

***Invasive procedures in the diagnosis
and treatment of liver diseases:
focal lesions***



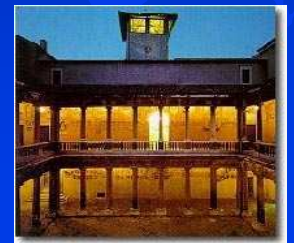
F. Farinati

Gastroenterologia, Padova

TACE: coming of age?



AISF 2005



TACE: LEVELS OF EVIDENCE

EBM

Degree of certainty

Methodology

A 1a

Metanalysis of homogenous RCTs

A 1b

Single RCT with accetable CI

A 1c

Metanalysis of homogenous cohort studies

B 2b

Cohort study or low quality RCT (f.u. <80% pts)

B 3a

Metanalysis with heterogeneity

B 3b

Case-control study

C 4

Observational or low quality case-control

D 5

Experts' opinion

TACE: coming of age?

- TACE IN INTERMEDIATE STAGE HCC***
- TACE IN THE NEO-ADJUVANT TREATMENT FOR OLTx***
- TACE COMBINED WITH LOCOREGIONAL TRANS-PARIETAL TREATMENTS (PEI-RFA)***

TACE: coming of age?

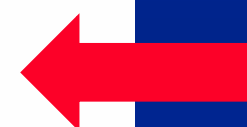
- TACE IN INTERMEDIATE STAGE HCC***
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Barcelona staging system(BCLC)

Table 2. Barcelona Clinic Liver Cancer Classification

Stage	Performance Status Test	Tumor Stage	Okuda Stage	Liver Function Status
A	0	Single	I-II	Child-Pugh A-B
B	0	Large multinodular	I-II	Child-Pugh A-B
C	1-2	Vascular invasion/ extrahepatic spread	I-II	Child-Pugh A-B
D	3-4	Any	III	Child-Pugh C

NOTE. Stage A and B, all criteria should be fulfilled; stage C and D, at least one criteria.

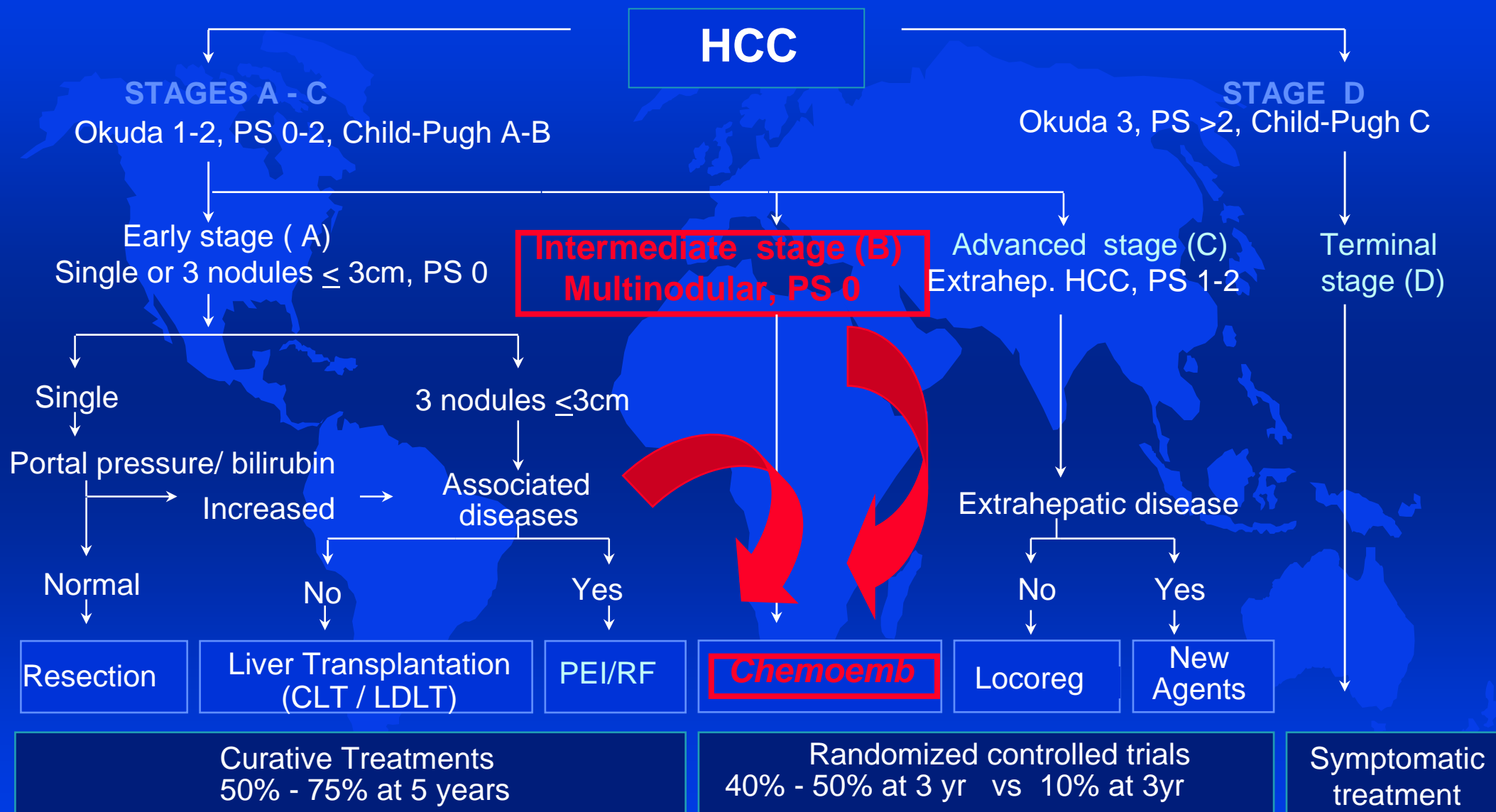


CLIP scoring system

<i>Variable</i>	<i>Score</i>		
	<i>0</i>	<i>1</i>	<i>2</i>
<i>Child-Pugh</i>	<i>A</i>	<i>B</i>	<i>C</i>
<i>Tumor morphology</i>	<i>Uninodular < 50%</i>	<i>Multinodular < 50%</i>	<i>Massive or > 50%</i>
<i>AFP (ng/dL)</i>	<i>< 400 ng/dL</i>	<i>> 400 ng/dL</i>	
<i>Portal thrombosis</i>	<i>No</i>	<i>Yes</i>	

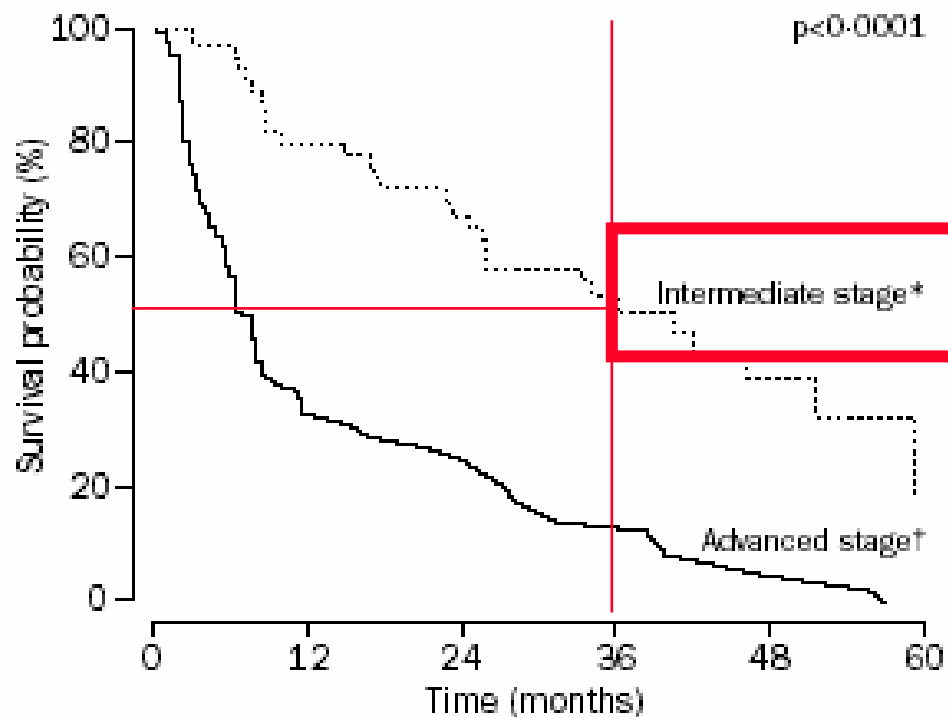
Score 0-2

BCLC Staging and Treatment Strategy



How long is untreated natural history in the intermediate stage?





Patients at risk	0	12	24	36	48	60
BCLC stage B	48	37	30	16	7	
BCLC stage C	54	17	9	2	–	

Figure 2: Survival, according to tumour stage, in untreated HCC BCLC=Barcelona-Clinic liver cancer. *Multinodular asymptomatic tumours, median survival 40 months. †Cancer-related symptoms, vascular invasion or extrahepatic spread, median survival 5.4 months. Reproduced from reference 56 with permission from The American Association for the Study of Liver Disease.

