Randomized Phase II Study Evaluating The Tolerability Of Adjuvant Docetaxelbased Chemotherapy For Completely Resected Stage IB-II Non-Small Cell Lung Cancer (NSCLC): TOLEDO trial

Lionel Bosquée on behalf of the TOLEDO-investigators





- Antwerps Kanker Register 2000:
  - 989 new cases of lung cancer in the province of Antwerp.
  - 83% were histologically of NSCLC type.
  - About one quarter (27%) of the NSCLCs were resected.
  - →≥ 200 resections for NSCLC / year in the province of Antwerp.

 Long-term survival for resected NSCLC remains disappointing:

- 5-year survival rates range from 73% for pathologic stage IA disease to 39% for pathologic pathologic stage IIB disease.
- Micrometastatic cancer cells are present in bone marrow of >30% of patients with operable NSCLC.
- The majority of relapses occur at distant sites.



Survival results of adjuvant chemotherapy from 5 recent randomised placebo-controlled trial

	Ν	Stage	# cycl	OS 5-yr	HR	p value
ALPI	1209	I-IIIA	3	+ 3%	.96	0.6
BLT	381	1-111	3	+0-1%	1	1
IALT	1867	I-IIIA	3-4	+ 4%	.86	<0.03
JBR.10	482	IB-II	4	+ 15%	.69	0.012
CALGB	344	IB	4	+ 12%*	.62	0.028

Furthermore, systematic reviews and meta-analysis confirm that adjuvant chemotherapy is associated with improved survival compared with surgical intervention alone



Sedrakyan ea. J Thorac Cardiovasc Surg 2004, 128: 414-419.

- Adjuvant chemotherapy is associated with significant morbidity / toxicity:
  - IALT: 22.6 % grade 4 toxicity
  - ALPI: 12 % grade 4 toxicity
  - BLT: overall 30% grade 3/4 toxicity (55% of patients receiving NP were reported as having grade 3 or 4 toxicity compared to 27% of the MIC and 17% of the MVP patients)
  - CALGB 9633: 28% grade 4 toxicity
  - NCIC trial: reported only combined grade 3 and 4 (i.e. 73% gr 3-4 neutropenia and 7% febrile neutropenia)



Drug delivery results of adjuvant chemotherapy from 5 recent randomised placebo-controlled trial

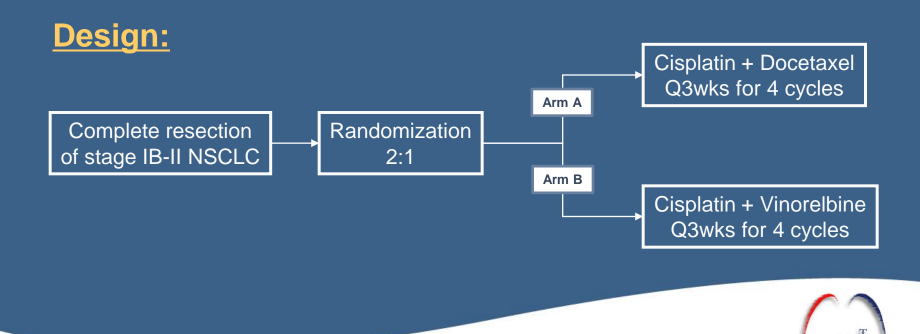
	Ν	Chemo	Chemo completion	PORT
ALPI	1209	MVdP	69%	43%
IALT	1867	P-Vb, P-Vd,	74%	23%
		P-Vrb or P-V	′P16	
Big Lung Trial	381	MIP, MVbP	64%	-
		P-Vb or P-Vo	d	
NCIC JBR.10	482	P-Vrb	<mark>65%</mark>	-
CALGB 9633	344	Cb-Pacli	85%	-



## **Toledo-trial: aim and design**

### <u> Aim:</u>

To evaluate the tolerability of four cycles of adjuvant docetaxel plus cisplatin in patients with completely resected stage IB/II NSCLC.



