recurrent Bone Pain after RT

	Oxford		
	LoE	GR	AGO
Single dose RT *	3b	C	++
Fractionated RT *	3b	C	++
Radionuclid therapy	3b	C	+
Magnetic resonance-guided focused ultrasound	1b	B	+
Radiofrequency ablation	4	C	+
Cryoablation	4	C	+

Recurrent bone pain in pre-irradiated parts of the skeleton

- Single dose RT *
- Fractionated RT *
- Radionuclid therapy
- Magnetic resonance-guided focused ultrasound
- Radiofrequency ablation
- Cryoablation

* Dosing and fractionation depending on location, interval from first RT, and dose and fractionation of first radiotherapy.

Side-Effects and Toxicity – Bisphosphonates (BP) and Denosumab (Db)

	<u>LoE</u>
<ul style="list-style-type: none"> ▪ Renal function deterioration due to IV-aminobisphosphonates 	1b
<ul style="list-style-type: none"> ▪ Osteonecrosis of the jaw (ONJ) mostly under IV-BP and denosumab therapy (1.3 % / 1.8 %) <ul style="list-style-type: none"> ▪ Association with (simultaneous) anti-angiogenetic therapies 	1b
<ul style="list-style-type: none"> ▪ Severe hypocalcemia (Dmab > BPs) 	1b
<ul style="list-style-type: none"> ▪ Acute Phase Reaction (IV Amino-BPs, Db) 10–30 % 	1b
<ul style="list-style-type: none"> ▪ Gastrointestinal side effects (oral BPs) 2–10 % 	1b
<ul style="list-style-type: none"> ▪ Atypical femur fractures (absolute risk of 11 per 10,000 person years of BP use) 	2b
<ul style="list-style-type: none"> ▪ Extremely rare: Uveitis / Scleritis under BP treatment 	4

Frequent side effects under treatment with BPs and Denosumab



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Drug	Acute phase-reaction	Kidney Tox.	Upper GI	Diarrhea	Osteo necrosis of the jaw	
Clodronate 1500 i.v.	0	+	0	0	0	Non-Amino.
Clodronate 1600 p.o.	0	0	+	+	0	Non-Amino.
Ibandronate 50 mg p.o.	0	0	+	0	0	Aminobisp.
Ibandronate 6 mg i.v.	+	0	0	0	+	Aminobisp.
Zoledronate 4 mg i.v. q4w oder q12w	+	+	0	0	+	Aminobisp.
Pamidronate 90 mg i.v.	+	+	0	0	+	Aminobisp.
Zoledronate 4 mg i.v. q6m	+	0	0	0	0	Aminobisp.
Denosumab 120 mg sc q4w	0	0	0	+	+	

Cave: Hypocalcemia under antiresorptive therapy in pts with bone metastases!

Recommendations for Prevention of Osteonecrosis of the Jaw (ONJ)

Oxford LoE: 4

GR: C

AGO: +

- During bisphosphonate or denosumab treatment, avoid any elective dental procedures involving jaw bone manipulations should be avoided during treatment with bisphosphonates or denosumab (**LoE 2b**)
- Optimize dental status before start of bisphosphonate or denosumab treatment (**LoE 2b**)
- Inform patients about ONJ risk and educate about early symptom reporting
- In case of high risk for ONJ, use oral bisphosphonate
- Good oral hygiene, limiting of alcohol intake and stopping smoking should be recommended

In adjuvant bisphosphonate therapy, ONJ was rare

Adjuvant Bone Targeted Therapy for Improvement of Prognosis



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- **Clodronate (oral)**
 - Postmenopausal patients
 - Premenopausal patients
- **Aminobisphosphonate (iv or oral)**
 - Postmenopausal patients
 - Premenopausal patients
- **Denosumab (6 x 120 mg/3–4w + 14 x 120 mg/3m)**
 - Postmenopausal patients Stage II and III
- **Denosumab (60 mg s.c. q6m)**
 - Postmenopausal patients undergoing AI therapy

	Oxford		
	LoE	GR	AGO
	1a	A	+
	1a	B	+/-
	1a	A	+
	1a	B	+/-
	1b	B	-
	1b ^a	B	+/-

Dosage of Adjuvant Bisphosphonates for Improvement of Survival

- **Non-Aminobisphosphonates:**
- **Clodronat po 1600 mg/d (Bonafos / Clodronic acid)**
- **Clodronat po 1040 mg/d (Ostac / Clodronic acid)**