**Survival in
untreated HCC
patients**

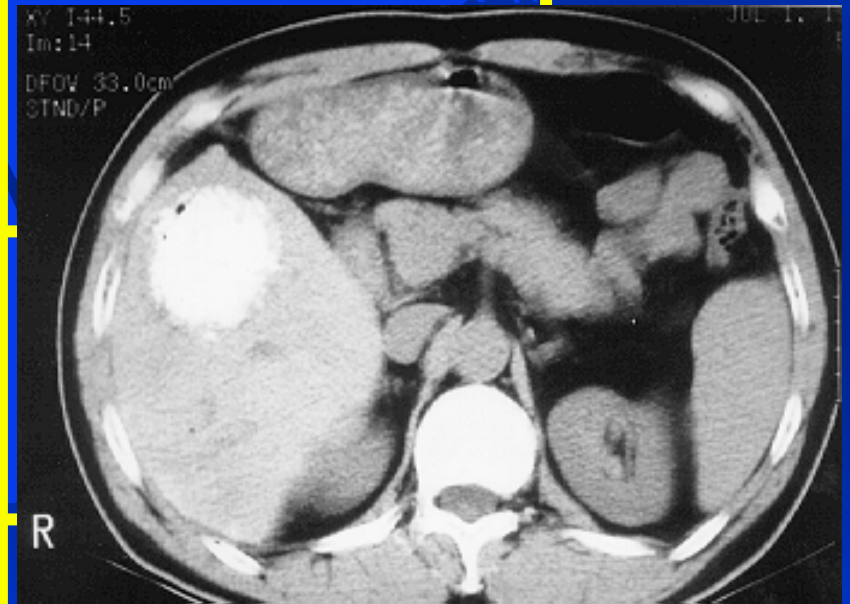
Llovet, Lancet 2003

Intermediate stage HCC ?

	<i>I year survival</i>	<i>II year survival</i>	<i>III year survival</i>
<i>BCLC I°</i>	<i>80%</i>	<i>65%</i>	<i>50%</i>
<i>BCLC II°</i>	<i>63%</i>	<i>27%</i>	<i>17%</i>
<i>ITALICA J.I.M.</i>	<i>47%</i>	<i>28%</i>	<i>14%</i>

HCC “advanced”: TACE

<i>EASL</i>	<i>AISF</i>
<i>Additional trials needed</i>	<i>Segmentary or sub-segmentary</i>
<i>Bruix, EASL 2001</i>	<i>Not adequate RCTs</i>



TACE: indications and contraindications

PRO:

- ◆ Single node / no surgery or PEI-RFA;
- ◆ Multiple nodes;
- ◆ Child A and B (?);
- ◆ Patent portal tract;
- ◆ No high grade varices.

CONTRA:

- ◆ Portal thrombosis/artero-portal fistula/blood flow inversion;
- ◆ vascular abnormalities;
- ◆ Child C;
- ◆ Severe discoagulopathy
- ◆ Renal failure
- ◆ Extra-hepatic metastases.

TACE: size of the problem

TREATMENT

SCREEN CHANCE

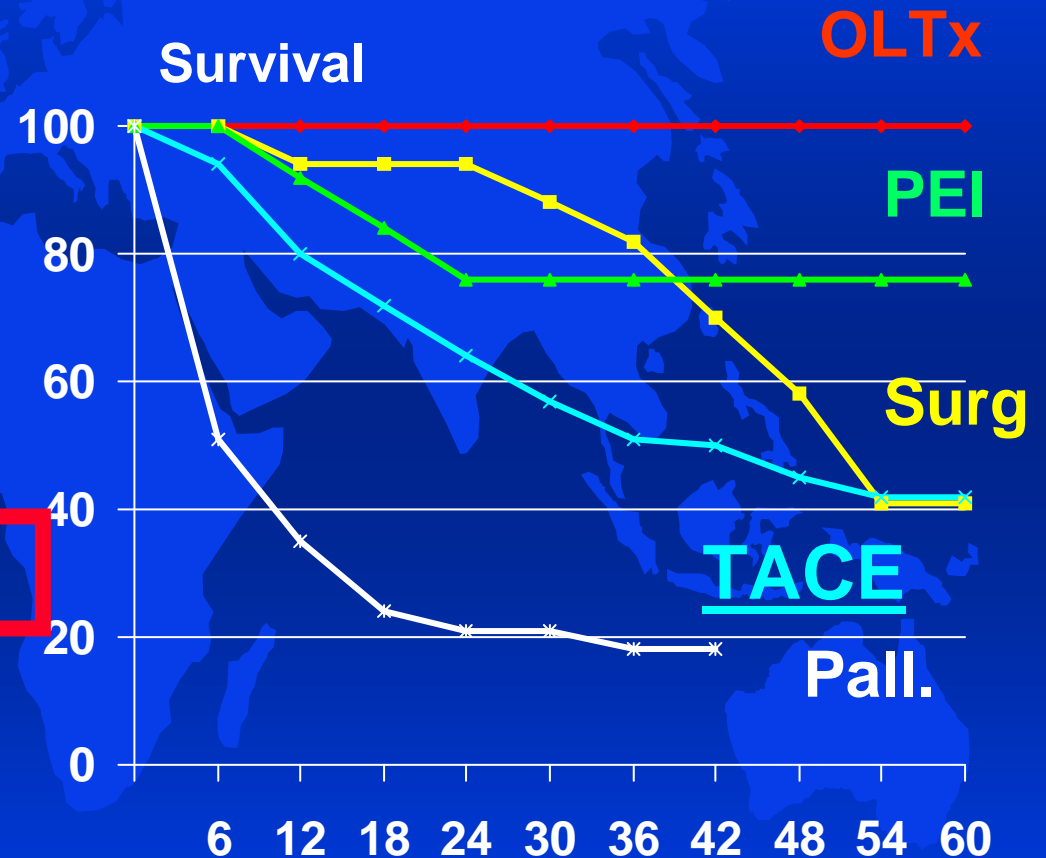
OTLx = 4 (5%) 0 (0%)

Surg = 14 (17%) 5 (6%)

PEI = 9 (11%) 4 (5%)

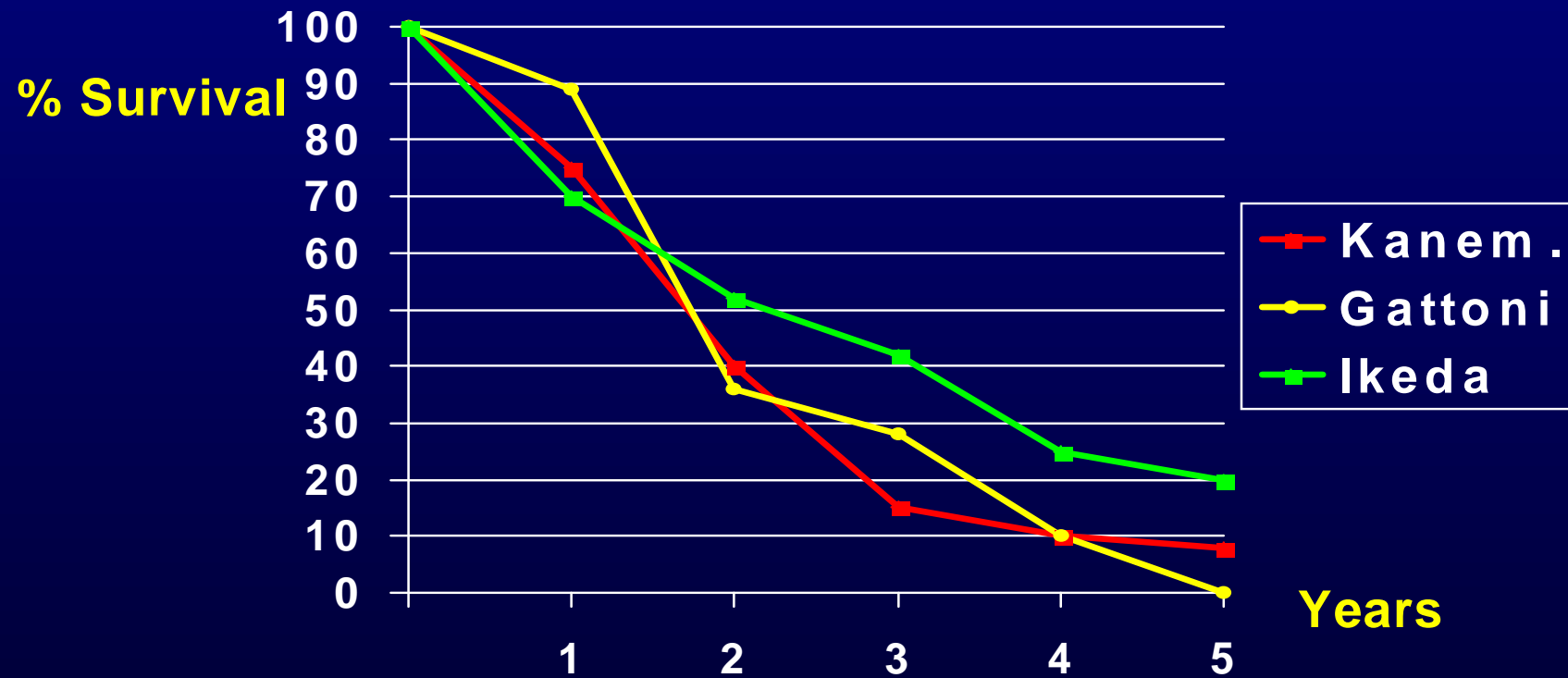
TACE = 48 (59%) 42 (51%)

Pall. = 6 (7%) 31 (38%)