# **Toledo-trial: endpoints**

#### Primary endpoints:

- 1. Success of delivery treatment
  - (To be considered as a "success" a patient should have received at least 3 cycles of chemotherapy, and have a relative dose intensity of 80% or above)
- 2. Toxicity

(Occurrence of any grade 4 non-haematological toxicity)

#### Secondary endpoints:

- 1. Overall toxicity
- 2. Progression free survival and overall survival



## **Toledo-trial: inclusion criteria**

- 1. Completely resected (R0\*) pathological stage IB or II NSCLC
- The first cycle of chemotherapy should be started within 60 days of resection
- 3. KS 70-100 (see appendix II)
- 4. Age: 18 75 year
- 5. Weight loss < 10% over previous 6 months
- 6. Adequate haematological function:
  - ANC > 1,5x 109/L
  - Platelets > 100x 109/L
  - Hgb > 10 g/dl

\*R0 resection: a resection is considered a complete resection if microscopic examination shows a radical resection of the primary tumor with tumor-free resction margins and if the highest prelevated mediastinal lymph node is tumor free



## **Toledo-trial: inclusion criteria**

- 7. Adequate renal and liver function:
  - Creatinine ≤ 1.5 mg/dL, or calculated creatinine clearance ≥ 60 ml/min (see appendix III for calc. creat. clearance)
  - Total bilirubin  $\leq$  ULN
  - Alkaline phosphatase ≤ 5.0x ULN
  - AST/ALT  $\leq 2.0x$  ULN
  - Serum calcium  $\leq 1.1 \times ULN$
- 8. Signed informed consent prior to beginning protocol specific procedures.
- 9. Women of childbearing potential must be nonpregnant, non-lactating and use adequate contraception during study treatment.



## **Toledo-trial: exclusion criteria**

- 1. Previous chemo- or radiotherapy for NSCLC
- 2. Bronchoalveolar cell subtype (ie those cases which show no stromal, pleural, or lymphatic invasion according to the WHO classification of 1999).
- 3. Second active primary malignancy (except basocellular CA of the skin, adequately treated CA in situ of the cervix, low-grade prostate cancer or other cancer from which the patient has been disease free for at least five years)
- 4. Serious concomitant medical disease (i.e. active infection, preexisting neuropathy, AMI less than 6 months old), immunosuppression or psychiatric disease that, in the opinion of the investigator, would compromise the safety of the patient or compromise the patient's ability to complete the study.
- 5. Pregnant or breast-feeding females.
- 6. Difficulties with adequate follow-up



## **Toledo-trial: randomization procedure**

- Patients will be randomized on a 2:1 basis, with stratification according to clinical stage: IB versus II (using two computer-generated randomization lists). Centralized randomization was used to conceal the allocation sequence to the investigator at time of enrolment.
- The Registration/Randomization center will inform the investigator of the treatment sequence: arm A or arm B. The study treatment had to start within 7 days from randomization and within 60 days of resection.
- The study was not blinded. Commercially available products were used.



## **Toledo-trial: treatment administration**

### <u>Arm A:</u>

cisplatin 75 mg/m2 on day 1 and docetaxel 75 mg/m2 on day 1 of each cycle

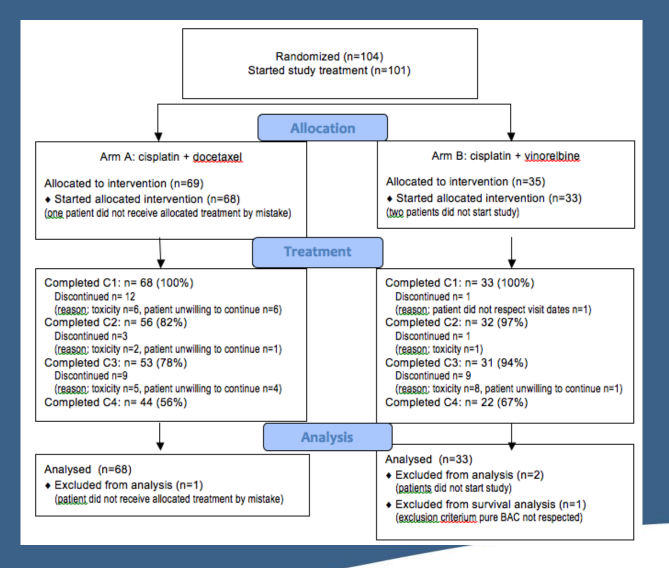
### <u>Arm B:</u>

cisplatin 80 mg/m2 on day 1 and vinorelbine 25 mg/m2 on day 1 and 8 of each cycle

- A cycle is defined as an interval of 21 days
- All patients will be treated with 4 cycles of adjuvant chemotherapy, unless there is the occurrence of unacceptable toxicity or early progression.



