

DECLARATION OF INTERESTS

None related to the presented work.

Icro Meattini declares occasional small fees received for advisory boards supported by Eli Lilly, Novartis, SeaGen, Gilead, Daiichi Sankyo, AstraZeneca, Pfizer.



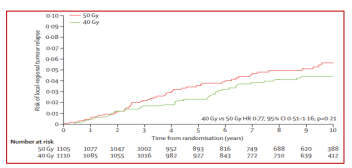
Hypofractionation

Partial Breast Irradiation
Omission of whole breast irradiation
Oligometastatic disease
Integration with systemic therapies

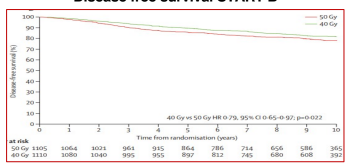
Moderate hypofractionation

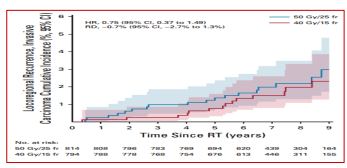
Efficacy

Cumulative risk of locoregional relapse

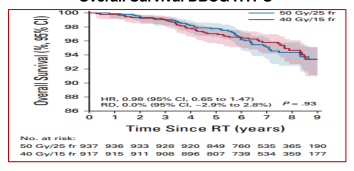


Disease free survival START B





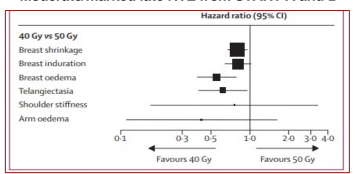
Overall Survival DBCG HYPO



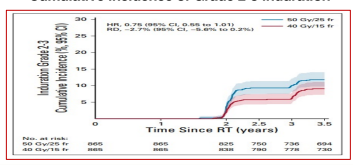
Moderate hypofractionation

Safety

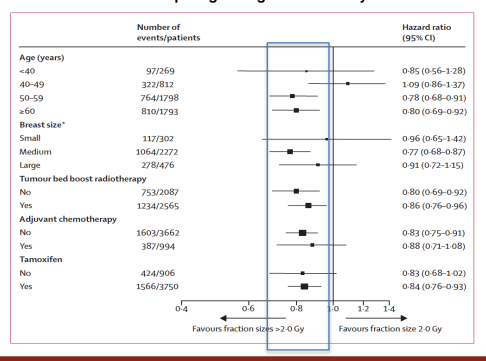
Moderate/marked late NTE from START A and B



Cumulative incidence of Grade 2-3 induration



Meta-analysis any moderate/marked physician-assessed NTE in breast comparing HF regimens *vs* 50 Gy/25



Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

Hypofractionation for early breast cancer

Ultra Hypofractionation (5-fraction)

FAST-Forward trial (n=4096) showed that ultra-hypofractionation (**26Gy in 5 fractions**) leads to **non-inferior local control** rates and **similar adverse event profile** as compared to **40Gy in 15 fractions over 3 weeks**

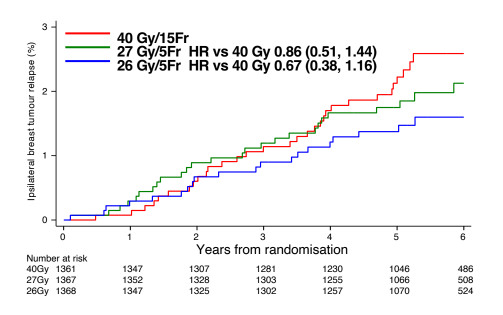
Median follow up 6 years

- → 5-fraction regimen non-inferior in terms of LR as compared to 40 Gy in 15 fractions (HR 0.67, 95%CI 0.38 to 1.16)
- → 5-year estimate 1.4% (26 Gy) *vs* 2.1% (40 Gy)
- → Late NTE as assessed by clinicians, patients, and photos similar for 26 Gy (HR 1.12, 95%Cl 0.94 to 1.34; p=0.20)



Hypofractionation for early breast cancer

Ultra Hypofractionation (5-fraction)

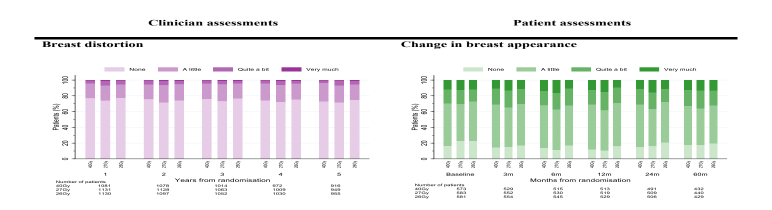


No. events		5yr estimate (95% CI)	Difference vs. 40 Gy (95% CI)	
40 Gy	31	2.1% (1.4, 3.1)	-	
27 Gy	27	1.7% (1.2, 2.6)	-0.3% (-1.0, <u>0.9</u>)	
26 Gy	21	1.4% (0.9, 2.2)	-0.7% (-1.3, <u>0.3</u>)	



Ultra Hypofractionation (5-fraction)

Clinician & patient assessments of adverse effects up to 5 years



At 5 years → any clinician-assessed moderate/marked AE: 10% in 40Gy vs 15% in 27Gy vs 12% in 26Gy

Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

Postmastectomy setting

Efficacy of hypofractionation

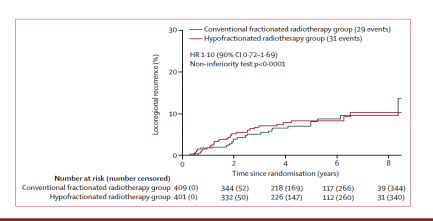
Hypofractionated versus conventional fractionated postmastectomy radiotherapy for patients with high-risk breast cancer: a randomised, non-inferiority, open-label, phase 3 trial



Shu-Lian Wang*, Hui Fang*, Yong-Wen Song, Wei-Hu Wang, Chen Hu, Yue-Ping Liu, Jing Jin, Xin-Fan Liu, Zi-Hao Yu, Hua Ren, Ning Li, Nina-Nina Lu, Yu Tana, Yuan Tana, Shu-Nan Qi, Guana-Yi Sun, Ran Pena, Shuai Li, Bo Chen, Yong Yana, Ye-Xiona Li

820 high-risk BC patients (2008-2016)

Hypofractionated PMRT (43.5Gy in 15 fractions) is as efficacious and safe as 50Gy in 25 fractions



Results

Median follow-up of 58.5 months

8.3% of 5-year cumulative incidence of LRR in the hypo-RT group *vs* **8.1%** in the CF-group (*p*<0.0001)



Postmastectomy setting

Safety of hypofractionation

No significant differences in acute and late toxicities

Grade 3 acute skin toxicity in **3% in the hypo-RT group** *vs* **8% in the CF-group** (*p*<0.0001)

Hypofractionated PMRT was non-inferior to CF-RT with similar toxicities in high-risk breast cancer

	Conventional fractionated radiotherapy group (n=409)	Hypofractionated radiotherapy group (n=401)	p value
Acute toxicity			
Skin toxicity		7···	<0.0001
Grade 1-2	357 (87%)	351 (89%)	
Grade 3	32 (8%)	14 (3%)	
Pneumonitis			0.278
Grade 1	62 (15%)	61 (15%)	
Grade 2	7 (2%)	14 (3%)	
Grade 3			
Late toxicity			
Skin toxicity			0.669
Grade 1-2	90 (22%)	86 (21%)	
Grade 3	0	1 (<1%)	
Lymphoedema			0.961
Grade 1-2	81 (20%)	78 (19%)	175
Grade 3	3 (1%)	3 (1%)	
Shoulder dysfunction			0.734
Grade 1-2	13 (3%)	7 (2%)	
Grade 3	1 (<1%)	1 (<1%)	
Lung fibrosis		2 · ·	0.081
Grade 1-2	42 (10%)	62 (15%)	
Grade 3	0	0	
Ischaemic heart disease			0.569
Grade 1-2	1 (<1%)	3 (1%)	
Grade 3	3 (1%)	4 (1%)	

RNI settingSafety of hypofractionation

Patient-assessed arm and shoulder effects according to +/- lymphatic RT

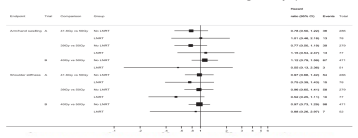


Table 3Physician-assessed moderate/marked normal tissue effects in the arm or shoulder following lymphatic radiotherapy in START-pilot, START-A and START-B.

Schedule	Total moderate/ marked events (n/total, %)	Estimated cumulative incidence by 5 years, % (95%CI)	Estimated cumulative incidence by 10 years, % (95%CI)	Hazard ratio (95% CI)	P- value ¹	Prevalence of moderate/ marked events at 5 years, n/total (%)	P- value ²	Prevalence of moderate/ marked events at 10 years, n/total (%)	P- value ²
Arm oedema									
START-B									
50 Gy	7/73 (9.6)	6.0 (2.3-15.3)	13.5 (6.4-27.0)	1		0/51 (0)		0/27 (0)	
40 Gy	3/81 (3.7)	2.8 (0.7-10.7)	4.7 (1.5-14.0)	0.42 (0.11-1.63)	0.21	2/57 (3.5)	0.50	0/20 (0)	-
Shoulder stiff	ness								
START-B									
50 Gy	4/73 (5.5)	2.9 (0.7-11.0)	8.2 (2.9-21.8)	1		1/51 (2.0)	>0.99	1/27 (3.7)	>0.99
40 Gy	3/81 (3.7)	3.1 (0.8-11.9)			0.72	1/57 (1.8)	>0.99	1/20 (5.0)	>0.99



Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

ESTRO-ACROP 2022 consensus statements

European Society for Radiotherapy and Oncology Advisory Committee in Radiation Oncology Practice consensus recommendations on patient selection and dose and fractionation for external beam radiotherapy in early breast

Marianne C Aznar Claus Bellin Adrian Murray Brunt Samantha Dicuanza Pierfrancesca Franco Mechthild Krause Maisead MacKenzi



Radiotherapy & Oncology

Whole breast irradiation

Moderate hypofractionated WBI should be offered regardless of:

- age at breast cancer diagnosis
- pathological tumour stage
- breast cancer biology
- surgical margins status
- tumour bed boost

- breast size
- invasive or pre-invasive DCIS disease
- oncoplastic breast conserving surgery
- use of systemic therapy

Ultra-hypofractionated WBI can be offered as:

- Standard of care
- Within a randomised controlled trial or prospective registration cohort

Nodal irradiation

- Moderate hypofractionation should be offered for RNI
- **Ultra-hypofractionation** should **not** be offered for RNI until ongoing trials results are reported



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Moderate hypofractionation can be offered:

- for chest wall irradiation without breast reconstruction
- for chest wall irradiation regardless of time and type of breast reconstruction

Ultra-hypofractionation for chest wall irradiation without breast reconstruction can be offered as:

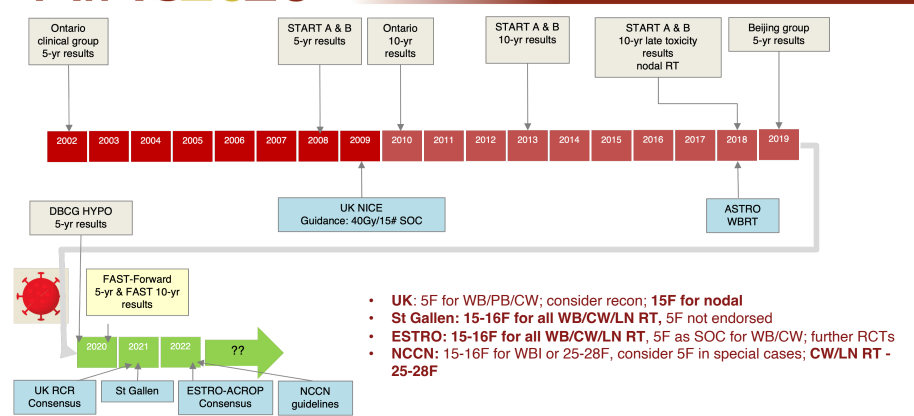
- Standard-of-care
- Within a randomised controlled trial or prospective registration cohort

Ultra-hypofractionation for chest wall irradiation after breast reconstruction can be offered within:

- A randomised controlled trial
- Prospective registration cohort



Radioterapia Oncologica: l'evoluzione al servizio dei pazienti





Hypofractionation

Partial Breast Irradiation

Omission of whole breast irradiation Oligometastatic disease Integration with systemic therapies

Primary endpoint: Ipsilateral breast tumour relapse

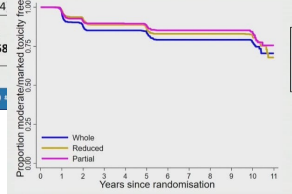
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	Whole	Reduced	Partial
	N=674	N=673	N=669
Number of IBTR events	17	11	17
KM 10 year cumulative	2.8%	1.9%	2.8%
IBTR estimate (95% CI)	(1.8, 4.5)	(1.1, 3.4)	(1.7, 4.5)
Hazard ratio		0.63	0.99
(95% CI)		(0.30, 1.35)	(0.51, 1.94
Absolute difference in IBTR rate compared with control group at 10 years (95% CI)		-1.02% (-1.97, 0.97)	-0.02% (-1.38, 2.58





Partial-breast radiotherapy after breast conservation surgery for patients with early breast cancer (UK IMPORT LOW trial): 5-year results from a multicentre, randomised, controlled, phase 3, non-inferiority trial

Time to any* moderate/marked clinician assessed breast NTE



10 year event-free estimates:

Whole: 70.5% (95%CI 64.1, 75.9)

Reduced: 75.3% (95%CI 69.6, 80.0) Partial: 75.6% (95%CI 69.1, 80.9)

NB. Clinician assessments of NTEs were conducted at years 1, 2, 5 and

then at 10 years



X f #ASTRO23

any=breast shrinkage, breast induration (index), breast induration (outside index), telangiectasia, breast oedema, other RT related adverse event

Partial breast irradiation

Trial design – APBI IMRT Florence (NCT 02104895)

APBI using IMRT 30 Gy in 5 Phase III trial non-consecutive (n=520 patients) Breast conserving 1:1 randomization surgery pT <25 mm Final surgical CF-WBI margins ≥5 mm 50 Gy in 25 Age >40 years 10 Gy in 5 boost

Primary endpoint

• IBTR

Follow Up

Secondary endpoints

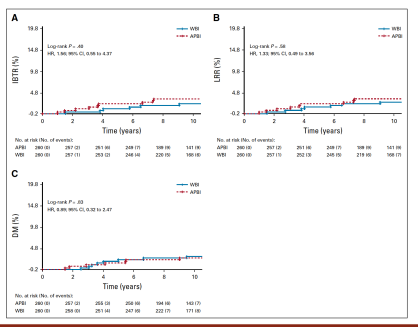
- OS and BCSS
- Contralateral breast cancer (CBC)
- Early and late toxicity
- Physician-rated cosmesis

Partial breast irradiation

Long term follow-up - APBI IMRT Florence (NCT 02104895)

Accelerated Partial-Breast Irradiation Compared With Whole-Breast Irradiation for Early Breast Cancer: Long-Term Results of the Randomized Phase III APBI-IMRT-Florence Trial