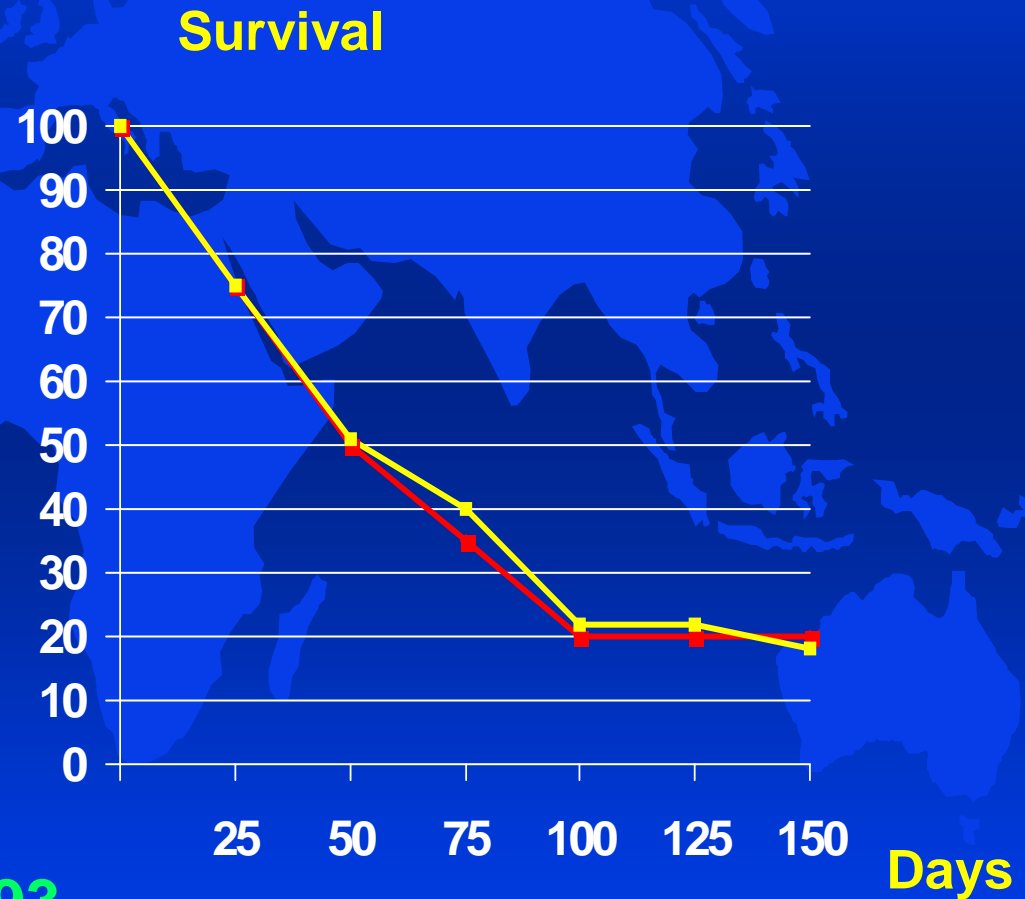
12% according to BRUIX, 2004

TACE: survival in open design studies



TACE: randomized studies

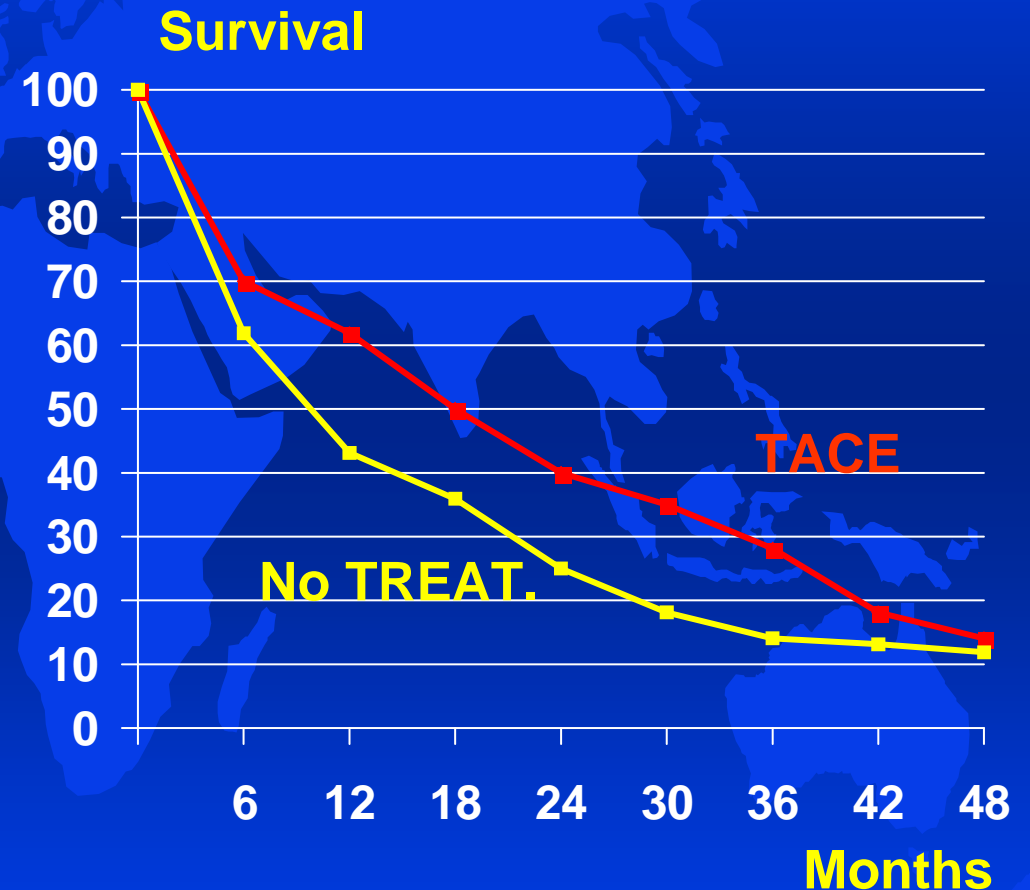
- ◆ No difference in survival between treated and untreated patients (short survival, no embolization, 75% Okuda II patients)



Madden, GUT 1993

TACE: randomized studies

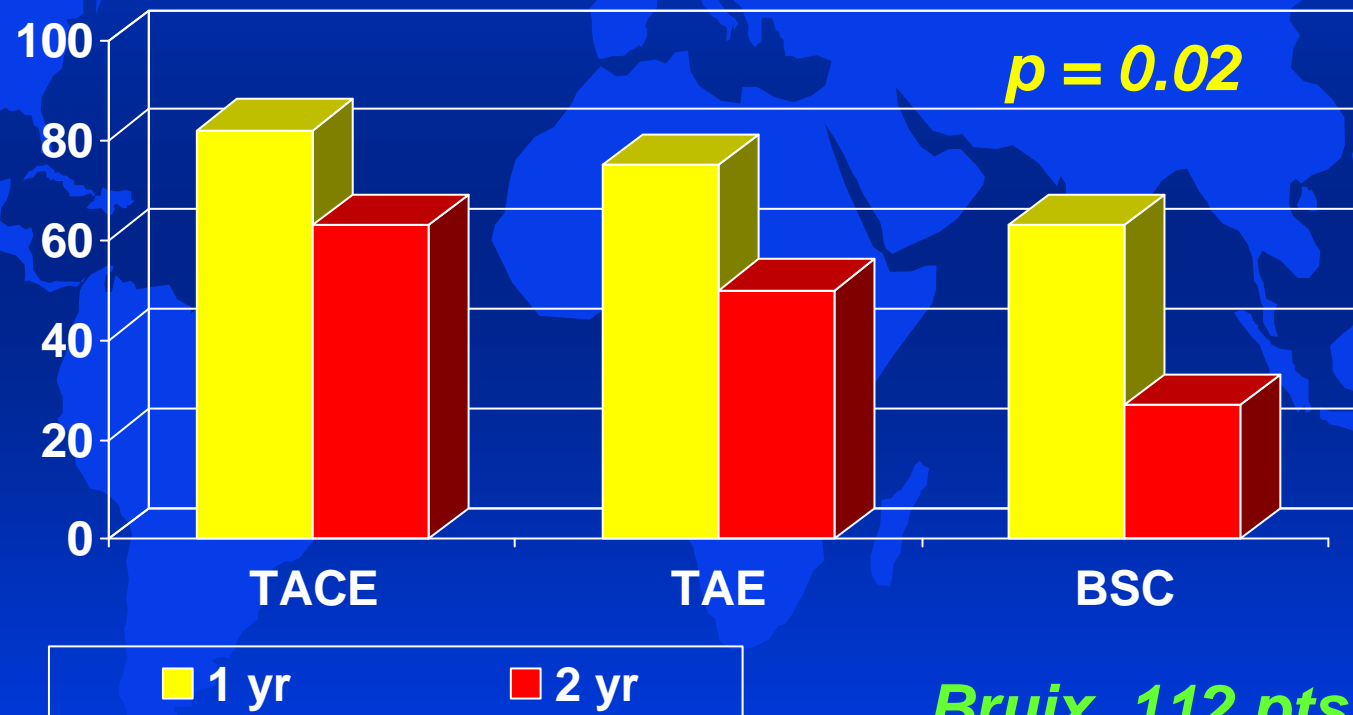
- ◆ 96 pts in 24 centers (4 pts/unit)
- ◆ 15 deaths in the first 2 months (Child A)
- ◆ 1 course/2months
- ◆ $p = 0.1$
- ◆ 53% of treated patients had tumor reduction, only 7% (versus 74%) had portal thrombosis.