## **Patient disposition**





# **Patient demographics**

	all patients	Cis-Doc	Cis-VRB
Ν	101	68	33
Age (Y):			
Median	61	61	62
(range)	(37-75)	(37-75)	(43-75)
Gender:			
Male	68	49	19
Female	33	19	14
Performance status:			
KPS 90-100	88	62	26
KPS 70-80	13	6	7
Lung function:			
FEV <sub>1</sub> (L): median	1,88	2,20	1,79
(range)	(1,02-4,60)	(1,02-4,60)	(1,10-2,39)
FEV <sub>1</sub> (%pred): median	66	70	60
(range)	(32-117)	(32-117)	(41-94)
DLCO (%pred): median	56	56	56
(range)	(20-130)	(20-125)	(35-130)
BSA (mg/m <sup>2</sup> ):			
Median	1,85	1,86	1,85
(range)	(1,41-2,66)	(1,49-2,66)	(1,41-2,29)



# Patient demographics

	all patients	Cis-Doc	Cis-VRB
N	all patients		
N	101	68	33
Interval between ourgany and			
Interval between surgery and randomization (in days):			
Median	38	36	40
	(14-69)	(14-69)	(25-69)
(range)	(14-09)	(14-09)	(20-09)
Type of surgery	- (0/)	~ (0/)	~ (0/)
Type of surgery	n (%)	n (%)	n (%)
pneumonectomy right lung	5 (5)	3 (4)	2 (6)
pneumonectomy left lung	14 (14)	11 (16)	3 (9)
lobectomy RUL	12 (12)	9 (13)	3 (9)
lobectomy RML	2 (2)	2 (3)	-
lobectomy RLL	14 (14)	9 (13)	5 (15)
lobectomy LUL	24 (24)	13 (19)	11 (33)
lobectomy LLL	14 (14)	10 (15)	4 (12)
bilobectomy RUL+RML	5 (5)	3 ( 4)	2 (6)
bilobectomy RML+RLL	2 (2)	1 (1)	1 (3)
bilobectomy RUL+RLL	1 (1)	1 (1)	-
bilobectomy RUL+RML+RLL	1 (1)	1 (1)	-
limited resection wedge	2 (2)	2 (3)	-
limited resection	1 (1)	-	1 (3)
segmentectomy			N 7
sleeve lobectomy RUL	2 (2)	2 (3)	-
pneumonectomy left	1 (1)		1 (3)
lung+partial thorax wall		-	
RUL +part of 2 ribs	1 (1)	1 (1)	-



# **Patient demographics**

	all patients	Cis-Doc	Cis-VRB
N	101	68	33
	(01)	(0/)	(01)
Histological tumor type	n (%)	n (%)	n (%)
adenocarcinoma	53 (52)	36 (53)	17 (52)
squamous cell	33 (33)	25 (37)	8 (24)
large cell	9 (9)	6 (9)	3 (9)
adenocarcinoma + squamous	1 (1)	1 (1)	-
adenocarcinoma + BAC	2 (2)	-	2 (6)
adenocarcinoma + large cell	1 (1)	-	1 (3)
muco-epidermoid	1 (1)	-	1 (3)
BAC	1 (1)	-	1 (3)
Histology differentiation			
well differentiated	12 (12)	8 (12)	4 (12)
moderately differentiated	49 (49)	34 (50)	15 (45)
poorly differentiated	32 (32)	23 (34)	9 (27)
unknown	8 (8)	3 (4)	5 (15)
Pathological TNM*			
T1N1M0	10 (10)	7 (10)	3 (9)
T2N0M0	56 (55)	37 (54)	19 (58)
T2N1M0	29 (29)	18 (26)	11 (33)
T3N0M0	6 (6)	6 (9)	-
Cancer stage*			
IB	56 (55)	37 (54)	19 (58)
IIA	12 (12)	8 (12)	4 (12)
IIB	33 (33)	23 (34)	10 (30)