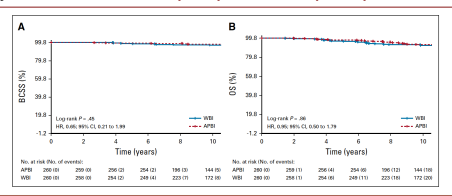
Icro Meattini, MD¹⁻²; Livia Marrazzo, MS²; Calogero Saieva, MD¹; Isacco Desideri, MD¹⁻²; Vieri Scotti, MD²; Gabriele Simontacchi, MD Pierbiugi Bomono, MD¹ (Barco Male Maria Mangoni, MD, PhD¹⁻²; Isivia Secciocatini, MD²; Sara Lucidi, MD¹; Lisa Paeletti, MD Massimillario Fambrini, MD¹, Marco Bernini, MD, PhD²; Liui Sanchex, MD²; Lorenzo Ozalesti, MD²; Lacopo, hori, MD²;





10-year IBTR 2.5% (WBI) vs 3.7% (APBI) (HR 1.56; P = 0.40)

10-year BCSS 96.7% (WBI) vs 97.8% (APBI) (HR 0.65; P = 0.45)



Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

Partial breast irradiation

Long term follow-up – APBI IMRT Florence (NCT 02104895)

Assessment	APBI (n = 246)	WBI (n = 260)	P
Acute period adverse events ^a			
None	194 (78.9)	87 (33.5)	.0001
Yes, any grade	52 (21.1)	173 (66.5)	_
Grade 1	47 (19.1)	75 (28.8)	.0001
Grade 2	5 (2.0)	81 (31.2)	_
Grade 3	_	17 (6.5)	_
Grade 4	_	_	_
Grade 0-1	241 (98.0)	162 (62.3)	.0001
Grade ≥ 2	5 (2.0)	98 (37.7)	.0001
Late period adverse events ^a			
None	235 (95.5)	182 (70.0)	.0001
Yes, any grade	11 (4.5)	78 (30.0)	.0001
Grade 1	11 (4.5)	71 (27.3)	.0001
Grade 2	_	7 (2.7)	
Grade 3	_		
Grade 4	_	_	_
Grade 0-1	246 (100)	253 (97.3)	.015
Grade ≥ 2	0	7 (2.7)	
Physician-rated cosmesis ^b			
Excellent	233 (94.7)	189 (72.7)	.0001
Good	13 (5.3)	66 (25.4)	_
Fair	_	5 (1.9)	_
Poor	_	_	_
Patient-rated cosmesis ^b			
Excellent	44 (17.9)	13 (5.1)	.0001
Good	200 (81.3)	209 (80.3)	
Fair	2 (0.8)	38 (14.6)	
Poor	_	_	

Accelerated Partial-Breast Irradiation Compared With Whole-Breast Irradiation for Early Breast Cancer: Long-Term Results of the Randomized Phase III APBI-IMRT-Florence Trial

Icro Meattini, MD^{1,2}; Livia Marrazzo, MS²; Calogero Saieva, MD³; Isacco Desideri, MD^{1,2}; Vieri Scotti, MD²; Gabriele Simontacchi, MD²; Pierluigi Bonomo, MD²; Daniela Greto, MD²; Monica Mangoni, MD, PhD^{1,2}; Silvia Scoccianti, MD²; Sara Lucidi, MD¹; Lisa Paoletti, MD⁴; Massimiliano Fambrini, MD^{1,2}; Marco Bemini, MD, PhD²; Luis Sanchez, MD²; Lorenzo Orzalesi, MD^{1,2}; Jacopo Nori, MD²; Simonetta Bianchi, MD^{1,2}; Stefania Pallotta. MS^{1,2}; and Lorenzo Livi. MD^{1,2}

APBI significantly favoured:

- acute and late adverse events
- both physician- and patient-rated cosmesis



Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

ESTRO-ACROP 2022 consensus statements

Partial breast irradiation

European Society for Radiotherapy and Oncology Advisory Committee in Radiation Oncology Practice consensus recommendations on patient selection and dose and fractionation for external beam radiotherapy in early breast cancer



European SocieTy for Radiotherapy & Oncology

Icro Meattini, Carlotta Becherini, Liesbeth Boersma, Orit Kaidar-Person, Gustavo Nader Marta, Angel Montero, Birgitte Vrou Offersen,
Marianne C Aznar, Claus Belka, Adrian Murray Brunt, Samantha Dicuorzo, Pierfrancesco Franco, Mechthild Krause, Mariead MacKenzie,
Tanja Marinko, Livia Marrazzo, Unica Ratosa, Astrid Schotlen, Elbizite Sentan, Shilay Sotabar, Philip Poertamas", Charlotte E Coles*

Low risk-features suitable for partial breast irradiation:

- luminal-like subtypes small tumour (≤3 cm)
- absence of lymph vascular space invasion
- non-lobular invasive carcinoma
- tumour grade 1-2
- low to intermediate grade DCIS (sized ≤2.5 cm with clear surgical margins ≥3 mm)
- age at diagnosis 50 years or more
- unicentric/unifocal lesion
- clear surgical margins (>2 mm)
- node negative (including isolated tumour cells)
- no use of primary systemic therapy/neoadjuvant chemotherapy





Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

ESTRO-ACROP 2022 consensus statements

Partial breast irradiation

European Society for Radiotherapy and Oncology Advisory Committee in Radiation Oncology Practice consensus recommendations on patient selection and dose and fractionation for external beam radiotherapy in early breast cancer