Group Etude Trait. Carcinome Hepatocell., NEJM 1995

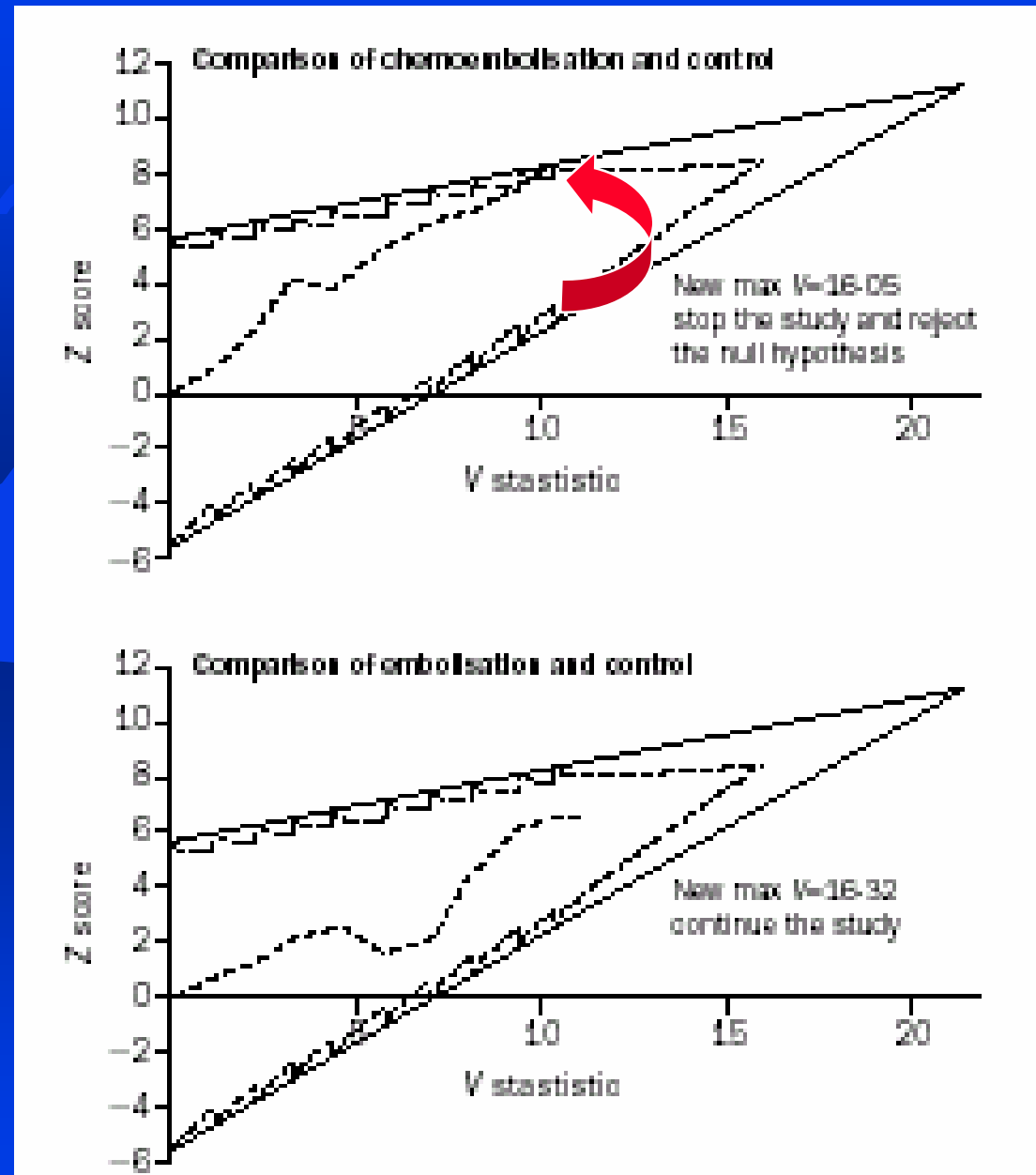
TACE: randomized studies

Survival



*Bruix, 112 pts,
Lancet, 2002*

Sequential design randomized trial



Evidence-based Practice

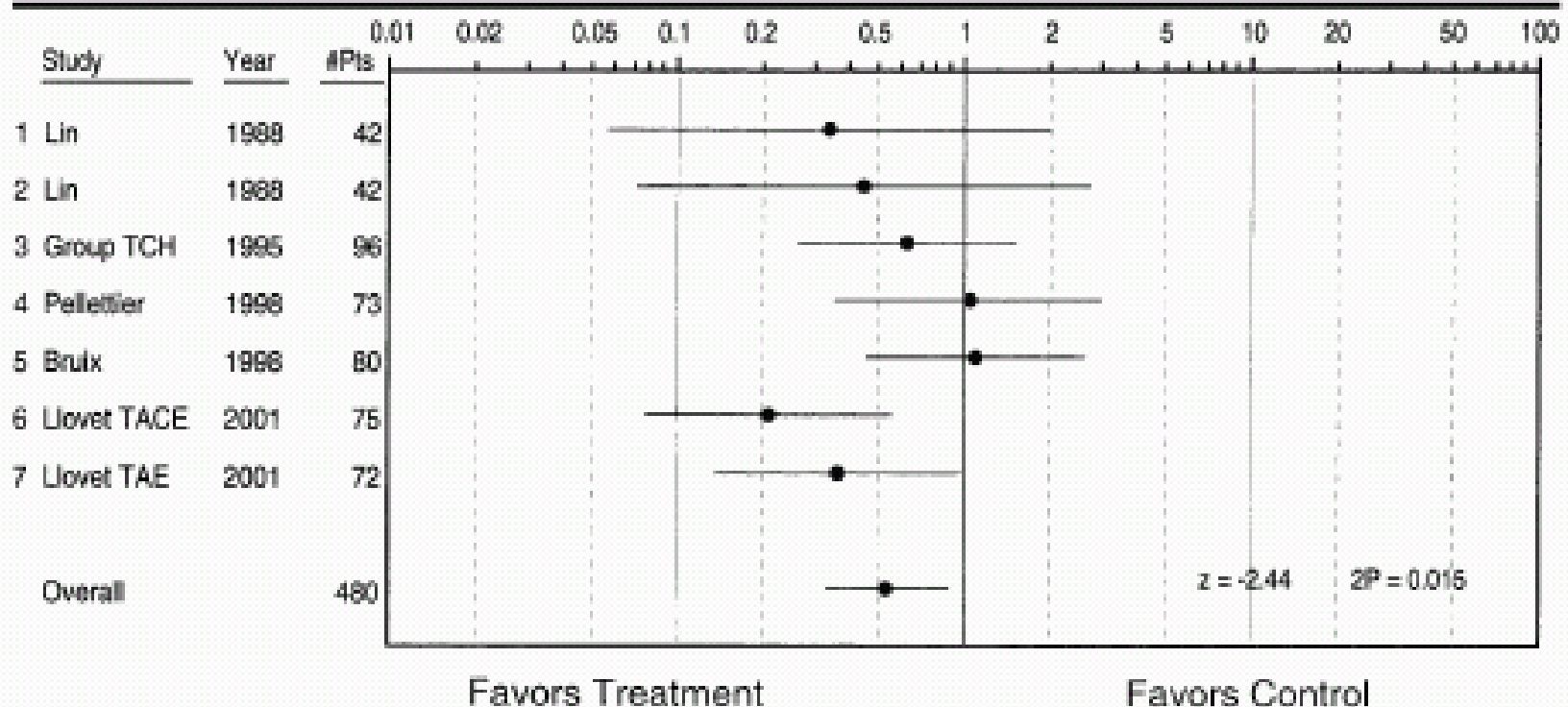
Radiology

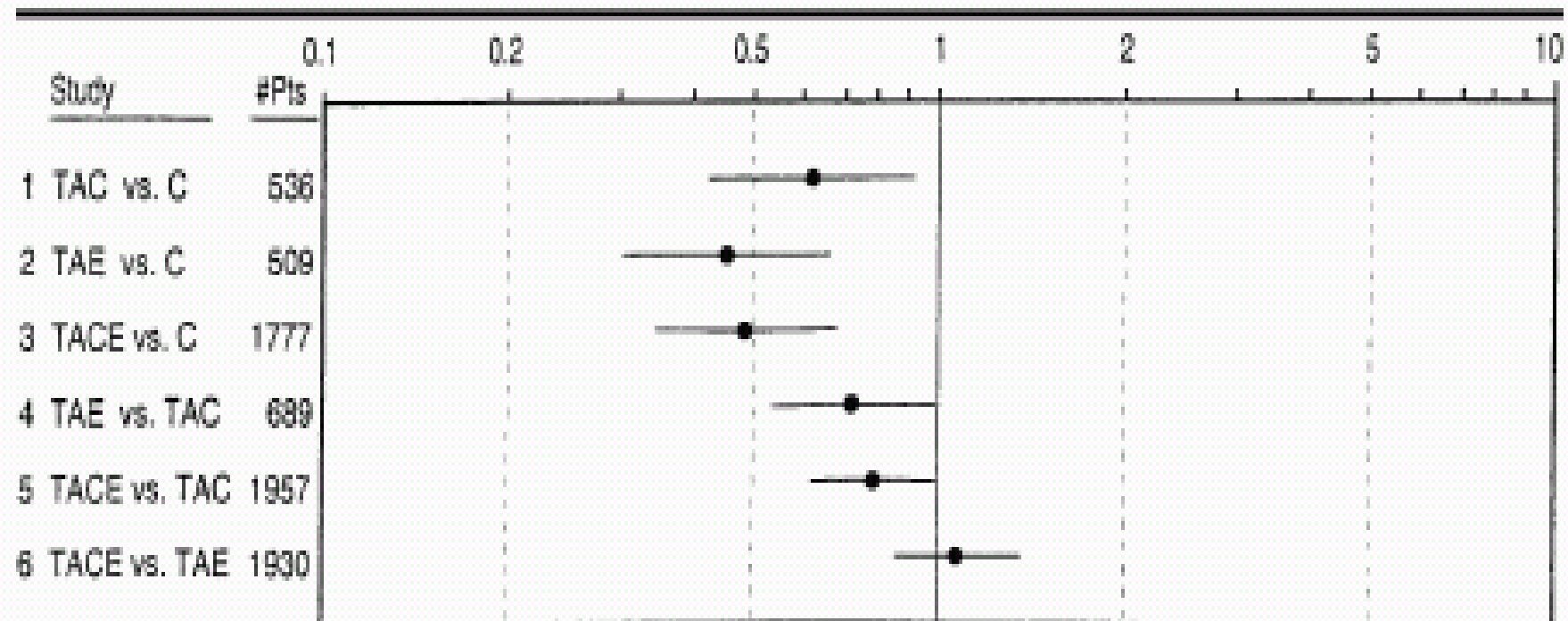
Calogero Cammà, MD
 Filippo Scephis, MD
 Ambrogio Orlando, MD
 Maddalena Albanese, MD
 Lillian Shahied, PhD
 Franco Trevisani, MD
 Pietro Andreone, MD
 Antonio Craxi, MD
 Mario Cottone, MD

Index terms:
 Efficacy study
 Liver neoplasms, 761.323
 Liver neoplasms, chemotherapeutic

Transarterial Chemoembolization for Unresectable Hepatocellular Carcinoma: Meta-Analysis of Randomized Controlled Trials¹

Radiology





The available evidence is sufficient to conclude that (a) chemoembolization significantly reduces overall 2-year mor-

tality in patients with unresectable HCC and (b) TACE was not more effective than TAE, which suggests that the addition of the chemotherapeutic agents currently used does not improve the benefit of therapy and emphasizes the need for more effective anticancer drugs. Future

Mean CR rate 6%

Systematic Review of Randomized Trials for Unresectable Hepatocellular Carcinoma: Chemoembolization Improves Survival