- **Aminobisphosphonates:**
- **Zoledronat iv 4 mg/6 m (Zometa / Zoledronic acid)**
- **Ibandronat po 50 mg/d (Bondronat / Ibandronic acid)**
- **Pamidronat po (orally not available in most countries)**
- **Risedronat po 35 mg/w (Actonel / Risedronic acid)**
- **Alendronat po 70 mg/w (Fosamax / Alendronic acid)**
- **Optimal duration yet to be defined; in adjuvant studies duration of BP treatment varied from 2–5 years**

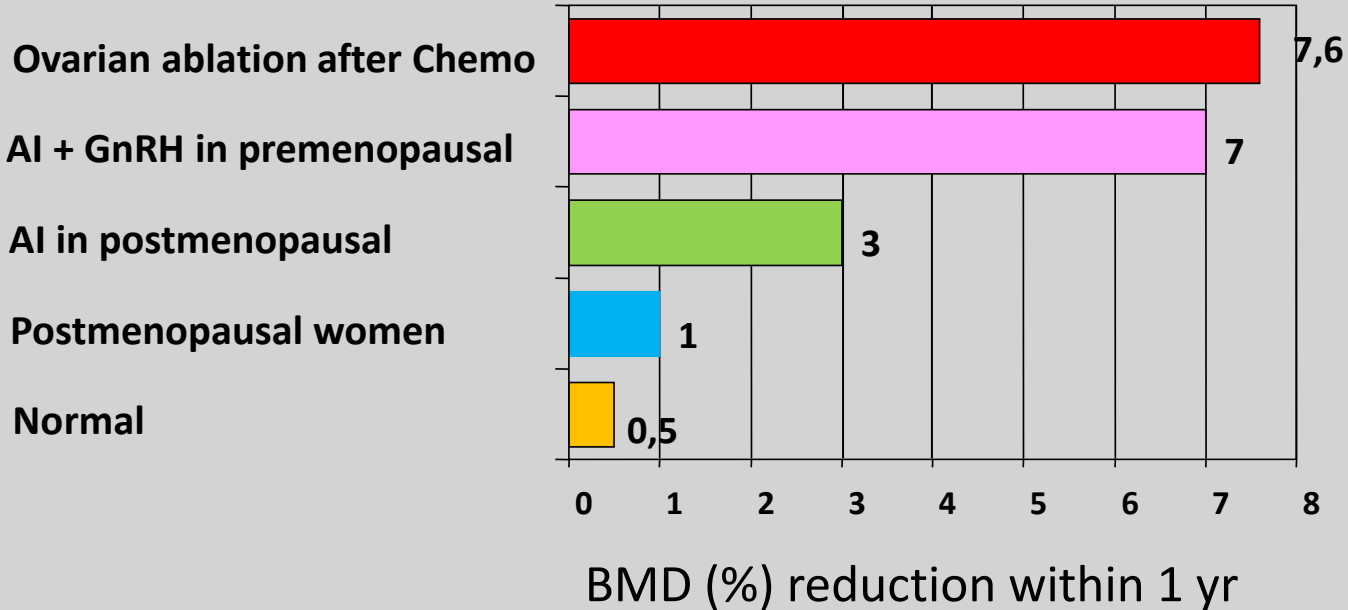
Aminobisphosphonates include:

Zoledronic acid (65 %), oral ibandronate (24 %), oral pamidronate (8 %), oral risedronate (2 %), oral alendronate (1 %) (data from EBCTCG-metaanalysis)