EUTRO European SocieTy for Radiotherapy & Oncology

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Partial breast irradiation-dose and fractionation:

- Moderate hypofractionation (40Gy in 15 fractions) and ultra hypofractionation (26–30Gy in 5 fractions) represent acceptable schedules for external beam partial breast irradiation
- Twice a day external beam partial breast irradiation dose and fractionations similar to those used in the RAPID trial should not be offered



Hypofractionation
Partial Breast Irradiation

Omission of whole breast irradiation

Oligometastatic disease Integration with systemic therapies

Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

The NEW ENGLAND JOURNAL of MEDICINE

ORIGINAL ARTICLE

Omitting Radiotherapy after Breast-Conserving Surgery in Luminal A Breast Cancer

T.J. Whelan, S. Smith, S. Parpia, A.W. Fyles, A. Bane, F.-F. Liu, E. Rakovitch, L. Chang, C. Stevens, J. Bowen, S. Provencher, V. Théberge, A.M. Mulligan, Z. Kos, M.A. Akra, K.D. Voduc, T. Hijal, I.S. Dayes, G. Pond, J.R. Wright, T.O. Nielsen, and M.N. Levine, for the LUMINA Study Investigators*

Whole breast irradiation (WBI) omission LUMINA trial

Prospective single-arm cohort study at 26 centres in Canada:

- Recruitment of **500 patients** from 2013-2017

Inclusion criteria

- Age ≥ 55 years
- Invasive breast carcinoma (NST, tubular, or mucinous)
- pT1a-c pN0 with a minimum of ASR 1 mm
- ER ≥ 1% PR ≥ 20% HER2 negative Ki67 ≤ 13.25%
- Planned ET for 5 years
- Omission of adjuvant RT

Statistical hypothesis

LR rate in the operated breast after 5 years <5%

Characteristic	All Patients (N=500)
Age	
Median (IQR) — yr	67.1 (62.9–71.6)
Distribution — no. (%)	
55 to <60 yr	61 (12)
60 to <65 yr	138 (28)
65 to <70 yr	136 (27)
70 to <75 yr	107 (21)
75 to <80 yr	42 (8)
≥80 yr	16 (3)
Tumor size	
Median (IQR) — cm	1.0 (0.7-1.4)
Distribution — no. (%)	
≤0.5 cm	39 (8)
0.5–1.0 cm	217 (43)
1.1–2.0 cm	244 (49)
Tumor grade — no. (%)	
1	330 (66)
2	170 (34)
Histologic cancer type — no. (%)
Ductal	437 (87)
Tubular	25 (5)
Mucinous	26 (5)
Other	12 (2)

Whole breast irradiation (WBI) omission

LUMINA trial

- LR rate after 5 years was **2.3%** (95%CI 1.2-4.1)
- Only **marginally lower** than the rate of contralateral second carcinomas (2.5% after 5 years)
- DFS and OS at 5 years were 89.9% and 97.2%, respectively

Authors' conclusion: The prospective and controlled nature of this study supports our conclusion that such patients are candidates for omission of radiotherapy



Whole breast irradiation (WBI) omission

Debunking

Key Details

- The **inclusion criteria for the study were quite broad**. The minimum age of 55 years was significantly lower than in most other studies of this kind; 40% <65 years old → particularly relevant due to **life expectancy** considerations
- The Ki67 value was determined centrally, and this was done for a good reason, as inter-rater reliability can be problematic.
 Indeed, 224 patients (30% of registered patients) were excluded in the screening after central testing due to their high Ki67 levels
- The **Ki67 cut-off of 13.25% is arbitrary** a large grey area between 5-30% was defined, where the use of gene expression analyses is recommended for luminal tumours
- Compliance with ET was significantly higher (82.7%) than expected outside of clinical studies
- The 5-year results of the LR rate **overlap with the CI** of comparable studies (i.e., PRIME II)



Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

Whole breast irradiation (WBI) omission

The wrong answer at the right question!

PRIME II study 10-year LR rates:

- All 9.5% (no RT) vs 0.9% (RT)

19.1% (no RT) vs 0.0% (RT) - ER-low

ER high was defined here as:

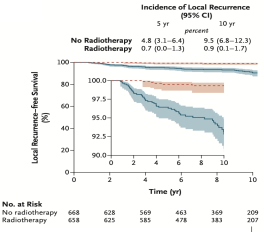
- ER ≥50% (!)
- Allred Score 7-8, or
- ER ≥20 fmol/mg (an obsolete method)

The NEW ENGLAND JOURNAL of MEDICINE

FEBRUARY 16, 2023

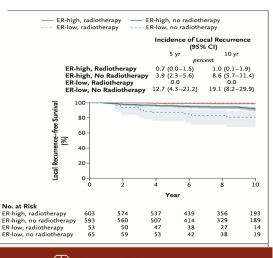
Breast-Conserving Surgery with or without Irradiation in Early Breast Cancer

Ian H. Kunkler, M.B., B.Chir., Linda J. Williams, Ph.D., Wilma J.L. Jack, M.B., Ch.B., David A. Cameron, M.D., and I. Michael Dixon, M.D.