Josep M. Llovet and Jordi Bruix for the Barcelona-Clinic Liver Cancer Group

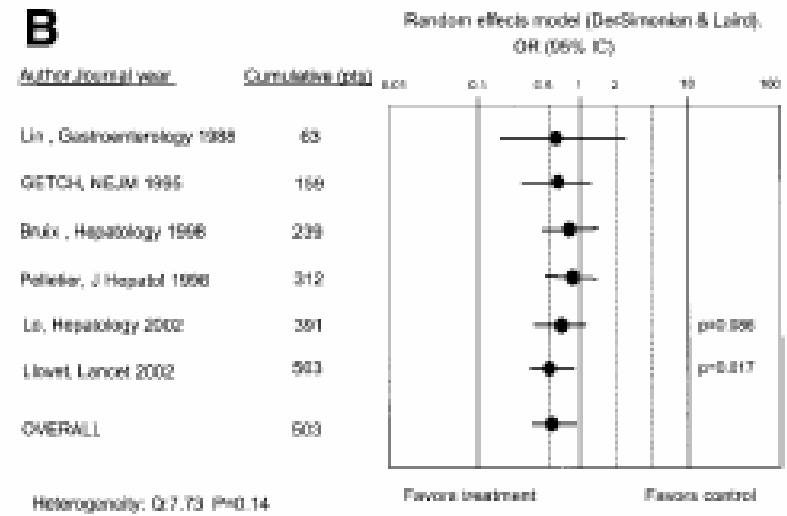
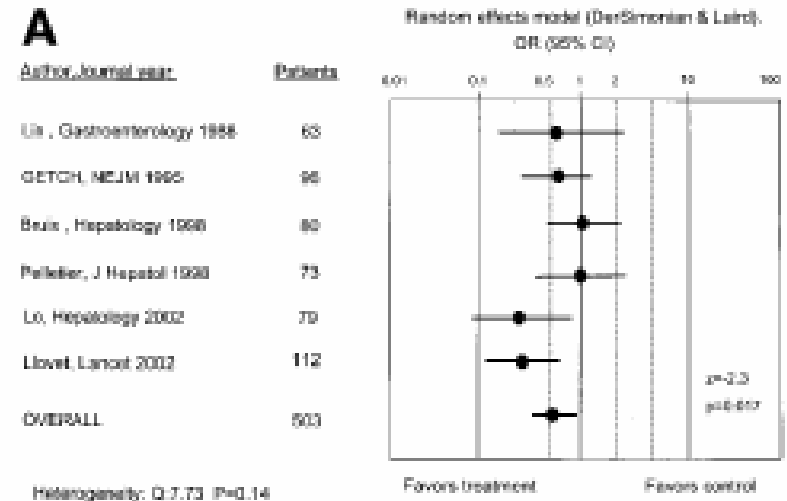


Fig. 2. (A) Meta-analysis of RCTs comparing 2-year survival with chemoembolization/embolization versus conservative management or suboptimal therapies for unresectable HCC (core group). Random effects model (OR, 0.53; 95% CI, 0.32-0.89; $P = .017$). (B) Cumulative meta-analysis according to time of publication.

Systematic Review of Randomized Trials for Unresectable Hepatocellular Carcinoma: Chemoembolization Improves Survival

Josep M. Llovet and Jordi Bruix for the Barcelona-Clinic Liver Cancer Group

Treatment vs no treatment (4 RCT, 367 pts)
 High quality trials (≥ 6 points) (5 RCT, 449 pts)
 Chemoembolization vs control (4 RCT, 323 pts)
 Embolization vs control (3 RCT, 215 pts)
 Treatment vs control- 1-yr survival (7 RCT, 545 pts)

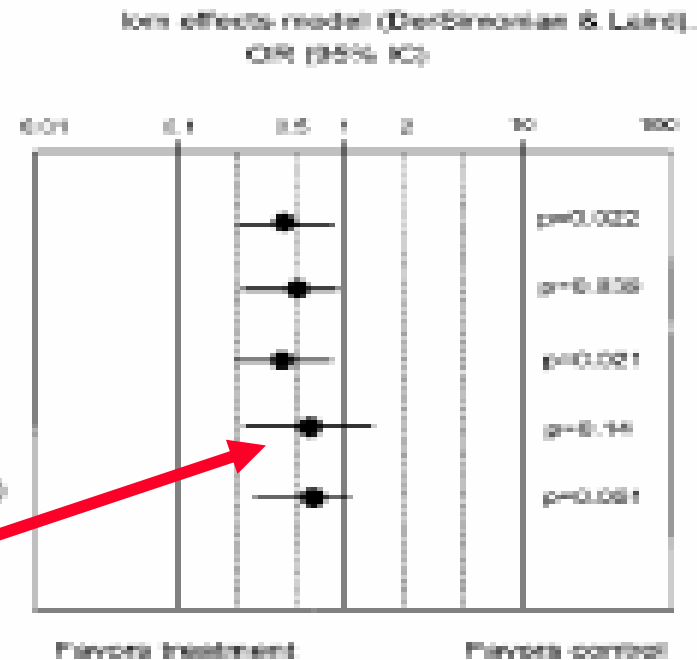


Fig. 3. Sensitivity meta-analysis of the core group (6 RCTs) reporting 2-year survival assessing embolization of RCTs with a control arm of conservative management (4 RCTs),^{28,29,22,22} the effect of chemoembolization (4 RCTs),^{29,21-22} embolization (3 RCTs),^{27,20,22} and high-quality trials (5 RCTs).²⁹⁻³² Sensitivity analysis including all studies reporting 1-year survival rates (7 RCTs).²⁷⁻³²

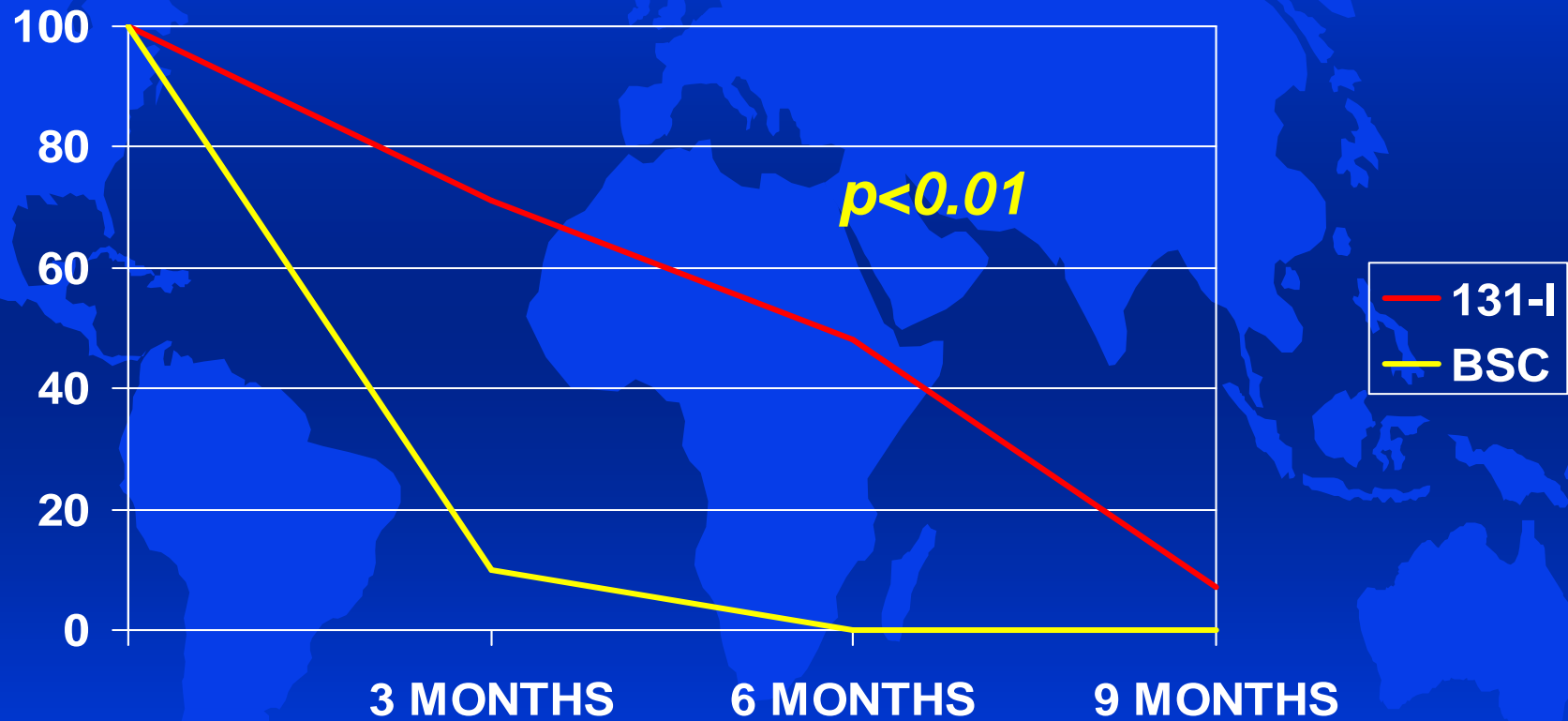
Table 4. Predictors of Survival in Patients With HCC Treated by Arterial Embolization/Chemoembolization, Multivariate Analysis

Author, journal (n)	Variables	P
Arterial embolization/chemoembolization		
Mondazzi et al., Hepatology 1994 ⁵¹ (n = 84)	Age	<.01
	Child-Pugh	<.05
	Serum bilirubin	<.025
	Tumor size	<.001
	Degree of lipiodol labelling	<.01
GETCH, N Eng J Med 1996 ¹⁴ (n = 96)	Karnofsky score	.004
	Ascites	<.001
	Serum albumin	.004
	Tumor type	.02
	Tumor mass	<.001
	Segmental portal obstruction	<.001
	Serum AFP	.009
Bruix et al., Hepatology 1998 ¹⁵ (n = 80)	Performance Status test	.005
	Serum bilirubin	.05
Llovet et al., Lancet 2002 ¹⁸ (n = 112)	Baseline variables (n = 112)	
	Treatment allocation	.02
	Inclusion of treatment response (n = 102)	
	Treatment response	.0007
Lo et al., Hepatology 2002 ¹⁷ (n = 79)	Constitutional syndrome	.04
	Treatment allocation	.006
	Portal vein thrombosis	.004
Arterial embolization and percutaneous ablation		
Tanaka et al., Cancer 1998 ⁵² (n = 83)	Child-Pugh	<.001
	Tumor diameter	.003

TACE: relevant problems

- ◆ **Schedule:** 1/ 6 weeks to 2-3 months, “a la demande” ?
- ◆ **Sections:** bilat. involvement, 1 course/lobe
- ◆ **Monitoring:** liver function, pancreatic damage, glucose homeostasis, thyroid function, US appearance (contrast media?).
- ◆ **Treatment:** antibiotics, insulin, drainage.
- ◆ **Stop:** progression, avascular lesions, complete response, liver failure
- ◆ **Morbidity (major):** 10% (ischemic cholecystitis, pancreatitis, liver abscess)
- ◆ **Mortality:** 1-5%
- ◆ **?????** Chemotherapeutic agent, embolizing tool.