\* according to 6<sup>th</sup> TNM classification

### Succes of treatment delivery (≥3 cycles with RDI ≥80%)

Succes of delivery	Cis-Doc		Cis-VRB
treatment	n = 68		n = 33
Succes cisplatinum: n (%)	52 (76)	Succes cisplatinum: n (%)	26 (79)
Succes docetaxel: n (%)	52 (76)	Succes vinorelbine: n (%)	25 (76)
Succes Cis-Doc: n (%)	52 (76)	Succes Cis-VRB: n (%)	23 (70)



# Reason for dose modification or delay

Cis-Doc 0					Cis-V	RB				
	Cycle 1	Cycle 2	Cycle 3	Cycle 4	Total	Cycle 1	Cycle 2	Cycle 3	Cycle 4	Total
Adverse event drug related:										
-hematological toxicity	-	4	3	-	7	-	11	8	14	33
-non-hematological toxicity	1	5	4	8	18	-	2	4	3	9
Adverse event not drug related	-	-	1	1	2	-	-	-	1	1
Patient unwilling/unable to continue	-	6	1	4	11	-	-	-	1	1
Organizational reason or other	-	2	4	3	9	2	3	1	-	6
Total	1	17	13	16	47	2	16	13	19	50



# Reason for withdrawal from chemotherapy

	Cis-Doc				Cis-VRB				
	Cycle 1	Cycle 2	Cycle 3	Cycle 4	Total	Cycle 2	Cycle 3	Cycle 4	Total
Adverse event drug related:									
-hematological toxicity	-	2	1	-	3	-	-	4	4
-non-hematological toxicity	1	3	1	5	10	-	1	3	4
-hematologic+non-hematologic	-	-	-	-	-	-	-	1	1
Patient unwilling/unable to continue	-	6	1	4	11	-	-	1	1
Investigator decision because patient did not respect visit dates	-	-	-	-	-	1	-	-	1
Total	1	11	3	9	24	1	1	9	11

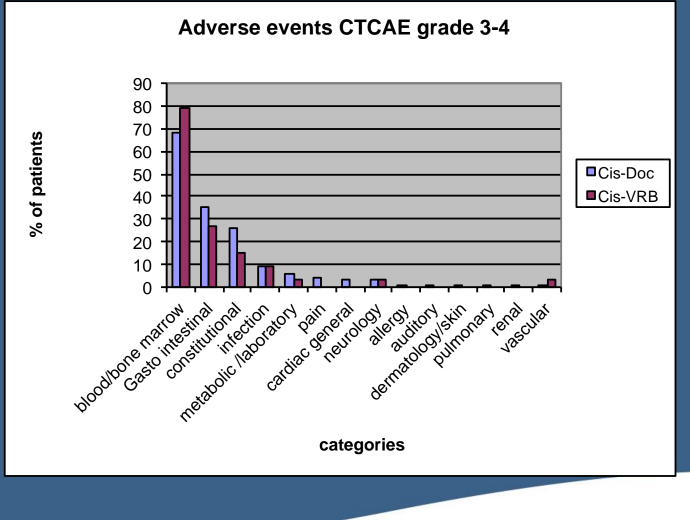


## **Grade 3-4 adverse events**

	Cis-Doo	c			Cis-VRE	3		
CTCAE categories	Grade	Grade	Grade		Grade	Grade	Grade	
	3	4	3-4		3	4	3-4	
	n	n	n	%	n	n	n	%
Blood/bone marrow	17	29	46	(68)	10	16	26	(79)
Gastro intestinal	24		24	(35)	9		9	(27)
Constitutional	18		18	(26)	5		5	(15)
Infection	6		6	(9)	3		3	(9)
Metabolic								
/laboratory	4		4	(6)	1		1	(3)
Pain	3		3	(4)	-		-	
Cardiac general	2		2	(3)	-		-	
Neurology	1	1	2	(3)	1		1	(3)
Allergy	1		1	(1)	-		-	
Auditory	1		1	(1)	-		-	
Dermatology/skin	1		1	(1)	-		-	
Pulmonary	1		1	(1)	-		-	
Renal	1		1	(1)	-		-	
Vascular	1		1	(1)	1		1	(3)
Hematological			46	(68)			26	(79)
Non hematological			36	(53)			14	(42)
Hematological + non hema	atological ev	ents	57	(84)			28	(85)