currence-free Survival

No radiotherapy Radiotherapy





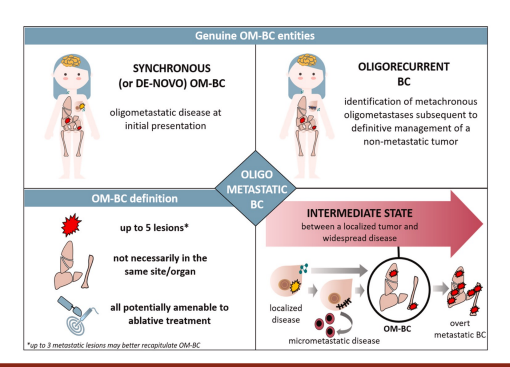
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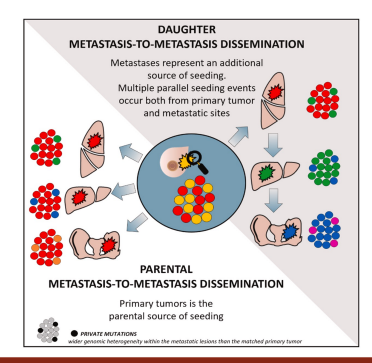
Oligometastatic disease

Integration with systemic therapies

Oligometastases

Definitions and Concepts





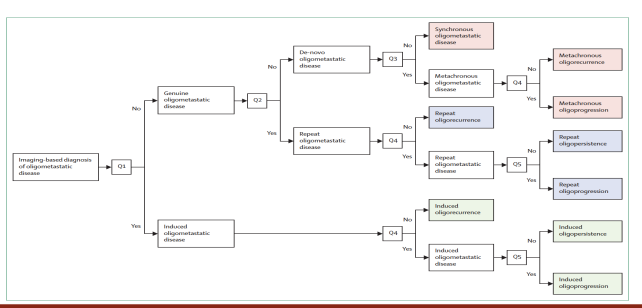
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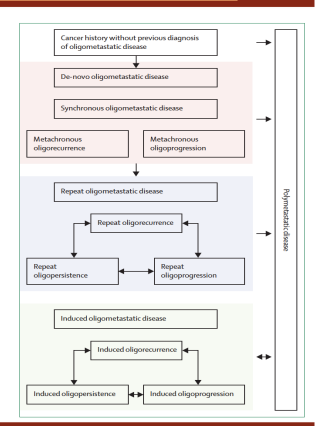
Oligometastases

Characterization and Classification

Characterisation and classification of oligometastatic disease: a European Society for Radiotherapy and Oncology and European Organisation for Research and Treatment of Cancer consensus recommendation

Matthias Guckenberger, Yalande Lievens, Angelique B Bouma, Laurence Collette, Andre Dekker, Nandita M deSouza, Anne-Marie C Dingemans, Beatrice Fournier, Coen Hurkmans, Frédric Lecowet, Icro Meattini, Alejandra Méndez Romero, Umberto Ricardi, Nicola S Russell, Daniel H Schanne, Marta Scorsetti, Bertrand Tombol, Dirk Verellen, Christine Verfaillie, Piet Ost





Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

Oligometastases

Use and evaluation of imaging methods in clinical trials

Review of the literature covering all aspects of oligometastatic breast cancer

35 experts from the EORTC Imaging and Breast Cancer Groups

Consensus recommendations:

- Oligometastatic breast cancer definition
- Optimal diagnostic pathways
- Clinical trials required to evaluate the effect of diagnostic imaging strategies and metastasis-directed therapies
- Strategies for the randomisation of imaging methods and therapeutic approaches in different subsets of patients

Designing clinical trials based on modern imaging and metastasis-directed treatments in patients with oligometastatic breast cancer: a consensus recommendation from the EORTC Imaging and Breast Cancer Groups



David Pasquier, Luc Bidaut, Daniela Elena Oprea-Lager, Nandita M deSouza, David Krug, Laurence Collette, Wolfgang Kunz, Yazid Belkacemi, Maria Grazia Bau, Caroline Caramella, Lioe-Fee De Geus-Oei, Alex De Caluwé, Christophe Deroose, Olivier Gheysens, Ken Herrmann, Isabelle Kindts, Michalis Kontos, Sherko Kümmel, Barbro Linderholm, Egesta Lopci, Icro Meattini, Ann Smeets, Orit Kaidar-Person, Philip Poortmans, Pelagia Tsoutsou, Nawale Hajjaji, Nicola Russell, Elżbieta Senkus, Jean-Noël Talbot, Lale Umutlu, Vincent Vandecaveye, Joost J C Verhoeff, Willemien Menke-van der Houven van Oordt, Helle D Zacho, Fatima Cardoso, Laure Fournier, Fredericke Van Dujinhoven, Frédéric E Lecouvet