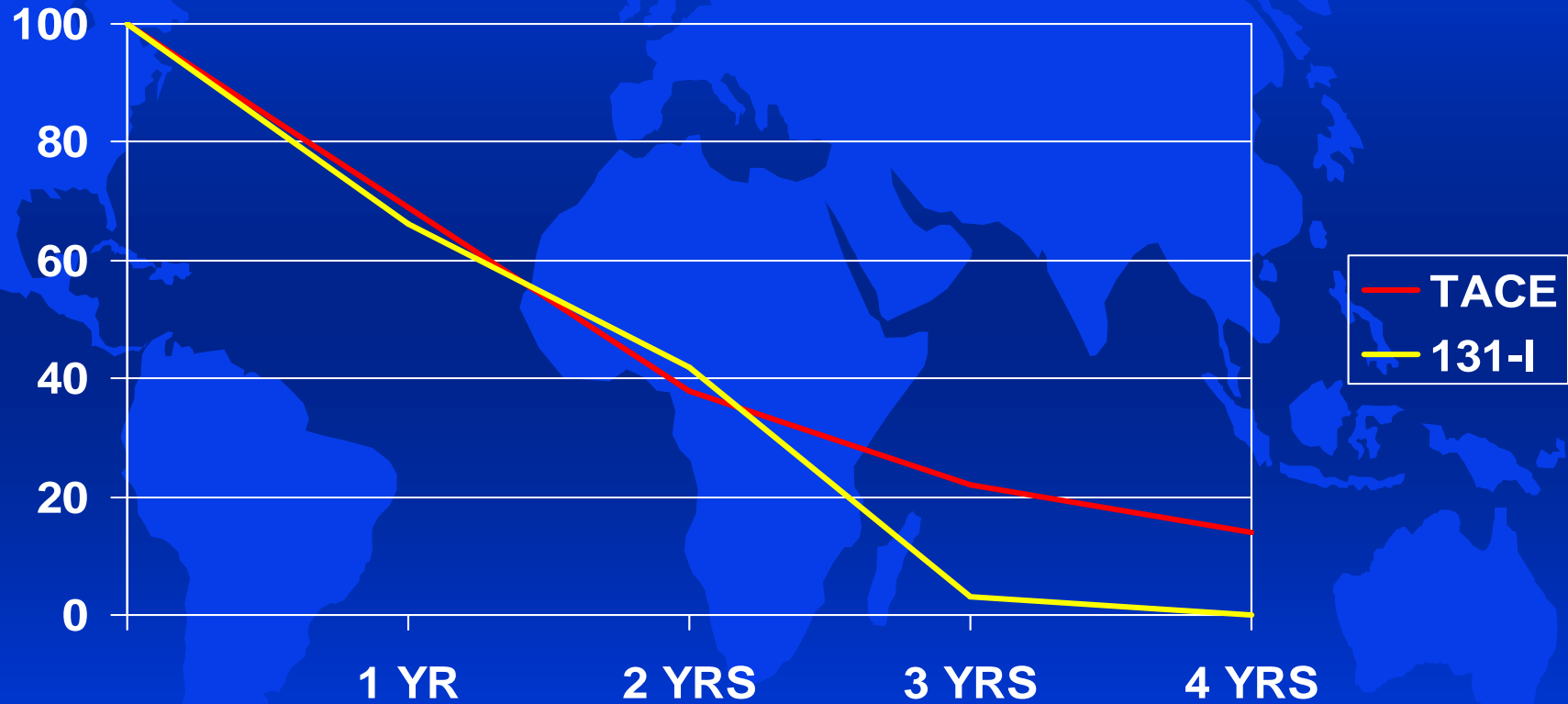
Radioactive-lipiodol Embolization (RCTs)



GOOD TOLERANCE!

Raoul, J.Nucl.Med., 1994

RADIO-ACTIVE LIPIODOL INJECTION (RCTs)



BETTER TOLERANCE!

Raoul, Hepatology, 1997

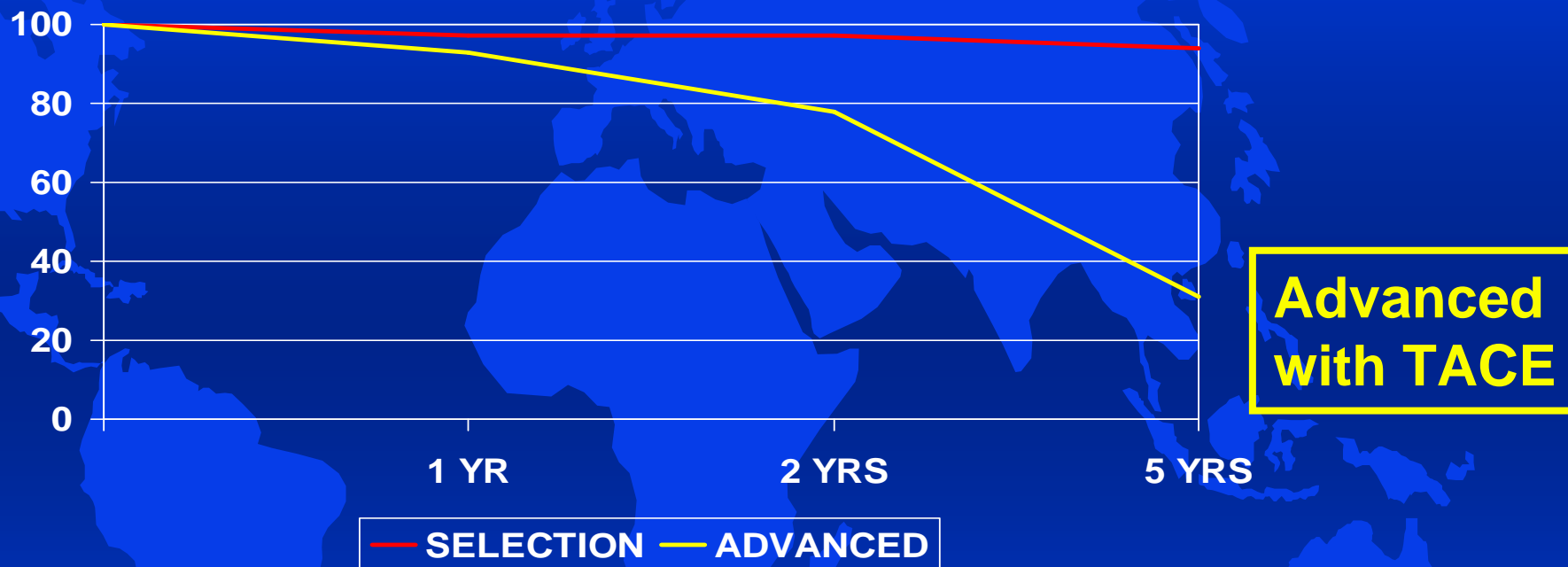
TACE: coming of age?

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TACE ad OLTx

- Multivariate analysis of predictor of survival in 67 HCC patients shows that neo-adjuvant chemoembolization reduces the risk of death (Shimoda, 2004)
- Aggressive ablation treatment.... optimizes the use of OLTx in HCC patients (Fisher 2004)

TACE ad OLTx



TACE is highly efficacious in preventing tumor progression and is associated with excellent outcome. No beneficial effect despite downstaging in advanced HCC

Graziadei, 2003

OLT x for HCC: The AISF experience

HCC transplanted: 1985-1999
587/3026 pts (20%)

Males: 505
Females: 82

M/F

HCC = 6.2

General=2.3

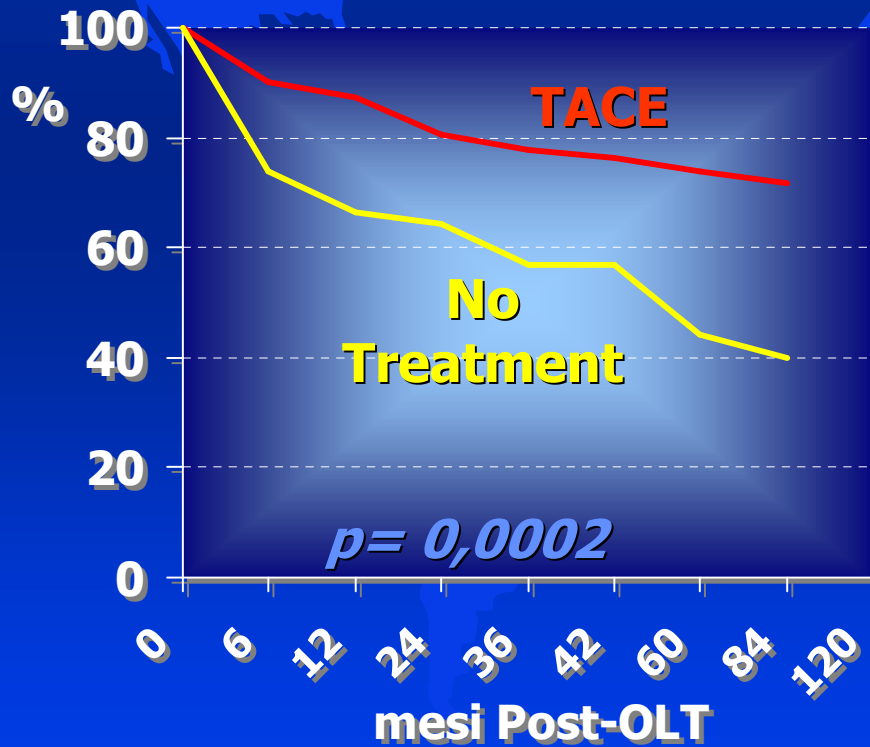
Mean age: 51,8 (range: 19-72)