## **Grade 3-4 adverse events**



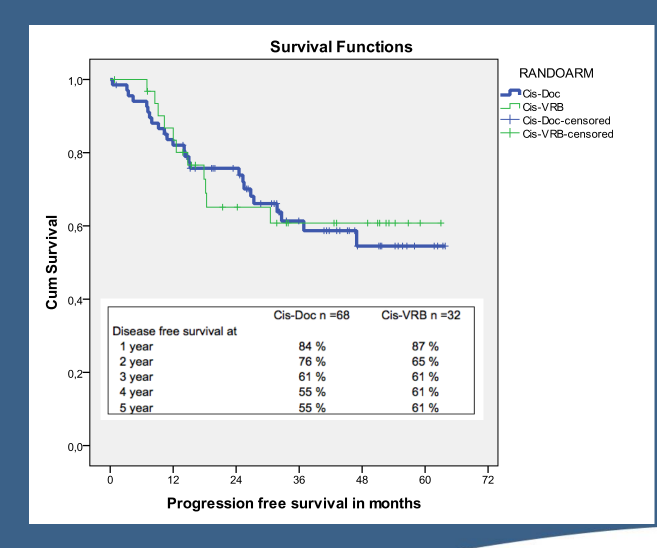


## Site of disease recurrence

	Cis-Doc	Cis-VRB
	n =68	n =32
Type of recurrence	n (%)	n (%)
Local (ipsilateral lung and/or hilum)	3 (4,4)	1 (3,1)
Regional mediastinum	1 (1,5)	1 (3,1)
Local + regional mediastinum	1 (1,5)	
Local + metastasis	2 (2,9)	2 (6,3)
Regional mediastinum + metastasis	1 (1,5)	1 (3,1)
Regional supraclavicular	1 (1,5)	1 (3,1)
Second primary tumor in contralateral lung	3 (4,4)	
Metastasis	11 (16,2)	4 (12,5)
Died from lung cancer		1 (3,1)
Died non lung cancer related	2 (2,9)	
Total number of recurrence	25 (36,8)	11 (34,4)