Imaging methods in clinical trials	Consensus and round
Use of imaging in trials	
["FFFDG-PET-CT (or ["F]FDG-PET-MRI) staging should be mandatory in trials enrolling oligometastatic disease breast cancer patients to ensure the true oligometastatic disease status	Consensus agreement=79% in round 1; absolute number=27; total number of responses=35; non-qualified=1
Whole-body MRI and diffusion-weighted imaging staging should be mandatory in trials enrolling patients with oligometastatic disease breast cancer to ensure the true oligometastatic disease status	Neither consensus nor (dis)agreement
Necessary evaluation of imaging in trials	
Prospective trials are needed to compare SIMs and MIMs for staging and response assessment in advanced breast cancer, including oligometastatic disease	Consensus agreement=86% in round 1; absolute number=30; total number of responses=35
Clinical trials aiming to compare SIMs and MIMs for staging and response assessment in advanced breast cancer should be designed in specific histological and breast cancer subtypes (eg., lobular cancer and triple negative)	Consensus agreement=87% in round 2 (74% in round 1); absolu number=28; total number of responses=33; non-qualified=1
The diagnostic performance of different MIMs (eg., PET-CT or PET-MRI, whole-body MRI, liver MRI, and ["F]NAF plus ["F]FDG-PET cocktail) deserves further comparisons in trials	Consensus agreement=89% in round 1; absolute number=31; total number of responses=35
The diagnostic performance of MIMs (eg. PET-CT or PET-MRI, whole-body-MRI, liver MRI, or brain MRI) should be compared in the different subtypes of breast cancer (eg. ductal, lobular, HR, and HER)	Consensus agreement=94% in round 1; absolute number=33; total number of responses=35
Diagnostic trials should further validate quantification with MIMs (ie, second order statistics) for turnour characterisation and prognostic purposes (in whole-body diffusion-weighted MRI and PET-CT (or PET-MRI))	Consensus agreement=91% in round 2 (74% in round 1); absolu number=29; total number of responses=33; non-qualified=1
Diagnostic trials should compare technical and diagnostic performance and robustness of MRI and diffusion-weighted imaging sequences from hybrid PET-MRI modallities and from stand-alone MRI, with the purpose of optimising and standardising technical and diagnostic performance across various instruments	Consensus agreement=90% in round 1; absolute number=28; total number of responses=35; non-qualified=4
HER2 PET-CT imaging is still experimental and is not recommended outside of clinical trials	Consensus agreement=78% in round 1; absolute number=25; total number of responses=35; non-qualified=3

Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

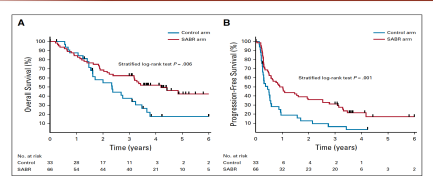
Oligometastases – Recent prospective studies SABR-COMET 51-month follow-up Update

palliative standard of care treatments alone

(control group)

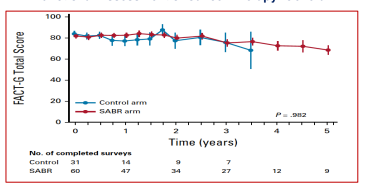
standard of care plus SBRT to all metastatic lesions

(SBRT group)



Kaplan-Meier plots for (A) OS and (B) PFS

Functional Assessment of Cancer Therapy: General



5-year OS rate **17.7%** (control) **vs 42.3%** (SABR)

5-year PFS rate not reached 3.2% (control) vs 17.3% (SABR)

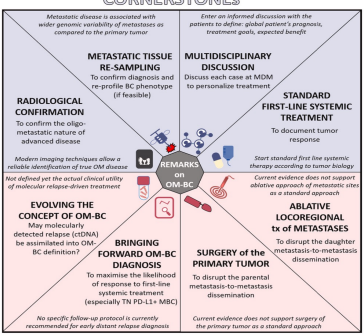
No new grade 2-5 adverse events and no differences in HRQoL between arms

Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

Oligometastases

Future directions

CORNERSTONES



RESEARCH AGENDA



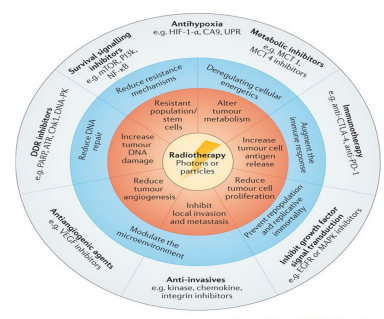
Hypofractionation
Partial Breast Irradiation
Omission of whole breast irradiation
Oligometastatic disease

Integration with systemic therapies

Integration of radiation therapy with targeted treatments for breast cancer

Treatment effectiveness Treatment safety





Nature Reviews | Clinical Oncology

Radioterapia Oncologica: 'evoluzione al servizio dei pazienti

Radiation and New Drugs

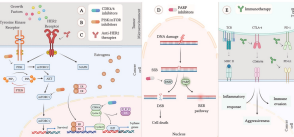
When is it concomitant?

RT is commonly considered given concurrent with systemic therapy when administered in a range < than 5 half-lives of the drug

Drug	Median Half-life	5 half-lives
Olaparib	15 hours	75 hours (≈ 3 days)
Lapatinib	24 hours	120 hours (≈ 5 days)
Abemaciclib	24.8 hours	124 hours (≈ 5 days)
Palbociclib	28.8 hours	144 hours (≈ 6 days)
Everolimus	30 hours	150 hours (≈ 6 days)
Ribociclib	29.7 – 54.7 hours	148.5 – 273.5 hours (≈ 6 - 11 days)
Talazoparib	90 hours	450 hours (≈ 19 days)
Trastuzumab-emtansine	96 hours	480 hours (≈ 20 days)
Trastuzumab-deruxtecan	168 hours	840 hours (≈ 35 days)
Trastuzumab	456 hours	2280 hours (≈ 95 days)
Bevacizumab	480 hours	2400 hours (≈ 100 days)
Nivolumab	578 hours	2890 hours (≈ 121 days)
Atezolizumab	648 hours	3240 hours (≈ 135 days)