"Monotematica AISF Liver Transplant Group"

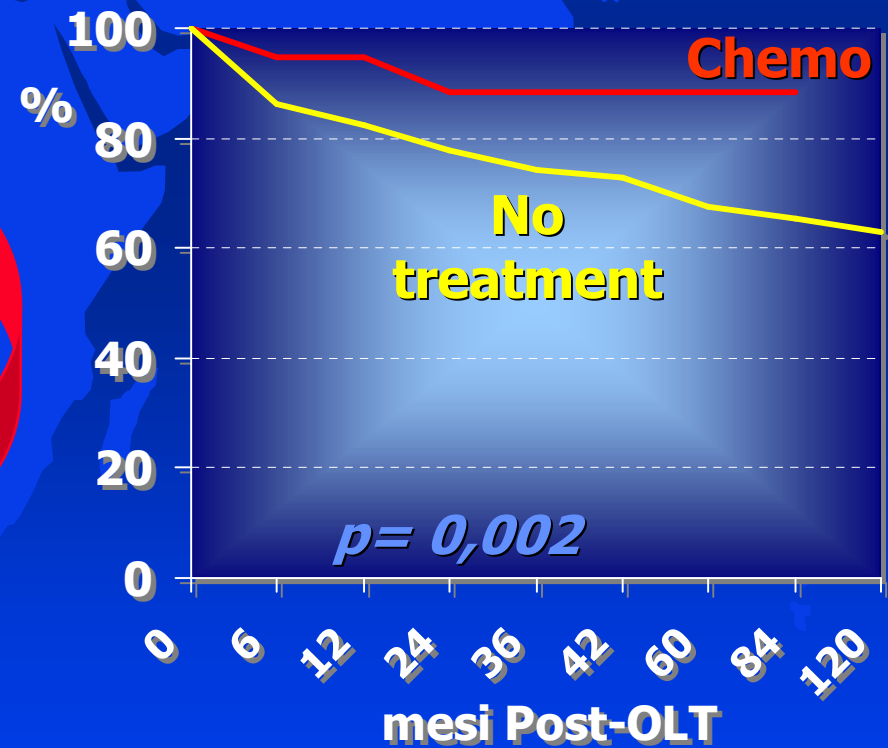
OLTx and HCC: Survival

(Kaplan-Meier survival analysis)

Neo-Adjuvant



Adjuvant



Partial Necrosis on Hepatocellular Carcinoma Nodules Facilitates Tumor Recurrence after Liver Transplantation

Matteo Ravaioli,¹ Gian Luca Grazi,¹ Giorgio Ercolani,¹ Michelangelo Fiorentino,² Matteo Cescon,¹ Rita Golfieri,³ Franco Trevisani,⁴ Walter Franco Grigioni,² Luigi Bolondi,⁵ and Antonio Daniele Pinna^{1,6}

Transplantation 2004

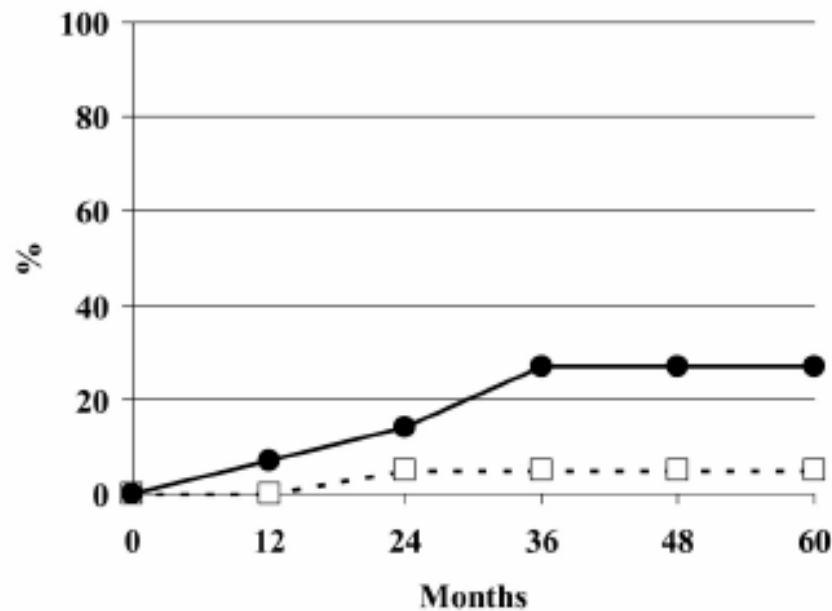


FIGURE 1. Cumulative incidence of recurrence according to the presence of partial necrosis (partial necrosis ●; no partial necrosis □).

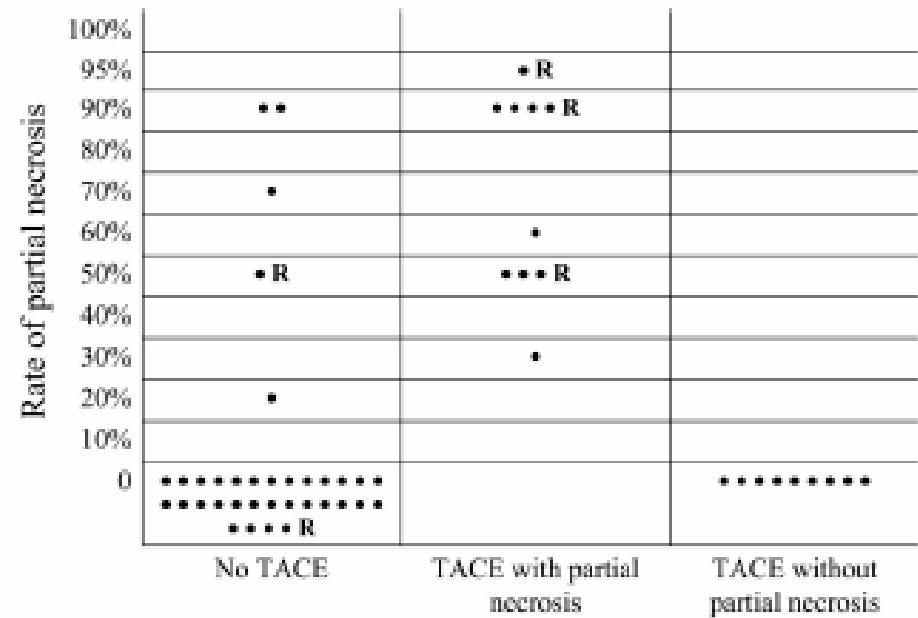


FIGURE 2. Rate of partial necrosis and Hepatocellular Carcinoma recurrences (R) distributed according to the transarterial chemoembolization (TACE) procedure (No. cases ●).

• BRIEF REPORTS •

Expression of plasma vascular endothelial growth factor in patients with hepatocellular carcinoma and effect of transcatheter arterial chemoembolization therapy on plasma vascular endothelial growth factor level

Xin Li, Gan-Sheng Feng, Chuan-Sheng Zheng, C

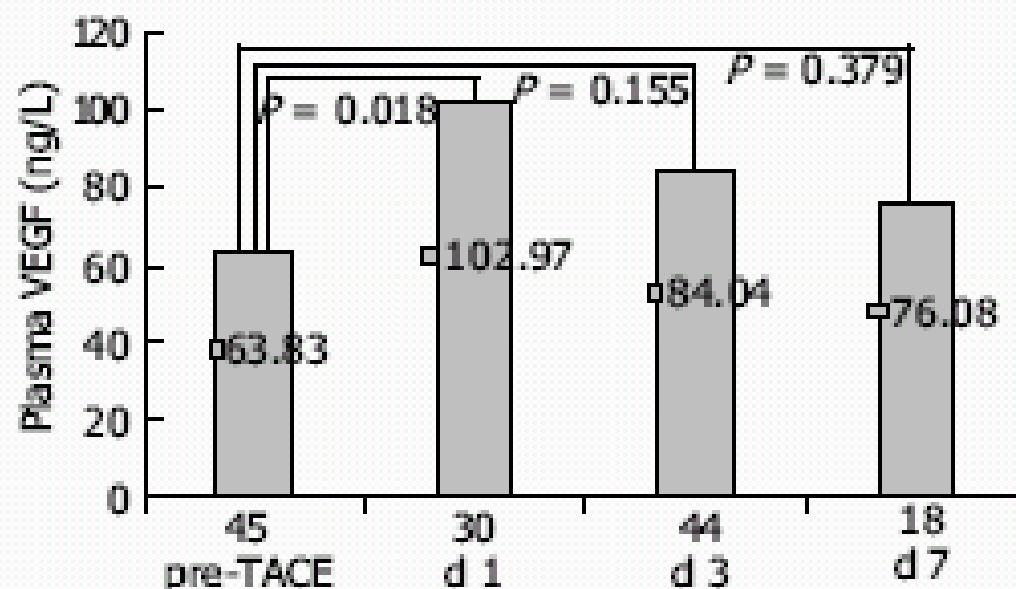


Figure 2 Effect of TACE on mean plasma vascular endothelial growth factor levels.

TACE: coming of age?