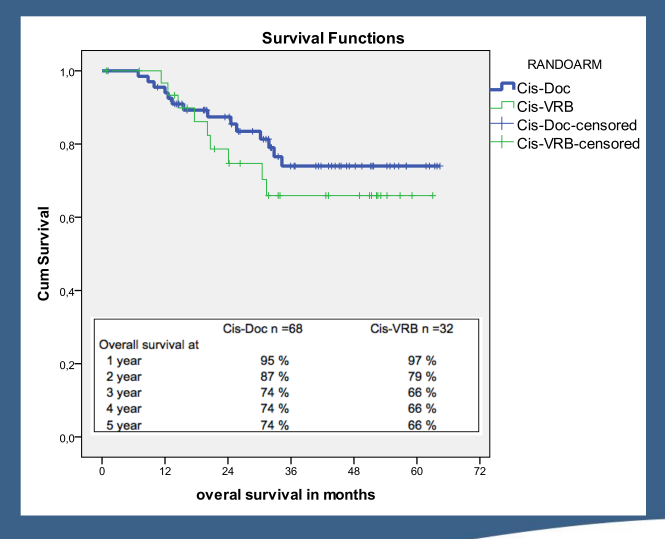


## **TOLEDO:** disease-free survival ~ arm



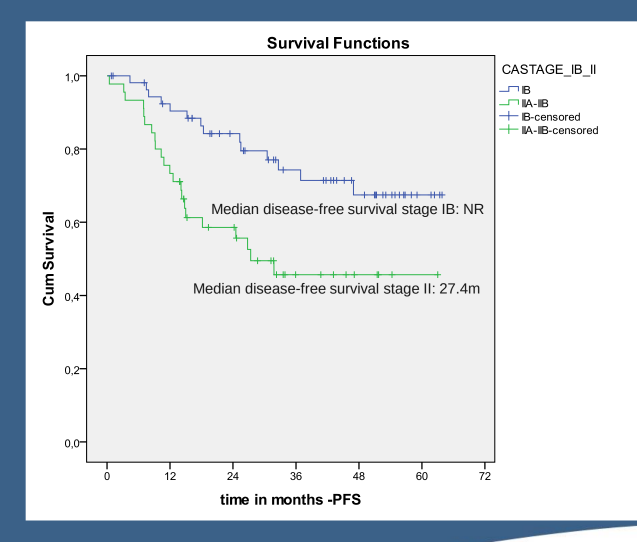


## **TOLEDO:** overall survival ~ arm



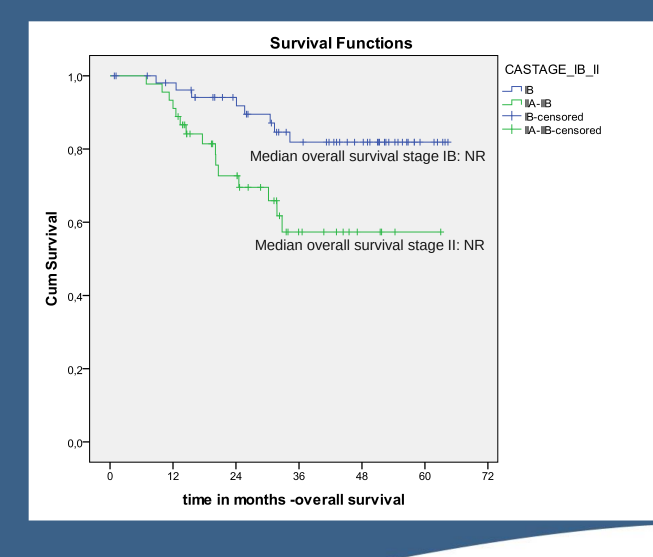


## **TOLEDO:** disease-free survival ~ stage





# **TOLEDO:** overall survival ~ stage





# **TOLEDO trial: conclusions**

- Adjuvant treatment with 3 week schedule Pt-TXT is :
  - feasible, tolerable
  - 1st endpoints reached: 76% delivery success
- 3 W schedule Adjuvant treatment PT- NVB shows
  - A better tolerability than a historical 4 w schedule
  - A 70% delivery success
- Data robust enough to propose a Random PhIII study based on DFS and Survival



### Investigational sites: 15 in Belgium - 3 in Holland

Site	Principal investigator	N° patients
Centre Hospitalier Universitaire de Liège, B	Dr L Bosquée	13
Amphia ziekenhuis Breda, NI	Dr J Aerts	12
ORBIS medisch centrum Geleen, NI	Dr F Peters	10
Universitair Ziekenhuis Antwerpen, B	Prof Dr P Germonpré	8
AZ Nikolaas, St Niklaas, B	Dr K Deschepper	8
ZNA Middelheim Antwerpen, B	Dr D Galdermans	7
AZ Turnhout, B	Dr P Driesen	7
AZ St Maarten Duffel Mechelen, B	Dr M Lambrechts	6
GZA St Vincentius, Antwerpen, B	Dr I Stappaerts	5
AZ Monica, Antwerpen Deurne, B	Dr T Huybrechs	5
Stedelijk ziekenhuis Aalst, B	Dr L Van Moorter	5
C.H.R de la Citadelle Liège, B	Dr F Bustin	4
GZA St Augustinus, Wilrijk, B	Dr D Verresen	3
AZ St Jozef Bornem Willebroek, B	Dr AM Morel	2
AZ St Elisabeth, Herentals, B	Dr Y Mentens	2
Ommelander ziekenhuisgroep Delfzijl Winschoten, NI	Dr R Pieterman	2
ZNA Jan Palfijn, Merksem, B	Dr C Van Schaardenburg	1
AZ OLV Aalst Asse, B	Dr P Vercauter	1