Radioterapia Oncologica: 'evoluzione al servizio dei pazienti

Preclinical and clinical findings



					voorolooox	Nucleus
Family drug	Drug	Preclinical Effectiveness	Clinical Toxic effect	LoE	Recommendation concomitant treatment	Drug 5-half-lives, days*
CDK4/6i	Palbociclib	Increased	Increased	4	Cautionary	5.8
	Ribociclib	Increased	Increased		Cautionary	6.7
	Abemaciclib	Increased	Increased		Cautionary	5
PI3Ki	Alpelisib	Increased	Uncertain	4	Unsuitable	1.9
mTORi	Everolimus	Increased	Increased		Unsuitable	6.2
Anti-HER	Trastuzumab	Increased	Safe	3	Suitable	175
	Pertuzumab	Increased	Safe		Suitable	90
	Lapatinib	Increased	Safe		Suitable	5
	T-DM1	Uncertain	Uncertain/Safe		Cautionary	20
PARPi	Olaparib	Increased	Increased	4	Unsuitable	3.1
	Talazoparib	Increased	Increased		Unsuitable	18.7
Immunotherapy	Atezolizumab	Uncertain	Safe	4	Suitable	135
	Pembrolizumab	Uncertain	Safe		Suitable	110

Abbreviations. CDK4/6i, cyclin-dependent kinase 4 and 6 inhibitors; PI3Ki, phosphatidyl-inositol-3-kinase inhibitors; mTORi, mammalian target of rapamycin inhibitors; HER, human epidermal growth factor receptor; PARPi, poly(ADP-ribose) polymerases inhibitors; LoE, level of evidence.

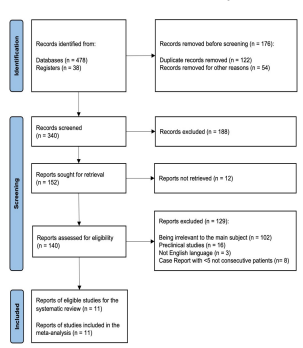
Level of Evidence followed the OCEBM Levels of Evidence Working Group. "The Oxford 2011 Levels of Evidence". Oxford Centre for Evidence-Based Medicine. https://www.cebm.net/wp-content/uploads/2014/06/CEBM-Levels-of-Evidence-2.1.pdf.

^{*} Radiation therapy is defined concomitant if a drug is administered within its 5-half-lives.

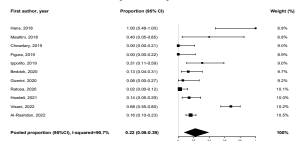
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CDK4/6 inhibitors and RT

Systematic Review and Meta-analyses

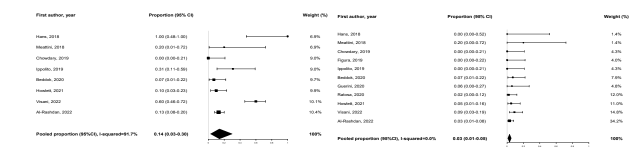


Any toxicity G3+



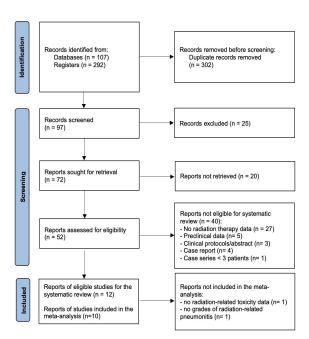
Haematological toxicity G3+

Non-haematological toxicity G3+

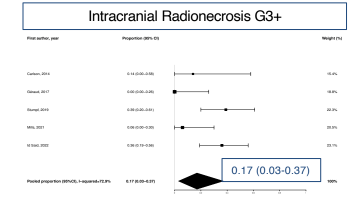


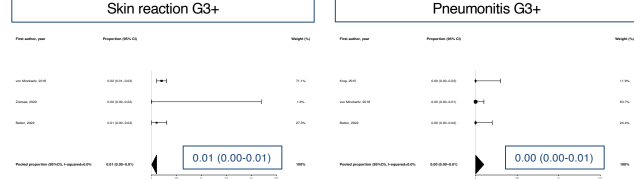
T-DM1 and RT

Systematic Review and Meta-analyses



Radioterapia Oncologica: l'evoluzione al servizio dei pazienti





Radioterapia Oncologica: l'evoluzione al servizio dei pazienti

Recommendations on integration of radiation therapy with targeted treatments for breast cancer consensus

Florence, Italy – June 16-17 #FlorenceBreast23



Recommendations on integration of radiation therapy with targeted treatments for breast cancer consensus meeting

Florence (IT), 16-17th June 2023

Grand Hotel Mediterraneo, Lungarno del Tempio, 44





Endorsed by







Radioterapia Oncologica: l'evoluzione al servizio dei pazient

Acknowledgments

Charlotte Elizabeth Coles Anna Kirby Philip Poortmans Orit Kaidar Person Birgitte Offersen Liesbeth Boersma

Livia Marrazzo

Marianne Camille Aznar

Viola Salvestrini

Luca Visani

Carlotta Becherini

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