- TACE IN INTERMEDIATE STAGE HCC***
- TACE IN THE NEO-ADJUVANT TREATMENT FOR OLTx***
- TACE + LOCOREGIONAL TRANS-PARIETAL TREATMENTS (PEI-RFA)***

Sequential transarterial chemoembolization and percutaneous acetic acid injection therapy versus repeated percutaneous acetic acid injection for unresectable hepatocellular carcinoma: a prospective study

T.-L. Huo^{1*}

† S.-D. Lee¹

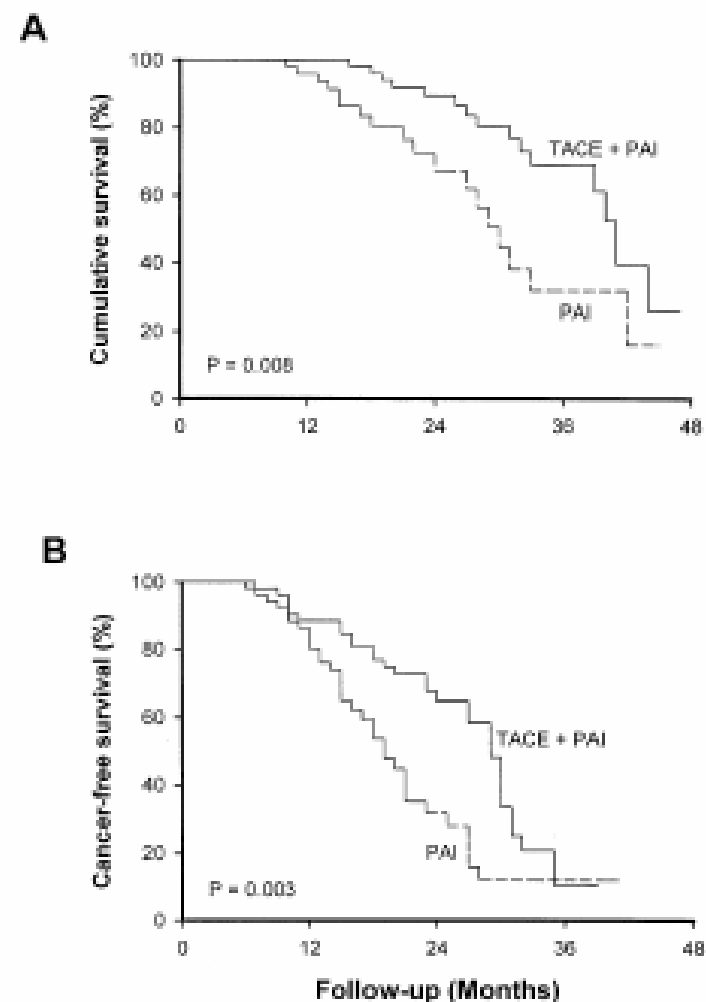
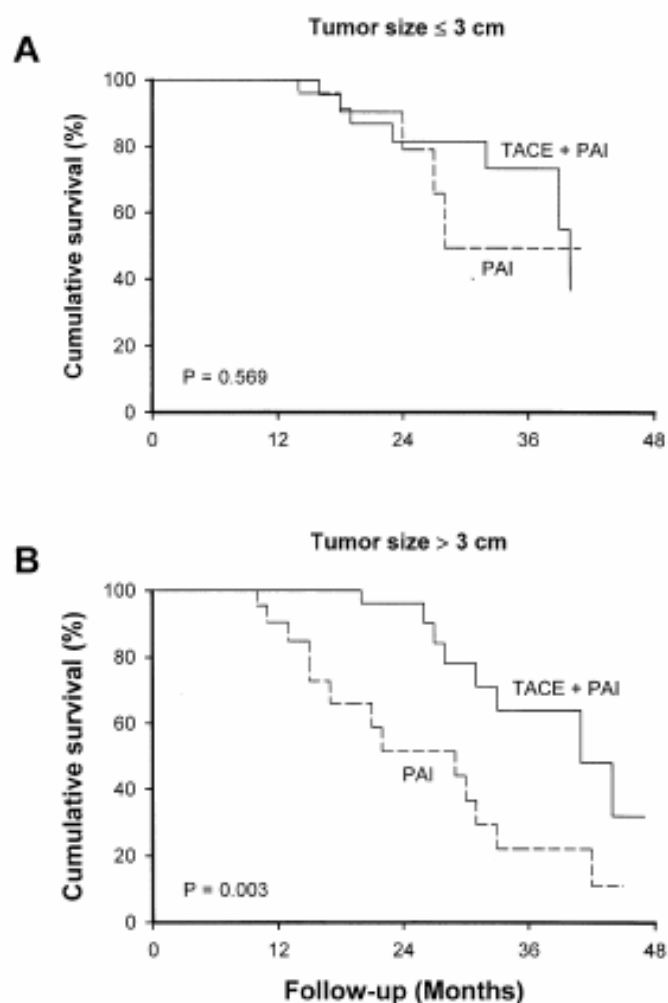


Figure 1. Comparison of (A) overall survival and (B) cancer-free survival between hepatocellular carcinoma patients treated with transarterial chemoembolization-percutaneous acetic acid injection (TACE-PAI) and PAI.

TACE-BASED COMBINATION TREATMENT IN LARGE HCC

- ◆ ***Randomized controlled trial in 53 HCC pts comparing TACE + PEI versus repeated TACE. Combined treatment....***
- ◆ ***Increases complete response***
- ◆ ***Reduces tumor recurrence***
- ◆ ***Correlates with longer survival ($p=0.1$)***
- ◆ ***Correlates with longer D.F. survival***

Bartolozzi, 1995

Combination Therapy with Transcatheter Arterial Chemoembolization and Percutaneous Ethanol Injection Compared with Percutaneous Ethanol Injection Alone for Patients with Small Hepatocellular Carcinoma

A Randomized Control Study

Masahiko Koda, M.D.
 Foshikazu Murawaki, M.D.
 Akemi Mitsuda, M.D.
 Genji Oyama, M.D.
 Ginya Okamoto, M.D.
 Foko Idobe, M.D.
 Takeaki Suou, M.D.
 Hirohisa Kawasaki, M.D.

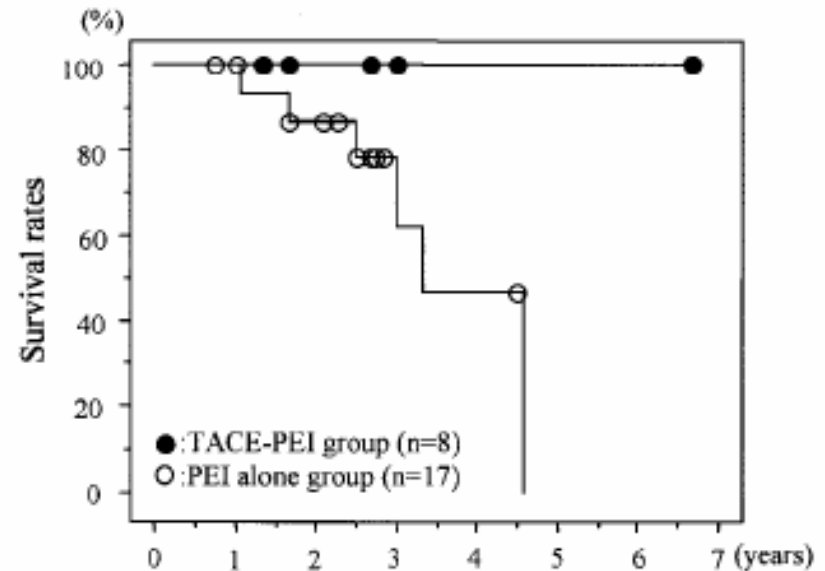
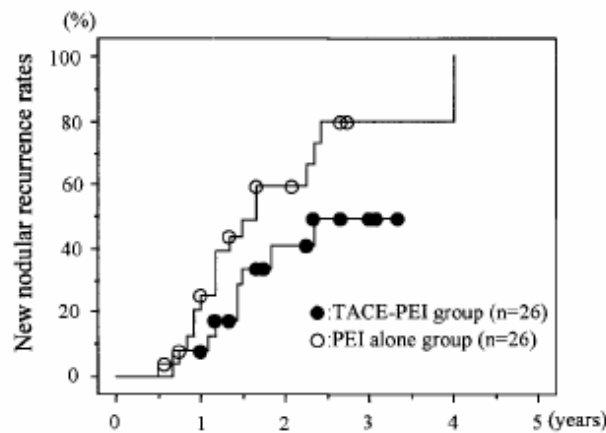
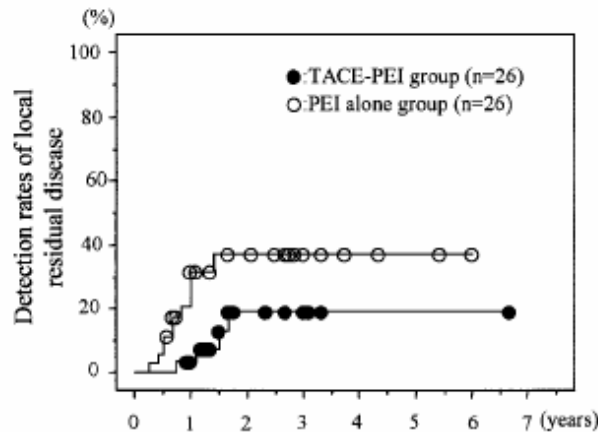


FIGURE 4. A comparison of the survival rates only in patients with hepatocellular carcinoma (HCC) tumors measuring < 2 cm in greatest dimension between the group that received transcatheter arterial chemoembolization (TACE) and percutaneous ethanol injection (PEI) (TACE-PEI group) and the group that received PEI alone (PEI alone group). The survival rate in the TACE-PEI group was higher than that in the PEI alone group ($P < 0.01$).

In summary....

Despite lack of standardization, TACE....

- ◆ *Is effective in prolonging survival in selected HCC patients in the intermediate stage (A1a)*
- ◆ *Is effective in reducing drop-out rate and increasing survival in HCC patients undergoing OLTx (B2b)*
- ◆ *When associated with percutaneous treatments, is effective in increasing response rate and prolonging survival in (large) HCC (A1b-B2b)*



Thanks !