Reduction in bone density of individual agents

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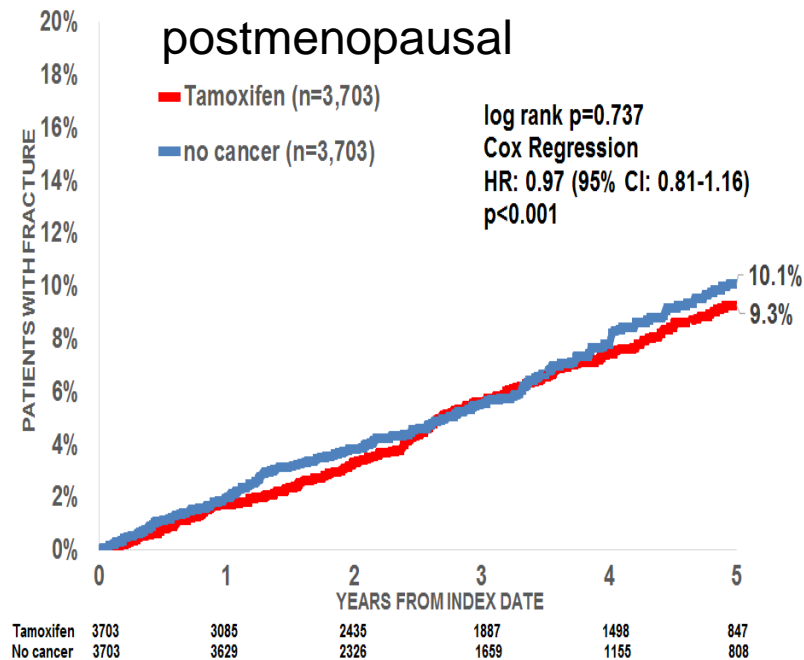
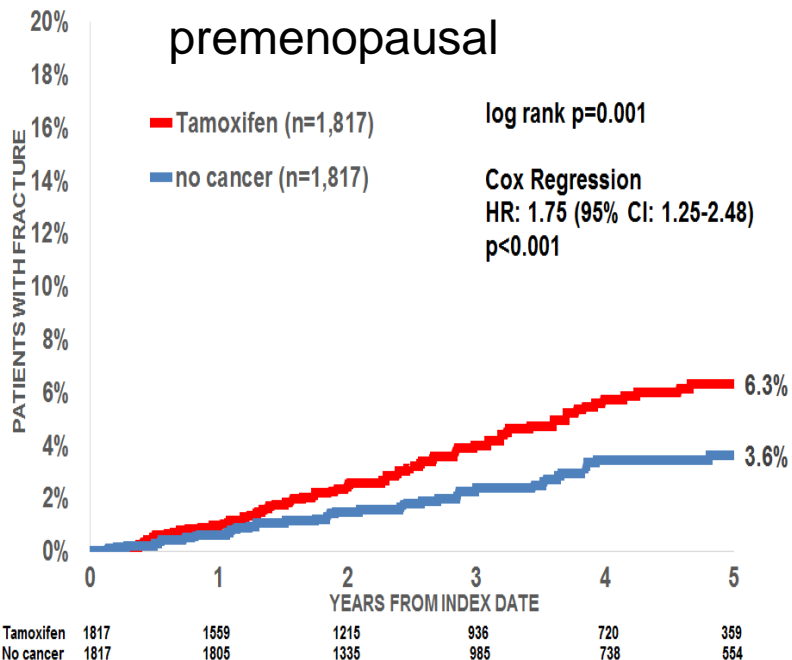


(1) Kanis JA Osteoporosis 22, 1997, (2) Gnant M SABCS 2004, (3) Shapiro CL, JCO 19:3305, 2001

Risk of osteoporosis and tamoxifen (fracture risk)

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Therapy and Prevention of Tumor Therapy-Induced Bone Loss / Osteoporosis



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	Oxford		
	LoE	GR	AGO
■ Bisphosphonates	1b	B	++
	1b	A	+
■ Denosumab	1b	B	++
	1b	A	+
■ Hormone replacement therapy	5	D	-
■ Clinical risk assessment for osteoporosis at baseline	5	D	++
■ DXA-Scan at baseline in pts with endocrine therapy and/or with premature menopause	5	D	+
■ Antiresorptive therapy in pts. with reduced bone density	5	D	++
■ Repeat DEXA-scan based on risk	5	D	+

Therapy and Prevention of Tumor Therapy-Induced Bone Loss / Osteoporosis



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Further recommendations (based on DVO-guidelines for treatment, diagnosis and prevention of osteoporosis)*

	Oxford		
	LoE	GR	AGO
■ Physical activity	4	C	++
■ Avoiding immobilisation	4	C	++
■ Calcium (1000–1500 mg/d)**	4	C	++
■ Vitamine D3 suppl. (800–2000 U/d or 20,000 U/w)	4	C	++
■ Cessation of smoking, reduction of alcohol	2b	B	++
■ Avoiding BMI < 20 mg/m²	3b	C	++
■ Antiresorptive therapy after discontinuation of Denosumab	4	C	+/-
■ Drugs approved for the treatment of osteoporosis in adults (see next slide)			

* http://www.dv-osteologie.org/dvo_leitlinien/dvo-leitlinie-2014; revised version expected in 2018
 ** if nutritional supply is insufficient, (in combination with Vit D3 only)

Effect of Denosumab Discontinuation

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FREEDOM / FREEDOM Extension Trial

N=1001, ≥ 2 dose of Denosumab or placebo, follow up ≥ 7 months after discontinuation treatment

Vertebral fracture rate per 100 participant year :

- 1.2 during denosumab therapy
- 7.1 after denosumab therapy
- 8.5 placebo

Non vertebral fracture rate per 100 participant year:

- 2.8 after denosumab vs. 3.8 placebo (n.s.)

Multiple vertebral fracture (% of all vertebral fractures):

60.7% after denosumab therapy vs. 38.7% placebo; p=0.049

Cummings SR et al. J Bone Miner Res 2017

Medical Treatment of Osteoporosis

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	Oxford		
	LoE	GR	AGO
▪ Alendronate 70 mg po/w*	1b	B	++
▪ Denosumab 60 mg sc/6m*	1b	B	++
▪ Ibandronate 150 mg po/m*	1b	B	++
▪ Ibandronat e 3 mg iv/3 m	1b	B	++
▪ Parathyroid hormone (1-84) 100 µg sc/d	1b	B	+
▪ Raloxifene 60 mg po/d (improves spine only)	1b	B	+/-
▪ Risedronate 35 mg po/w*	1b	B	++
▪ Strontium ranelate 2 g po/d**	1b	B	+
▪ Teriparatide (1-34) 20 µg sc/d	1b	B	+
▪ Zoledronate 5 mg iv/12 m*	1b	B	++

* Drugs tested in clinical studies with breast cancer patients and tumor therapy-induced osteoporosis

** Elevated risk of myocardial infarction. Substance restricted to postmenopausal pats. with severe osteoporosis and high risk of fractures

<http://www.dv-osteologie.org/uploads/Leitlinie%202014/DVO-Leitlinie%20Osteoporose%202014Rev%20Kitteltaschenversion%2015.12.2014.pdf>
Revised version expected in 2018



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TABELLE 4.2.: INDIKATION FÜR EINE MEDIKAMENTÖSE OSTEOPOROSETHERAPIE NACH RISIKOPROFIL in Abhängigkeit von Geschlecht, Lebensalter, DXA-Knochendichte und weiteren Risikofaktoren.¹

Lebensalter in Jahren		T-Score (Nur anwendbar auf DXA-Werte. Die Wirksamkeit einer medikamentösen Therapie ist für periphere Frakturen bei einem T-Score > -2,0 nicht sicher belegt.)				
Frau	Mann ²	-2,0 bis -2,5	-2,5 bis -3,0	-3,0 bis -3,5	-3,5 bis -4,0	< -4,0
50-60	60-70	Nein	Nein	Nein	Nein	Ja
60-65	70-75	Nein	Nein	Nein	Ja	Ja
65-70	75-80	Nein	Nein	Ja	Ja	Ja
70-75	80-85	Nein	Ja	Ja	Ja	Ja
>75	>85	Ja	Ja	Ja	Ja	Ja

¹ Alternative Risikomodellierungen können bei Bedarf vergleichend zu Rate gezogen werden (siehe Langfassung).
² bei Verwendung eines männlichen Referenzkollektivs für die T-Scores

Therapieindikation auch schon bei um 1,0 höherem T-Score^{3,4}, wenn:

- Glukokortikoide oral ≥ 2,5 mg und < 7,5 mg Prednisolonäquivalent tgl. (außer bei rheumatoider Arthritis +0,5)
- Diabetes mellitus Typ 1
- ≥ 3 niedrigtraumatische Frakturen in den letzten 10 Jahren im Einzelfall (mit Ausnahme von Finger-, Zehen-, Schädel- und Knöchelfrakturen)